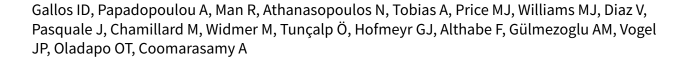


Cochrane Database of Systematic Reviews

Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis (Review)



Gallos ID, Papadopoulou A, Man R, Athanasopoulos N, Tobias A, Price MJ, Williams MJ, Diaz V, Pasquale J, Chamillard M, Widmer M, Tunçalp Ö, Hofmeyr GJ, Althabe F, Gülmezoglu AM, Vogel JP, Oladapo OT, Coomarasamy A. Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis. *Cochrane Database of Systematic Reviews* 2018, Issue 12. Art. No.: CD011689. DOI: 10.1002/14651858.CD011689.pub3.

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[Intervention Review]

Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis

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ABSTRACT

Background

Postpartum haemorrhage (PPH) is the leading cause of maternal mortality worldwide. Prophylactic uterotonic agents can prevent PPH, and are routinely recommended. The current World Health Organization (WHO) recommendation for preventing PPH is 10 IU (international units) of intramuscular or intravenous oxytocin. There are several uterotonic agents for preventing PPH but there is still uncertainty about which agent is most effective with the least side effects. This is an update of a Cochrane Review which was first published in April 2018 and was updated to incorporate results from a recent large WHO trial.

Objectives

To identify the most effective uterotonic agent(s) to prevent PPH with the least side effects, and generate a ranking according to their effectiveness and side-effect profile.

Search methods

We searched the Cochrane Pregnancy and Childbirth's Trials Register, ClinicalTrials.gov, the World Health Organization (WHO) International Clinical Trials Registry Platform (ICTRP) (24 May 2018), and reference lists of retrieved studies.



Selection criteria

All randomised controlled trials or cluster-randomised trials comparing the effectiveness and side effects of uterotonic agents with other uterotonic agents, placebo or no treatment for preventing PPH were eligible for inclusion. Quasi-randomised trials were excluded. Randomised trials published only as abstracts were eligible if sufficient information could be retrieved.

Data collection and analysis

At least three review authors independently assessed trials for inclusion and risk of bias, extracted data and checked them for accuracy. We estimated the relative effects and rankings for preventing PPH ≥ 500 mL and PPH ≥ 1000 mL as primary outcomes. Secondary outcomes included blood loss and related outcomes, morbidity outcomes, maternal well-being and satisfaction and side effects. Primary outcomes were also reported for pre-specified subgroups, stratifying by mode of birth, prior risk of PPH, healthcare setting, dosage, regimen and route of administration. We performed pairwise meta-analyses and network meta-analysis to determine the relative effects and rankings of all available agents.

Main results

The network meta-analysis included 196 trials (135,559 women) involving seven uterotonic agents and placebo or no treatment, conducted across 53 countries (including high-, middle- and low-income countries). Most trials were performed in a hospital setting (187/196, 95.4%) with women undergoing a vaginal birth (71.5%, 140/196).

Relative effects from the network meta-analysis suggested that all agents were effective for preventing PPH \geq 500 mL when compared with placebo or no treatment. The three highest ranked uterotonic agents for prevention of PPH \geq 500 mL were ergometrine plus oxytocin combination, misoprostol plus oxytocin combination and carbetocin. There is evidence that ergometrine plus oxytocin (RR 0.70, 95% CI 0.59 to 0.84, moderate certainty), carbetocin (RR 0.72, 95% CI 0.56 to 0.93, moderate certainty) and misoprostol plus oxytocin (RR 0.70, 95% CI 0.58 to 0.86, low certainty) may reduce PPH \geq 500 mL compared with oxytocin. Low-certainty evidence suggests that misoprostol, injectable prostaglandins, and ergometrine may make little or no difference to this outcome compared with oxytocin.

All agents except ergometrine and injectable prostaglandins were effective for preventing PPH \geq 1000 mL when compared with placebo or no treatment. High-certainty evidence suggests that ergometrine plus oxytocin (RR 0.83, 95% CI 0.66 to 1.03) and misoprostol plus oxytocin (RR 0.88, 95% CI 0.70 to 1.11) make little or no difference in the outcome of PPH \geq 1000 mL compared with oxytocin. Low-certainty evidence suggests that ergometrine may make little or no difference to this outcome compared with oxytocin meanwhile the evidence on carbetocin was of very low certainty. High-certainty evidence suggests that misoprostol is less effective in preventing PPH \geq 1000 mL when compared with oxytocin (RR 1.19, 95% CI 1.01 to 1.42). Despite the comparable relative treatment effects between all uterotonics (except misoprostol) and oxytocin, ergometrine plus oxytocin, misoprostol plus oxytocin combinations and carbetocin were the highest ranked agents for PPH \geq 1000 ml

Misoprostol plus oxytocin reduces the use of additional uterotonics (RR 0.56, 95% CI 0.42 to 0.73, high certainty) and probably also reduces the risk of blood transfusion (RR 0.51, 95% CI 0.37 to 0.70, moderate certainty) when compared with oxytocin. Carbetocin, injectable prostaglandins and ergometrine plus oxytocin may also reduce the use of additional uterotonics but the certainty of the evidence is low. No meaningful differences could be detected between all agents for maternal deaths or severe morbidity as these outcomes were rare in the included randomised trials where they were reported.

The two combination regimens were associated with important side effects. When compared with oxytocin, misoprostol plus oxytocin combination increases the likelihood of vomiting (RR 2.11, 95% CI 1.39 to 3.18, high certainty) and fever (RR 3.14, 95% CI 2.20 to 4.49, moderate certainty). Ergometrine plus oxytocin increases the likelihood of vomiting (RR 2.93, 95% CI 2.08 to 4.13, moderate certainty) and may make little or no difference to the risk of hypertension, however absolute effects varied considerably and the certainty of the evidence was low for this outcome.

Subgroup analyses did not reveal important subgroup differences by mode of birth (caesarean versus vaginal birth), setting (hospital versus community), risk of PPH (high versus low risk for PPH), dose of misoprostol (≥ 600 mcg versus < 600 mcg) and regimen of oxytocin (bolus versus bolus plus infusion versus infusion only).

Authors' conclusions

All agents were generally effective for preventing PPH when compared with placebo or no treatment. Ergometrine plus oxytocin combination, carbetocin, and misoprostol plus oxytocin combination may have some additional desirable effects compared with the current standard oxytocin. The two combination regimens, however, are associated with significant side effects. Carbetocin may be more effective than oxytocin for some outcomes without an increase in side effects.

PLAIN LANGUAGE SUMMARY

Which drug is best for reducing excessive blood loss after birth?

What is the issue?



The aim of this Cochrane Review was to find out which drug is most effective in preventing excessive blood loss at childbirth and has the least side effects. We collected and analysed all the relevant studies to answer this question (date of search: 24 May 2018).

Why is this important?

Excessive bleeding after birth is the most common reason why mothers die in childbirth worldwide. Although most women will have moderate bleeding at birth, others may bleed excessively, and this can pose a serious risk to their health and life. To reduce excessive bleeding at birth, the routine administration of a drug to contract the uterus (uterotonic) has become standard practice across the world.

Different drugs given routinely at birth have been used for reducing excessive bleeding. They include oxytocin, misoprostol, ergometrine, carbetocin, injectable prostaglandins and combinations of these drugs, each with different effectiveness and side effects. Some of the side effects identified include: vomiting, high blood pressure and fever. Currently, oxytocin is recommended as the standard drug to reduce excessive bleeding. We analysed all the available evidence to compare the effectiveness and side-effect profiles for each drug.

What evidence did we find?

We found 196 studies involving 135,559 women. We compared seven uterotonic agents against each other and against women receiving no uterotonic. Studies were conducted across 53 countries. In most studies women were giving birth normally and in a hospital.

The analysis suggests that all drugs are effective for preventing blood loss that equals or exceeds 500 mL when compared with no routine uterotonic treatment. Compared with oxytocin (the standard recommended drug), the three best drugs for this outcome were a combination of ergometrine plus oxytocin, carbetocin, and a combination of misoprostol plus oxytocin. We found the other drugs misoprostol, injectable prostaglandins, and ergometrine may make little or no difference to this outcome compared with oxytocin.

All drugs except ergometrine and injectable prostaglandins are effective for preventing blood loss that equals or exceeds 1000 mL when compared with no treatment. Ergometrine plus oxytocin and misoprostol plus oxytocin make little or no difference in this outcome compared with oxytocin. It is uncertain whether carbetocin and ergometrine alone make any difference to this outcome. However, misoprostol is less effective in preventing blood loss that equals or exceeds 1000 mL compared with oxytocin.

Misoprostol plus oxytocin reduces the use of additional uterotonics and probably also reduces the risk of blood transfusion when compared with oxytocin. Carbetocin, injectable prostaglandins and ergometrine plus oxytocin may also reduce the use of additional uterotonics but the certainty of the evidence is low. No meaningful differences could be detected between all agents for maternal deaths or severe birth complication as these are rare in such studies.

The two combinations of drugs were associated with important side effects. When compared with oxytocin, women receiving misoprostol plus oxytocin combination are more likely to suffer vomiting and fever. Women receiving ergometrine plus oxytocin are also more likely to suffer vomiting and may make little or no difference to the risk of hypertension, however the certainty of the evidence was low for this outcome.

The analyses gave similar results irrespective of whether women were giving birth normally or by caesarean, in a hospital or in the community, were at high or low risk for bleeding excessively after birth, whether they received a high or a low dose of misoprostol and whether they received a bolus or an infusion of oxytocin or both.

What does this mean?

All agents were generally effective for preventing excessive bleeding when compared with no uterotonic drug treatment. Ergometrine plus oxytocin combination, carbetocin, and misoprostol plus oxytocin combination may have some additional benefits compared with the current standard oxytocin. The two combination drugs, however, are associated with significant side effects that women might find disturbing compared with oxytocin. Carbetocin may have some additional benefits compared with oxytocin and appears to be without an increase in side effects.

SUMMARY OF FINDINGS

Summary of findings for the main comparison. PPH >= 500 mL

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxytocin

Outcome: PPH ≥ 500 mL

Utero- tonic	Direct evi	idence	Indirect 6	evidence	NMA evid	ence	Anticipated ab	r NMA estimate	
agent(s)	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with in- tervention
Carbe- tocin	0.75 (0.58 to 0.98)	⊕⊕⊕⊚ MODER- ATE ^a	0.59 (0.31 to 1.12)	31 to LOW b	0.72 (0.56 to	MODER-	145 per 1000	104 per 1000	41 fewer per 1000 (from 64 fewer to 10 fewer)
	0.007	AIL	,		0.93)	AIL	Vaginal birth: 122 per 1000	Vaginal birth:87 per 1000	Vaginal birth: 34 fewer per 1000 (from 54 fewer to 9 fewer)
							Caesarean birth: 604 per 1000	Caesarean birth: 435 per 1000	Caesarean birth: 169 fewer per 1000 (from 266 fewer to 42 fewer)
Miso- prostol	1.08 (0.94 to 1.24)	⊕⊕⊝⊝ LOW <i>d</i>	1.07 (0.83 to 1.39)	⊕⊝⊝⊝ VERY LOW ^e	1.08 (0.96 to 1.22)	⊕⊕⊚⊝ LOW ^f	145 per 1000	157 per 1000	12 more per 1000 (4 fewer to 32 more)
	1.2 1)		1.33)	LOW	1.22)		Vaginal birth:122 per 1000	Vaginal birth:132 per 1000	Vaginal birth:10 more per 1000 (4 fewer to 27 more)
							Caesarean birth: 604 per 1000	Caesarean birth: 652 per 1000	Caesarean birth: 48 more per 1000 (18 fewer to 133 more)
In- jectable	0.84 (0.26 to	⊕⊕⊝⊝ LOW <i>9</i>	1.08 (0.72 to	⊕⊝⊝⊝ VERY	1.05 (0.73 to	⊕⊕⊙⊝ LOW ^f	145 per 1000	152 per 1000	7 more per 1000
prostagla	2.71)	2011 0	1.62)	LOW e	1.51)				(39 fewer to 74 more)

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							Vaginal birth: 122 per 1000	Vaginal birth:128 per 1000	Vaginal birth:6 more per 1000 (33 fewer to 62 more)
							Caesarean birth: 604 per 1000	Caesarean birth: 634 per 1000	Caesarean birth: 30 more per 1000 (163 fewer to 308 more)
Er- gometrine	1.31 (0.86 to	⊕⊝⊝⊝ VERY	0.96 (0.70 to	⊕⊕⊝⊝ LOW <i>i</i>	1.09 (0.85 to	⊕⊕⊝⊝ LOW <i>j</i>	145 per 1000	158 per 1000	13 more per 1000
gomeanic	1.99)	LOW h	1.31)	LOW	1.39)	LOW			(22 fewer to 57 more)
							Vaginal birth: 122 per 1000	Vaginal birth: 133 per 1000	Vaginal birth: 11 more per 1000
									(18 fewer to 48 more)
							Caesarean birth: 604 per 1000	Caesarean birth: 610 per 1000	Caesarean birth: 6 more per 1000 (91 fewer to 236 more)
Er- gometrine plus	0.72 (0.57 to 0.91)	⊕⊕⊕⊝ MODER- ATE ^k	0.69 (0.54 to 0.90)	⊕⊕⊝⊝ LOW ^b	0.70 (0.59 to 0.84)	⊕⊕⊕⊝ MODER- ATE ^Ç	145 per 1000	101 per 1000	44 fewer per 1000 (59 fewer to 23 fewer)
oxy- tocin	0.01,	ALL	0.507		0.0 1)	ALE	Vaginal birth: 122 per 1000	Vaginal birth: 85 per 1000	Vaginal birth: 37 fewer per 1000 (50 fewer to 20 few- er)
							Caesarean birth: 604 per 1000	Caesarean birth: 423 per 1000	Caesarean birth: 181 fewer per 1000 (248 fewer to 97 fewer)
Miso- pros- tol plus	0.71 (0.59 to 0.85)	⊕⊕⊝⊝ LOW [[]	0.79 (0.35 to 1.77)	⊕⊕⊝⊝ LOW ^j	0.70 (0.58 to 0.86)	⊕⊕⊝⊝ LOW ^m	145 per 1000	101 per 1000	44 fewer per 1000 (61 fewer to 20 fewer)
oxy- tocin	0.03)		1.11)		0.00/		Vaginal birth: 122 per 1000	Vaginal birth: 85 per 1000	Vaginal birth: 37 fewer per 1000 (51 fewer to 17 few- er)
							Caesarean birth: 604 per 1000	Caesarean birth: 423 per 1000	Caesarean birth: 181 fewer per 1000 (254 fewer to 85 fewer)

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin **CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -1 due to severe unexplained statistical heterogeneity
- b Indirect evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity
- C Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity, incoherence, or imprecision)
- d Direct evidence downgraded -2 due to multiple crucial limitations in study design
- e Indirect evidence downgraded -3 due to multiple crucial limitations in study design, severe unexplained statistical heterogeneity and serious imprecision
- f Network evidence downgraded -2 due to low certainty direct evidence (no intransitivity or incoherence, network estimate remains imprecise)
- 9 Direct evidence downgraded -2 due to multiple limitations in study design and serious imprecision
- h Direct evidence downgraded -3 due to multiple crucial limitations in study design and serious imprecision
- i Indirect evidence downgraded -2 due to severe unexplained statistical heterogeneity, multiple limitations in study design and serious imprecision
- j Network evidence downgraded -2 due to low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- k Direct evidence downgraded -1 due to multiple limitations in study design
- Direct evidence downgraded -2 due to multiple limitations in study design and strong suspicion of publication bias
- m Network evidence downgraded -2 due to low certainty direct and indirect evidence (no intransitivity, incoherence, or imprecision)

Summary of findings 2. PPH >= 1000 mL

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxvtocin

Outcome: PPH ≥ 1000 mL

Utero- tonic	Direct evidence		Indirect evidence		NMA evidence		Anticipated absolute effects for NMA estimate		
agent(s)	RR (95%	Certain- ty	RR (95%	Certain- ty	RR (95%	Certain- tv	Risk with oxytocin	Risk with in- tervention	Risk difference with in- tervention
	CI)	•,	CI)	•,	CI)	•,	CA, to c	ter remain	to vention

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Trusted evidence.
Informed decisions.
Better health.

									(other utero- tonics)	
Carbe- tocin		0.73 (0.45 to	⊕⊕⊝⊝ LOW a	0.30 (0.13 to	⊕⊕⊝⊝ LOW ^b	0.87 (0.62 to	⊕⊝⊝⊝ VERY LOW ^ç	37 per 1000	32 per 1000	5 fewer per 1000 (from 14 fewer to 8 more)
		1.19)		0.72)		1.21)	LOW	Vaginal birth: 30 per 1000	Vaginal birth: 26 per 1000	Vaginal birth: 4 fewer per 1000 (11 fewer to 6 more)
								Caesarean birth: 33 per 1000	Caesarean birth: 116 per 1000	Caesarean birth: 17 fewer per 1000 (from 51 fewer to 28 more)
Miso- prostol		1.26 (1.11 to	⊕⊕⊕⊕ HIGH	1.23 (0.92 to	⊕⊕⊕⊝ MODER- ATE ^d	1.19 (1.01 to	⊕⊕⊕⊕ HIGH ^e	37 per 1000	44 per 1000	7 more per 1000 (0 fewer to 16 more)
		1.43)		1.64)	ATE	1.42)		Vaginal birth: 30 per 1000	Vaginal birth: 36 per 1000	Vaginal birth: 6 more per 1000 (0 fewer to 13 more)
								Caesarean birth: 133 per 1000	Caesarean birth: 158 per 1000	Caesarean birth: 25 more per 1000 (1 more to 56 more)
In- jectable prostagla	ar	1.43 (0.20 to	⊕⊝⊝⊝ VERY LOW ^f	0.74 (0.31 to	⊕⊝⊝⊝ VERY LOW ^g	0.88 (0.41 to	⊕⊝⊝⊝ VERY LOW ^h	37 per 1000	33 per 1000	4 fewer per 1000 (22 fewer to 33 more)
		10.31)		1.72)		1.89)		Vaginal birth: 30 per 1000	Vaginal birth: 27 per 1000	Vaginal birth: 3 fewer per 1000 (18 fewer to 27 more)
								Caesarean birth: 133 per 1000	Caesarean birth: 118 per 1000	Caesarean birth: 15 fewer per 1000
Er- gometrir	16	1.30 (0.52 to	⊕⊝⊝⊝ VERY LOW ^f	0.61 (0.22 to	⊕⊕⊝⊝ LOW ^j	0.94 (0.48 to	⊕⊕⊝⊝ LOW <i>j</i>	37 per 1000	35 per 1000	2 fewer per 1000 (19 fewer to 31 more)
		3.27)	LOVV	1.67)		1.84)		Vaginal birth: 30 per 1000	Vaginal birth: 28 fewer per 1000	Vaginal birth: 2 fewer per 1000 (16 fewer to 25 more)
								Caesarean birth: 133 per 1000	Caesarean birth: 122 per 1000	Caesarean birth: 8 fewer per 1000 (69 fewer to 112 more)

Er- gometrine plus	Ę		⊕⊕⊕⊕ HIGH	1.07 (0.75 to	⊕⊕⊕⊝ MODER- ATE ^k	0.83 (0.66 to	⊕⊕⊕⊕ HIGH ^e	37 per 1000	31 per 1000	6 fewer per 1000 (13 fewer to 1 more)
oxy- tocin		0.93)		1.54)	AIL"	1.03)		Vaginal birth: 30 per 1000	Vaginal birth: 25 per 1000	Vaginal birth: 5 fewer per 1000 (10 fewer to 1 more)
								Caesarean birth: 133 per 1000	Caesarean birth: 124 per 1000	Caesarean birth: 9 few- er per 1000 (45 fewer to 4 more)
Miso- pros- tol plus		0.87 (0.69 to	⊕⊕⊕⊚ MODER-	1.17	⊕⊕⊕⊕ HIGH		⊕⊕⊕⊕ HIGH ^e	37 per 1000	31 per 1000	6 fewer per 1000 (13 fewer to 1 more)
tot ptus			ATE /	(0.47 to		(0.70 to				
oxy- tocin		1.09)	ATE [[]	(0.47 to 2.86)		(0.70 to 1.11)		Vaginal birth: 30 per 1000	Vaginal birth: 25 per 1000	Vaginal birth: 5 fewer per 1000 (10 fewer to 1 more)

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin CI: Confidence interval; RR: Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -2 due to serious imprecision and strong suspicion of publication bias
- *b* Indirect evidence downgraded -2 due to very serious imprecision
- c Network evidence downgraded -2 due to low certainty direct and indirect evidence, and -1 due to incoherence between the direct and indirect estimates (no intransitivity, network estimate remains imprecise)
- d Indirect evidence downgraded -1 due to serious imprecision
- e Network evidence not downgraded due to high certainty indirect evidence (no intransitivity, incoherence, or imprecision)

g Indirect evidence downgraded -3 due to multiple limitations in study design and very serious imprecision

h Network evidence downgraded -3 due to very low certainty direct and indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)

¹ Indirect evidence downgraded -2 due to multiple limitations in study design and serious imprecision

J Network evidence downgraded -2 due to low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)

k Indirect evidence downgraded -1 due to multiple limitations in study design

¹ Direct evidence downgraded -1 due to serious imprecision

Summary of findings 3. Additional uterotonics

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxytocin Outcome: use of additional uterotonics **Setting:** hospital or community setting

Utero- tonic	Direct evi	dence	Indirect e	vidence	NMA evid	ence	Anticipated ab	solute effects for	NMA estimate
agent(s)	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with in- tervention
Carbe- tocin	0.48 (0.34 to	⊕⊕⊝⊝ LOW <i>a</i>	0.35 (0.22 to	⊕⊕⊝⊝ LOW <i>b</i>	0.45 (0.34 to	⊕⊕⊝⊝	135 per 1000	61 per 1000	74 fewer per 1000 (89 few- er to 55 fewer)
	0.68)		0.57)		0.59)		Vaginal birth: 116 per 1000	Vaginal birth: 52 per 1000	Vaginal birth: 64 fewer per 1000 (77 fewer to 48 few- er)
							Caesarean birth: 304 per 1000	Caesarean birth: 137 per 1000	Caesarean birth: 167 fewer per 1000 (201 fewer to 125 fewer)
Miso- prostol	1.01 (0.85 to	⊕⊕⊝⊝ LOW <i>a</i>	1.18 (0.81 to	⊕⊕⊝⊝ LOW ^b	1.04 (0.88 to	⊕⊕⊝⊝ LOW c	135 per 1000	140 per 1000	5 more per 1000 (16 fewer to 32 more)
	1.20)		1.73)		1.24)		Vaginal birth: 116 per 1000	Vaginal birth: 121 per 1000	Vaginal birth: 5 more per 1000 (14 fewer to 28 more)
							Caesarean birth: 304 per 1000	Caesarean birth: 316 per 1000	Caesarean birth: 12 more per 1000 (36 fewer to 73 more

In- jectable prostaglar	0.29 (0.09 to	⊕⊕⊝⊝ LOW <i>d</i>	0.78 (0.38 to	⊕⊕⊝⊝	0.55 (0.31 to	⊕⊕⊝⊝ LOW ¢	135 per 1000	74 per 1000	61 fewer per 1000 (93 fewer to 5 fewer)	
prostugiui		0.94)		1.59)		0.96)		Vaginal birth: 116 per 1000	Vaginal birth: 64 per 1000	Vaginal birth: 52 fewer per 1000 (80 fewer to 5 fewer)
								Caesarean birth: 304 per 1000	Caesarean birth: 167 per 1000	Caesarean birth: 137 fewer per 1000 (210 fewer to 12 fewer)
Er- gometrine		1.46 (0.61 to	⊕⊝⊝⊝ VERY LOW ^f	0.83 (0.55 to	⊕⊝⊝⊝ VERY LOW 9		⊕⊝⊝⊝ VERY LOW ^h	135 per 1000	131 per 1000	4 fewer per 1000 (42 fewer to 49 more)
		3.48)	LOW	1.26)	LOWS	1.36)	LOW"	Vaginal birth: 116 per 1000	Vaginal birth: 113 per 1000	Vaginal birth: 3 fewer per 1000 (36 fewer to 42 more)
							Caesarean birth: 304 per 1000	Caesarean birth: 295 per 1000	Caesarean birth: 9 fewer per 1000 (94 fewer to 109 more)	
Er- gometrine plus		0.79 (0.59 to	⊕⊝⊝⊝ VERY LOW ^f	0.57 (0.40 to	⊕⊕⊝⊝ LOW <i>b</i>	0.65 (0.50 to	⊕⊕⊝⊝	135 per 1000	89 per 1000	46 fewer per 1000 (66 fewer to 20 fewer)
oxy- tocin		1.07)		0.81)		0.85)		Vaginal birth: 116 per 1000	Vaginal birth: 77 per 1000	Vaginal birth: 39 fewer per 1000 (57 fewer to 17 few- er)
								Caesarean birth: 304 per 1000	Caesarean birth: 201 per 1000	1Caesarean birth: 03 fewer per 1000 (149 fewer to 46 fewer)
Miso- pros- tol plus		0.54 (0.44 to	⊕⊕⊕⊕ HIGH	0.68 (0.31 to	⊕⊕⊝⊝ LOW <i>b</i>	0.56 (0.42 to	⊕⊕⊕⊕ HIGH ⁱ	135 per 1000	77 per 1000	58 fewer per 1000 (76 fewer to 35 fewer)
oxy- tocin		0.67)		1.51)		0.73)		Vaginal birth: 116 per 1000	Vaginal birth: 66 per 1000	Vaginal birth: 50 fewer per 1000 (65 fewer to 30 few- er)
								Caesarean birth: 304 per 1000	Caesarean birth: 173 per 1000	Caesarean birth: 131 fewer per 1000 (170 fewer to 79 fewer)

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the Carbetocin, Misoprostol, Injectable prostaglandins, Ergometrine, Ergometrine plus oxytocin (Syntometrine®), Misoprostol plus oxytocin

Uterotonic agents for preventing postpartum haemorrhage:

groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin **CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity
- b Indirect evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity
- C Network evidence downgraded -2 due to low certainty direct and indirect evidence (no intransitivity, incoherence or serious imprecision)
- dDirect evidence downgraded -2 due to multiple crucial limitations in study design
- e Indirect evidence downgraded -2 due to multiple limitations in study design, severe unexplained statistical heterogeneity and serious imprecision
- f Direct evidence downgraded -3 due to multiple limitations in study design, severe unexplained statistical heterogeneity and serious imprecision
- 9 Indirect evidence downgraded -3 due to multiple limitations in study design, severe unexplained statistical heterogeneity and serious imprecision
- h Network evidence downgraded -3 due to very low certainty direct and indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- ⁱ Network evidence not downgraded due to high certainty direct evidence (no intransitivity, incoherence, or imprecision)

Summary of findings 4. Blood transfusion

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin

Comparison (reference): oxvtocin Outcome: blood transfusion

Utero- tonic	Dir	Direct evidence Indirect evider		vidence	nce NMA evidence		Anticipated absolute effects for NMA estimate			
agent(s)	RR (95 CI)	5% ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with in- tervention	

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Carbe- tocin	0.68 (0.38 to 1.22)	⊕⊕⊕⊝ MODER- ATE ^a	0.62 (0.21 to 1.85)	⊕⊕⊚⊚ LOW <i>b</i>	0.81 (0.49 to 1.32)	⊕⊕⊕⊝ MODER- ATE [¢]	22 per 1000	18 per 1000	4 fewer per 1000 (11 fewer to 7 more)
	1.22)	AIE	1.05)		1.32)	AIE	Vaginal birth:	Vaginal birth:	Vaginal birth: 3 fewer per
							15 per 1000	12 per 1000	1000 (5 fewer to 4 more)
							Caesarean birth:	Caesarean birth:66 per	Caesarean birth:15 fewer per 1000 (41 fewer to 26
							81 per 1000	1000	more)
Miso- prostol	0.81 (0.65 to	⊕⊕⊕⊝ MODER- ATE ^a	1.02 (0.59 to 1.77)	⊕⊕⊝⊝ LOW <i>d</i>	0.88 (0.68 to 1.13)	⊕⊕⊕⊝ MODER- ATE [¢]	22 per 1000	19 per 1000	3 fewer per 1000 (7 fewer to 3 more)
	1.00)	AIE "	1.77		1.13)	AIL	Vaginal birth: 15 per 1000	Vaginal birth: 13 per 1000	Vaginal birth: 2 fewer per 1000 (5 fewer to 2 more)
							Caesarean birth: 81 per 1000	Caesarean birth: 71 per 1000	Caesarean birth: 10 few- er per 1000 (26 fewer to 11 more)
In- jectable prostaglar	1.01 (0.04 to 23.65)	⊕⊝⊝⊝ VERY LOW ^e	0.49 (0.16 to 1.52)	⊕⊝⊝⊝ VERY LOW ^f	0.66 (0.25 to 1.72)	⊕⊝⊝⊝ VERY LOW g	22 per 1000	15 per 1000	7 fewer per 1000 (17 fewer to 16 more)
Fr. com Sim.	25.05)	LOW	1.52)	LOW	1112)	LOW	Vaginal birth: 15 per 1000	Vaginal birth: 10 per 1000	Vaginal birth: 5 fewer per 1000 (11 fewer to 11 more)
							Caesarean birth: 81 per 1000	Caesarean birth: 56 per 1000	Caesarean birth: 28 few- er per 1000 (61 fewer to 58 more)
Er- gometrine	1.44 (0.20 to 10.23)	⊕⊝⊝⊝ VERY LOW ^h	1.01 (0.38 to 2.68)	⊕⊕⊝⊝ LOW ⁱ	1.11 (0.54 to 2.28)	⊕⊕⊝⊝ LOW <i>j</i>	22 per 1000	24 per 1000	2 more per 1000 (10 fewer to 28 more)
	10.23)	LOW"	2.00)		2.20)		Vaginal birth: 15 per 1000	Vaginal birth: 17 per 1000	Vaginal birth: 2 more per 1000 (7 fewer to 19 more)
							Caesarean birth: 81 per 1000	Caesarean birth: 90 per 1000	Caesarean birth: 9 more per 1000 (37 fewer to 104 more)
Er- gometrine plus	0.88 (0.53 to 1.44)	⊕⊕⊝⊝ LOW ^k	0.64 (0.41 to 1.00)	⊕⊕⊝⊝ LOW ⁱ	0.77 (0.58 to 1.03)	⊕⊕⊝⊝ LOW <i>j</i>	22 per 1000	17 per 1000	5 fewer per 1000 (9 fewer to 1 more)

oxy- tocin							Vaginal birth: 15 per 1000	Vaginal birth: 12 per 1000	Vaginal birth: 3 fewer per 1000 (6 fewer to 0 fewer)
							Caesarean birth: 81 per 1000	Caesarean birth: 63 per 1000	Caesarean birth: 18 few- er per 1000 (33 fewer to 2 more)
Miso- pros- tol plus	0.50 (0.37 to 0.67)	⊕⊕⊝⊝ LOW [[]	0.77 (0.27 to 2.26)	⊕⊕⊕⊚ MODER- ATE ^m	0.51 (0.37 to 0.70)	⊕⊕⊕⊚ MODER- ATE [¢]	22 per 1000	11 per 1000	11 fewer per 1000 (14 fewer to 7 fewer)
oxy- tocin	0.017		2.20)	AIL	0.10)	AIL -	Vaginal birth: 15 per 1000	Vaginal birth: 8 per 1000	Vaginal birth: 7 fewer per 1000 (9 fewer to 5 fewer)
							Caesarean birth: 81 per 1000	Caesarean birth: 42 per 1000	Caesarean birth: 39 few- er per 1000 (50 fewer to 24 fewer)

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin **CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -1 due to serious imprecision
- b Indirect evidence downgraded -2 due to very serious imprecision
- c Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity or incoherence, network estimate remains imprecise)
- d Indirect evidence downgraded -2 due to multiple limitations in study design and strong suspicion of publication bias
- ^e Direct evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- f Indirect evidence downgraded -3 due to multiple crucial limitations in study design and very serious imprecision
- 9 Network evidence downgraded -3 due to very low certainty direct and indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- h Direct evidence downgraded -3 due to multiple limitations in study design, severe unexplained statistical heterogeneity and very serious imprecision

j Network evidence downgraded -2 due to low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)

k Direct evidence downgraded -2 due to multiple limitations in study design and serious imprecision

¹ Direct evidence downgraded -2 due to multiple limitations in study design and strong suspicion of publication bias

m Indirect evidence downgraded -1 due to multiple limitations in study design and serious imprecision

Summary of findings 5. Vomiting

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxytocin

Outcome: vomiting

	Utero- tonic		dence	Indirect e	Indirect evidence		ence	Anticipated absolute effects for NMA estimate			
agent	s)	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with in- tervention	
Carbe tocin		0.90 (0.53 to 1.50)	⊕⊕⊕⊝ MODER- ATE ^a	1.00 (0.51 to 1.95)	⊕⊕⊝⊝ LOW <i>b</i>	0.93 (0.64 to 1.35)	⊕⊕⊕⊚ MODER- ATE [¢]	28 per 1000	26 per 1000	2 fewer per 1000 (10 fewer to 10 more)	
								Vaginal birth: 13 per 1000	Vaginal birth: 12 per 1000	Vaginal birth: 1 fewer per 1000 (5 fewer to 5 more)	
								Caesarean birth: 97 per 1000	Caesarean birth: 91 per 1000	Caesarean birth: 6 fewer per 1000 (34 fewer to 35 more)	
Miso- prosto	ı	1.51 (1.19 to 1.91)	⊕⊕⊕⊕ HIGH	2.73 (1.66 to 4.50)	⊕⊕⊝⊝ LOW ^d	1.63 (1.25 to 2.14)	⊕⊕⊕⊚ MODER- ATE ^e	28 per 1000	46 per 1000	18 more per 1000 (7 more to 32 more)	
		1.31)		1.50)		2.1 1)	AIL	Vaginal birth: 13 per 1000	Vaginal birth: 21 per 1000	Vaginal birth: 8 more per 1000 (3 more to 15 more)	
								Caesarean birth: 97 per 1000	Caesarean birth: 158 per 1000	Caesarean birth: 61 more per 1000 (24 more to 111 more)	

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In- jectable prostaglar	2.48 (0.57 to 10.73)	⊕⊝⊝⊝ VERY LOW ^f	4.07 (1.93 to 8.60)	⊕⊝⊝⊝ VERY LOW ^g	3.76 (1.90 to 7.42)	⊕⊕⊝⊝ LOW <i>h</i>	28 per 1000	105 per 1000	77 more per 1000 (25 more to 180 more)
prostagtar	10.73)	LOW	8.60)	LOW9	1.42)		Vaginal birth: 13 per 1000	Vaginal birth: 49 per 1000	Vaginal birth: 36 more per 1000 (12 more to 83 more)
							Caesarean birth: 97 per 1000	Caesarean birth: 365 per 1000	Caesarean birth: 268 more per 1000 (87 more to 623 more)
Er-	3.83	00 00	1.83	⊕⊕⊝⊝ 	2.36	⊕⊕⊕⊝ MODED	28 per 1000	66 per 1000	38 more per 1000
gometrine	(1.10 to 13.28)	LOW ⁱ	(1.18 to 2.84)	LOW <i>j</i>	(1.56 to 3.55)	MODER- ATE ^k			(16 more to 71 more)
							Vaginal birth: 13 per 1000	Vaginal birth: 31 per 1000	Vaginal birth: 18 more per 1000 (7 more to 33 more)
							Caesarean birth: 97 per 1000	Caesarean birth: 229 per 1000	Caesarean birth: 132 more per 1000 (54 more to 247 more)
Er- gometrine plus	3.05 (1.76 to 5.29)	⊕⊕⊕⊝ MODER- ATE [[]	2.77 (1.75 to 4.38)	⊕⊕⊝⊝ LOW <i>d</i>	2.93 (2.08 to 4.13)	⊕⊕⊕⊝ MODER- ATE ^m	28 per 1000	82 per 1000	54 more per 1000 (30 more to 88 more)
oxy- tocin	3.23)	AIE	4.30)		4.13)	ATE.	Vaginal birth: 13 per 1000	Vaginal birth: 38 per 1000	Vaginal birth: 25 more per 1000 (14 more to 41 more)
							Caesarean birth: 97 per 1000	Caesarean birth: 284 per 1000	Caesarean birth: 187 more per 1000 (105 more to 304 more)
Miso- pros- tol plus	2.24 (1.52 to 3.31)	⊕⊕⊕⊕ HIGH	1.48 (0.52 to 4.27)	⊕⊝⊝⊝ VERY LOW ^g	2.11 (1.39 to 3.18)	⊕⊕⊕⊕ HIGH ⁿ	28 per 1000	59 per 1000	31 more per 1000 (11 more to 61 more)
oxy- tocin	3.31)		T. Z. I)	LOWY	3.10)		Vaginal birth: 13 per 1000	Vaginal birth: 27 per 1000	Vaginal birth: 14 more per 1000 (5 more to 28 more)
							Caesarean birth: 97 per 1000	Caesarean birth: 205 per 1000	Caesarean birth: 108 more per 1000 (38 more to 211 more)

Uterotonic agents for preventing postpartum haemorrhage: a

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin **CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -1 due to serious imprecision
- b Indirect evidence downgraded -2 due to very serious imprecision
- ^c Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity or incoherence, network estimate remains imprecise)
- d Indirect evidence downgraded -2 due to multiple crucial limitations in study design and serious imprecision
- e Network evidence not initially downgraded given high certainty direct evidence; however, downgraded -1 due to incoherence between the direct and indirect estimates (no intransitivity or imprecision)
- f Direct evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- 9 Indirect evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- h Network evidence initially downgraded -3 due to very low certainty direct and indirect evidence, however upgraded +1 due to precision of network estimate (when direct and indirect were both imprecise; no intransitivity or incoherence)
- ⁱ Direct evidence downgraded -2 due to multiple crucial limitations in study design
- J Indirect evidence downgraded -2 due to multiple limitations in study design and serious imprecision
- k Network evidence initially downgraded -2 due to low certainty direct and indirect evidence, however upgraded +1 due to precision of network estimate (when direct and indirect were both imprecise; no intransitivity or incoherence)
- United Evidence downgraded -1 due to multiple limitations in study design
- m Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity, incoherence, or serious imprecision)
- ⁿ Network evidence not downgraded due to high certainty direct evidence (no intransitivity, incoherence, or serious imprecision)

Summary of findings 6. Hypertension

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxvtocin

Outcome: hypertension

Setting: hospital or community setting Cochrane Library

Utero- tonic	Direct evi	dence	Indirect e	vidence	NMA evid	ence	Anticipated a	bsolute effects f	or NMA estimate
agent(s)	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with intervention
Carbe- tocin	Not re- ported	_	1.24	⊕⊝⊝⊝ VERY	1.24	⊕⊝⊝⊝ VERY	82 per 1000	102 per 1000	20 more per 1000
toem	by in- cluded		(0.28 to 5.56)	LOW a	(0.28 to 5.56)	LOW b			(59 fewer to 374 more)
	studies		,		,		Vaginal birth: 76 per 1000	Vaginal birth: 94 per 1000	Vaginal birth: 18 more per 1000 (55 fewer to 347 more)
							Caesarean birth: 167 per 1000	Caesarean birth: 207 per 1000	Caesarean birth: 40 more per 1000 (120 fewer to 762 more)
Miso- prostol	3.64 (0.60 to	⊕⊝⊝⊝ VERY LOW [¢]	1.01 (0.28 to	⊕⊕⊝⊝ LOW <i>d</i>	1.50 (0.49 to	⊕⊕⊝⊝ LOW e	82 per 1000	123 per 1000	41 more per 1000 (42 fewer to 296 more)
	22.27)	LOW	3.65)		4.61)		Vaginal birth: 76 per 1000	Vaginal birth: 114 per 1000	Vaginal birth: 38 more per 1000 (39 fewer to 274 more)
							Caesarean birth: 167 per 1000	Caesarean birth: 250 per 1000	Caesarean birth: 83 more per 1000 (85 fewer to 603 more)
In- jectable prostaglai	Not re- ported by in-	_	1.40 (0.09 to	⊕⊝⊝⊝ VERY LOW ^a	1.40 (0.09 to	⊕⊝⊝⊝ VERY LOW ^b	82 per 1000	115 per 1000	33 more per 1000 (75 fewer to 1000 more)
prostugiui	cluded studies		20.66)	LOW	20.66)	LOW	Vaginal birth: 76 per	Vaginal birth: 106 per 1000	Vaginal birth: 30 more per 1000
							1000		(69 fewer to 1000 more)
							Caesarean birth: 167 per 1000	Caesarean birth: 234 per 1000	Caesarean birth: 67 more per 1000 (152 fewer to 1000 more)

Er- gometrine	13.39 (2.01 to 89.44)	⊕⊕⊝⊝ LOW ^f	12.42 (0.91 to 168.67)	⊕⊝⊝⊝ VERY LOW <i>g</i>	8.54 (2.12 to 34.48)	⊕⊕⊝⊝ LOW <i>h</i>	82 per 1000	700 per 1000	618 more per 1000 (92 more to 2745 more)
	65.44)		100.07)		34.40)		Vaginal birth: 76 per 1000	Vaginal birth: 649 per 1000	Vaginal birth: 573 more per 1000 (85 more to 1000 more)
							Caesarean birth: 167 per 1000	Caesarean birth: 1000 per 1000	Caesarean birth: 1000 more per 1000 (187 more to 1000 more)
Er- gometrine	2.00 (0.29 to	⊕⊕⊚⊚ LOW ⁱ	5.16 (0.63 to	⊕⊝⊝⊝ VERY	2.48 (0.89 to	⊕⊕⊝⊝ LOW ^k	82 per 1000	203 per 1000	121 more per 1000 (9 fewer to 482 more)
plus oxy- tocin	13.97)		42.13)	LOW ^j	6.88)		Vaginal birth: 76 per 1000	Vaginal birth: 188 per 1000	Vaginal birth: 112 more per 1000 (8 fewer to 447 more)
							Caesarean birth: 167 per 1000	Caesarean birth: 414 per 1000	Caesarean birth: 247 more per 1000 (18 fewer to 982 more)
Miso- pros- tol plus oxy- tocin	Not re- ported by in- cluded studies	-	Not re- ported by in- cluded studies	-	Not re- ported by in- cluded studies	-	see com- ment*	see com- ment**	see comment***

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

^{*} No included studies or there are no event in included studies to estimate the baseline risk

^{**} Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin

^{***}Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin CI: Confidence interval; RR: Risk ratio.

- a Indirect evidence downgraded -3 due to multiple limitations in study design, very serious imprecision and severe unexplained statistical heterogeneity
- b Network evidence downgraded -3 due to very low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- ^c Direct evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- d Indirect evidence downgraded -2 due to multiple limitations in study design, serious imprecision and severe unexplained statistical heterogeneity
- e Network evidence downgraded -2 due to low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- f Direct evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity
- 9 Indirect evidence downgraded -3 due to multiple crucial limitations in study design and severe unexplained statistical heterogeneity
- h Network evidence downgraded -3 due to very low certainty indirect evidence (no intransitivity, incoherence or imprecision; although CI is wide there is a clear increase in this outcome for ergometrine)
- ⁱ Direct evidence downgraded -2 due to severe unexplained statistical heterogeneity and serious imprecision
- J Indirect evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- k Network evidence downgraded -2 due to low certainty direct evidence (no intransitivity or incoherence, network estimate remains imprecise)

Summary of findings 7. Fever

Patient or population: women in the third stage of labour

Interventions: carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine®), misoprostol plus oxytocin

Comparison (reference): oxytocin

Outcome: fever

Setting: hospital or community setting

Utero- tonic	Direct evi	dence	Indirect e	vidence	NMA evid	ence	Anticipated ab	solute effects for	r NMA estimate
agent(s)	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	RR (95% CI)	Certain- ty	Risk with oxytocin	Risk with in- tervention (other utero- tonics)	Risk difference with in- tervention
Carbe- tocin	1.58 (0.27 to	⊕⊕⊕⊝ MODER-	0.77	⊕⊕⊝⊝ 	1.07	⊕⊕⊕⊝ MODER-	29 per 1000	31 per 1000	2 more per 1000
tocin	9.35)	ATE a	(0.18 to 3.42)	LOW ^b	(0.43 to 2.69)	ATE c			(17 fewer to 49 more)
			,		,		Vaginal birth: 24 per 1000	Vaginal birth: 26 per 1000	Vaginal birth: 2 more per 1000
									(14 fewer to 41 more)
							Caesarean birth: 55 per 1000	Caesarean birth: 59 per 1000	Caesarean birth: 4 more per 1000 (31 fewer to 93 more)

Cochrane
Library

Trusted evidence. Informed decisions. Better health.

Uterotoni Copyright	Miso- prostol	3.75 (2.73 to	⊕⊕⊝⊝ LOW <i>d</i>	6.49 (2.24 to	⊕⊕⊕⊝ MODER- ATE ^e	3.87 (2.90 to	⊕⊕⊕⊚ MODER- ATE [¢]	29 per 1000	112 per 1000	83 more per 1000 (55 more to 121 more)
c agents for I		5.15)		18.76)	AIL	5.16)	AIL-	Vaginal birth: 24 per 1000	Vaginal birth: 93 per 1000	Vaginal birth: 69 more per 1000 (46 more to 100 more)
preventing pouthors. Cochi								Caesarean birth: 55 per 1000	Caesarean birth: 213 per 1000	Caesarean birth: 158 more per 1000 (105 more to 229 more)
ostpar ane Da	In- jectable	2.00	⊕⊝⊝⊝ VERY	0.96	⊕⊕⊝⊝	1.12	00 00	29 per 1000	32 per 1000	3 more per 1000
tum h	prostaglar	(0.18 to 21.71)	LOW f	(0.24 to 3.87)	LOW ^b	(0.33 to 3.86)	LOW9			(19 fewer to 83 more)
aemorrh:				5.5.,		3.33,		Vaginal birth: 24 per 1000	Vaginal birth: 27 per 1000	Vaginal birth: 3 more per 1000
age: a										(16 fewer to 69 more)
Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis (Review) Copyright © 2018 The Authors. Cochrane Database of Systematic Reviews published by John Wiley & Sons, Ltd. on behalf of The Cochrane								Caesarean birth: 55 per 1000 (for cae- sarean birth)	Caesarean birth: 61 per 1000 (for cae- sarean birth)	Caesarean birth: 6 more per 1000 (37 fewer to 153 more)
analys ed by J	Er-	2.97	⊕ ⊝⊝⊝	0.63	00 00	0.77	0 000	29 per 1000	22 per 1000	7 fewer per 1000
is (Rev	gometrine	(0.97 to 9.05)	VERY LOW ^f	(0.35 to 1.16)	LOW ^h	(0.44 to 1.35)	VERY LOW ⁱ			(16 fewer to 10 more)
iew) iley & Son		3.03)		1.10)		1.55)		Vaginal birth: 24 per 1000	Vaginal birth: 18 per 1000	Vaginal birth: 6 fewer per 1000
s, Ltd.										(13 fewer to 8 more)
on behalf of								Caesarean birth: 55 per 1000	Caesarean birth: 42 per 1000	Caesarean birth: 13 few- er per 1000 (31 fewer to 18 more)
The Co	Er-	1.08	⊕⊕ ⊝⊝	0.54	⊕⊕⊝⊝ . ••••/•	0.70	00 00	29 per 1000	20 per 1000	9 fewer per 1000
chrane	gometrine plus	(0.48 to 2.43)	LOW j	(0.22 to 1.32)	LOW k	(0.35 to 1.42)	LOW [[]			(19 fewer to 12 more)
	tocin	2.10)		1.52)		±. 1∠j		Vaginal birth: 24 per 1000	Vaginal birth: 17 per 1000	Vaginal birth: 7 fewer per 1000
20										(16 fewer to 10 more)

							Caesarean birth: 55 per 1000	Caesarean birth: 42 per 1000	Caesarean birth: 13 few- er per 1000 (31 fewer to 19 more)
Miso- pros- tol plus	2.99 (2.00 to	⊕⊕⊕⊝ MODER- ATE <i>m</i>	5.43 (1.48 to	⊕⊕⊝⊝ LOW <i>n</i>	3.14 (2.20 to	⊕⊕⊕⊝ MODER- ATE Ø	29 per 1000	91 per 1000	62 more per 1000 (35 more to 101 more)
oxy- tocin	4.45)	AIE'''	19.95)		4.49)	AIE	Vaginal birth: 24 per 1000	Vaginal birth: 75 per 1000	Vaginal birth: 51 more per 1000 (29 more to 84 more)
							Caesarean birth: 55 per 1000	Caesarean birth: 173 per 1000	Caesarean birth: 118 more per 1000 (66 more to 192 more)

The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin groups in the network meta-analysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin groups (and their 95% confidence interval) are based on the assumed risk in the oxytocin group and the relative effect of individual uterotonic when compared with oxytocin (and its 95% CI) derived from the network meta-analysis.

- * No included studies or there are no event in included studies to estimate the baseline risk
- ** Absolute risk with uterotonic cannot be estimated in the absence of absolute risk with oxytocin
- ***Risk difference cannot be estimated in the absence of absolute risks with intervention and oxytocin **CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- ^a Direct evidence downgraded -1 due to serious imprecision
- b Indirect evidence downgraded -2 due to multiple limitations in study design, severe unexplained statistical heterogeneity and serious imprecision
- ^c Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity or incoherence, network estimate remains imprecise)
- d Direct evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity
- ^e Indirect evidence downgraded -1 due to multiple limitations in study design and serious imprecision
- f Direct evidence downgraded -3 due to multiple limitations in study design and very serious imprecision
- 9 Network evidence downgraded -2 due to low certainty indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)
- h Indirect evidence downgraded -2 due to multiple limitations in study design, severe unexplained statistical heterogeneity and strong suspicion of publication bias. The indirect estimate is imprecise, however the effect estimates for the two head-to-head comparisons in the dominant first-order loop were not imprecise, so we have not downgraded for imprecision

¹ Network evidence initially downgraded -2 due to low certainty indirect evidence; however, downgraded further -1 due to incoherence between the direct and indirect estimates (no intransitivity. Network estimate is imprecise, unlike indirect evidence, however no further downgrade considered because certainty already very low)

j Direct evidence downgraded -2 due to very serious imprecision

k Indirect evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity. The indirect estimate is imprecise, however the effect estimates for the two head-to-head comparisons in the dominant first-order loop were not imprecise, so we have not downgraded for imprecision

Network evidence downgraded -2 due to low certainty direct and indirect evidence (no intransitivity or incoherence, network estimate remains imprecise)

m Direct evidence downgraded -1 due to multiple limitations in study design

n Indirect evidence downgraded -2 due to multiple limitations in study design and severe unexplained statistical heterogeneity

^o Network evidence downgraded -1 due to moderate certainty direct evidence (no intransitivity, incoherence or imprecision)



BACKGROUND

Description of the condition

An estimated 303,000 women died during childbirth in 2015 (Alkema 2016). Postpartum haemorrhage (PPH) accounted for up to a third of all these maternal deaths (Say 2014). Almost all deaths occurred in low- or middle-income countries. Even when death from PPH is avoided, the need for blood transfusion, hysterectomy and additional intervention place a huge burden on women's health and health services (Penney 2007; Souza 2013).

The third stage of labour, defined as the period of time from birth until the delivery of the placenta, and the immediate postpartum period are the most hazardous periods of childbirth due to the risk of PPH. The World Health Organization (WHO) defines PPH as when the blood loss after birth equals or exceeds 500 mL in the first 24 hours (WHO 2012). The most common cause of PPH is uterine atony (failure of the uterus to contract after birth). Even though risk factors for adverse maternal outcomes from severe haemorrhage have been identified (Souza 2013), often PPH is unpredictable as it occurs in the absence of identifiable clinical or historical risk factors (Combs 1991). Therefore, effective prevention of PPH is advocated for all women during childbirth (WHO 2012). The administration of uterotonic agents routinely in the third stage of labour is the key intervention that prevents PPH, although there is uncertainty about which agent may be the most effective.

Description of the intervention

The administration of uterotonic agents to prevent PPH is part of the active management of the third stage of labour (Begley 2015). The active management of the third stage of labour refers to the administration of a uterotonic agent, early cord clamping, and controlled cord traction until delivery of the placenta. In 2012, a WHO guideline panel revisited the evidence underpinning each component of active management of the third stage of labour and considered the use of uterotonics as the main intervention within this package (WHO 2012).

How the intervention might work

Several different uterotonic agents have been used for preventing PPH. These agents include ergometrine, misoprostol, carbetocin, oxytocin, injectable prostaglandins (such as carboprost and sulprostone) and the combinations of agents such as misoprostol plus oxytocin and ergometrine plus oxytocin.

Oxytocin

Oxytocin (Syntocinon®) is the most widely used uterotonic agent. At low doses, it produces rhythmic uterine contractions that are indistinguishable in frequency, force and duration from those observed during spontaneous labour, but at higher dosages, it causes sustained uterine contractions (MEDICINES.ORG.UK). It has a short half-life, approximately three to five minutes, and can be used as an infusion to maintain uterine contraction. When used intramuscularly, the latent phase lasts three to seven minutes, but produces a longer-lasting clinical effect of up to one hour (MEDICINES.ORG.UK). However, oxytocin cannot be used orally. It is unstable in ambient temperatures and it requires a cold chain through storage and transport. It should also not be given intravenously as a large bolus, because it can cause severe hypotension (Thomas 2007). Because of its anti-diuretic effect,

water intoxication can occur with prolonged infusion of oxytocin (MEDICINES.ORG.UK).

Ergometrine

Ergometrine and methylergometrine are ergot alkaloids that increase the uterine muscle tone by causing sustained uterine contractions. They have a latent phase of two to five minutes after intramuscular injection and the plasma half-life is 30 to 120 minutes (de Groot 1998). After intravenous administration, the onset of action is one minute or less and the duration of action is 45 minutes (although rhythmic contractions may persist for up to three hours). However, ergometrine and methylergometrine have an unpredictable bioavailability, which prevents oral use of the agent and requires protection from light, and storage at a temperature between 2° and 8°C to prolong shelf life (de Groot 1996a). They are vasoconstrictive and are contraindicated in women with hypertensive or cardiovascular disorders (MEDICINES.ORG.UK).

Misoprostol

Misoprostol is a prostaglandin E1 analogue, which is licensed for the prevention and treatment of gastric ulcers. It is well known for its off-label use as a uterotonic agent (Tuncalp 2012). It is water-soluble and heat stable (Davies 2001). It is absorbed nine to 15 minutes after sublingual, oral, vaginal, and rectal use. The half-life is about 20 to 40 minutes. Oral and sublingual routes have the advantage of rapid onset of action, while the vaginal and rectal routes result in prolonged activity and greater bioavailability (Schaff 2005).

Injectable prostaglandins

Prostaglandin preparations are available in injectable forms and the most commonly used agents are carboprost tromethamine (Hemabate), an analogue of 15-methyl-prostaglandin F2a, and sulprostone, which is a PGE2 analogue. After intramuscular administration, the time to peak plasma concentration is between 15 and 60 minutes. The half-life is about eight minutes. They require storage at a temperature between 2° and 8°C to prolong shelf life (MEDICINES.ORG.UK). They both enhance uterine contractility and cause vasoconstriction in postpartum women (MEDICINES.ORG.UK). However, they are not contraindicated in hypertensive women (MEDICINES.ORG.UK). In the management of the third stage of labour, injectable prostaglandins have been mainly used for intractable PPH as a last resort when other measures fail. Important disadvantages of injectable prostaglandins have been their cost and availability.

Carbetocin

Carbetocin is a newer long-acting synthetic analogue of oxytocin with agonist properties. After intravenous injection, it produces sustained uterine contractions within two minutes, lasting for approximately six minutes followed by rhythmic contractions for 60 minutes (Hunter 1992). When carbetocin is administered by an intramuscular injection, the sustained uterine contractions last for approximately 11 minutes and the rhythmic contractions for 120 minutes (Hunter 1992). A heat stable carbetocin is now available and has been evaluated against oxytocin in a large randomised trial (Widmer 2018). Carbetocin also appears to have a favourable side-effect profile (Su 2012).



Combination agents

The use of combinations of uterotonic agents is also popular and the most commonly used agent is ergometrine plus oxytocin (Syntometrine ®). This is a fixed-combination agent containing 5 international units (IU) of oxytocin and 500 mcg of ergometrine. Intramuscular injection is the recommended route (MEDICINES.ORG.UK). When used intramuscularly, the latent period for the occurrence of the uterine response is about 2.5 minutes and the uterotonic effects last for around three hours. Another combination that has been investigated is misoprostol plus oxytocin. This combination is not in synthetic (fixed-drug) or naturally occurring forms.

The WHO recommends that all women giving birth should be offered uterotonics during the third stage of labour for the prevention of PPH; oxytocin (intramuscular/intravenous, 10 IU is the uterotonic agent of choice (WHO 2012). Other injectable uterotonics and misoprostol are recommended as alternatives for the prevention of PPH in settings where oxytocin is not available.

Why it is important to do this review

The individual uterotonics described above have been compared in existing Cochrane Reviews and all comparisons are based on trials that directly compared one uterotonic against another uterotonic agent in head-to-head trials (Begley 2015; Liabsuetrakul 2018; McDonald 2004; Su 2012; Tuncalp 2012; Westhoff 2013). The existing Cochrane Reviews have variable eligibility criteria for study inclusion, uterotonic agent comparisons and outcomes. In the absence of a single randomised controlled trial comparing all available uterotonic agents, uncertainty remains over their relative effectiveness and ranking. When multiple interventions are available, a network meta-analysis is better placed for synthesising and interpreting the wider picture of the evidence and to understand the relative effects of all available interventions. Network meta-analysis has advantages over conventional pairwise meta-analysis, as the technique uses both direct and indirect evidence in a single coherent analysis to improve certainty about all possible treatment comparisons. Indirect evidence is obtained when the relative effectiveness of two competing interventions is inferred through a common comparator, even though this pair may not have been compared directly (Caldwell 2005; Lumley 2002).

This is an update of a review first published in April 2018. It has been updated to incorporate results from a large WHO trial (Widmer 2018) and a number of other large recently published trials.

OBJECTIVES

To identify the most effective uterotonic agent(s) to prevent postpartum haemorrhage (PPH) with the least side effects, and generate a ranking according to their effectiveness and side-effect profile.

METHODS

Criteria for considering studies for this review

Types of studies

All randomised controlled trials or cluster-randomised trials comparing the effectiveness and side effects of uterotonic agents with other uterotonic agents, placebo or no treatment for preventing postpartum haemorrhage (PPH) were eligible for

inclusion. Quasi-randomised trials were excluded. Randomised trials published only as abstracts were eligible if sufficient information could be retrieved.

Types of participants

The review included studies of women in the third stage of labour following a vaginal or caesarean birth in hospital or community settings.

Types of interventions

Trials were eligible if they administered uterotonic agents of any dosage, route or regimen systemically at birth for preventing PPH, and compared them with other uterotonic agents, placebo or no treatment. Trials evaluating uterotonic agents not administered systemically, such as intrauterine administration, or not immediately after birth, or exclusively comparing different dosages, routes or regimens of the same uterotonic agent were excluded. We included trials in which non-pharmacologic cointerventions such as controlled cord traction, cord clamping, or uterine massage was performed as a randomised intervention in all arms of the trial and the effects of such co-interventions were tested through a sensitivity analysis.

We classified agents into single agents including oxytocin, carbetocin, injectable prostaglandins (carboprost tromethamine, sulprostone), misoprostol, ergometrine (included also ergonovine, methylergonovine), and combination agents including ergometrine plus oxytocin (Syntometrine ® as a fixed-combination agent containing 5 international units (IU) of oxytocin and 500 mcg of ergometrine, any oxytocin dose and route when combined with any dose and route of ergometrine, ergonovine, or methylergonovine), and misoprostol plus oxytocin (any oxytocin dose and route when combined with any dose and route of misoprostol).

For this review, we assumed that any woman who meets the inclusion criteria is, in principle, equally likely to be randomised to any of the eligible uterotonic agents.

Types of outcome measures

We estimated the relative effects and rankings of the competing interventions according to the following outcomes.

Primary outcomes

The primary outcomes of the review were:

- 1. PPH \geq 500 mL; and
- 2. PPH ≥ 1000 mL.

Secondary outcomes

The secondary outcomes of the review were:

- 1. maternal deaths;
- 2. severe maternal morbidity: intensive care admissions;
- 3. severe maternal morbidity: shock (as defined by the trialists);
- 4. additional uterotonics;
- 5. blood transfusion;
- 6. mean volumes of blood loss (mL);
- change in haemoglobin measurements before versus after birth (g/L);



- 8. breastfeeding at hospital discharge;
- 9. nausea;
- 10.vomiting;
- 11.hypertension;
- 12.headache;
- 13.fever (>= 38°C);
- 14.shivering;
- 15.abdominal pain;
- 16.diarrhoea;
- 17.maternal sense of well-being (as defined by the trialists);
- 18.maternal satisfaction (as defined by the trialists).

Search methods for identification of studies

Electronic searches

We searched Cochrane Pregnancy and Childbirth's Trials Register by contacting their Information Specialist (24 May 2018).

The Register is a database containing over 25,000 reports of controlled trials in the field of pregnancy and childbirth. It represents over 30 years of searching. For full current search methods used to populate Pregnancy and Childbirth's Trials Register including the detailed search strategies for CENTRAL, MEDLINE, Embase and CINAHL; the list of handsearched journals and conference proceedings, and the list of journals reviewed via the current awareness service, please follow this link

Briefly, Cochrane Pregnancy and Childbirth's Trials Register is maintained by their Information Specialist and contains trials identified from:

- monthly searches of the Cochrane Central Register of Controlled Trials (CENTRAL);
- 2. weekly searches of MEDLINE (Ovid);
- 3. weekly searches of Embase (Ovid);
- 4. monthly searches of CINAHL (EBSCO);
- handsearches of 30 journals and the proceedings of major conferences;

6. weekly current awareness alerts for a further 44 journals plus monthly BioMed Central email alerts.

Search results are screened by two people and the full text of all relevant trial reports identified through the searching activities described above is reviewed. Based on the intervention described, each trial report is assigned a number that corresponds to a specific Pregnancy and Childbirth review topic (or topics), and is then added to the Register. The Information Specialist searches the Register for each review using this topic number rather than keywords. This results in a more specific search set that has been fully accounted for in the relevant review sections (Included studies; Excluded studies; Studies awaiting classification; Ongoing studies).

In addition, we searched ClinicalTrials.gov and the WHO International Clinical Trials Registry Platform (ICTRP) for unpublished, planned and ongoing trial reports using the terms given in Appendix 1 (24 May 2018).

Searching other resources

We retrieved additional relevant references cited in papers identified through the above search strategy and we did search for the full texts of trials initially identified as abstracts. For randomised trials published only as abstracts, we sought information from primary authors to investigate whether these studies met our eligibility criteria before including them. Trials that compared at least two of the agents were eligible and we searched for all possible comparisons. We did not apply any language or date restrictions.

Data collection and analysis

Selection of studies

Three review authors retrieved and independently assessed for inclusion all the potential studies we identified (IDG, AP, NA). We resolved any disagreements through discussion or, if required, in consultation with a third person (AC). We created a study flow diagram to map out the number of records identified, included and excluded (Figure 1).



Figure 1. Study flow diagram.

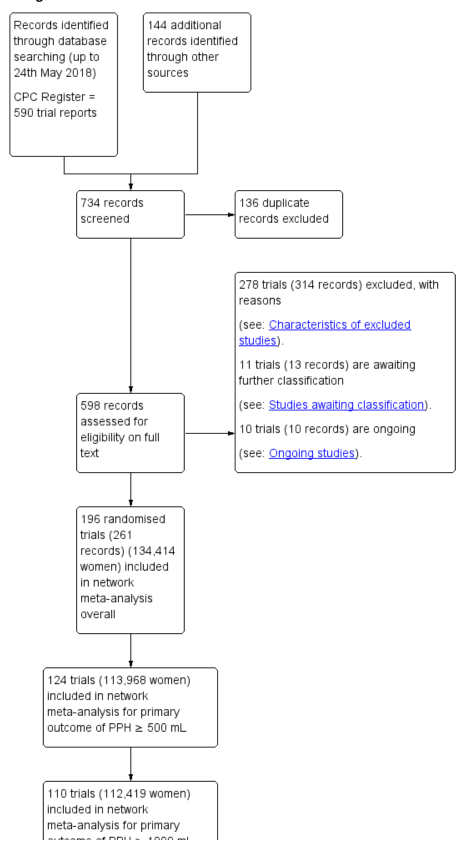




Figure 1. (Continued)

meta-analysis for primary outcome of PPH ≥ 1000 mL

Data extraction and management

We designed an electronic form on ©Microsoft Access to extract data. For eligible studies, at least three review authors independently extracted the data using a blank electronic form (IDG, AP, NA, RM, OT). We resolved discrepancies through discussion or, if required, we consulted another person (AC). We entered data into STATA and Review Manager software (RevMan 2014) and checked for accuracy. When information was unclear, we attempted to contact authors of the original reports to provide further details. The following data were extracted.

Outcome data

From each included study we extracted: the number of participants, the gestational age and the parity of participants, and any exclusion criteria. We also extracted: the interventions being compared, and their respective primary and secondary outcomes. All relevant arm level data were extracted (e.g. number of events and number of patients for binary outcomes and means and standard deviations per study arm for continuous outcomes).

Data on potential effect modifiers

From each included study we extracted the following study, intervention and population characteristics that may act as effect modifiers:

- 1. mode of delivery (vaginal or caesarean birth);
- prior risk of PPH (as defined by trialists and categorised as low, high, mixed or not stated);
- dosage, regimen, and route of administration (sublingual, subcutaneous, intramuscular, rectal, oral, intravenous bolus and/or infusion); and
- 4. setting of the study (community or hospital).

Other data

From each included study we extracted the following additional information:

- 1. country or countries in which the study was performed;
- 2. date of publication and dates of recruitment;
- 3. type of publication (full-text publication, abstract publication, unpublished data); and
- 4. trial registration reference.

Assessment of risk of bias in included studies

At least three (IDG, AP, NA, RM) review authors independently assessed the risk of bias for each study using the criteria outlined in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). Any disagreements were resolved by discussion or by involving another assessor (AC).

(1) Random sequence generation (checking for possible selection bias)

Studies were excluded if found to be at high risk for bias for random sequence generation (any non-random process, e.g. odd or even date of birth; hospital or clinic record number). We described for each included study the method used to generate the allocation sequence in sufficient detail to allow an assessment of whether it should produce comparable groups.

We assessed the methods as:

- low risk of bias (any truly random process, e.g. random number table; computer random number generator); or
- · unclear risk of bias.

(2) Allocation concealment (checking for possible selection bias)

We described for each included study the method used to conceal allocation to interventions prior to assignment and assessed whether intervention allocation could have been foreseen in advance of, or during recruitment, or changed after assignment.

We assessed the methods as:

- low risk of bias (e.g. telephone or central randomisation; consecutively numbered sealed opaque envelopes);
- high risk of bias (open random allocation; unsealed or nonopaque envelopes, alternation; date of birth); or
- unclear risk of bias.

(3.1) Blinding of participants and personnel (checking for possible performance bias)

We described for each included study the methods used, if any, to blind study participants and personnel from knowledge of which intervention a participant received. We considered that studies were at low risk of bias if they were blinded, or if we judged that the lack of blinding would be unlikely to have affected the results.

We assessed the methods as:

- low, high or unclear risk of bias for participants; and
- low, high or unclear risk of bias for personnel.

(3.2) Blinding of outcome assessment (checking for possible detection bias)

We described for each included study the methods used, if any, to blind outcome assessors from knowledge of which intervention a participant received.

We assessed methods used to blind outcome assessment as:

· low, high or unclear risk of bias.



(4) Incomplete outcome data (checking for possible attrition bias due to the amount, nature and handling of incomplete outcome data)

We described for each included study the completeness of data including attrition and exclusions from the analysis. We stated whether attrition and exclusions were reported and the numbers included in the analysis at each stage (compared with the total randomised participants), reasons for attrition or exclusion where reported, and whether missing data were balanced across groups or were related to outcomes. Where sufficient information was reported, or supplied by the trial authors, we re-included missing data in the analyses. We assessed methods to handle incomplete outcome data as:

- low risk of bias (e.g. no missing outcome data; missing outcome data balanced across groups and less than 10% of missing outcome data):
- high risk of bias (e.g. numbers or reasons for missing data imbalanced across groups; 'as treated' analysis done with substantial departure of intervention received from that assigned at randomisation or more than 10% of missing outcome data); or
- unclear risk of bias.

(5) Selective reporting (checking for reporting bias)

We described for each included study how we investigated the possibility of selective outcome reporting bias and what we found.

We assessed the methods as:

- low risk of bias (where it is clear that all of the study's prespecified outcomes and all expected outcomes of interest to the review have been reported);
- high risk of bias (where not all the study's pre-specified outcomes have been reported; one or more reported primary outcomes were not pre-specified; outcomes of interest are reported incompletely and so cannot be used; study fails to include results of a key outcome that would have been expected to have been reported); or
- unclear risk of bias.

(6) Other bias (checking for bias due to problems not covered by (1) to (5) above)

We described for each included study any important concerns about other possible sources of bias, such as the source of funding and potential conflicts of interest.

We assessed these interests as:

- low risk of other bias (public funding or no funding and no significant conflicts of interest identified);
- high risk of other bias (industry funding or significant conflicts of interest identified); or
- unclear risk of other bias.

Another source of bias was generated by the method of measuring blood loss. We assessed the method described in each study and classified it as at:

- low risk of other bias (objective measurements such as weighing sponges, measurements in drapes, volumetric assessment, tagged red cells, etc);
- high risk of other bias (subjective measurement such as clinical or visual estimates); or
- unclear risk of other bias (unspecified methods of measurement).

(7) Overall risk of bias

We made explicit judgements about whether studies were at high risk of bias, according to the criteria given in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). For our primary outcomes, we combined quality items and judged trials as "low risk of bias" if they were double-blinded, had allocation concealment and with little loss to follow-up (less than 10%). Trials were judged as "intermediate risk of bias" if they demonstrated adequate allocation concealment, with assessor blinding and little loss to follow-up (less than 10%). Alternatively, trials were considered to be at "high risk of bias". We explored the impact of the level of bias through undertaking sensitivity analyses - see Sensitivity analysis for information about how the risk of bias was incorporated in the sensitivity analysis.

Summary of findings

The 'Summary of findings' tables present evidence comparing all other uterotonic agents with a reference comparator, oxytocin. Each table describes key features of the evidence relating to a single outcome, and there is one table for each of our seven most important outcomes in accordance with the GRADE approach. These include PPH \geq 500 mL, PPH \geq 1000 mL, blood transfusion, additional uterotonics, vomiting, hypertension and fever.

We used the GRADE working group's approach (Brignardello-Petersen 2018; Puhan 2014) for rating the certainty of the network meta-analysis effect estimates for all the comparisons and all outcomes. We appraised the certainty of the direct, indirect, and network evidence sequentially (in this order). First, we assessed the certainty of the direct evidence (where available) for a given outcome, and rated the evidence using the standard GRADE approach based on consideration of: study design limitations (risk of bias); inconsistency; imprecision; indirectness and publication bias (Higgins 2011). On the network diagram for all the comparisons and all outcomes we display the certainty of the direct evidence. Then we rated the certainty of the indirect evidence for the same outcome, and this was determined based on the lower of the certainty ratings of the two arms forming the dominant 'first-order' loop in the network diagram for this outcome. Our final step was to determine the quality of network evidence based on: (i) the higher certainty rating of the direct and indirect evidence, (ii) whether the relevant network diagram exhibited 'intransitivity', i.e. whether all the comparisons contributing data to the estimate were directly consistent with the PICO question, (iii) consideration of coherence between direct and indirect effect estimates, and (iv) precision of the network effect estimate. Where the network estimate was precise, and the direct and/or indirect evidence contributing to the certainty ratings were not, the certainty of the network evidence was upgraded by one level for precision. At each of these stages, two review authors (MJW, VD, JP, MC) independently appraised the certainty ratings for the direct, indirect and network evidence. Disagreements between authors were resolved through discussion



and consultation with a third review author (OTO, JPV) where necessary.

The quality of network evidence for each outcome was rated as 'high', 'moderate', 'low' or 'very low' in accordance with the GRADE approach:

- High certainty: we are very confident that the true effect lies close to that of the estimate of the effect;
- Moderate certainty: we are moderately confident in the effect estimate. The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different;
- Low certainty: our confidence in the effect estimate is limited. The true effect may be substantially different from the estimate of the effect; and
- Very low certainty: we have very little confidence in the effect estimate. The true effect is likely to be substantially different from the estimate of effect.

For ease of comparison when interpreting the relative effects of all uterotonic agents, the 'Summary of findings' tables include the effect estimate and certainty judgements for each of the direct evidence, indirect evidence and the network metaanalysis, describing all the findings for a single outcome in each table. The anticipated absolute effects are also included, based on the network effect estimate for each agent/agent combination in comparison with oxytocin. The assumed risks in the oxytocin group are based on weighted means of baseline risks from the studies with oxytocin arms in the network metaanalysis. The corresponding risks in the carbetocin, misoprostol, injectable prostaglandins, ergometrine, ergometrine plus oxytocin (Syntometrine ®), misoprostol plus oxytocin groups (and their 95% confidence interval (CI)) are based on the assumed risk in the oxytocin group and the relative effect of the individual uterotonic when compared with oxytocin (and its 95% CI) as derived from the network meta-analysis. The baseline risks differed significantly by the mode of birth subgroups, so the anticipated absolute effects are presented separately for vaginal and caesarean births based on the weighted means of baseline risks according to these modes of birth.

Measures of treatment effect

Relative treatment effects

We summarised relative treatment effects for dichotomous outcomes as risk ratios (RR) and for continuous outcomes as mean difference (MD) with 95% CIs (Dias 2013). These are summarised in forest plots displaying the results from pairwise, indirect and network (combining direct and indirect) analyses for the comparisons of uterotonic agents versus placebo or no treatment and the comparisons of uterotonic agents versus oxytocin. All other comparisons are available from Appendix 2.

Relative treatment ranking

We estimated the cumulative probabilities for each uterotonic agent being at each possible rank and obtained a treatment hierarchy using the surface under the cumulative ranking curve (SUCRA); the larger the SUCRA the higher its rank among all available agents (Salanti 2011). The probabilities to rank the treatments are estimated under a Bayesian model with flat priors, assuming that the posterior distribution of the parameter estimates is approximated by a normal distribution with mean and variance equal to the frequentist estimates and variance-

covariance matrix. Rankings are constructed drawing 1000 samples from their approximate posterior density. For each draw, the linear predictor is evaluated for each study, and the largest linear predictor is noted (White 2011).

Unit of analysis issues

Cluster-randomised trials

For a cluster-randomised trial included in this review (Stanton 2013), we used the unadjusted standard errors as the clusters and the Intracluster Correlation Co-efficient (ICC) was small (ICC = 0.012). Another cluster-randomised trial (Chandhiok 2006) did not report the ICC and the ICC from Stanton 2013 was used. Chandhiok 2006 was reduced it to its effective sample size taking into account the design effects as described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We considered it reasonable to combine the results from the cluster-randomised and the individually-randomised trials as there was little heterogeneity between the study designs and any interaction between the relative effects of agents and the choice of randomisation unit was considered to be unlikely. The effect of the unit of randomisation was also assessed in sensitivity analysis (Higgins 2011).

Multi-arm trials

Multi-arm trials were included and we accounted for the correlation between the effect estimates in the network meta-analysis. We treated multi-arm studies as multiple independent comparisons in pairwise meta-analyses and these were not combined in any analysis.

Dealing with missing data

For included studies, we noted the levels of attrition. We explored the impact of including studies with high levels of missing data in the overall assessment of treatment effect by using sensitivity analysis. For all outcomes, we carried out analyses, as far as possible, on an intention-to-treat basis, i.e. we included all participants randomised to each group in the analyses, and all participants were analysed in the group to which they were allocated, regardless of whether or not they received the allocated intervention. We used the number randomised minus any participants whose outcomes were known to be missing as the denominator for each outcome in each trial.

Assessment of clinical and methodological heterogeneity within treatment comparisons

To evaluate the presence of clinical heterogeneity, we described the study population characteristics across all included trials. We assessed the presence of clinical heterogeneity by comparing these characteristics.

Assessment of intransitivity across treatment comparisons

In this context we expect that the intransitivity assumption holds assuming the following: 1) the common treatment used to compare different uterotonics indirectly is similar when it appears in different trials (e.g. oxytocin is administered in a similar way in oxytocin versus misoprostol trials and in oxytocin versus oxytocin plus ergometrine trials); 2) all pairwise comparisons do not differ with respect to the distribution of effect modifiers (e.g. the design and study characteristics of oxytocin versus misoprostol trials are similar to oxytocin versus oxytocin plus ergometrine trials). The assumption of intransitivity was evaluated epidemiologically by



comparing the clinical and methodological characteristics of sets of studies from the various treatment comparisons.

Assessment of reporting biases

If there were 10 or more studies in the meta-analysis, we investigated reporting biases (such as publication bias) using funnel plots. The funnel plots were assessed visually for asymmetry. We also assessed potential reporting bias for the primary outcomes by assessing the sensitivity of results to exclusion of studies with fewer than 400 participants.

Data synthesis

Methods for direct treatment comparisons

Initially, we performed pairwise meta-analyses using a randomeffects model in Stata and Review Manager software (RevMan 2014) for every treatment comparison with at least two studies (DerSimonian 1986).

Methods for indirect and network comparisons

We initially generated and assessed the network diagrams to determine if a network meta-analysis is feasible. Then we performed the network meta-analysis within a frequentist framework using multivariate meta-analysis estimated by restricted maximum likelihood. All analyses were done using Stata statistical software, release 15 (StataCorp, College Station, TX). We used the network suite of Stata commands designed for this purpose (White 2012; White 2015).

Assessment of statistical heterogeneity

Assumptions when estimating the heterogeneity

In pairwise meta-analyses, we estimated the heterogeneity for each comparison. In network meta-analysis we assumed a common estimate for the heterogeneity variance across all of the different comparisons.

Measures and tests for heterogeneity

We assessed statistically the presence of heterogeneity within each pairwise comparison for the primary outcomes using the I^2 statistic that measures the percentage of variability that cannot be attributed to random error (Higgins 2002). The certainty of the evidence was downgraded for inconsistency where $I^2 \geq 60\%$. The assessment of statistical heterogeneity in the entire network was based on the magnitude of the heterogeneity variance parameter estimated from the multivariate meta-analysis.

Assessment of statistical inconsistency

To check the assumption of consistency in the entire network we used the "design-by- treatment" interaction model as described by Higgins (Higgins 2012). This method accounts for a different source of inconsistency that can occur when studies with different designs (two-arm trials versus three-arm trials) give different results as well as disagreement between direct and indirect evidence. Using this approach we inferred about the presence of inconsistency from any source in the entire network based on a Chi² test.

Investigation of heterogeneity and inconsistency

Where we found important heterogeneity and/or inconsistency, we explored the possible sources for primary outcomes. Where

sufficient studies were available, we performed multivariate metaanalyses for subgroups and sensitivity analyses by using potential effect modifiers as possible sources of inconsistency and/or heterogeneity.

Subgroup analysis

For the primary outcomes we carried out the following prespecified subgroup analyses.

- Population: prior risk of PPH (high versus low), mode of delivery (vaginal versus caesarean birth), setting (hospital versus community).
- 2. Intervention: dose of misoprostol (≥ 600 mcg versus < 600 mcg), and regimen of oxytocin (bolus versus bolus plus infusion versus infusion only).

We assessed subgroup differences by firstly comparing the network diagram for each subgroup. Next, we performed a pairwise and network meta-analysis for each subgroup and we compared their relative treatment effects and their relative treatment ranking. We examined the subgroups for qualitative interactions where the direction of effect could be reversed, that is if an intervention was beneficial in one subgroup but harmful in another.

Sensitivity analysis

For the primary outcomes we performed sensitivity analysis for the following.

- 1. Risk of bias (restricted to low risk of bias studies only): studies are ranked as 'low risk of bias' if they are double-blinded, and have allocation concealment with little loss to follow-up (less than 10%). The concealed studies with assessor blinding and little loss to follow-up (less than 10%) are ranked as 'intermediate risk of bias' and the rest as 'high risk of bias'. We considered that assessor blinding was likely to be very important, in order to eliminate any risk of bias in subjective measurements or estimates of blood loss (not all studies measure this outcome objectively). We considered protocol publication in advance of the results to be an unsuitable criterion for sensitivity analyses, because protocol publication only became widespread in recent years.
- 2. Funding source (restricted to studies with funding source at low risk of bias (public or no funding)).
- 3. Whether an objective method of outcome assessment was employed (restricted to studies with an objective method of measuring blood loss). Objective methods of blood loss measurement were considered to be all methods that employed a measurement of the blood loss. This is in contrast to subjective methods where a healthcare professional is estimating the blood loss, usually visually.
- 4. Trial size (restricted to large studies (> 400 participants), in recognition of the greater likelihood for small studies than large or multi-centre studies to suffer publication bias). In terms of trial size, there is evidence that smaller studies can exaggerate estimated benefits (Nüesch 2010). However, the cutoff for deciding the definition of a small study can vary between research topics. For this topic, it appears that trials with more than 400 participants are more likely to be of higher quality, prospectively registered and overall at low risk of bias.



- Removing trials that also randomised participants to cointerventions such as uterine massage or controlled cord traction.
- 6. Removing trials with more than 10% missing data.
- 7. Removing trials published before 1990.
- 8. Randomisation unit (restricted to individually-randomised trials and removing cluster-randomised trials).
- Choice of relative effect measure (risk ratio (RR) versus odds ratio (OR)).

10.Use of fixed-effect versus random-effects model.

Differences were assessed by evaluating the relative effects and assessment of model fit.

RESULTS

Description of studies

Results of the search

The results of the search are summarised in the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) flow diagram (Figure 1).

The search of Cochrane Pregnancy and Childbirth's (CPC) Trials Register on 24 May 2018 retrieved in total 590 records. We retrieved a further 144 records from additional author searches and manual searching of reference lists for a total of 734 available records. From these, we excluded 136 records as duplicates. We examined the full text of 598 records and included in the network meta-analysis 196 randomised trials (reported in 261 publications).

We contacted the authors from 98 randomised trials for additional data or clarifications. We were able to obtain additional data from trial authors for 39 randomised trials (Characteristics of included studies and Appendix 2). We excluded 278 trials (reported in 314 publications) (Characteristics of excluded studies), 11 trials (reported in 13 publications) could not be classified (Studies awaiting classification) and 10 trials were still ongoing (Ongoing studies).

Included studies

The network meta-analysis includes 196 randomised trials involving 135,559 women. Most studies were reported in English; 11 translations were obtained (six Spanish, two French, two Turkish and one Chinese). The studies were conducted across 53 countries (including high-, middle- and low-income countries) and often involved more than one country. A number of multi-arm trials were identified: two five-arm trials, eight four-arm trials and 22 three-arm trials. The median size of the trials was around 213 participants (interquartile range (IQR) 123 to 529).

Most trials (95.4%, 187/196) were performed in a hospital setting, seven were performed in a community setting (3.6%), one (0.5%) in a mixed setting and one (0.5%) of unspecified setting. The majority of the trials included women undergoing a vaginal birth (71.5%, 140/196), and 53 trials (27%) involved women undergoing elective or emergency caesareans. Only two (1%) trials included women undergoing either a vaginal birth or caesarean and in one trial (0.5%) the mode of birth was not specified. Women included in the trials were judged to be at high risk for postpartum haemorrhage (PPH) in 66 of 196 trials (33.7%), low risk in 52 trials (26.5%) and 68 trials (34.7%) included women both at high or low risk for PPH. The risk for PPH was not specified in 10 trials (5.1%).

Women with a singleton pregnancy only were recruited in 124 trials (63.3%), 36 trials (18.4%) included women with either singleton or multiple pregnancies and only one trial (0.5%) included women with twin pregnancies only. Thirty-five trials (17.8%) did not specify. Six trials (3.1%) included only nulliparous or primigravida women, one trial included only multiparous women (0.5%), 108 trials (55.1%) included both nulliparous and multiparous women of all parities, and 81 trials (41.3%) did not specify the parity of the women included in the trials. Exclusion criteria varied significantly and usually encompassed women with significant medical comorbidities.

Across all 196 trials (412 trial arms) in the network metaanalysis, the following agents were used either as intervention or comparison:

- 1. 137 trial arms (33.3%) used oxytocin;
- 2. 96 trial arms (23.3%) used misoprostol;
- 3. 39 trial arms (9.5%) used ergometrine;
- 4. 35 trial arms (8.5%) used ergometrine plus oxytocin;
- 5. 33 trial arms (8%) used carbetocin;
- 6. 29 trial arms (7%) used placebo or no treatment;
- 7. 26 trial arms (6.3%) used misoprostol plus oxytocin;
- 8. 17 trial arms (4.1%) used injectable prostaglandins.

See Characteristics of included studies for details.

Excluded studies

We excluded 278 trials (for details see Characteristics of excluded studies). The most common reasons for exclusion were because trials were comparing exclusively doses or routes of the same uterotonic agents, trials that were quasi randomised or trials investigating ineligible interventions such as tranexamic acid.

Risk of bias in included studies

We present summaries of the methodological quality of the included studies for each of the domains we assessed across all studies (Figure 2) and for each included study (Figure 3).



Figure 2. 'Risk of bias' graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.

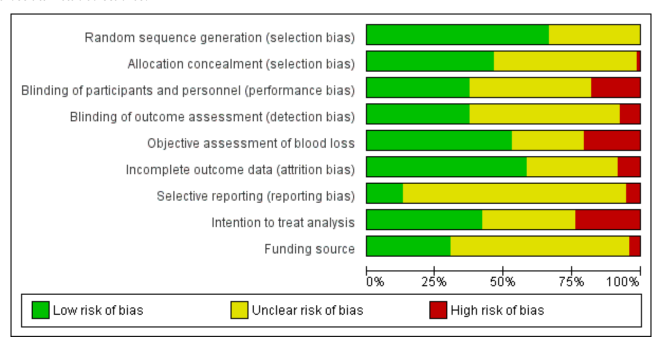




Figure 3. 'Risk of bias' summary: review authors' judgements about each risk of bias item for each included study.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Objective assessment of blood loss	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Intention to treat analysis	Funding source
Abdel-Aleem 1993	•	?	?	?	•	?	?	?	?
Abdel-Aleem 2010	•	•	?	?	•	•	?	•	•
Acharya 2001	?	•	?	?	•	•	?	•	?
Adanikin 2012	•	•	•	•	?	•	?	•	?
Adanikin 2013	•	•	•	•	•	•	?	•	?
Afolabi 2010	•	?	?	?	•	•	?	•	?
Ahmed 2014	?	?	?	?	?	?	?	?	?
Al-Sawaf 2013	?	?	?	?	•	•	?	•	?
Alwani 2014	•	?	?	?	?	?	?	?	•
Amant 1999	•	•	•	•	?	•	?	•	?
Amin 2014	?	?	?	?	•	?	?	?	?
Askar 2011	•	•	•	•	•	•	?	•	?
Asmat 2017	•	•	?	?	?	?	?	?	?
Attilakos 2010	•	•	•	•	•	•	•	•	•
Atukunda 2014	•	•	•	•	•	•	•	•	•
Badejoko 2012	•	•	•	•	•	•	?	•	•
Balki 2008	•	•	•	•	•	•	?	•	•
Bamigboye 1998a	•	•	•	?	•	•	?	•	?
Bamigboye 1998b	•	•	?	?	•	?	?	•	•
Barton 1996	?	?	?	?	?	?	?	?	?



Figure 3. (Continued)

B-st-11 4000	•	_	_	•	_	_	_	_	_
Barton 1996	?	?	?	?	?	?	?	?	?
Baskett 2007	•	•	•	•	•	•	?	•	•
Begley 1990	•	•	•		•	•	?	•	•
Begum 2015	?	?	?	?	?	?	?	?	?
Bellad 2012	•	•	•	•	•	•	?	•	•
Benchimol 2001	•	?	?	?	•	•	?	•	?
Bhatti 2014	?	?	?	?	•	?	?	?	?
Bhullar 2004	•	•	?	?	•	•	?	•	?
Biswas 2007	?	?	?	?	•	?	?	?	?
Borruto 2009	?	?	?	?	•	•	?	?	
Boucher 1998	?	?	•	•	•	•	?		
Boucher 2004	•	?	•	•	?	•	?	•	•
Bugalho 2001	?	?	•	?	•	•	•	•	•
Butwick 2010	•	?	•	?	•	•	?	•	•
Caliskan 2002	•	•	•	•	•	•	?	•	?
Caliskan 2003	•	•	•	•	•	•	?	•	?
Carbonell 2009	•	•	?	?	•	•	?	•	•
Carrillo-Gaucin 2016	?	?	?	?	?	•	?	?	?
Cayan 2010	?	?	?	?	?	•	?	•	?
Chalermpolprapa 2010	?	?	?	?	?	?	?	?	?
Chandhiok 2006	?	•	?	?	•	•	?	•	•
Chaudhuri 2010	•	•	•	•	•	•	•	•	?
Chaudhuri 2012	•	•	•	•	•	•	•	•	?
Chaudhuri 2015	•	•	•	•	•	•	•	•	?
Chaudhuri 2016	•	•	•	•	•	•	•	•	?
Chhabra 2008	•	?	?	?	?	?	?	?	?
Choy 2002	•	•	•	•	•	•	?	•	?
Chua 1995	•	?	?	?	•	?	?	•	?
Cook 1999	•	•	•	•	?	•	?	•	?
Dabbaghi Gale 2012	?	?	?	?	?	?	?	?	?
Dansereau 1999	•	?	•	•	?	•	?	•	•
	'	'	'		'	'	'	'	



Figure 3. (Continued)

Dansereau 1999										
Del Angel-Garcia 2006 Derman 2004 Del Angel-Garcia 2006 Del Angel-Garcia 2007 Del Angel-Garcia 2006 Del Angel-Garcia 2007 Del Angel-	Dansereau 1999	•	?	•	•	?	•	?	•	
Del Angel-Garcia 2006 Derman 2006 Derman 2006 Diallo 2017 Diallo 2017 Diop 2016 Docherty 1981 Dutta 2016 Cite Refaera 2007 El Behery 2015 El Sankwy 2013 El-Refaev 2000 Elsedeek 2012 El Tahan 2012 Del Angel-Garcia 2006 Fakour 2015 Fakour 2016 Cite Refaev 2007 Fakour 2016 Fakour 2016 Fararjeh 2003 Fararjeh 2003 Fararjeh 2003 Fararyeh 2016 Fararyeh 2016 Fararyeh 2016 Fararyeh 2016 Fararyeh 2016 Fararyeh 2016 Fararyeh 2017 Fararyeh 2017 Fararyeh 2018 Fararyeh 2011 Fararyeh 2011 Fararyeh 2012 Fararyeh 2013 Fararyeh 2014 Fararyeh 2015 Fararyeh 2014 Fararyeh 2015 Fararyeh 2017 Far	Dasuki 2002	?	?	?	?	?	?	?	?	?
Derman 2006 Dhananjaya 2014 Diallo 2017 Diallo 2017 Diop 2016 Docherty 1981 Dutta 2016 Deffekhari 2009 El Behery 2015 Elbohoty 2016 El-Refaev 2000 El-Refaev 2000 Elsedeek 2012 El Tahan 2012 El Tahan 2012 Enakpene 2007 Ezeama 2014 Fahmy 2016 Deffekhari 2007 Ezeama 2014 Fahmy 2016 Fahmy 2017 Fawole 2011 Fawo	de Groot 1996	•	•	•	?	•	•	?	•	?
Dhananjaya 2014 7 7 7 7 7 7 7 7 7	Del Angel-Garcia 2006	?	?	?	?	?	?	?	?	?
Diallo 2017 Diop 2016 Diop 2016	Derman 2006	•	•	•	•	•	•	•	•	•
Diop 2016	Dhananjaya 2014	?	?	?	?	•	?	?	?	?
Docherty 1981 Dutta 2016 Eftekhari 2009 El Behery 2015 El Behery 2016 El Refaey 2000 El-Refaey 2000 El Sadeek 2012 El Tahan 2012 Enakpene 2007 Ezeama 2014 Fahrmy 2016 Fahrmy 2	Diallo 2017	•	•	?	•	•	?	?	?	•
Dutta 2016	Diop 2016	•	•	•	•	•	•	•	•	•
Effekhari 2009	Docherty 1981	?	?	?	?	?	?	?	?	?
El Behery 2015 Elbohoty 2016 Elbohoty 2016 El-Refaey 2000	Dutta 2016	?	?	?	?	?	?	?	?	?
Elbohoty 2016	Eftekhari 2009	?	?	•	?	•	?	•	?	?
Elgafor El Sharkwy 2013 El-Refaey 2000 El-Refaey 2000 Elsedeek 2012 Elsedeek 2012 El Tahan 2012 Enakpene 2007 Ezeama 2014 Ezeama 2014 Enahmy 2015 Fahmy 2016 Fakour 2013 Refararjeh 2003 Refararjeh 2	El Behery 2015	•	•	•	•		?	?	?	?
El-Refaey 2000 Elsedeek 2012 El Tahan 2012 El Tahan 2012 Enakpene 2007 Ezeama 2014 Fahmy 2015 Fahmy 2016 Favor 2013 Fararjeh 2003 Favor 2014 Fawzy 2012 Fawzy 2012 Faxel 2013 Faxel 2013 Fekih 2009 Fekih 2009 Fuks 2014 Fuks 2014 Payson Aller	Elbohoty 2016	•	•	•	•	•	•	•	•	?
Elsedeek 2012	Elgafor El Sharkwy 2013	•	•	•	•	•	•	?	•	?
El Tahan 2012 Enakpene 2007 Ezeama 2014 Fahmy 2015 Fahmy 2016 Fakour 2013 Favole 2011 Fawzy 2012 Fazel 2013 Fazel 2013 Fakour 2012 Fu 2003 Fuks 2014 Fuks 2014 Fuks 2014 Fakour 2014 Faxel 2014 Fuks 2014	El-Refaey 2000	•	•	•	•	•	•	?	•	?
Enakpene 2007 Ezeama 2014 Fahmy 2015 Fahmy 2016 Fakour 2013 Fararjeh 2003 Fawzy 2012 Fazel 2013 Fekih 2009 Fuks 2014 Fuks 2014 Fuks 2014 Fuks 2014 Fuks 2014 Fixed 2014 Fixed 2014 Fuks 2014 Fixed 2014 Fixe	Elsedeek 2012	•	?	•	•	•	•	?	•	•
Ezeama 2014 Fahmy 2015 Fahmy 2016 Fakour 2013 Fararjeh 2003 Fawzy 2012 Fazel 2013 Fekih 2009 Fuks 2014 Fuks 2014 Fuks 2014 Fuks 2014 Fakour 2014 Faxole 2014 Faxole 2014 Faxole 2016 Faxole 2017 Faxole 2018 Faxole 2018 Faxole 2018 Faxole 2019 Fax	El Tahan 2012	•	•	•	•	•	•	?	•	•
Fahmy 2015 Fahmy 2016 Pahmy 2016 Pahmy 2016 Pakour 2013 Pararjeh 2003 Fawole 2011 Fawzy 2012 Pazel 2013 Fekih 2009 Paxel 2012 Paxel 2014 Paxel 2014 Paxel 2014 Paxel 2014 Paxel 2014	Enakpene 2007	•	•	?	?	•	•	?	•	•
Fahmy 2016	Ezeama 2014	•	•	•	•	•	•	•	•	•
Fakour 2013 ? ? • • ? ? ? ? ? ? ? ? ? Fararjeh 2003 • ? ? ? • • • ? ?	Fahmy 2015	•	•	?	?	•	?	?	?	?
Fararjeh 2003	Fahmy 2016	•	?	?	?	?	?	?	?	?
Fawole 2011 Fawzy 2012 ? ? • • • ? ? ? ? ? ? ? ? ? ? ? ? ? ?	Fakour 2013	?	?	•	•	?	?	?	?	?
Fawzy 2012 ? ? • • • ? ? ? ? ? Fazel 2013 • ? ? ? • • ? ? ? • • ? ? Fekih 2009 • • ? ? • • ? ? • ? Fenix 2012 • ? • • • ? ? • ? ? ? ? ? ? ? ? ? ? ?	Fararjeh 2003	•	?	?	?	•	•	?	•	?
Fazel 2013	Fawole 2011	•	•	•	•	•	•	?	?	•
Fekih 2009	Fawzy 2012	?	?	•	•	•	?	?	?	?
Fenix 2012	Fazel 2013	•	?	?	?	•	?	?	?	•
Fu 2003 ? ? ? ? • ? ? ? ? ? ? ? ? ? ? ? ? ? ?	Fekih 2009	•	•	?	?	•	•	?	•	?
Fuks 2014 ? ? ? ? ? ? ? ? ?	Fenix 2012	•	?	•	•	•	•	?	•	?
	Fu 2003	?	?	?	?	•	?	?	?	?
Garg 2005 ? ? ? ? ? ? ? ?	Fuks 2014	?	?	?	?	?	?	?	?	?
	Garg 2005	?	?	?	?	?	?	?	?	?



Figure 3. (Continued)

I									
Garg 2005	?	?	?	?	?	?	?	?	?
Gavilanes 2016	•	?		?	•	?	?	?	?
Gerstenfeld 2001	•	•	•	•	•	•	?		?
Gore 2017	?	?	?	?	•	?	?	?	•
Gulmezoglu 2001	•	•	•	•	•	•	•	•	•
Gupta 2006	•	?	•	•	•	•	?	•	?
Hamm 2005	•	•	•	•	?	•	?	•	?
Harriott 2009	•	?	•	•	•	•	?	•	•
Hernandez-Castro 2016	•	•	?	?	•	•	•	•	?
Hofmeyr 1998	•	•	•	•	•	•	?	•	•
Hofmeyr 2001	•	•	•	•	•	•	?	•	•
Hofmeyr 2011	•	•	•	•	•	•	•	•	•
Hoj 2005	•	•	•	•	•	•	?	•	•
Hong 2007	?	?	?	?	?	?	?	?	?
Humera 2016	?	?	?	?	•	?	?	?	•
ls 2012	?	?	?	?	?	?	?	?	?
Jago 2007	•	?	?	?	?	•	?	•	?
Jangsten 2011	•	•	•	•	•	•	?	•	•
Jans 2017	•	•	•	•	•	•	?	•	•
Jerbi 2007	?	?	?	?	?	•	?	•	?
Jirakulsawas 2000	?	?	?	?	?	?	?	?	?
Kabir 2015	•	?	?	?	•	•	?	•	?
Karkanis 2002	•	•	•	•	?	•	?	•	•
Kerekes 1979	?	?	?	?	?	•	?	•	?
Khan 1995	?	•	•	•	•	•	?	•	?
Khurshid 2010	•	?	?	?	•	?	?	?	?
Koen 2016	•	•	•	•	•	•	•	•	?
Kumar 2016	?	•	?	?	•	?	?	•	?
Kumru 2005	?	?	?	?	•	?	?	?	?
Kundodyiwa 2001	•	•	•	•	•	•	?	•	?
Kushtagi 2006	?	?	?	?	•	?	?	?	?



Figure 3. (Continued)

Kushtagi 2006	?	?	?	?	•	?	?	?	?
Lam 2004	•	?	•	?	•	?	?	?	?
Lamont 2001	?	?	•	•	?	?	?	•	?
Lapaire 2006	•	•	•	•	•	•	•	•	•
Leung 2006	•	•	•	•	•	•	•	•	•
Lokugamage 2001	•	•	•	?	•	•	?	•	•
Lumbiganon 1999	•	•	•	•	•	•	?	•	•
Maged 2016	•	?	?	?	•	•	?	•	?
Maged 2017	•	?	?	?	•	•	?	•	?
Malik 2018	?	?	?	?	•	?	?	?	?
Mannaerts 2018	•	?	•	•	?	•	•	?	?
McDonald 1993	•	•	•	•	•	•	?	•	•
Mitchell 1993	?	•	•	•	•	•	?	•	•
Mobeen 2011	•	•	•	•	•	•	•	•	•
Modi 2014	?	?	?	?	•	?	?	?	•
Moertl 2011	•	?	•	•	•	•	•	•	•
Mohamed 2015	•	?	?	?	•	?	?	?	?
Moir 1979	?	?	?	?	•	•	?	•	?
Moodie 1976	?	?	?	?	•		?	•	?
Mukta 2013	?	?	?	?	?	•	?	•	?
Musa 2015	•	?	•	•	•	•	?	•	•
Nankaly 2016	?	?	•	•	?	?	?	?	?
Nasr 2009	•	•	•	•	•	•	?	•	?
Nayak 2017	?	?	?	?	•	?	?	?	?
Nellore 2006	?	?	?	?	?	?	?	?	?
Ng 2001	•	•	•	•	•	•	?	•	?
Ng 2004	?	?	?	?	?	?	?	?	?
Ng 2007	•	•	•	•	•	•	•	•	?
Nirmala 2009	•	?	?	?	•	•	?	•	?
Nordstrom 1997	•	•	•	•	•	•	?	•	•
Nuamsiri 2016	•	•	•	•	•	•	?	•	?

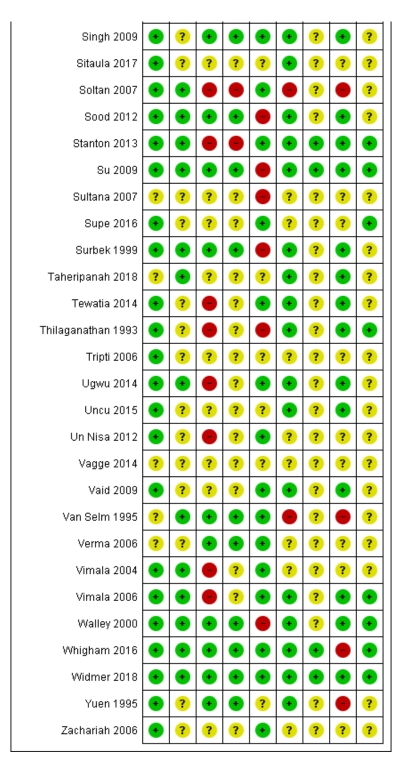


Figure 3. (Continued)

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Orji 2008		_	_	_	_	_		_	_	
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Quibel 2016 • • • • • • • • • • • • • • • • • • •	Poeschmann 1991	?	•	•	•	•	•	?	•	?
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Reyes 2011 ? ? ? ? ? ?	Reddy 2001	?	?	?	?	?	?	?	?	?
Rogers 1998	Reyes, Gonzalez 2011	•	•	•	•	?	•	?	•	?
Rosseland 2013 Sadiq 2011 Samimi 2013 Shady 2017 Shrestha 2011 Rosseland 2013 Procedure of the content of	Reyes 2011	?	?	?	?	?	•	•	•	•
Sadiq 2011	Rogers 1998	•	•	•	•	•	•	?	•	•
Samimi 2013	Rosseland 2013	•	•	•	•	•	•	•	•	•
Samimi 2013	Sadiq 2011	•	?	•	?	?	•	?	•	•
Shady 2017		•		•			•		•	
Shrestha 2011 • ? • ? • ? ? ?	Shady 2017	•		?	?	•	?	?	?	
		•	?			•	_		?	
	Singh 2009	•	?	•	•	•	•	?	•	?



Figure 3. (Continued)



Allocation

Trials with evidence of inadequate random sequence generation were excluded from this review. As a result 130 of 196 included trials (66.3%) were found to have used an adequate method generating the random sequence and were at low risk of bias. However, 66 trials (33.7%) did not report the method used in sufficient detail and the risk of bias was judged to be unclear. Ninety of 196 trials (45.9%)

reported adequate methods for allocation concealment and were judged to be at low risk of bias. Only three trials (1.5%) showed evidence of inadequate allocation concealment and 103 trials (52.6%), did not provide enough information to assess allocation concealment and the risk of bias was judged to be unclear.



Blinding

In total, 73 of 196 trials (37.2%) reported adequate methods for blinding both participants and personnel to treatment allocation. Thirty-five trials (17.9%) were judged to be at high risk of bias for blinding of participants and personnel. Eighty-eight trials (44.9%) did not provide enough information to assess the blinding of participants and personnel and the risk of bias was judged to be unclear. Seventy-three of 196 trials (37.2%) reported adequate methods for blinding the assessment of the primary outcomes. Fifteen trials (7.7%) were judged to be at high risk of bias for blinding the assessment of the primary outcomes. There were 108 trials (55.1%) that did not provide enough information for blinding the assessment of the primary outcomes and the risk of bias was judged to be unclear.

Incomplete outcome data

There were 114 of 196 trials (58.1%) that were judged to be at a low risk of bias. In these trials, missing outcome data were less than 10% for the primary outcomes of the review and balanced in numbers across intervention groups with similar reasons for missing data across groups. In 16 trials (8.2%), more than 10% of patients dropped out or were not analysed as per the "intention-to-treat" principles following randomisation, indicating a high risk of bias. Sixty-six trials (33.7%) did not provide enough information to assess so that it was uncertain whether or not the handling of incomplete data was appropriate and the risk of bias was judged to be unclear in these trials.

Selective reporting

Only 25 of 196 trials (12.8%) pre-specified all outcomes in publicly available study protocols and were judged to be at low risk of bias. Ten trials (5.1%) did not report all pre-specified outcomes as reported in their published protocols or methodology within the main report and were judged to be at high risk of bias for selective reporting. For most trials (161 trials; 82.1%), we were unable to identify a published protocol and the risk of bias was judged to be unclear.

Other potential sources of bias

Eighty-two of 196 trials (41.8%) analysed data by the intention-to-treat principle and were judged to be at low risk of bias. Forty-seven trials (24%) did not analyse data by the intention-to-treat principle and were judged to be at high risk of bias. For 67 trials (34.2%), we were unable to identify whether data were analysed by the intention-to-treat principle and the risk of bias was judged to be unclear.

We found that 59 of 196 trials (30.1%) were either conducted with public or no funding, and declared that they had no potential

conflicts of interest. Eight trials (4.1%) were judged to be at high risk of bias as they were funded directly by the manufacturer of the drug under investigation. There were 129 trials (65.8%) that did not provide enough information to assess the source of funding or potential conflicts of interest and the risk of bias was judged to be unclear.

Among all the studies, 103 of 196 trials (52.6%) reported relatively objective methods for measuring blood loss such as weighing sponges, measurements in drapes or volumetric assessment and were judged to be at low risk of bias. The studies that did not measure blood loss as this was not an outcome of interest were also considered at low risk of bias. Forty-one trials (20.9%) were judged to be at high risk of bias for measuring blood loss as they used subjective measurement such as clinical or visual estimates. Fifty-two trials (26.5%) did not provide enough information to assess the method for measuring blood loss, and the risk of bias was judged to be unclear.

For the purpose of sensitivity analysis we analysed how many trials were judged to be at low, intermediate or high overall risk of bias. For PPH ≥ 500 mL, 38 of 124 trials (30.6%) were found to be at low overall risk of bias. Eighty-six of 124 trials (69.4%) were judged to be at high risk of bias as they were judged to be either at high risk or unclear risk of bias for at least one of the domains mentioned above. There were no trials judged as intermediate risk of bias - see Sensitivity analysis for information about how this risk of bias has impacted the results.

Effects of interventions

See: Summary of findings for the main comparison PPH >= 500 mL; Summary of findings 2 PPH >= 1000 mL; Summary of findings 3 Additional uterotonics; Summary of findings 4 Blood transfusion; Summary of findings 5 Vomiting; Summary of findings 6 Hypertension; Summary of findings 7 Fever

Please note that all of the analyses presented in the Data and analyses section relate to the 'direct evidence' and were used as per our methods to grade the evidence. The results from Data and analyses were also used to check the direction of effect in the subgroups and not to formally check for subgroup effects using the interaction test. These results are not described.

The following section presents the results as reported in all of the figures (Figure 4 to Figure 5). The figures present the results from the network diagrams, the forest plots with the pairwise, indirect and network (combining direct and indirect) effect estimates and the cumulative rankograms for all the outcomes with available data. The figures present the results for different uterotonics in comparison to placebo or no treatment and different uterotonics in comparison to the reference uterotonic agent oxytocin. All other comparisons are available from Appendix 2.



Figure 4. Network Diagram for PPH ≥ 500 mL. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

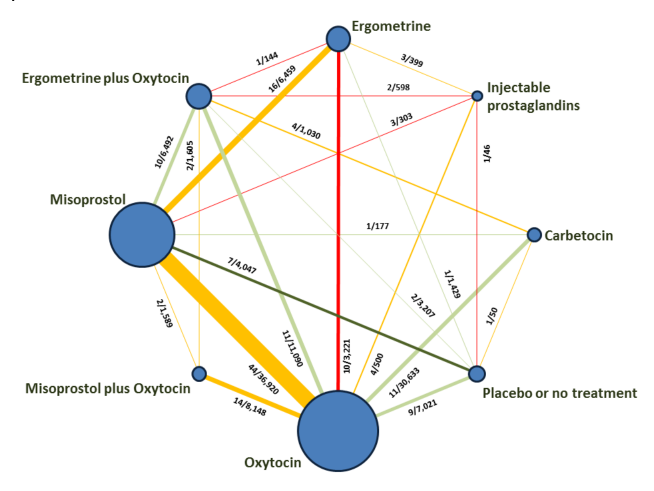
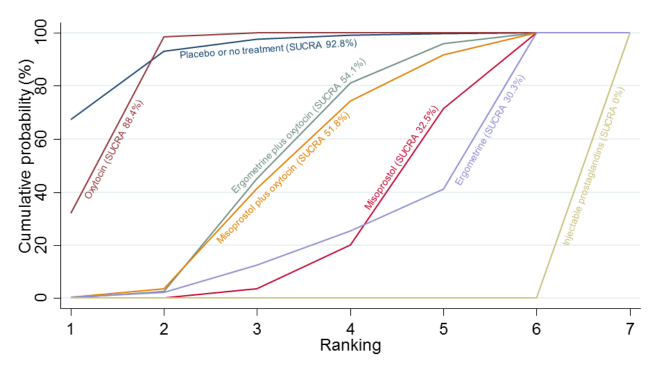




Figure 5. Cumulative rankograms comparing each of the uterotonic agents for diarrhoea. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Primary outcomes

Postpartum haemorrhage (PPH) ≥ 500 mL

The network diagram for PPH ≥ 500 mL is presented in Figure 4. Oxytocin was the most frequently investigated uterotonic agent (88 of 124 trials, 71%) for this outcome (Figure 4).

Relative effects from the network meta-analysis of 124 trials suggested that all agents were effective for preventing PPH ≥ 500 mL when compared with placebo or no treatment (Figure 6). When compared with oxytocin, ergometrine plus oxytocin combination, carbetocin, and misoprostol plus oxytocin combination were more effective in preventing PPH ≥ 500 mL. When compared with oxytocin, moderate-certainty evidence suggests that carbetocin

(risk ratio (RR) 0.72, 95% confidence interval (CI) 0.56 to 0.93) and ergometrine plus oxytocin (RR 0.70, 95% CI 0.59 to 0.84) probably reduce PPH \geq 500 mL, while low-certainty evidence suggests that misoprostol plus oxytocin (RR 0.70, 95% CI 0.58 to 0.86) may reduce PPH \geq 500 mL. Low-certainty evidence suggests that misoprostol, injectable prostaglandins, and ergometrine may make little or no difference to this outcome compared with oxytocin (Summary of findings for the main comparison). Based on these results, about 122 per 1000 women given oxytocin for a vaginal birth would experience a PPH of \geq 500 mL compared with 85 given ergometrine plus oxytocin combination, 87 given carbetocin, and 85 given misoprostol plus oxytocin (Summary of findings for the main comparison).

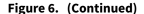


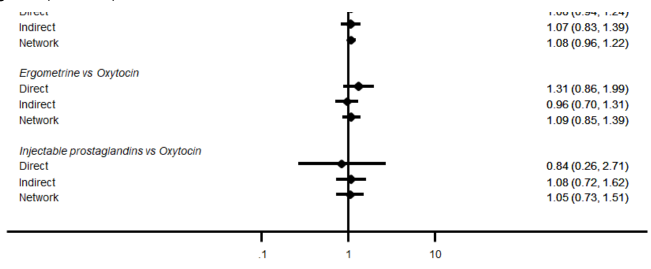
Collaboration.

Figure 6. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for prevention of PPH ≥ 500 mL.

Comparison	RR (95% CI)
Oxytocin vs Placebo or no treatment	
Direct	0.61 (0.52, 0.71)
Indirect -	0.57 (0.43, 0.74)
Network •	0.58 (0.49, 0.70)
Ergometrine+Oxytocin vs Placebo or no treatment	
Direct -	0.37 (0.30, 0.46)
Indirect -	0.42 (0.33, 0.55)
Network -	0.41 (0.33, 0.51)
Carbetocin vs Placebo or no treatment	
Direct	0.75 (0.30, 1.85)
Indirect	0.40 (0.29, 0.55)
Network ——	0.42 (0.31, 0.57)
Misoprostol+Oxytocin vs Placebo or no treatment	0.44 (0.24, 0.52)
Network	0.41 (0.31, 0.53)
Misoprostol vs Placebo or no treatment	
Direct ←	0.75 (0.59, 0.94)
Indirect •	0.57 (0.45, 0.73)
Network •	0.63 (0.52, 0.76)
Ergometrine vs Placebo or no treatment	
Direct	0.24 (0.14, 0.42)
Indirect	0.73 (0.55, 0.98)
Network	0.63 (0.48, 0.84)
Injectable prostaglandins vs Placebo or no treatment	
Direct	0.55 (0.22, 1.35)
Indirect	0.62 (0.41, 0.93)
Network	0.61 (0.42, 0.90)
Ergometrine+Oxytocin vs Oxytocin	
Direct	0.72 (0.57, 0.91)
Indirect	0.69 (0.54, 0.90)
Network	0.70 (0.59, 0.84)
Carbetocin vs Oxytocin	
Direct	0.75 (0.58, 0.98)
Indirect	0.59 (0.31, 1.12)
Network	0.72 (0.56, 0.93)
Misoprostol+Oxytocin vs Oxytocin	
Direct	0.71 (0.59, 0.85)
Indirect	0.79 (0.35, 1.77)
Network	0.70 (0.58, 0.86)
Misoprostol vs Oxytocin	
Misoprostol vs Oxytocin Direct	1.08 (0.94, 1.24) 1.07 (0.83, 1.39)



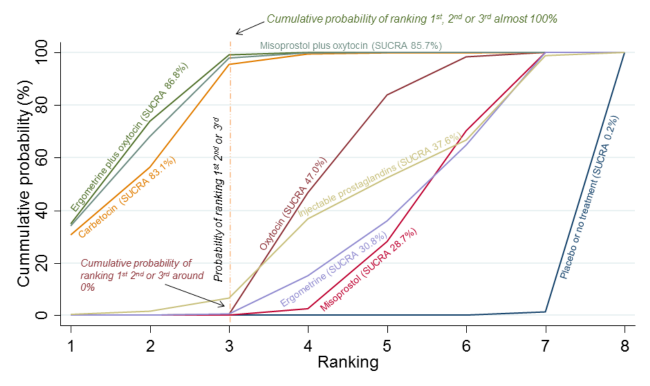




The cumulative probabilities for each agent being at each possible rank for preventing PPH ≥ 500 mL are shown in Figure 7. Treatment hierarchies are presented with the surface under the cumulative ranking curve (SUCRA); the larger the SUCRA the higher its rank among all available agents. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. A SUCRA of 100% means the uterotonic agent is the best and

a SUCRA of 0% means the agent is the worst. The highest ranked agents were ergometrine plus oxytocin combination (SUCRA 86.8%), misoprostol plus oxytocin combination (SUCRA 85.7%) and carbetocin (SUCRA 83.1%). Oxytocin ranked fourth (47%) followed by injectable prostaglandins (SUCRA 37.6%), ergometrine (SUCRA 30.8%), misoprostol (SUCRA 28.7%) and placebo or no treatment (SUCRA 0.2%).

Figure 7. Cumulative rankograms comparing each of the uterotonic agents for prevention of PPH ≥ 500 mL. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.

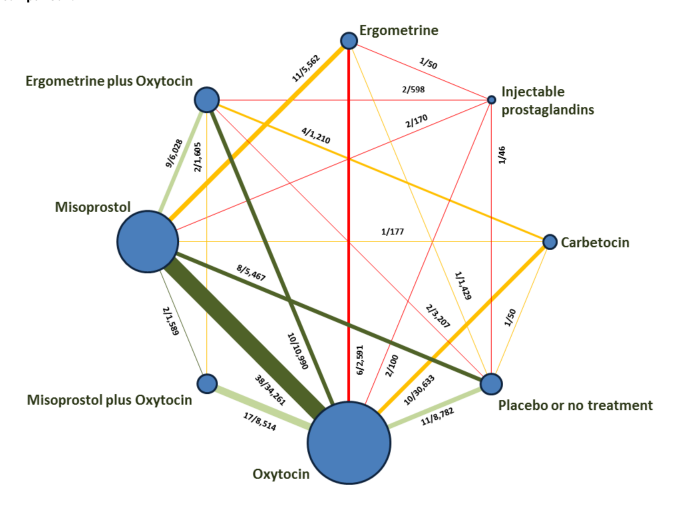




Postpartum haemorrhage (PPH) ≥ 1000 mL

The network diagram for PPH \geq 1000 mL is presented in Figure 8. Oxytocin was the most frequently investigated uterotonic agent (72.7%, 80 of 110 trials) for this outcome (Figure 8).

Figure 8. Network Diagram for PPH ≥ 1000 mL. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.



Relative effects from the network meta-analysis of 110 trials suggested that all agents except ergometrine and injectable prostaglandins were effective for preventing PPH \geq 1000 mL when compared with placebo or no treatment (Figure 9). No differences were observed in the effects of uterotonic agents compared with the reference uterotonic agent oxytocin for PPH \geq 1000 mL. High-certainty evidence suggests that misoprostol plus oxytocin (RR 0.88, 95% CI 0.70 to 1.11) and ergometrine plus oxytocin (RR 0.83, 95% CI 0.66 to 1.03) make little or no difference to PPH \geq 1000 mL when compared with oxytocin. In absolute terms, these results

suggest that about 30 per 1000 women given oxytocin for a vaginal birth would experience PPH \geq 1000 mL, compared with 26 given misoprostol plus oxytocin and 25 given ergometrine plus oxytocin. Low-certainty evidence suggests that ergometrine (RR 0.94, 95% CI 0.48 to 1.84) may make little or no difference to this outcome when compared with oxytocin. The evidence for carbetocin and injectable prostaglandins was uncertain. High-certainty evidence suggests that misoprostol is less protective against PPH \geq 1000 mL when compared with oxytocin (RR 1.19, 95% CI 1.01 to 1.42) (Summary of findings 2).



Figure 9. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for prevention of PPH ≥ 1000 mL.

Comparison	RR (95% CI)
Oxytocin vs Placebo or no treatment	
Direct +	0.61 (0.52, 0.73)
Indirect -	0.56 (0.42, 0.75)
Network -	0.59 (0.50, 0.70)
Ergometrine+Oxytocin vs Placebo or no treatment	
Direct	0.44 (0.18, 1.05)
Indirect -	0.50 (0.38, 0.66)
Network	0.49 (0.38, 0.63)
Carbetocin vs Placebo or no treatment	0.50 (0.07. 0.70)
Network	0.52 (0.37, 0.73)
Misoprostol+Oxytocin vs Placebo or no treatment	
Network -	0.52 (0.39, 0.69)
Misoprostol vs Placebo or no treatment	
Direct -	0.73 (0.56, 0.95)
Indirect	0.68 (0.51, 0.90)
Network -	0.71 (0.59, 0.85)
	(0.00, 0.00)
Ergometrine vs Placebo or no treatment	0.00 (0.04.0.70)
Direct	0.09 (0.01, 0.72)
Indirect	0.70 (0.34, 1.43)
Network	0.56 (0.28, 1.10)
Injectable prostaglandins vs Placebo or no treatment	
Direct	0.36 (0.04, 3.24)
Indirect	0.54 (0.24, 1.22)
Network	0.52 (0.24, 1.13)
Ergometrine+Oxytocin vs Oxytocin	
Direct	0.73 (0.57, 0.93)
L.	
Indirect	1.07 (0.75, 1.54)
Network	0.83 (0.66, 1.03)
Carbetocin vs Oxytocin	
Direct	0.73 (0.45, 1.19)
Indirect ——	0.30 (0.13, 0.72)
Network	0.87 (0.62, 1.21)
Misoprostol+Oxytocin vs Oxytocin	
Direct -	0.87 (0.69, 1.09)
Indirect	1.17 (0.47, 2.86)
_	
Network	0.88 (0.70, 1.11)
Misoprostol vs Oxytocin	
Direct	1.26 (1.11, 1.43)
Indirect +-	1.23 (0.92, 1.64)
Network	1.19 (1.01, 1.42)
Ergometrine vs Oxytocin	
Direct	1.30 (0.52, 3.27)
Indirect	0.61 (0.22, 1.67)
munect	
Network	0.94 (0.48, 1.84)



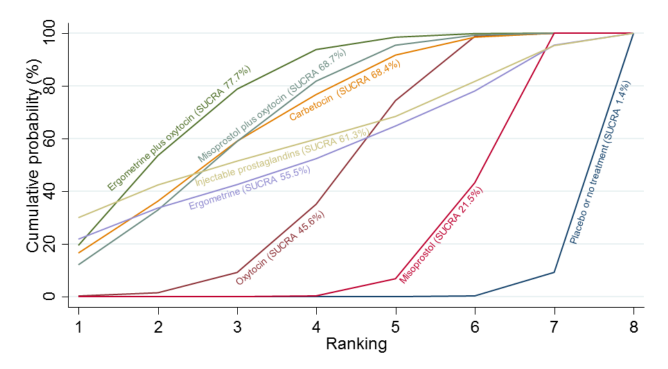
Figure 9. (Continued)



Despite the comparable relative treatment effects between all uterotonics (except misoprostol) and oxytocin, cumulative probabilities for each agent being at each possible rank for PPH ≥ 1000 mL are shown in Figure 10. Ergometrine plus oxytocin (SUCRA 77.7%), misoprostol plus oxytocin (SUCRA 68.7%) combinations

and carbetocin (SUCRA 68.4%) were the highest ranked agents. Oxytocin ranked sixth (45.6%) after injectable prostaglandins (SUCRA 61.3%) and ergometrine (SUCRA 55.5%). Misoprostol was seventh (SUCRA 21.5%) ranking higher than placebo or no treatment (1.4%).

Figure 10. Cumulative rankograms comparing each of the uterotonic agents for prevention of PPH ≥ 1000 mL. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Secondary outcomes

Maternal death

The network diagram for maternal death is presented in Figure 11. Relative effects from the network meta-analysis of 59 trials suggested that no meaningful differences could be detected between all uterotonic agents for maternal deaths as this outcome

was rare (14 deaths across all trials were reported) (Figure 12). When compared with oxytocin, carbetocin (RR 2.00, 95% CI 0.37 to 10.92) and misoprostol (RR 0.62, 95% CI 0.14 to 2.74) probably make little or no difference to maternal death. Network relative effects were not estimable for the comparisons of other uterotonics with oxytocin (Figure 12).



Figure 11. Network Diagram for maternal death. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

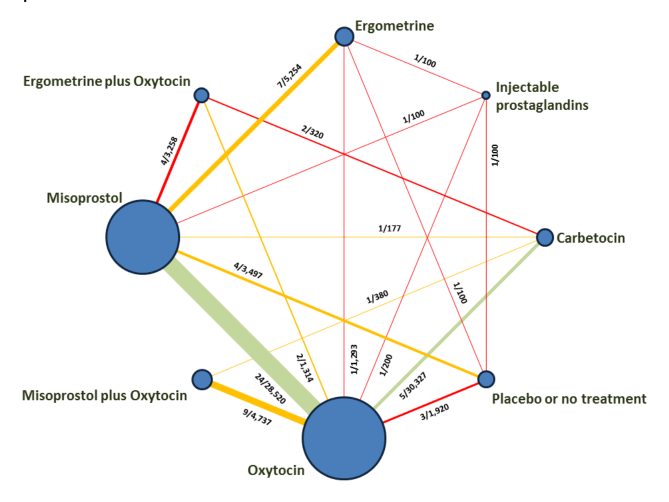




Figure 12. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for prevention of maternal death.

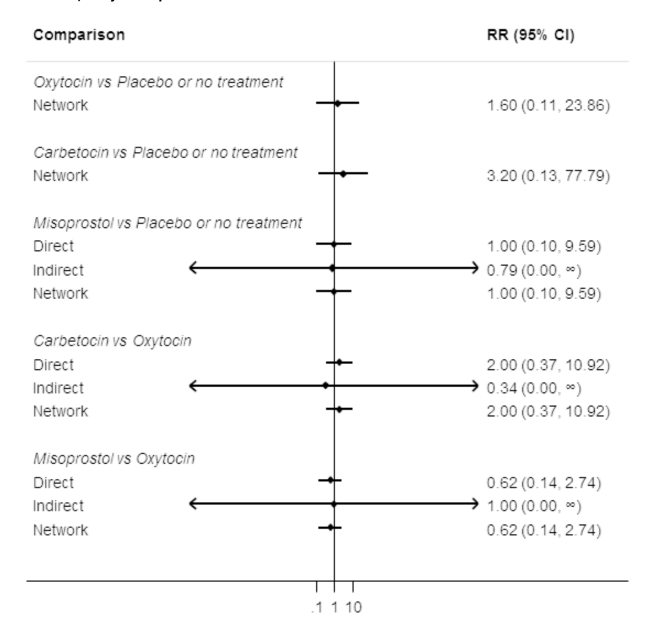
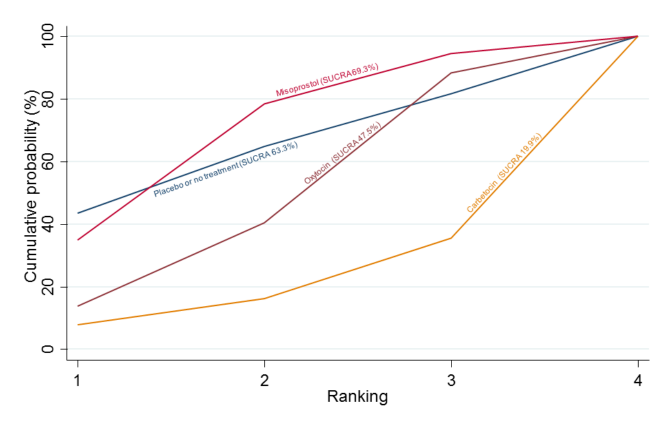


Figure 13 shows the cumulative probabilities for each agent being at each possible rank for maternal death. No reliable ranking could

be derived for this outcome because of the rarity of maternal deaths.



Figure 13. Cumulative rankograms comparing each of the uterotonic agents for prevention of maternal death. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Severe maternal morbidity: intensive care admissions

The network diagram for intensive care admissions as an outcome of severe morbidity is presented in Figure 14. Relative effects from the network meta-analysis of 21 trials for the various

comparisons suggested that there were no detectable differences among uterotonic agents for intensive care admissions as this outcome was rare. This outcome was not reported for any trial involving injectable prostaglandins (Figure 15).



Figure 14. Network Diagram for severe maternal morbidity: intensive care admissions. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

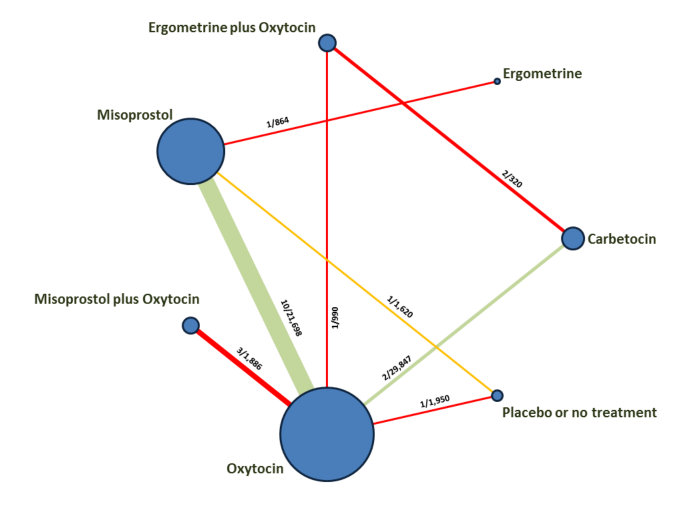




Figure 15. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for prevention of severe maternal morbidity: intensive care admissions.

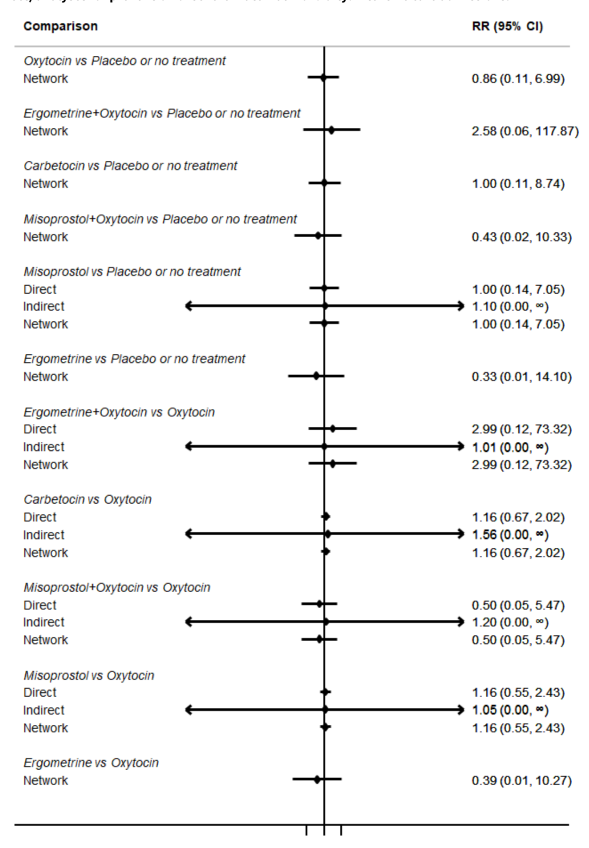




Figure 15. (Continued)

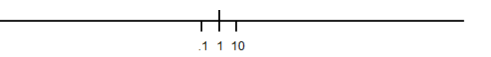
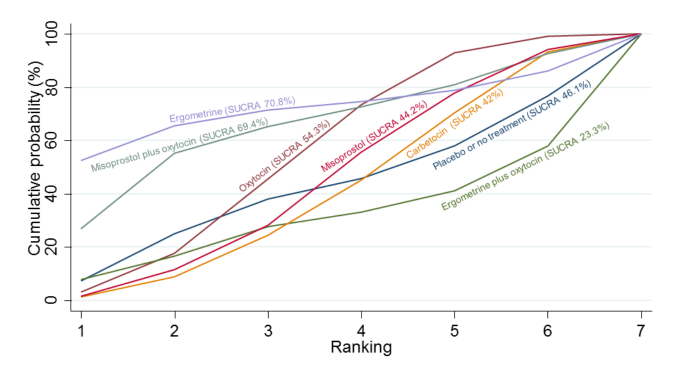


Figure 16 shows the cumulative probabilities for each agent being at each possible rank for intensive care admissions. The ranking for all agents was not clear for this outcome due to limited data.

Figure 16. Cumulative rankograms comparing each of the uterotonic agents for prevention of severe maternal morbidity: intensive care admissions. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Severe maternal morbidity: shock

There were no trials reporting shock as an outcome of severe maternal morbidity.

Additional uterotonics

The network diagram for the use of additional uterotonics is presented in Figure 17. Relative effects from the network metaanalysis of 142 trials suggested that all agents were effective at reducing the use of additional uterotonics when compared with placebo or no treatment (Figure 18). High-certainty evidence suggests that misoprostol plus oxytocin (RR 0.56, 95% CI 0.42 to 0.73) reduces the use of additional uterotonics when compared with oxytocin (Summary of findings 3). Based on these results, about 116 per 1000 women given oxytocin for a vaginal birth would require the administration of additional uterotonic agents, compared with 66 given misoprostol plus oxytocin. There is low-certainty evidence that carbetocin (RR 0.45, 95% CI 0.34 to 0.59), injectable prostaglandins (RR 0.55, 95% CI 0.31 to 0.96) and ergometrine plus oxytocin (RR 0.65, 95% CI 0.50 to 0.85) may also reduce the use of additional uterotonics compared with oxytocin. It is uncertain whether ergometrine reduces use of additional uterotonics because the certainty of this evidence is very low (Summary of findings 3).



Figure 17. Network Diagram for additional uterotonics. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

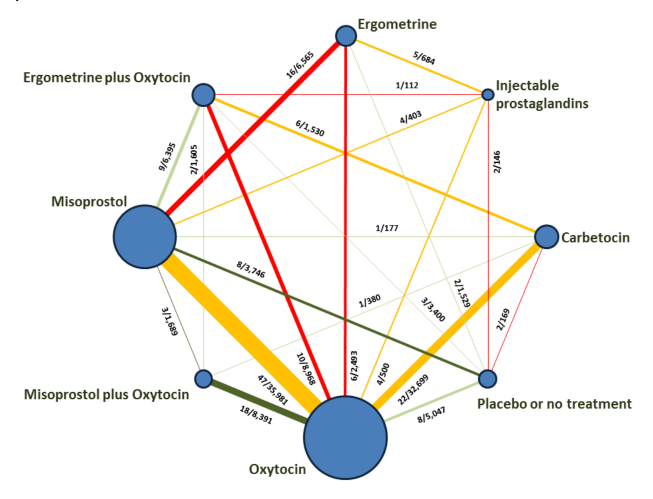




Figure 18. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for additional uterotonics.

Comparison	RR (95% CI)
Oxytocin vs Placebo or no treatment	
Direct	0.43 (0.32, 0.58
Indirect	0.43 (0.29, 0.63
Network	0.42 (0.32, 0.56
Helifoli	0.42 (0.02, 0.00
Ergometrine+Oxytocin vs Placebo or no treatment	0.40/0.45 0.04
Direct	0.19 (0.15, 0.24
Indirect	0.32 (0.22, 0.47
Network	0.28 (0.20, 0.39
Carbetocin vs Placebo or no treatment	
Direct	0.19 (0.12, 0.32
Indirect	0.19 (0.13, 0.29
Network ——	0.19 (0.13, 0.27
Misoprostol+Oxytocin vs Placebo or no treatment	
Network	0.24 (0.16, 0.35
Misoprostol vs Placebo or no treatment	
Direct	0.67 (0.52, 0.87
Indirect	0.36 (0.25, 0.53
Network	0.44 (0.33, 0.59
Ergometrine vs Placebo or no treatment	
Direct	0.18 (0.09, 0.37
Indirect ——•	0.47 (0.30, 0.74
Network	0.41 (0.27, 0.61
Injectable prostaglandins vs Placebo or no treatment	
Direct	0.66 (0.21, 2.09
Indirect	0.19 (0.10, 0.37
Network ——	0.23 (0.13, 0.43
Ergometrine+Oxytocin vs Oxytocin	
Direct	0.79 (0.59, 1.07
Indirect	0.57 (0.40, 0.81
Network	0.65 (0.50, 0.85
Carbetocin vs Oxytocin	
Direct	0.48 (0.34, 0.68
Indirect	0.35 (0.22, 0.57
Network	0.45 (0.34, 0.59
Misoprostol+Oxytocin vs Oxytocin	
Direct	0.54 (0.44, 0.67
Indirect	0.68 (0.31, 1.51
Network	0.56 (0.42, 0.73
Misoprostol vs Oxytocin	
Direct	1.01 (0.85, 1.20
Indirect	1.18 (0.81, 1.73
Network	1.04 (0.88, 1.24
Ergometrine vs Oxytocin	
Direct	1 46 (0 64 2 49
	1.46 (0.61, 3.48
Indirect	0.83 (0.55, 1.26
Network	0.97 (0.69, 1.36
Injectable prostaglandins vs Oxytocin	
Direct Direct	0.29 (0.09.0.94



Figure 18. (Continued)

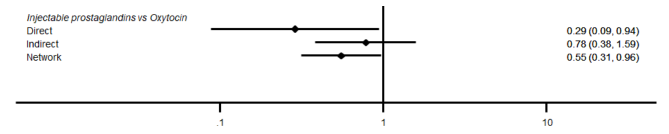
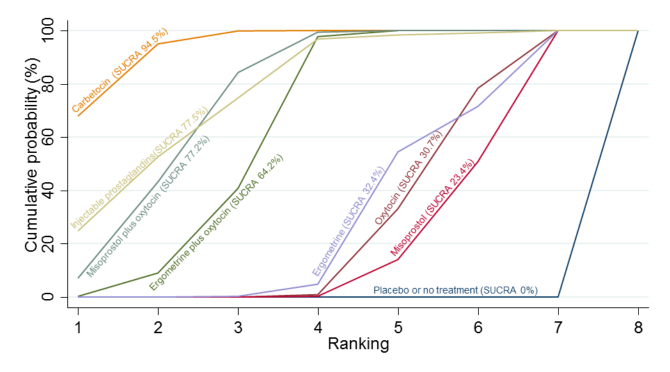


Figure 19 shows the cumulative probabilities for each agent being at each possible rank for the use of additional uterotonics. The highest ranked agents were carbetocin (SUCRA 94.5%), injectable prostaglandins (77.5%), misoprostol plus oxytocin (SUCRA 77.2%)

and ergometrine plus oxytocin (SUCRA 64.2%). Oxytocin was ranked sixth (SUCRA 30.7%) behind ergometrine (SUCRA 32.4%). The lowest ranked agents were misoprostol (SUCRA 23.4%), and placebo or no treatment (SUCRA 0%).

Figure 19. Cumulative rankograms comparing each of the uterotonic agents for additional uterotonics. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Blood transfusion

The network diagram for blood transfusion is presented in Figure 20. Relative effects from the network meta-analysis of 124 trials suggested that all agents except ergometrine and injectable prostaglandins were effective for preventing blood transfusion when compared with placebo or no treatment (Figure 21). Moderate-certainty evidence suggests that misoprostol plus oxytocin probably prevents the need for blood transfusion when compared with oxytocin (RR 0.51, 95% CI 0.37 to 0.70). This suggests that whilst around 15 per 1000 women would require a blood

transfusion when given oxytocin for a vaginal birth, about 8 per 1000 women would need a transfusion with misoprostol plus oxytocin. Moderate-certainty evidence suggests that carbetocin (RR 0.81, 95% CI 0.49 to 1.32) and misoprostol (RR 0.88, 95% CI 0.68 to 1.13) make little or no difference to the need for blood transfusion when compared with oxytocin. Low-certainty evidence suggests that ergometrine (RR 1.11, 95% CI 0.54 to 2.28) and ergometrine plus oxytocin (RR 0.77, 95% CI 0.58 to 1.03) may make little or no difference to this outcome when compared with oxytocin. The evidence for injectable prostaglandins is uncertain (Summary of findings 4).



Figure 20. Network Diagram for blood transfusion. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

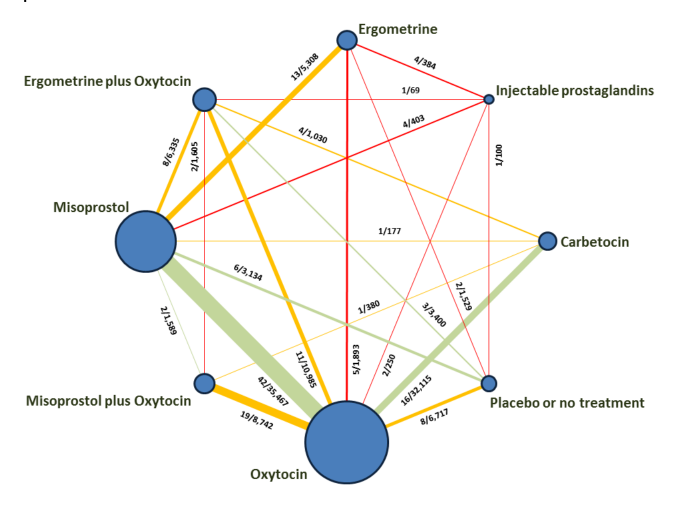




Figure 21. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for blood transfusion.

Comparison	RR (95% CI)
Oxytocin vs Placebo or no treatment	
Direct	0.75 (0.51, 1.12)
Indirect —	0.42 (0.23, 0.75)
Network	0.60 (0.41, 0.87)
Ergometrine+Oxytocin vs Placebo or no treatment	
Direct	0.34 (0.18, 0.66)
Indirect	0.58 (0.35, 0.98)
Network	0.46 (0.31, 0.69)
Carbetocin vs Placebo or no treatment	
Network	0.48 (0.26, 0.89)
Misoprostol+Oxytocin vs Placebo or no treatment	
Network	0.30 (0.19, 0.50)
Misoprostol vs Placebo or no treatment	
Direct	0.46 (0.15, 1.47)
Indirect	0.53 (0.34, 0.84)
Network ——	0.52 (0.35, 0.80)
Ergometrine vs Placebo or no treatment	
Direct	0.34 (0.04, 3.28)
Indirect	0.72 (0.31, 1.67)
Network	0.66 (0.30, 1.45)
Injectable prostaglandins vs Placebo or no treatment	
Network	0.39 (0.14, 1.08)
Ergometrine+Oxytocin vs Oxytocin	
Direct	0.88 (0.53, 1.44)
Indirect	0.64 (0.41, 1.00)
Network -	0.77 (0.58, 1.03)
Carbetocin vs Oxytocin	
Direct	0.68 (0.38, 1.22)
Indirect	0.62 (0.21, 1.85)
Network	0.81 (0.49, 1.32)
Misoprostol+Oxytocin vs Oxytocin	
Direct ←	0.50 (0.37, 0.67)
Indirect	0.77 (0.27, 2.26)
Network ——	0.51 (0.37, 0.70)
Misoprostol vs Oxytocin	
Direct	0.04 /0.05 4.00\



Figure 21. (Continued)

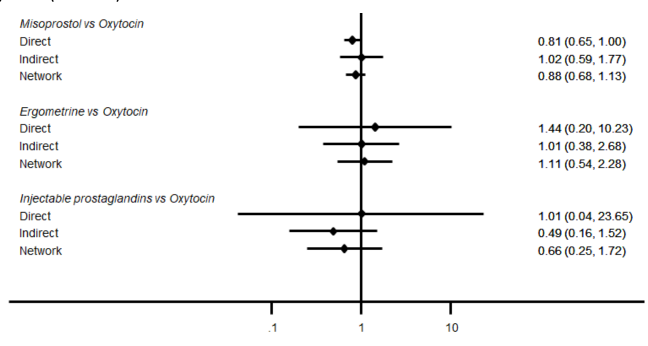
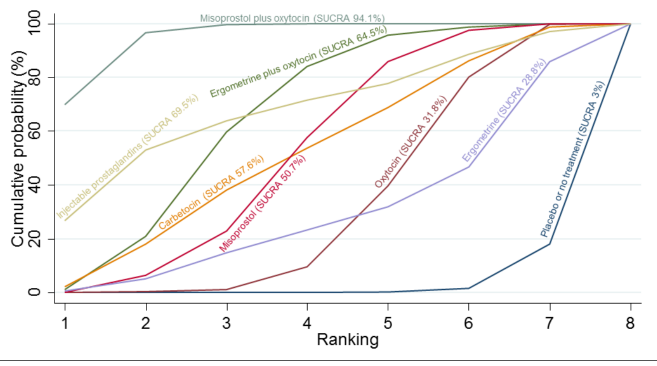


Figure 22 shows the cumulative probabilities for each agent being at each possible rank for preventing blood transfusion. The highest ranked agents were misoprostol plus oxytocin (SUCRA 94.1%), injectable prostaglandins (SUCRA 69.5%) and ergometrine plus

oxytocin (SUCRA 64.5%). Oxytocin was ranked sixth (SUCRA 31.8%) behind carbetocin (SUCRA 57.6%) and misoprostol (SUCRA 50.7%), but higher than ergometrine (SUCRA 28.8%) and placebo or no treatment (SUCRA 3%).

Figure 22. Cumulative rankograms comparing each of the uterotonic agents for blood transfusion. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.





Mean volumes of blood loss

The network diagram for blood loss (mL) as a continuous outcome is presented in Figure 23. Relative effects from the network metaanalysis of 136 trials suggested that all agents are effective for reducing blood loss as a continuous outcome when compared with placebo or no treatment (Figure 24). When compared with oxytocin, moderate-certainty evidence suggests that blood loss is probably on average reduced among women receiving misoprostol plus oxytocin (mean difference (MD) 88.31 mL lower, 95% CI 127.08 mL lower to 49.54 mL lower), and low-certainty evidence suggests that it may be reduced among women receiving carbetocin (MD 81.39 mL lower, 95% CI 119.91mL lower to 42.87 mL lower). Low-certainty evidence suggests that there may be little or no difference between ergometrine (MD 4.82 mL higher, 95% CI 28.00 mL lower to 37.64 mL higher) and oxytocin for this outcome. The effects of misoprostol, injectable prostaglandins and ergometrine plus oxytocin were unclear because the certainty of the evidence was very low (Figure 24).

Figure 23. Network Diagram for mean blood loss (mL). The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

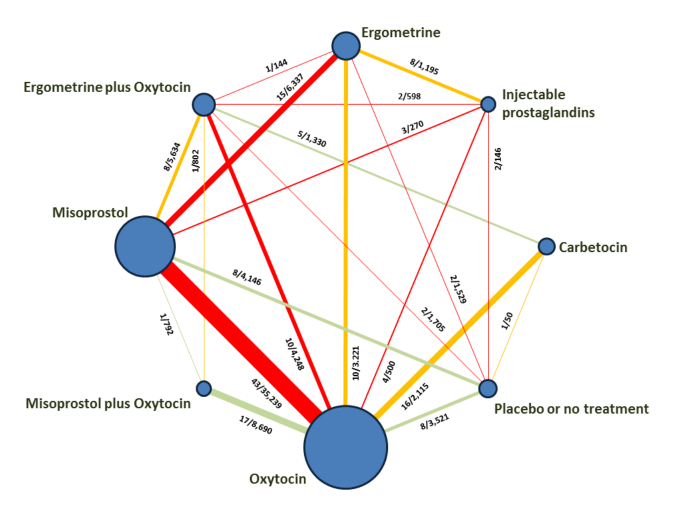




Figure 24. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for mean blood loss (mL).

Comparison	Mean Differences (95% CI)
Oxytocin vs Placebo or no treatment	
Direct •	-118.52 (-141.40, -95.64)
Indirect	-27.19 (-79.51, 25.14)
Network -	-56.98 (-98.15, -15.82)
Ergometrine+Oxytocin vs Placebo or no treatment	
Direct	-35.02 (-101.63, 31.59)
Indirect ——	-93.76 (-147.66, -39.86)
Network ——	-82.24 (-130.59, -33.89)
Carbetocin vs Placebo or no treatment	
Direct	-274.00 (-591.60, 43.60)
Indirect ——	-135.26 (-190.82, -79.71)
Network ——	-138.37 (-193.24, -83.50)
Misoprostol+Oxytocin vs Placebo or no treatment	
Network	-145.29 (-201.55, -89.03)
Misoprostol vs Placebo or no treatment	
Direct	-42.07 (-52.47, -31.68)
Indirect ——	-88.91 (-148.00, -29.81)
Network -	-66.33 (-106.96, -25.69)
Ergometrine vs Placebo or no treatment	
Direct	-50.64 (-119.92, 18.65)
Indirect	-52.53 (-105.09, 0.02)
Network ——	-52.16 (-99.32, -4.99)
Injectable prostaglandins vs Placebo or no treatment	
Direct	-95.17 (-296.09, 105.75)
Indirect	-91.44 (-155.66, -27.22)
Network —	-87.43 (-144.93, -29.93)
Ergometrine+Oxytocin vs Oxytocin	
Direct -	-10.31 (-40.32, 19.70)
Indirect -	-34.53 (-79.23, 10.17)
Network	-25.26 (-59.15, 8.64)
Carbetocin vs Oxytocin	
Direct	-92.73 (-154.97, -30.49)
Indirect	-68.57 (-147.48, 10.33)
Network -	-81.39 (-119.91, -42.87)
Misoprostol+Oxytocin vs Oxytocin	
Direct	-87.26 (-157.83, -16.69)
Indirect	-65.33 (-288.87, 158.20)
Network -	-88.31 (-127.08, -49.54)
Misoprostol vs Oxytocin	
Direct	-8.90 (-23.45, 5.65)
Indirect	-6.35 (-52.97, 40.26)
Network	-9.34 (-31.08, 12.39)
Ergometrine vs Oxytocin	
Direct	8.09 (-17.83, 34.00)
Indirect	3.07 (-39.95, 46.09)
Network	4.82 (-28.00, 37.64)
Injectable prostaglandins vs Oxytocin	
Direct	-15.83 (-152.28, 120.62)
Indirect	_35 N5 /_Q1 18 21 NQ\



Figure 24. (Continued)

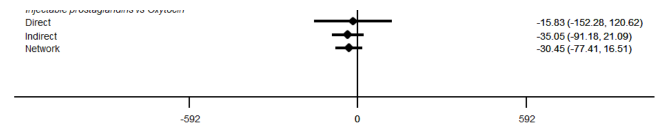
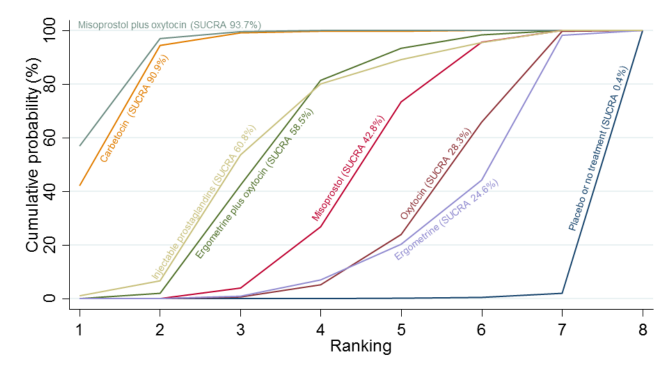


Figure 25 shows the cumulative probabilities for each agent being at each possible rank for preventing blood loss (mL) as a continuous outcome. The highest ranked agents were misoprostol plus oxytocin (SUCRA 93.7%), carbetocin (SUCRA 90.9%), injectable prostaglandins (SUCRA 60.8%) and ergometrine plus oxytocin

(SUCRA 58.5%). Oxytocin was ranked sixth (SUCRA 28.3%) behind misoprostol (SUCRA 42.8%). The lowest ranked agents were ergometrine (SUCRA 24.6%) and placebo or no treatment (SUCRA 0.4%)

Figure 25. Cumulative rankograms comparing each of the uterotonic agents for mean blood loss (mL). Ranking indicates the cumulative probability of being the best, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Change in haemoglobin

The network diagram for the change in haemoglobin measurements before versus after birth (g/L) is presented in Figure 26. Relative effects from the network meta-analysis of 86 trials suggested that all agents except ergometrine and the injectable prostaglandins were effective for reducing the change in haemoglobin measurements when compared with placebo or no treatment (Figure 27). There is low-certainty evidence to suggest that the mean change in haemoglobin level before versus after birth

may be lower among women receiving misoprostol plus oxytocin (MD 2.53 g/L lower, 95% CI 3.80 g/L lower to 1.26 g/L lower) and carbetocin (MD 2.18 g/L lower, 95% CI from 3.57 g/L lower to 0.79 g/L lower) compared with those receiving oxytocin. Low-certainty evidence suggests that there may be little or no difference between ergometrine (MD 0.98 g/L higher, 95% CI from 0.74 g/L lower to 2.69 g/L higher); or ergometrine plus oxytocin (MD 1.07 g/L lower, 95% CI 2.38 g/L lower to 0.25 g/L higher) and oxytocin for this outcome. The effects of misoprostol and injectable prostaglandins were unclear because the certainty of the evidence was very low (Figure 27).



Figure 26. Network Diagram for change in haemoglobin measurements before and after birth (g/L). The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

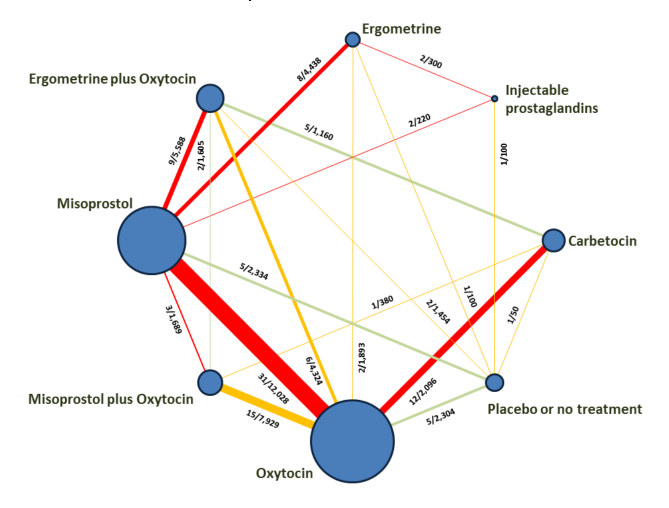




Figure 27. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for change in haemoglobin measurements before and after birth (g/L).

Comparison	Mean Differences (95% C
Oxytocin vs Placebo or no treatment	
Direct	-2.68 (-4.47, -0.89)
Indirect	-1.68 (-3.99, 0.62)
Network	-2.14 (-3.87, -0.41)
THE WORK	2.14 (0.01, 0.41)
Ergometrine+Oxytocin vs Placebo or no treatment	277/272
Direct	-3.57 (-6.50, -0.63)
Indirect	-3.09 (-5.34, -0.84)
Network ———	-3.21 (-5.13, -1.29)
Carbetocin vs Placebo or no treatment	
Direct	-3.40 (-7.23, 0.43)
Indirect — •	-4.47 (-6.71, -2.23)
Network ———	-4.33 (-6.42, -2.23)
Misoprostol+Oxytocin vs Placebo or no treatment	
Network	-4.67 (-6.77, -2.57)
Misoprostol vs Placebo or no treatment	
Direct — •	-2.53 (-3.53, -1.52)
Indirect	-2.53 (-4.94, -0.11)
Network	-2.22 (-3.94, -0.50)
Ergometrine vs Placebo or no treatment	
Direct ◆	-0.50 (-0.58, -0.42)
Indirect	-1.34 (-3.83, 1.15)
Network	-1.17 (-3.37, 1.04)
Injectable prostaglandins vs Placebo or no treatment	
Direct -	0.90 (0.56, 1.24)
Indirect —	-3.10 (-7.00, 0.80)
Network	-1.54 (-4.59, 1.52)
Ergometrine+Oxytocin vs Oxytocin	
Direct	-2.23 (-5.24, 0.77)
Indirect	-0.39 (-2.07, 1.29)
Network	
Network	-1.07 (-2.38, 0.25)
Carbetocin vs Oxytocin	
Direct	-1.66 (-3.81, 0.50)
Indirect	-3.27 (-5.69, -0.84)
Network	-2.18 (-3.57, -0.79)
Misoprostol+Oxytocin vs Oxytocin	
Direct —	-2.59 (-3.70, -1.48)
Indirect •	-2.18 (-5.85, 1.50)
Network ——	-2.53 (-3.80, -1.26)
Misoprostol vs Oxytocin	
Direct	-0.14 (-0.74, 0.47)
Indirect	0.03 (-2.08, 2.14)
Network ——	-0.08 (-0.97, 0.82)
Francontrino us Ovutagia	
Ergometrine vs Oxytocin	0.40/0.00 4.40
Direct	0.42 (-0.30, 1.13)
Indirect	1.20 (-0.78, 3.17)
	0.98 (-0.74, 2.69)
Network	0.90 (-0.74, 2.09)
Injectable prostaglandins vs Oxytocin	0.30 (-0.14, 2.03)



Figure 27. (Continued)

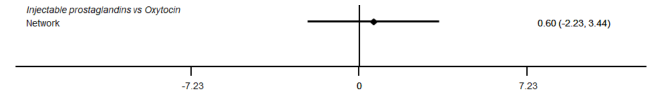
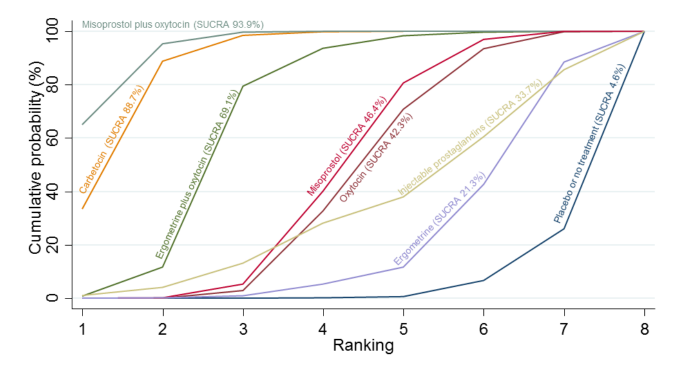


Figure 28 shows the cumulative probabilities for each agent being at each possible rank for change in haemoglobin measurements before versus after birth (g/L). The highest ranked agents were misoprostol plus oxytocin (93.9%), carbetocin (88.7%), and

ergometrine plus oxytocin (69.1%). Oxytocin ranked fifth (SUCRA 42.3%) behind misoprostol (SUCRA 46.4%), but ranked better than injectable prostaglandins (SUCRA 33.7%), ergometrine (SUCRA 21.3%) and placebo or no treatment (SUCRA 4.6%).

Figure 28. Cumulative rankograms comparing each of the uterotonic s for change in haemoglobin measurements before and after birth (g/L). Ranking indicates the cumulative probability of being the best, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Breastfeeding at hospital discharge

The network diagram for breastfeeding at hospital discharge is presented in Figure 29. Relative effects from the network meta-analysis of six trials suggested that there were no detectable differences among oxytocin, carbetocin, ergometrine plus oxytocin for breastfeeding at hospital discharge when compared with placebo or no treatment. HIgh-certainty evidence suggests that ergometrine plus oxytocin (RR 0.99, 95% CI 0.96 to 1.03) makes little or no difference to the proportion of women who are breastfeeding

at the time of discharge from hospital when compared with oxytocin. In absolute terms, these results suggest that about 849 per 1000 women given oxytocin for vaginal birth would be breastfeeding at discharge, compared to 841 per 1000 women with ergometrine plus oxytocin. The findings for carbetocin were unclear, because we found the evidence to be of very low certainty. There were no clear findings relating to any other uterotonics as the outcome was not reported in any of the included trials involving misoprostol, injectable prostaglandins, ergometrine and misoprostol plus oxytocin (Figure 30).



Figure 29. Network Diagram for breastfeeding at discharge. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

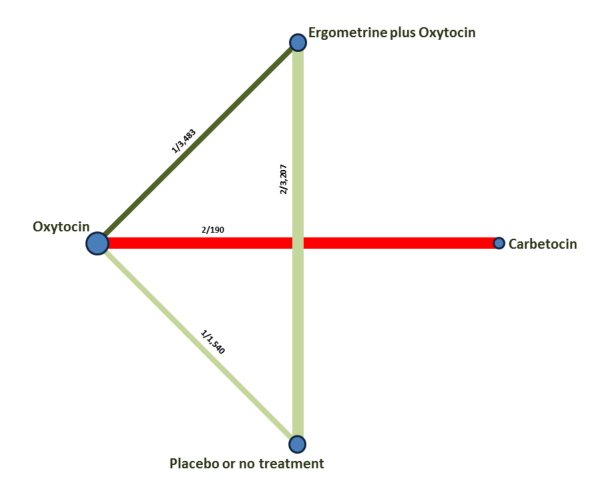




Figure 30. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for breastfeeding at discharge.

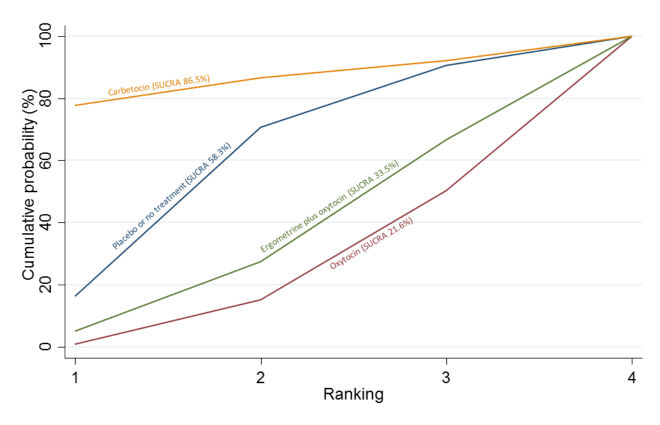
Comparison	RR (95% CI)
Oxytocin vs Placebo or no treatment	
Direct	1.00 (0.95, 1.05)
Indirect	1.04 (0.99, 1.09)
Network	1.02 (0.98, 1.06)
Ergometrine+Oxytocin vs Placebo or no treatment	
Direct	1.03 (0.99, 1.07)
Indirect	0.98 (0.93, 1.04)
Network	1.01 (0.97, 1.05)
Carbetocin vs Placebo or no treatment	
Network	0.96 (0.87, 1.06)
Ergometrine+Oxytocin vs Oxytocin	
Direct	0.99 (0.96, 1.01)
Indirect	1.03 (0.97, 1.10)
Network	0.99 (0.96, 1.03)
Carbetocin vs Oxytocin	
Direct	0.94 (0.86, 1.03)
Indirect <	→ 0.95 (0.00, ∞)
Network	0.94 (0.86, 1.03)
	10
.1 1	10

Figure 31 shows the cumulative probabilities for each agent being at each possible rank for breastfeeding at hospital discharge. The

ranking for all agents was not clear for this outcome due to limited data.



Figure 31. Cumulative rankograms comparing each of the uterotonic agents for breastfeeding at discharge. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Side effects

Nausea

The network diagram for nausea is presented in Figure 32. Relative effects from the network meta-analysis of 100 trials suggest that ergometrine and ergometrine plus oxytocin are worse than placebo or no treatment in causing nausea (Figure 33). When compared with oxytocin, there is high-certainty evidence to suggest that women receiving ergometrine plus oxytocin (RR 2.03, 95% CI 1.47 to 2.80) and misoprostol plus oxytocin (RR 1.88, 95% CI 1.14 to 3.09) are more likely to experience nausea and moderate-certainty evidence that women receiving misoprostol (RR 1.41, 95% CI 1.10 to 1.81), ergometrine (RR 2.40, 95% CI 1.65 to 3.49) or injectable

prostaglandins (RR 2.25, 95% CI 1.16 to 4.39) are more likely to experience nausea than women receiving oxytocin alone (Figure 33). Based on these results, about 86 per 1000 women given oxytocin for a vaginal birth would experience nausea, compared with 175 given ergometrine plus oxytocin, 162 given misoprostol plus oxytocin, 121 given misoprostol, 206 given ergometrine, and 193 given injectable prostaglandins. Low-certainty evidence suggests that carbetocin may make little or no difference to experience of nausea among women when compared with oxytocin (RR 1.00, 95% CI 0.71 to 1.41). With carbetocin, the anticipated absolute effect is the same as oxytocin, with 86 per 1000 women experiencing nausea.



Figure 32. Network Diagram for nausea. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

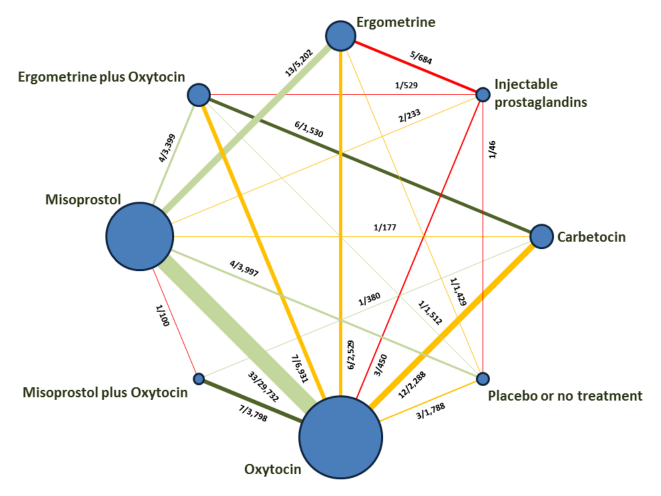




Figure 33. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for nausea.

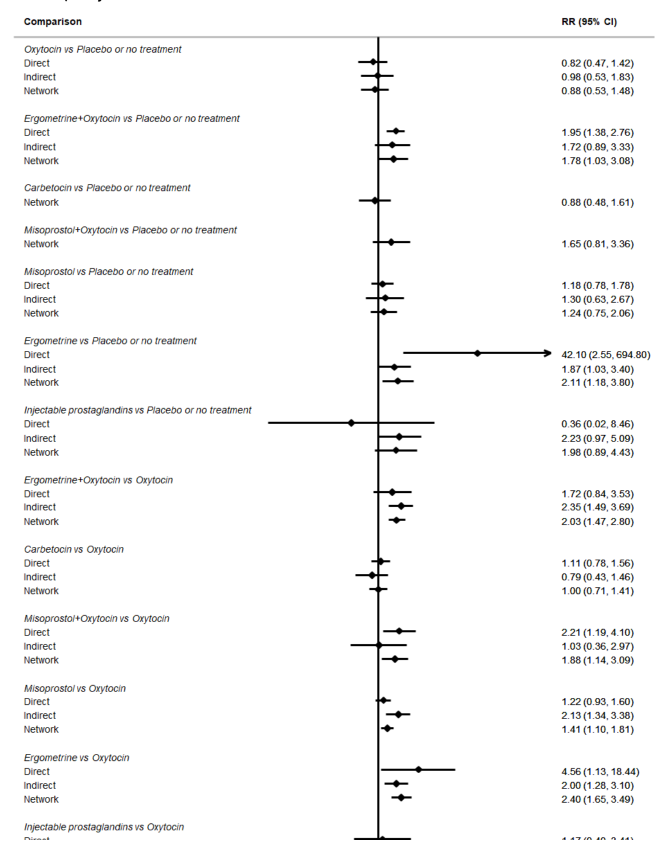




Figure 33. (Continued)

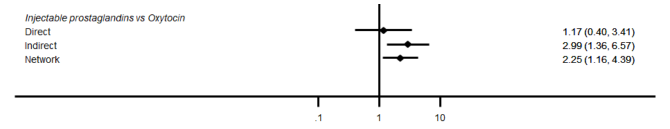
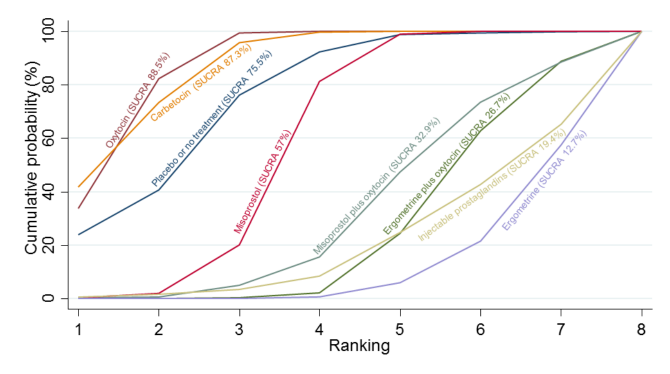


Figure 34 shows the cumulative probabilities for each agent being at each possible rank for causing nausea. The highest ranked agents with which women are less likely to experience nausea are oxytocin (SUCRA 88.5%), carbetocin (SUCRA 87.3%) and placebo or no treatment (SUCRA 75.5%). These are followed by misoprostol

(SUCRA 57%) and misoprostol plus oxytocin (SUCRA 32.9%). The lowest ranked agents are ergometrine plus oxytocin (SUCRA 26.7%), injectable prostaglandins (SUCRA 19.4%) and ergometrine (SUCRA 12.7%).

Figure 34. Cumulative rankograms comparing each of the uterotonic agents for nausea. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Vomiting

The network diagram for vomiting is presented in Figure 35. Relative effects from the network meta-analysis of 110 trials suggested that ergometrine, injectable prostaglandins, misoprostol plus oxytocin and ergometrine plus oxytocin are worse than placebo or no treatment in causing vomiting (Figure 36). When compared with oxytocin, there is evidence that all agents besides carbetocin increase the incidence of vomiting. High-certainty evidence suggests misoprostol plus oxytocin combination (RR 2.11, 95% CI 1.39 to 3.18) increases the likelihood of vomiting, while moderate-certainty evidence suggests that ergometrine plus

oxytocin (RR 2.93, 95% CI 2.08 to 4.13), misoprostol (RR 1.63, 95% CI 1.25 to 2.14), and ergometrine (RR 2.36, 95% CI 1.56 to 3.55) probably increase the likelihood of vomiting. These results suggest that 13 per 1000 women given oxytocin experience vomiting, compared to 12 per 1000 with carbetocin, 27 with misoprostol plus oxytocin, 38 with ergometrine plus oxytocin, 21 with misoprostol, and 31 with ergometrine. Low-certainty evidence also suggests that injectable prostaglandins (RR 3.76, 95% CI 1.90 to 7.42) may increase women's experience of vomiting. Moderate-certainty evidence suggests that carbetocin probably makes little or no difference to women's experience of vomiting compared with oxytocin (RR 0.93, 95% CI 0.64 to 1.35) (Summary of findings 5).



Figure 35. Network Diagram for vomiting. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

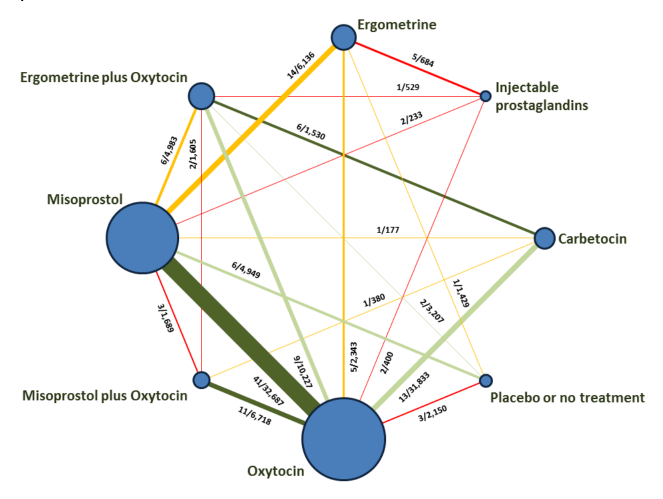




Figure 36. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for vomiting.

Comparison		RR (95% CI)
Oxytocin vs Placebo or no treatment		
Direct		1.40 (0.44, 4.41)
Indirect	+	0.93 (0.53, 1.64)
Network	+	0.98 (0.58, 1.66)
Ergometrine+Oxytocin vs Placebo or no treatment		
Direct		2.15 (1.46, 3.18)
Indirect	—	3.66 (1.80, 7.44)
Network	🛨	2.88 (1.73, 4.78)
Carbetocin vs Placebo or no treatment		
Network	†	0.91 (0.49, 1.68)
Misoprostol+Oxytocin vs Placebo or no treatment		
Network	—	2.07 (1.08, 3.97)
Misoprostol vs Placebo or no treatment		
Direct	 • -	1.41 (0.92, 2.16)
Indirect	 •-	1.48 (0.73, 3.03)
Network	→	1.61 (0.98, 2.62)
Ergometrine vs Placebo or no treatment		
Direct	─	2 5.67 (1.52, 432.78)
Indirect		2.10 (1.15, 3.81)
Network	 	2.31 (1.28, 4.18)
Injectable prostaglandins vs Placebo or no treatment		
Network	—	3.69 (1.65, 8.26)
Ergometrine+Oxytocin vs Oxytocin		
Direct	 → -	3.05 (1.76, 5.29)
Indirect	→	2.77 (1.75, 4.38)
Network	+	2.93 (2.08, 4.13)
Carbetocin vs Oxytocin		
Direct		0.90 (0.53, 1.50)
Indirect	+	1.00 (0.51, 1.95)
Network	+	0.93 (0.64, 1.35)
Misoprostol+Oxytocin vs Oxytocin		
Direct	→	2.24 (1.52, 3.31)
Indirect		1.48 (0.52, 4.27)
Network	-	2.11 (1.39, 3.18)
Misoprostol vs Oxytocin		
Di	I.a.	1 5 1 / 1 1 1 1 1 1 1



Figure 36. (Continued)

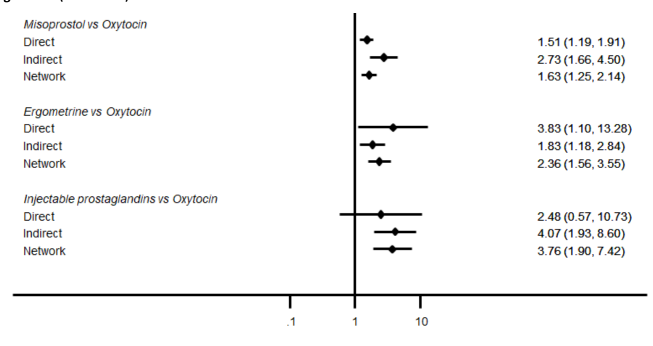
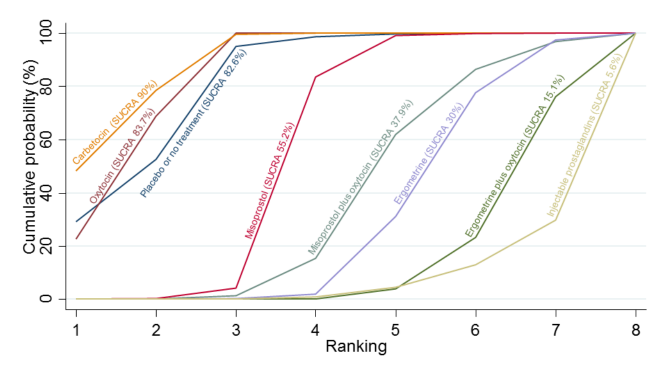


Figure 37 shows the cumulative probabilities for each agent being at each possible rank for causing vomiting. The highest ranked agents were carbetocin (SUCRA 90%), oxytocin (SUCRA 83.7%) and placebo or no treatment (SUCRA 82.6%). These are followed by

misoprostol (SUCRA 55.2%) and misoprostol plus oxytocin (SUCRA 37.9%). The lowest ranked agents were ergometrine (SUCRA 30%), ergometrine plus oxytocin (SUCRA 15.1%) and injectable prostaglandins (SUCRA 5.6%).

Figure 37. Cumulative rankograms comparing each of the uterotonic agents for vomiting. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.





Hypertension

The network diagram for hypertension is presented in Figure 38. Relative effects from the network meta-analysis of 20 trials suggest that ergometrine is worse than placebo or no treatment in causing hypertension (Figure 39). Low-certainty evidence suggests that ergometrine (RR 8.54, 95% CI 2.12 to 34.48) may increase the risk of hypertension when compared with oxytocin, whereas misoprostol

(RR 1.50, 95% 0.49 to 4.61) and ergometrine plus oxytocin (RR 2.48, 95% CI 0.89 to 6.88) may make little or no difference to this outcome. The baseline risk of hypertension for women receiving oxytocin is 76 per 1000 women. Taking into account the very wide 95% CIs, this suggests that the range of possible true effects varies substantially for each agent. It is uncertain whether carbetocin or injectable prostaglandins increase hypertension because the certainty of evidence was very low (Summary of findings 6).

Figure 38. Network Diagram for hypertension. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

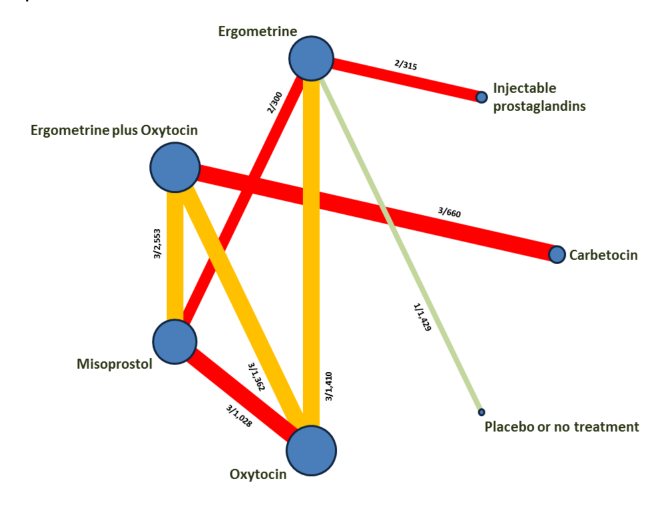




Figure 39. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for hypertension.

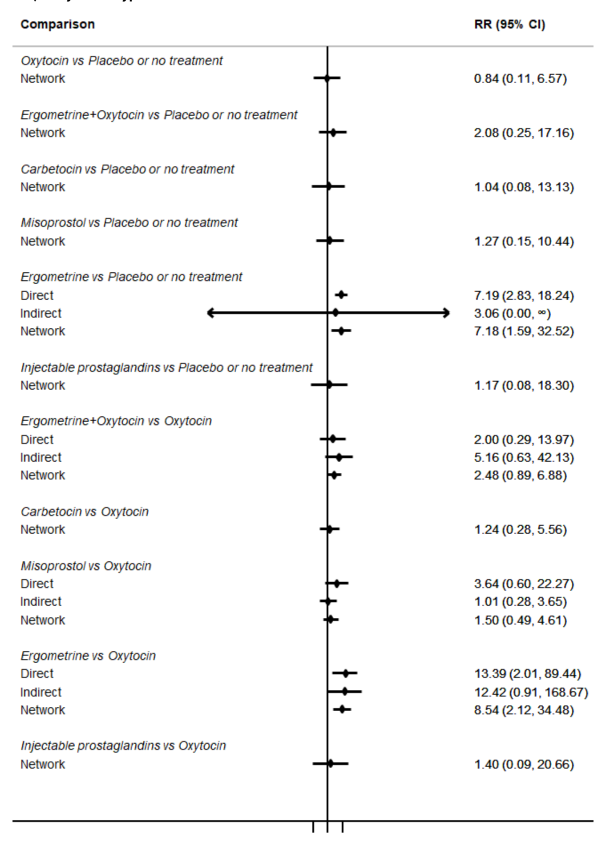




Figure 39. (Continued)

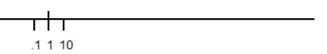
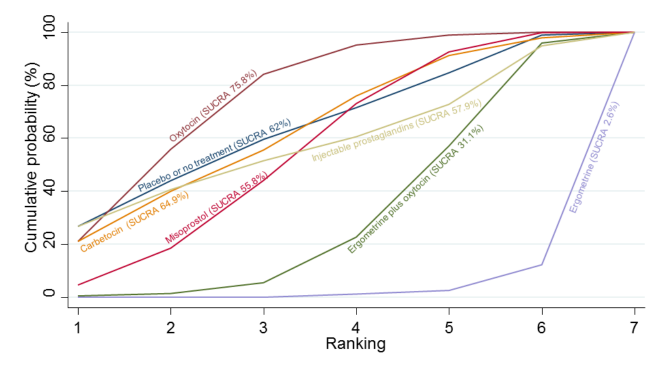


Figure 40 shows the cumulative probabilities for each agent being at each possible rank for causing hypertension. The lowest ranked agents were ergometrine (SUCRA 2.6%) and ergometrine plus

oxytocin (31.1%). The rest of the agents were of comparable ranking. There were no trials involving misoprostol plus oxytocin so this agent could not be ranked.

Figure 40. Cumulative rankograms comparing each of the uterotonic agents for hypertension. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Headache

The network diagram for headache is presented in Figure 41. Relative effects from the network meta-analysis of 57 trials suggested that ergometrine is worse than placebo or no treatment in causing headache (Figure 42). When compared with oxytocin, there is low-certainty evidence to suggest that women receiving ergometrine (RR 1.89, 95% CI 1.02 to 3.50) may be more likely

to experience headache (Figure 42), with 167 per 1000 women given oxytocin experiencing headache compared to 316 with ergometrine. Low-certainty evidence also suggests that carbetocin, misoprostol, and misoprostol plus oxytocin may make little or no difference to experience of headache when compared with oxytocin. It is uncertain whether injectable prostaglandins impact on women's experience of headache because the certainty of evidence was very low.



Figure 41. Network Diagram for headache. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

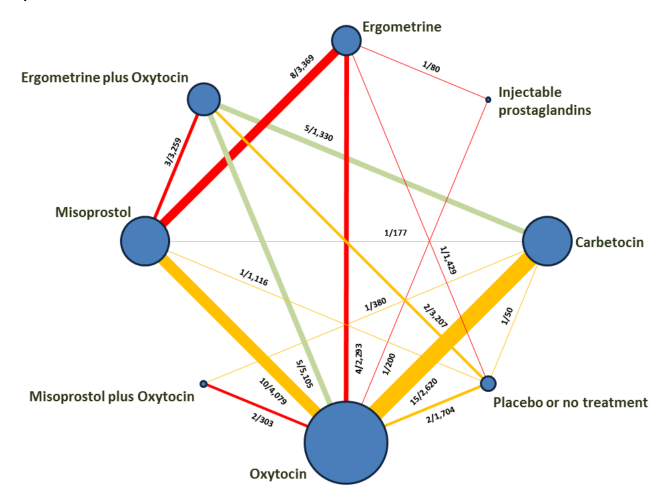




Figure 42. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for headache.

Comparison		RR (95% CI)
Oxytocin vs Placebo or no treatment		
Direct		1.56 (0.52, 4.74)
Indirect	- •	1.40 (0.59, 3.31)
Network	 • -	1.45 (0.74, 2.81)
Ergometrine+Oxytocin vs Placebo or no treatment		
Direct	 • -	1.65 (0.78, 3.48)
Indirect	+•	1.51 (0.60, 3.82)
Network	 • -	1.57 (0.80, 3.06)
Carbetocin vs Placebo or no treatment		
Direct	- •	5.00 (0.25, 99.16)
Indirect	- •	1.32 (0.62, 2.80)
Network	 	1.35 (0.65, 2.82)
Misoprostol+Oxytocin vs Placebo or no treatment		
Network		2.14 (0.52, 8.77)
Misoprostol vs Placebo or no treatment		
Direct		0.94 (0.32, 2.77)
Indirect	+•	1.62 (0.72, 3.66)
Network	+•	1.41 (0.71, 2.81)
Ergometrine vs Placebo or no treatment		
Direct		7.19 (0.37, 138.91)
Indirect		2.53 (1.04, 6.16)
Network	——	2.73 (1.15, 6.50)
Injectable prostaglandins vs Placebo or no treatment		
Network		2.55 (0.43, 14.99)
Ergometrine+Oxytocin vs Oxytocin		
Direct	+•-	1.26 (0.79, 1.99)
Indirect		0.87 (0.48, 1.58)
Network	+	1.08 (0.73, 1.61)
Carbetocin vs Oxytocin		
Direct	- ♦+	0.84 (0.63, 1.12)
Indirect	+•-	1.46 (0.66, 3.25)
Network	+	0.94 (0.66, 1.33)
Misoprostol+Oxytocin vs Oxytocin		
Direct		1.26 (0.26, 6.23)
Indirect	- •	1.90 (0.27, 13.36)
Network		1.48 (0.42, 5.18)
Misoprostol vs Oxytocin		
Direct	- • -	0.88 (0.54, 1.42)
Indirect		1.19 (0.60, 2.33)
Network	+	0.98 (0.68, 1.40)
Ergometrine vs Oxytocin	_	
Direct	<u> </u>	5.63 (0.93, 33.96)
Indirect		1.34 (0.65, 2.76)
Network		1.89 (1.02, 3.50)



Figure 42. (Continued)

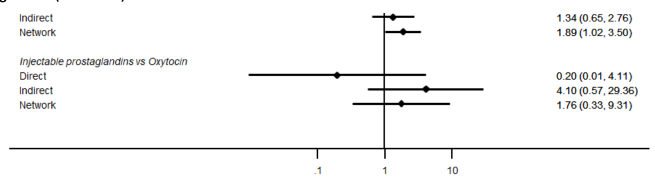
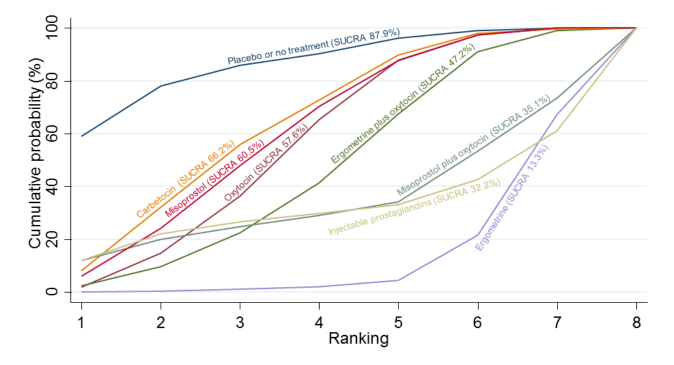


Figure 43 shows the cumulative probabilities for each agent being at each possible rank for causing headache. The highest ranked intervention was placebo or no treatment (SUCRA 87.9%), carbetocin (SUCRA 66.2%), misoprostol (SUCRA 60.5%)

and oxytocin (SUCRA 57.6%). The lowest ranked agents were ergometrine plus oxytocin (SUCRA 47.2%), misoprostol plus oxytocin (SUCRA 35.1%), injectable prostaglandins (SUCRA 32.2%) and ergometrine (SUCRA 13.3%).

Figure 43. Cumulative rankograms comparing each of the uterotonic agents for headache. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Fever

The network diagram for fever is presented in Figure 44. Relative effects from the network meta-analysis of 83 trials suggested that misoprostol and misoprostol plus oxytocin are worse than placebo or no treatment in causing fever (Figure 45). Moderate-certainty evidence suggests that misoprostol (RR 3.87, 95% CI 2.90 to 5.16) and misoprostol plus oxytocin (RR 3.14, 95% CI 2.20 to 4.49) probably increase the occurrence of fever when compared with oxytocin. Moderate-certainty evidence suggests that carbetocin (RR

1.07, 95% CI 0.43 to 2.69) probably makes little or no difference to women's experience of fever. These results suggests that 24 per 1000 women given oxytocin would experience fever, compared to 93 with misoprostol, 75 with misoprostol plus oxytocin and 26 with carbetocin. Low-certainty evidence suggests that injectable prostaglandins (RR 1.12, 95% CI 0.33 to 3.86) and ergometrine plus oxytocin (RR 0.70, 95% CI 0.35 to 1.42) may make little or no difference to this outcome, when compared with oxytocin. Evidence regarding the comparative effect of ergometrine on this



outcome is uncertain because the certainty of the evidence was very low (Summary of findings 7).

Figure 44. Network Diagram for fever. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

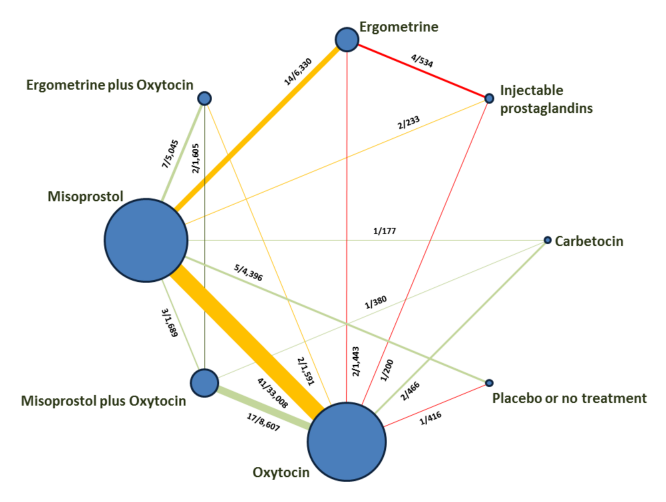




Figure 45. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for fever.

Comparison		RR (95% CI)
Oxytocin vs Placebo or no treatment Network	+	1.06 (0.51, 2.21)
Ergometrine+Oxytocin vs Placebo or no treatment Network	-	0.74 (0.28, 1.95)
Carbetocin vs Placebo or no treatment Network	-	1.14 (0.36, 3.59)
Misoprostol+Oxytocin vs Placebo or no treatment Network		3.33 (1.49, 7.41)
Misoprostol vs Placebo or no treatment Direct Indirect Network		4.09 (2.01, 8.32) 0.49 (0.00, 202.19) 4.10 (2.09, 8.05)
Ergometrine vs Placebo or no treatment Network		0.82 (0.35, 1.89)
Injectable prostaglandins vs Placebo or no treatment Network	-	1.19 (0.30, 4.77)
Ergometrine+Oxytocin vs Oxytocin Direct Indirect Network		1.08 (0.48, 2.43) 0.54 (0.22, 1.32) 0.70 (0.35, 1.42)
Carbetocin vs Oxytocin Direct Indirect Network		1.58 (0.27, 9.35) 0.77 (0.18, 3.42) 1.07 (0.43, 2.69)
Misoprostol+Oxytocin vs Oxytocin Direct Indirect Network	-	2.99 (2.00, 4.45) 5.43 (1.48, 19.95) 3.14 (2.20, 4.49)
Misoprostol vs Oxytocin Direct Indirect Network	<u>+</u>	3.75 (2.73, 5.15) 6.49 (2.24, 18.76) 3.87 (2.90, 5.16)
Ergometrine vs Oxytocin Direct Indirect Network Injectable prostaglandins vs Oxytocin	•	2.97 (0.97, 9.05) 0.63 (0.35, 1.16) 0.77 (0.44, 1.35)
injectable prostaglandins vs Oxytocin	I	



Figure 45. (Continued)

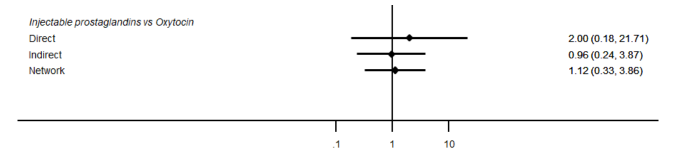
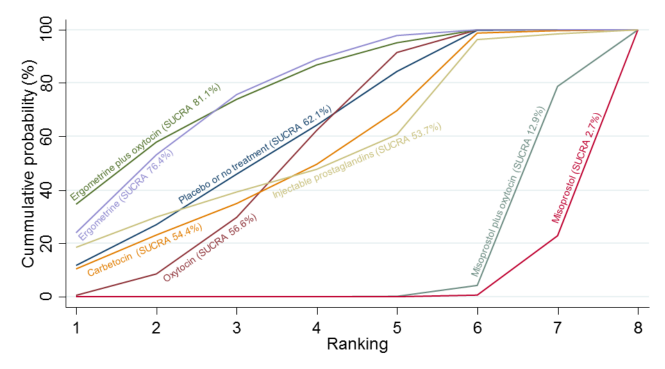


Figure 46 shows the cumulative probabilities for each agent being at each possible rank for causing fever. The lowest ranked agents were misoprostol (SUCRA 2.7%) and misoprostol plus oxytocin

(SUCRA 12.9%). The rest of the agents were similar in ranking to the placebo or no treatment group.

Figure 46. Cumulative rankograms comparing each of the uterotonic agents for fever. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Shivering

The network diagram for shivering is presented in Figure 47. Relative effects from the network meta-analysis of 109 trials suggested that misoprostol and misoprostol plus oxytocin are worse than placebo or no treatment in causing shivering (Figure 48). When compared with oxytocin, there is moderate-certainty evidence to suggest that women receiving misoprostol plus oxytocin (RR 3.62, 95% CI 2.59 to 5.05) are probably more likely to experience shivering (Figure 48). In absolute terms, whereas

89 per 1000 women given oxytocin would experience shivering with oxytocin, 322 would experience shivering with misoprostol plus oxytocin. Low-certainty evidence suggests that misoprostol (RR 4.18, 95% CI 3.34 to 5.23) may also increase the experience of shivering. Moderate-certainty evidence suggests that ergometrine plus oxytocin (RR 1.38, 95% CI 0.86 to 2.22) probably makes little or no difference to shivering when compared with oxytocin. Likewise, low-certainty evidence suggests that carbetocin and injectable prostaglandins may make little or no difference to this outcome when compared with oxytocin (Figure 48).



Figure 47. Network Diagram for shivering. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

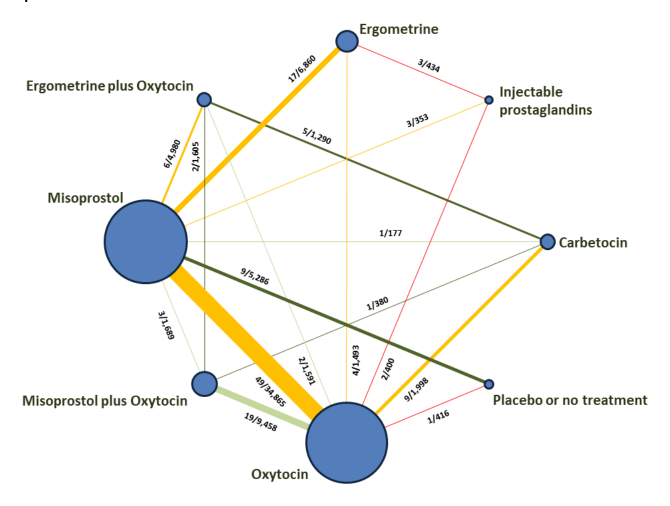




Figure 48. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for shivering.

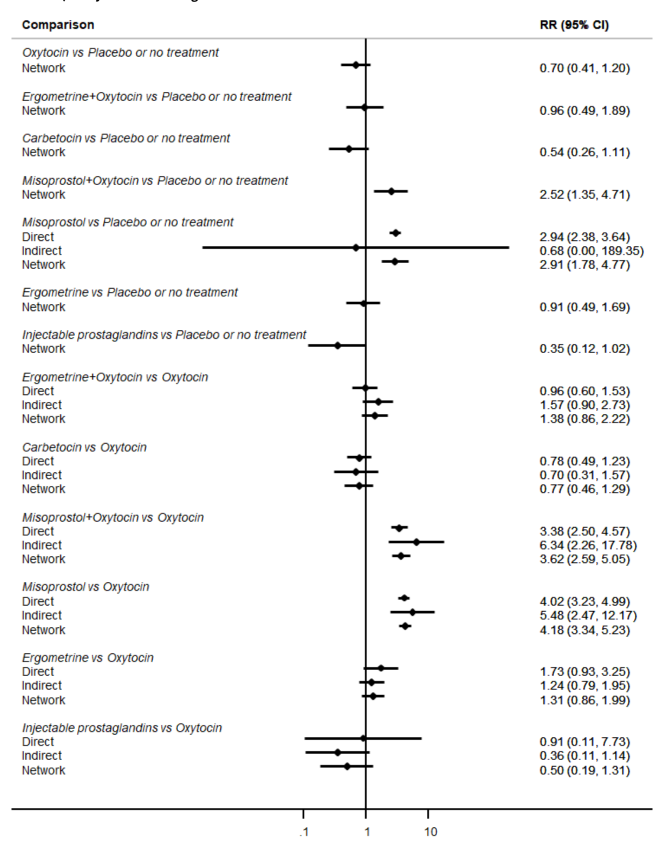




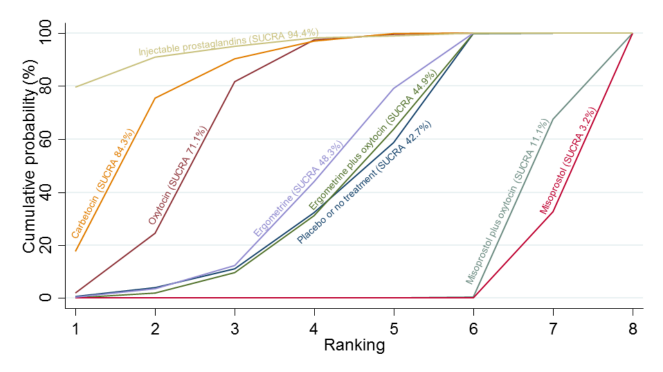
Figure 48. (Continued)

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Figure 49 shows the cumulative probabilities for each agent being at each possible rank for causing shivering. The highest ranked agents are injectable prostaglandins (SUCRA 94.4%), carbetocin (SUCRA 84.3%) and oxytocin (SUCRA 71.1%). These are followed

by ergometrine (SUCRA 48.3%), ergometrine plus oxytocin and placebo or no treatment (SUCRA 42.7%). The lowest ranked agents were misoprostol plus oxytocin (SUCRA 11.1%) and misoprostol (SUCRA 3.2%).

Figure 49. Cumulative rankograms comparing each of the uterotonic agents for shivering. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Abdominal pain

The network diagram for abdominal pain is presented in Figure 50. Relative effects from the network meta-analysis of 32 trials suggested that ergometrine is worse than placebo or no treatment in causing abdominal pain (Figure 51). High-certainty evidence suggests that misoprostol and misoprostol plus oxytocin make

little or no difference to women's experience of abdominal pain, when compared with oxytocin. Low-certainty evidence suggests that ergometrine plus oxytocin probably make little or no difference to women's experience of abdominal pain compared with oxytocin. The effects of injectable prostaglandins and ergometrine were uncertain as the certainty of evidence was very low (Figure 51).



Figure 50. Network Diagram for abdominal pain. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

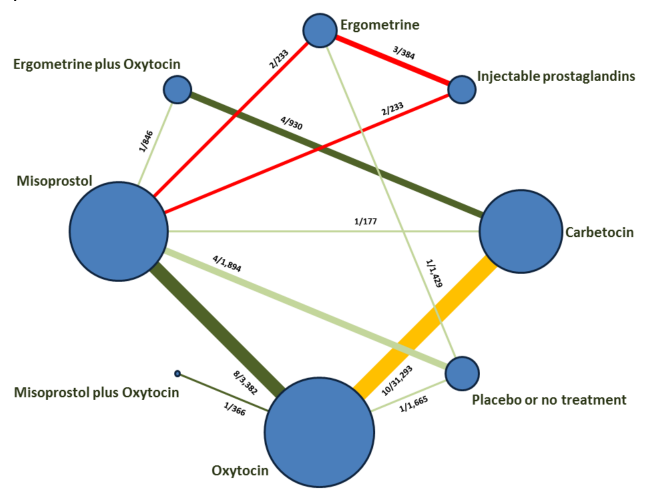




Figure 51. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for abdominal pain.

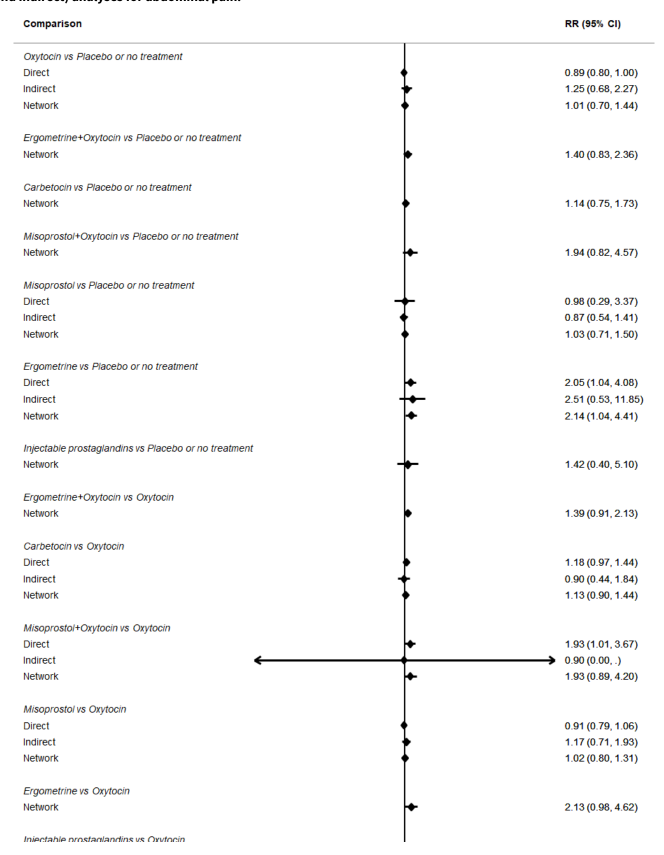




Figure 51. (Continued)

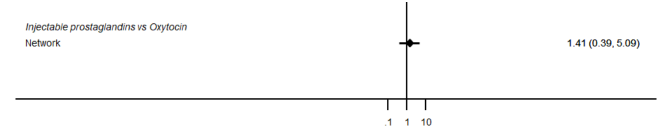
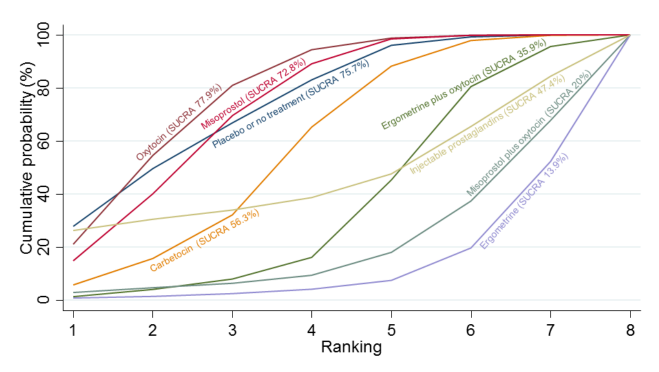


Figure 52 shows the cumulative probabilities for each agent being at each possible rank for causing abdominal pain. The highest ranked agent was oxytocin (SUCRA 77.9%), misoprostol (SUCRA 72.8%) and placebo or no treatment (SUCRA 75.7%). These were

followed by carbetocin (SUCRA 56.3%), injectable prostaglandins (SUCRA 47.4%) and ergometrine plus oxytocin (SUCRA 35.9%). The lowest ranked agents are misoprostol plus oxytocin (SUCRA 20%) and ergometrine (SUCRA 13.9%).

Figure 52. Cumulative rankograms comparing each of the uterotonic agents for abdominal pain. Ranking indicates the cumulative probability of being the best agent, the second best, the third best, etc. The x axis shows the relative ranking and the y-axis the cumulative probability of each ranking. We estimate the SUrface underneath this Cumulative RAnking line (SUCRA); the larger the SUCRA the higher its rank among all available agents.



Diarrhoea

The network diagram for diarrhoea is presented in Figure 53. Relative effects from the network meta-analysis of 55 trials suggested that misoprostol, ergometrine and injectable prostaglandins are worse than placebo or no treatment in causing diarrhoea (Figure 54). High-certainty evidence shows that misoprostol (RR 2.24, 95% CI 1.64 to 3.05) and misoprostol plus oxytocin (RR 1.82, 95% CI 1.12 to 2.98) increase the likelihood of diarrhoea when compared with oxytocin (Figure 54), Moderate-

certainty evidence suggests that ergometrine plus oxytocin (RR 1.80, 95% CI 1.18 to 2.75) and injectable prostaglandins (RR 23.41, 95% CI 11.03 to 49.70) probably increase the likelihood of diarrhoea, when compared with oxytocin (Figure 54). These results suggest that 11 women per 1000 given oxytocin for vaginal birth would experience diarrhoea, compared to 25 per 1000 women with misoprostol, 23 with misoprostol plus oxytocin, 20 with ergometrine plus oxytocin and 254 with injectable prostaglandins. There is also low-certainty evidence that ergometrine (RR 2.51, 95% CI 1.20 to 5.26) may increase diarrhoea (Figure 54).



Figure 53. Network Diagram for diarrhoea. The nodes represent an intervention and their size is proportional to the number of trials comparing this intervention to any other in the network. The lines connecting each pair of interventions represent a direct comparison and are drawn proportional to the number of trials making each direct comparison. Numbers on the lines represent the number of trials and participants for each comparison. The colour of the line is green for high-certainty evidence; light green for moderate-certainty evidence; orange for low-certainty evidence and red for very low-certainty evidence. Multi-arm trials contribute to more than one comparison.

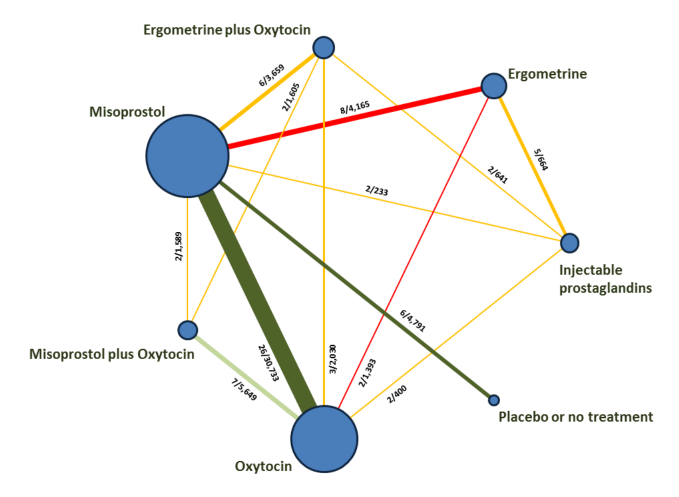




Figure 54. Forest plot with relative risk ratios and 95% CIs from pairwise, indirect and network (combining direct and indirect) analyses for diarrhoea.

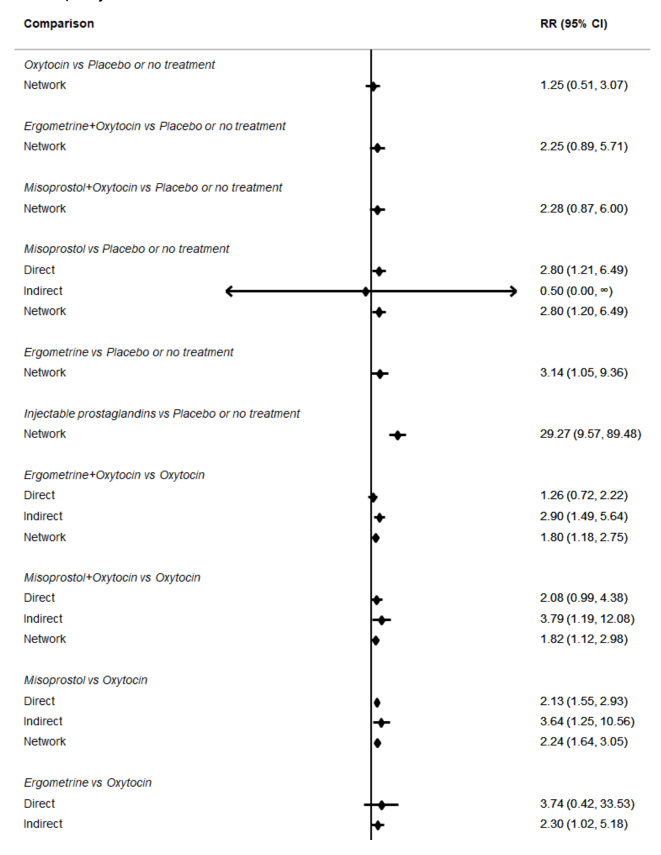




Figure 54. (Continued)

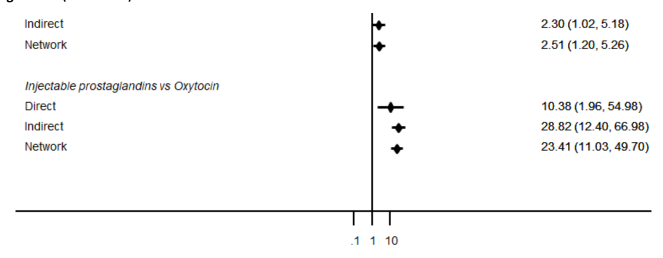


Figure 5 shows the cumulative probabilities for each agent being at each possible rank for causing diarrhoea. The highest ranked agents were placebo or no treatment (SUCRA 92.8%) and oxytocin (SUCRA 88.4%). These were followed by ergometrine plus oxytocin (SUCRA 54.1%), misoprostol plus oxytocin (SUCRA 51.8%). The lowest ranked agents are misoprostol (SUCRA 32.5%), ergometrine (SUCRA 30.3%) and injectable prostaglandins (SUCRA 0%).

Maternal sense of well-being

In total there were two trials reporting outcomes relevant to maternal sense of well-being. However, because of the heterogeneous ways maternal sense of well-being was defined in these trials a decision was made not to perform a meta-analysis.

This outcome was reported in two different ways by one trial comparing oxytocin with no treatment (Jans 2017). Low-certainty evidence suggests that the use of prophylactic oxytocin may make little or no difference to women's experience of less energy than before birth at three months postpartum (RR 1.02, 95% CI 0.93 to 1.13), or to experience of fatigue at three months postpartum (RR 0.99, 95% CI 0.95 to 1.04), Analysis 1.19.

This outcome was reported in eight different ways by another trial comparing ergometrine plus oxytocin with no treatment (Rogers 1998), Analysis 6.19. Low-certainty evidence suggests that prophylactic ergometrine plus oxytocin may make little or no difference to women's general health at six weeks postpartum when compared to no treatment (RR 0.99, 95% CI 0.71 to 1.37). Moderate-certainty evidence suggests that prophylactic ergometrine plus oxytocin probably makes little or no difference to women's exhaustion since birth when compared to no treatment (RR 0.95, 95% CI 0.79 to 1.15). Low-certainty evidence suggests that prophylactic ergometrine plus oxytocin may make little or no difference to women's exhaustion at six weeks postpartum when compared to no treatment (RR 0.95, 95% CI 0.74 to 1.21). Moderatecertainty evidence suggests that prophylactic ergometrine plus oxytocin probably makes little or no difference to women's blues at six weeks postpartum when compared to no treatment (RR 0.93, 95% CI 0.83 to 1.04). Low-certainty evidence suggests that prophylactic ergometrine plus oxytocin may make little or no difference to women experiencing depression at six weeks postpartum when compared to no treatment (RR 1.22, 95% CI 0.84 to 1.78). Low-certainty evidence suggests that prophylactic ergometrine plus oxytocin may make little or no difference to women looking for help for depression at six weeks postpartum when compared to no treatment (RR 1.05, 95% CI from 0.82 to 1.35). It is uncertain whether prophylactic ergometrine plus oxytocin reduces admissions to hospital for depression at six weeks postpartum when compared to no treatment because the certainty of this evidence is very low (RR 3.06, 95% CI 0.12 to 75.06). Moderate-certainty evidence suggests that prophylactic ergometrine plus oxytocin probably makes little or no difference to women reporting health problems at six weeks postpartum when compared to no treatment (RR 0.95, 95% CI 0.90 to 1.01).

Maternal satisfaction

In total, there were five trials reporting outcomes relevant to maternal satisfaction. However, because of the heterogeneous ways maternal satisfaction was defined in these trials a decision was made not to perform a meta-analysis.

This outcome was reported in three different ways by one trial comparing oxytocin with no treatment (Jangsten 2011). Moderate-certainty evidence suggests that the use of prophylactic oxytocin may make little or no difference to women's perception of whether management of the birth positively influenced the childbirth experience for mothers (RR 1.01, 95% CI 0.89 to 1.15); or made little or no difference to the mother's childbirth experience (RR 1.02, 95% CI 0.90 to 1.15). Low-certainty evidence suggests that the use of prophylactic oxytocin may make little or no difference to the extent to which women perceive that management of the birth negatively influenced the childbirth experience for mothers (RR 0.73, 95% CI 0.47 to 1.13), Analysis 1.20.

This outcome was reported in two different ways by one trial comparing ergometrine plus oxytocin with no treatment (Rogers 1998). Moderate-certainty evidence suggests that prophylactic ergometrine plus oxytocin probably makes little or no difference to satisfaction with third-stage management when compared to no treatment (RR 1.03, 95% CI from 1.00 to 1.05). Moderate-certainty evidence suggests that prophylactic ergometrine plus oxytocin probably decreased women feeling in control during third stage



when compared to no treatment (RR 0.95, 95% CI 0.91 to 0.99), Analysis 6.20.

This outcome was reported in four different ways by one trial comparing misoprostol with oxytocin (Diop 2016). Moderate-certainty evidence suggests that prophylactic misoprostol, when compared to oxytocin, probably makes little or no difference to women being satisfied or very satisfied with the uterotonic agent they received (RR 1.01, 95% CI 1.00 to 1.02), or that women would make a complaint about, or have problems with, the uterotonic agent they received (RR 0.36, 95% CI 0.20 to 0.64), or that women would take the specific uterotonic agent again after subsequent deliveries (RR 1.01, 95% CI 1.00 to 1.02), or that women would recommend the specific uterotonic agent to a friend (RR 1.01, 95% CI 1.00 to 1.02), Analysis 8.20.

This outcome was reported by one trial comparing ergometrine plus oxytocin with misoprostol using an eight-item Client Satisfaction Questionnaire (Ng 2007). Moderate-certainty evidence suggests that prophylactic ergometrine plus oxytocin, when compared to misoprostol, probably makes little or no difference to women being satisfied with the uterotonic agent they received (MD 0.6 lower, 95% CI 1.22 lower to 0.02 higher).

Subgroup analyses

We carried out subgroup analyses for PPH ≥ 500 mL and PPH ≥ 1000 mL by mode of birth (caesarean versus vaginal birth), setting (hospital versus community), risk of PPH (high versus low risk for PPH), dose of misoprostol (≥ 600 mcg versus < 600 mcg), and regimen of oxytocin (bolus versus bolus plus infusion versus infusion only). The network diagrams for all subgroups are available from Appendix 2. Relative effects and cumulative probabilities for each agent being at each possible rank from the network meta-analysis for each subgroup are also available from Appendix 2. Subgroup analyses did not reveal important subgroup differences for any of the subgroups.

Sensitivity analyses

We carried out pre-specified sensitivity analyses by restricting our analyses to studies at low risk of bias, studies at low risk of bias in terms of funding sources, to studies that used an objective method of measuring blood loss and large trials with more than 400 participants. Details of these analyses are available from Appendix 2. Sensitivity analyses were also performed according to the choice of relative effect measure (risk ratio (RR) versus odds ratio (OR)) and the statistical model (fixed-effect versus random-effects model). Further sensitivity analyses identified during the review process were performed by removing trials published earlier than 1990, cluster trials, removing trials with a high level of missing data and removing trials where participants were also randomised to coagents such as uterine massage or early controlled cord traction, or both. The sensitivity analyses show that the overall results are not affected by the abovementioned criteria or decisions.

DISCUSSION

Summary of main results

This network meta-analysis of 196 randomised trials (135,559 women) shows that all uterotonic agents are effective in preventing postpartum haemorrhage (PPH) ≥ 500 mL when compared with placebo or no treatment. The three highest ranked uterotonic

agents were ergometrine plus oxytocin combination, carbetocin and misoprostol plus oxytocin combination. Ergometrine plus oxytocin combination and carbetocin are probably more effective uterotonic agents for preventing PPH ≥ 500 mL than oxytocin. Misoprostol plus oxytocin may also be more effective but the certainty of the evidence is low. Misoprostol, injectable prostaglandins, and ergometrine have comparable relative effects to oxytocin for preventing PPH ≥ 500 mL but again the certainty of the evidence is low.

This network meta-analysis shows all agents except ergometrine and injectable prostaglandins were effective for preventing PPH ≥ 1000 mL when compared with placebo or no treatment. Misoprostol plus oxytocin and ergometrine plus oxytocin combinations make little or no difference to PPH ≥ 1000 mL when compared with oxytocin. Ergometrine also may make little or no difference to this outcome when compared with oxytocin but the evidence was of low certainty. The evidence for carbetocin and injectable prostaglandins was of very low quality. Misoprostol is less effective against PPH ≥ 1000 mL when compared with oxytocin. Despite the comparable relative treatment effects between all uterotonics (except misoprostol) and oxytocin, ergometrine plus oxytocin, misoprostol plus oxytocin combinations and carbetocin were the highest ranked agents for PPH ≥ 1000 mL.

Misoprostol plus oxytocin reduces the use of additional uterotonics and probably also reduces the risk of blood transfusion when compared with oxytocin. Carbetocin, injectable prostaglandins and ergometrine plus oxytocin may also reduce the use of additional uterotonics but the certainty of the evidence is low. No meaningful differences could be detected between all agents for maternal deaths or severe morbidity as these outcomes were so rare in the included randomised trials where they were reported.

The two combination regimens were associated with important side effects. When compared with oxytocin, misoprostol plus oxytocin combination increases the likelihood of vomiting and fever. No included studies reported on hypertension for misoprostol plus oxytocin versus oxytocin. Ergometrine plus oxytocin probably increases the likelihood of vomiting and may make little or no difference to the risk of hypertension, however absolute effects varied considerably and the certainty of the evidence was low for this outcome.

Subgroup analyses did not reveal important subgroup differences by mode of birth (caesarean versus vaginal birth), setting (hospital versus community), risk of PPH (high versus low risk for PPH), dose of misoprostol (≥ 600 mcg versus < 600 mcg) and regimen of oxytocin (bolus versus bolus plus infusion versus infusion only).

Overall completeness and applicability of evidence

This network meta-analysis provides the relative effectiveness of all agents used for the prevention of PPH in a coherent and methodologically robust way across important clinical outcomes by combining both direct and indirect evidence, thus increasing the statistical power and confidence in the results. We found that most of the included trials reported our primary outcomes and most of the secondary outcomes. This increased the power across most of our analyses and contributed to the consistency in the ranking across all blood loss outcomes. We were thorough in our evaluation of the important potential treatment effect modifiers (mode of birth, prior risk of PPH, healthcare setting, dose, route and



regimen of the agents). We did not encounter important differences in the distribution of the effect modifiers between the different comparisons. In addition, the ranking of the agents in each of the subgroups was comparable with the overall ranking. The results of the network meta-analyses were mostly consistent and where there was significant inconsistency this was likely due to unstable estimates from single studies.

Many trials excluded women with significant comorbidities and at very high risk for PPH. Women recruited to the included studies were predominantly delivered at more than 37 weeks of gestation. Most of the trials were carried out in hospital settings and with women having a vaginal birth. For women having a vaginal birth, uterotonic agent administration used to be a component of the active management of the third stage of labour, alongside controlled cord traction and early cord clamping. The most upto-date guidelines from the WHO (WHO 2012), place emphasis on the administration of a uterotonic agent as the main aspect within this package for prevention of PPH. These guidelines state that early cord clamping is generally not advised, whilst controlled cord traction is optional where skilled birth attendants are present (WHO 2012). Rankings of the available agents were similar in subgroups where trials included either only women having a vaginal birth or only women undergoing a caesarean section. Evidently, uterine tone plays a major role in PPH at caesarean section, with a relative reduction of PPH \geq 500 mL similar to the reduction seen in women undergoing vaginal births when more effective agents are used. The ranking is relevant to women at either high or low risk for PPH in hospital settings. There were not enough trials to be able to recommend a ranking in community settings, even though a similar ranking in terms of effectiveness can be expected.

The dosages, regimens and routes of administration for the most effective uterotonic agents varied. In most of the studies investigating this agent, carbetocin was administered as a single intravenous bolus of 100 mcg or intramuscularly. The combination of ergometrine plus oxytocin was usually administered intramuscularly combining 500 mcg of ergometrine plus 5 IU (international units) of oxytocin. Misoprostol plus oxytocin combinations varied greatly, with some studies administering an intravenous infusion of 20 IU of oxytocin and 400 mcg of misoprostol sublingually, or 200 mcg of misoprostol sublingually, others administering an intravenous bolus of oxytocin of 10 IU plus 400 mcg misoprostol sublingually, while others administered an intravenous infusion of 10 IU of oxytocin and 400 mcg of misoprostol rectally. There were also several other ways of administering the misoprostol plus oxytocin combination described (see Characteristics of included studies).

Quality of the evidence

We recognise that there is no single established approach for assessing the certainty of the effect estimates generated by the network meta-analysis. We applied the rigorous method for appraising quality of network evidence as proposed by the GRADE Working group. Overall, the evidence presented varied widely in quality, and our confidence in the effect estimates ranged from very low to high certainty. When we compared oxytocin with all the other uterotonic agents and agent combinations, most individual outcomes included a range in quality of evidence across the different interventions, and this was equally true for our most important outcomes. Our reasons for downgrading the evidence also varied across comparisons and outcomes.

Summarising the quality of the evidence for the seven most important outcomes (also described in the summary of findings), for PPH ≥ 500 mL, moderate-certainty evidence pointed to the probable superiority of both carbetocin and ergometrine plus oxytocin when compared with oxytocin alone; in both cases this evidence was downgraded due to some concerns regarding risk of bias and unexplained statistical heterogeneity in both the direct and indirect comparisons contributing most weight to the network estimate, however the direct and indirect effect estimates were coherent with one another.

For PPH ≥ 1000 mL, we had high-certainty evidence that there is little difference between oxytocin and the uterotonic combinations of ergometrine plus oxytocin and misoprostol plus oxytocin, whereas misoprostol is slightly worse. For carbetocin and injectable prostaglandins, very low-certainty evidence suggested unclear effects due to imprecision, strong suspicion of publication bias for the direct evidence, and incoherence between the direct and indirect effect estimates.

We had high-certainty evidence that misoprostol plus oxytocin reduces the need for additional uterotonic agents compared to oxytocin alone, and low-certainty evidence suggested that carbetocin, injectable prostaglandins and ergometrine plus oxytocin may all also reduce the use of additional uterotonics. The low-certainty findings were all downgraded due to risk of bias and unexplained statistical heterogeneity, with findings on injectable prostaglandins and ergometrine plus oxytocin also being downgraded for imprecision. Moderate-certainty evidence suggested misoprostol plus oxytocin may reduce the need for blood transfusion compared to oxytocin, with the evidence being downgraded due to imprecision.

The quality of the evidence on side effects was also somewhat varied, although with less variation within the evidence for each individual outcome. For vomiting, most of the findings were high or moderate certainty, with the main reasons for downgrading being concerns about risk of bias and imprecision. For hypertension, the evidence was all low or very low quality due to wideranging concerns about risk of bias, imprecision and unexplained statistical heterogeneity. The evidence on fever ranged from moderate to very low certainty, and we had most confidence in the finding that misoprostol alone or in combination with oxytocin led to a considerable increase in this outcome for women. We downgraded these findings due to concerns about risk of bias, severe unexplained statistical heterogeneity and imprecision.

Potential biases in the review process

Several authors have been involved in one or more previous or ongoing trials related to the use of uterotonics for the prevention of PPH that could be eligible for inclusion in this review. They did not participate in any decisions regarding these trials (i.e. assessment for inclusion/exclusion, trial quality, data extraction) for the purposes of this review – these tasks were carried out by other members of the team who were not directly involved in the trials.

The quality of the evidence was assessed by a team of authors based in different countries. Before we could GRADE the network meta-analysis evidence, we had to determine the methodology for this process because there is no well-established approach or accompanying tools such as software. All GRADE assessments



were undertaken by one individual (MJW, VD, MC or JP) and then re-assessed independently by another of those four authors, in consultation with OTO and JPV where additional decision-making was required.

The earliest included trial was conducted in 1976 (Moodie 1976), and in the decades since then, the clinical care and the clinical response to PPH may have improved. These temporal changes could have contributed to heterogeneity and increased the uncertainty of findings. However, we carried out a sensitivity analysis by removing trials published before 1990 and this did not vary the ranking of the agents. As objective methods of measuring blood loss became increasingly available this could perhaps have also led to apparent changes in reported blood loss. The trials included in the review recruited women with varied clinical characteristics, and it is important to consider this when interpreting results. The inclusion criteria were not always reported in detail and, when they were, these varied across trials. Further heterogeneity may also be present in the overall analysis related to the dose, route or regimen of the uterotonic agents. Even though we did not observe subgroup effects when we examined the dose of misoprostol or regimen of oxytocin administration, we were not able to perform subgroup analyses for every single increment in dosage or route of administration. Lastly, not all trials reported data on side effects, hence these analyses were often underpowered.

Agreements and disagreements with other studies or reviews

In this update of the review first published in April 2018, we have incorporated results from a large WHO trial (Widmer 2018) and overall, 56 new trials involving 46,612 women. The conclusions remain largely the same. The results for the primary outcome of PPH ≥ 500 mL were similar to the previously published review (Gallos 2018), although the quality of the evidence for carbetocin has changed from 'very low-' to 'moderate-certainty evidence' for this outcome, due to the addition of data from three studies including approximately 30,000 women. For the primary outcome of PPH ≥ 1000 mL, none of the agents is significantly more effective when compared with the reference uterotonic agent oxytocin. In the previous version of the review, high-quality evidence suggested that ergometrine plus oxytocin was more effective in reducing PPH ≥ 1000 mL in comparison to oxytocin. For all other outcomes (blood transfusion; additional uterotonics; and side effects), the results are largely the same.

Our results agree with existing Cochrane Reviews (Begley 2015; Liabsuetrakul 2018; McDonald 2004; Su 2012; Tuncalp 2012; Westhoff 2013) that focus on the comparison of a uterotonic agent versus another (direct comparisons). However, this network meta-analysis has several more studies than included in the previous reviews because of its nature of comparing all available uterotonic agents in one single analysis and because it is the most up-to-date including recently published trials. Hence, some estimates differ slightly, as expected.

AUTHORS' CONCLUSIONS

Implications for practice

Collaboration.

The current WHO recommendation on the choice of uterotonics for preventing postpartum haemorrhage (PPH) is 10 IU of intramuscular or intravenous oxytocin (WHO 2012). We found that

oxytocin has substantial desirable effects compared with placebo or no treatment and trivial side effects. As a result, the balance of effects is expected to favour oxytocin. A problem with oxytocin, though, is that it needs to be kept refrigerated (2 °C to 8 °C) to maintain its potency. Several studies have demonstrated that oxytocin loses potency if stored at room temperature for too long or at higher temperatures, making its use difficult in low-resource settings (Hogerzeil 1993; WHO 1993).

We found that ergometrine plus oxytocin combination (Syntometrine ®), misoprostol plus oxytocin combination and carbetocin have additional desirable effects compared with oxytocin, whereas misoprostol, injectable prostaglandins and ergometrine have no additional benefits compared with oxytocin. However, these uterotonic agents with the exception of carbetocin also have substantial undesirable effects as they increase the likelihood of side effects compared with oxytocin.

While the combination of ergometrine plus oxytocin may be more effective than oxytocin alone for some desirable outcomes, this combination also increases important side effects for the woman. Notably, caution should be exercised when using ergot derivatives for PPH prevention as these drugs have clear contraindications in women with underlying hypertensive or cardiovascular disorders. Thus, it is probably safer to avoid the use of ergot derivatives containing uterotonics in unscreened populations.

It is important to note that although the combination of misoprostol plus oxytocin may be more effective than oxytocin alone for some desirable outcomes, this combination also increases important side effects for the woman. In addition, as misoprostol and oxytocin are not available as a fixed drug combination (like Syntometrine®), and the two agents have to be administered through separate routes (parenteral and oral/rectal), the application of this combination may be less feasible in routine clinical settings compared with using either oxytocin or misoprostol as a single uterotonic agent. Therefore, the care provider and the parturient woman may need to carefully balance the additional benefits of a combination of misoprostol and oxytocin (over either of these agents alone) with the drawbacks (including side effects, and the challenges and inconvenience) of using two drugs through separate routes before using this combination.

There is evidence that carbetocin may be more effective than oxytocin for some desirable outcomes but with a comparable side-effect profile when compared with oxytocin. While this risk-benefit balance appears to favour carbetocin, carbetocin is more expensive and currently not widely available. A room temperature stable formulation of carbetocin is also now available, which could make it an attractive option for settings where maintaining the cold chain for storage and transport of oxytocin is problematic, if the cost limitations can be addressed. Nonetheless, despite the unit cost of carbetocin being higher than oxytocin it may still be cost-effective in high-income settings such as the UK where the cost of caring for PPH and its complication is substantial (Gallos 2018).

Before making decisions, policymakers would need to balance the desirable and undesirable effects of the range of effective uterotonics presented with their available resources and other contextual issues. An economic assessment would need to assess the consequences of various single or combination uterotonic agents compared with their current standard, with consideration of differences between their effects (benefits and harms), supply



costs, and other resource requirements (staffing and training, equipment and infrastructure, staff time, supplies, supervision and monitoring). Other important considerations for decision-making include the potential impact of introducing or scaling up the uterotonic on health equity, acceptability to key stakeholders and feasibility of using these uterotonics in routine clinical practice.

Implications for research

There is still uncertainty around the best doses and routes of administration for each of the uterotonic agents. For oxytocin for example, there are uncertainties around the optimal dose at caesarean section, whether it should be administered intravenously or intramuscularly and whether it should be administered as an intravenous bolus or infusion. The current network meta-analysis analyses the effectiveness and side effects for the various agents grouping together all doses and routes of the agents analysing them at an aggregate level only. The current network meta-analysis cannot answer if a specific dose or route for any of the agents is preferred as it excludes trials that have compared different doses or routes of the same agents. We propose to update our existing network meta-analysis by adding evidence from all the trials comparing the various doses and routes for all available agents. We wish to analyse each agent by disaggregating the various doses and routes available and then analyse in the context of the network. In this way we plan to make use of both existing direct evidence and indirect evidence from the whole network. This approach potentially can give us answers about preferred doses and routes for each of the agents and identify research gaps in the evidence base.

Consultation with our consumer group demonstrated the need for more research into outcomes identified as priorities for women and their families, such as women's views regarding the agents used, severe maternal morbidity such as shock, and breastfeeding at discharge. To date, trials have rarely investigated these outcomes. Consumers also considered the side effects of uterotonic agents to be important but these were often not reported.

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REFERENCES

References to studies included in this review

Abdel-Aleem 1993 (published data only)

Abdel-Aleem H, Abol-Oyoun EM, Moustafa SA, Kamel HS, Abdel-Wahab HA. Carboprost trometamol in the management of the third stage of labor. *International Journal of Gynecology & Obstetrics* 1993;**42**:247-50.

Abdel-Aleem 2010 (published data only)

Abdel-Aleem H, Singata M, Abdel-Aleem M, Mshweshwe N, Williams X, Hofmeyr GJ. Uterine massage to reduce postpartum hemorrhage after vaginal delivery. *International Journal of Gynecology & Obstetrics* 2010;**111**(1):32-6.

Acharya 2001 {published data only}

Acharya G, Al-Sammarai MT, Patel N, Al-Habib A, Kiserud T. A randomized, controlled trial comparing effect of oral misoprostol and intravenous syntocinon on intra-operative blood loss during cesarean section. *Acta Obstetricia et Gynecologica Scandinavica* 2001;**80**(3):245-50.

Adanikin 2012 (published data only)

Adanikin AI, Orji EO, Adanikin PO, Olaniyan O. Comparative study of rectal misoprostol to oxytocin infusion in preventing postpartum haemorrhage post-caesarean section. *International Journal of Gynecology & Obstetrics* 2012;**119**(Suppl 3):S825.

* Adanikin AI, Orji EO, Fasubaa OB, Onwudiegwu U, Ijarotimi OA, Olaniyan O. The effect of post-cesarean rectal misoprostol on intestinal motility. *International Journal of Gynecology and Obstetrics* 2012;**119**(2):159-62.

Orji EO, Adanikin AI. Prospective randomised double blind study on the effect of post-caesarean rectal misoprostol on intestinal motility. *International Journal of Gynecology and Obstetrics* 2012;**119**(Suppl 3):S446-S447.

Orji EO, Adanikin AO. The effect of post-caesarean rectal misoprostol on intestinal motility. *BJOG: an international journal of obstetrics and gynaecology* 2013;**120**(Suppl s1):21.

Adanikin 2013 (published data only)

Adanikin AI, Orji E, Adanikin PO, Olaniyan O. Comparative study of rectal misoprostol to oxytocin infusion in preventing postpartum haemorrhage after caesarean section. *Nepal Journal of Obstetrics and Gynaecology* 2013;**8**(2):34-7.

Afolabi 2010 {published data only}

Afolabi EO, Kuti O, Orji EO, Ogunniyi SO. Oral misoprostol versus intramuscular oxytocin in the active management of the third stage of labour. *Singapore Medical Journal* 2010;**51**(3):207-11.

Ahmed 2014 (published data only)

Ahmed WA, Ibrahim ZM, Mostafa I, Kishk EA, Elbahie MA. Safety and efficacy of carbetocin in hypertensive pregnant women undergoing cesarean delivery. *Journal of Maternal-Fetal & Neonatal Medicine* 2014;**27**(Suppl 1):49.

Al-Sawaf 2013 {published data only}

Al-Sawaf A, El-Mazny A, Shohayeb A. A randomised controlled trial of sublingual misoprostol and intramuscular oxytocin for prevention of postpartum haemorrhage. *Journal of Obstetrics and Gynaecology* 2013;**33**(3):277-9.

Alwani 2014 (published data only)

Alwani M, Singh S, Thakur R, Mishra S. A randomized study comparing rectally administered misoprostol after spinal anesthesia versus intramuscular oxytocin for prevention of postpartum hemorrhage in caesarean section. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2014;**3**(3):512-5.

Amant 1999 {published data only}

Amant F, Spitz B, Timmerman D, Corresmans A, Van Assche FA. Misoprostol compared with methylergometrine for the prevention of postpartum haemorrhage: a double-blind randomised trial. *British Journal of Obstetrics and Gynaecology* 1999;**106**:1066-70.

Amin 2014 (published data only)

Amin N. Prophylactic use of misoprostol in management of third stage of labour and prevention of atonic uterus. *Journal of Postgraduate Medical Institute* 2014;**28**(2):196-200.

Askar 2011 {published data only}

Askar AA, Ismail MT, El-Ezz AA, Rabie NH. Carbetocin versus syntometrine in the management of third stage of labor following vaginal delivery. *Archives of Gynecology and Obstetrics* 2011;**284**(6):1359-65.

Asmat 2017 {published data only}

Asmat R, Ashraf T, Asmat F, Asmat S, Asmat N. Effectiveness of per rectal misoprostol versus intramuscular oxytocin for prevention of primary postpartum haemorrhage. *Journal of the College of Physicians and Surgeons Pakistan* 2017;**27**(1):13-7.

Attilakos 2010 {published data only}

Attilakos G, Psaroudakis D, Ash J, Buchanan R, Winter C, Donald F, et al. Can a new oxytocin analogue reduce the need for additional oxytocics after caesarean section? The results of a double-blind randomised trial. *Archives of Disease in Childhood. Fetal and Neonatal Edition* 2008;**93**(Suppl 1):Fa51.

* Attilakos G, Psaroudakis D, Ash J, Buchanan R, Winter C, Donald F, et al. Carbetocin versus oxytocin for the prevention of postpartum haemorrhage following caesarean section: the results of a double-blind randomised trial. *BJOG: an international journal of obstetrics and gynaecology* 2010;**117**(8):929-36.

Attilakos G, Psaroudakis D, Ash J, Buchanan R, Winter C, Donald F, et al. The haemodynamic effects of oxytocin and carbetocin following caesarean section: the results of a doubleblind randomised study. *BJOG: an international journal of obstetrics and gynaecology* 2008;**115**(s1):140-1.

Attilakos G, Psaroudakis D, Ash J, Buchanen R, Winter C, Draycott T. Low recruitment rate for a drug trial in obstetrics:



an effect of the publicity following the TGN1412 clinical trial at the PAREXEL Research Unit in Northwick Park Hospital? [abstract]. 31st British International Congress of Obstetrics and Gynaecology; 2007 July 4-6; London, UK. 2007:110.

Atukunda 2014 (published data only)

Atukunda EC, Siedner MJ, Obua C, Mugyenyi GR, Twagirumukiza M, Agaba AG. Sublingual misoprostol versus intramuscular oxytocin for prevention of post-partum haemorrhage in Uganda: a randomised, controlled, non-inferiority trial. *Lancet* 2014;**384**(Suppl 1):S3.

* Atukunda EC, Siedner MJ, Obua C, Mugyenyi GR, Twagirumukiza M, Agaba AG. Sublingual misoprostol versus intramuscular oxytocin for prevention of postpartum hemorrhage in Uganda: a double-blind randomized non-inferiority trial. *PLOS Medicine* 2014;**11**(11):e1001752.

Badejoko 2012 {published data only}

Badejoko OO, Ijarotimi AO, Awowole IO, Loto OM, Badejoko BO, Olaiya DS, et al. Adjunctive rectal misoprostol versus oxytocin infusion for prevention of postpartum hemorrhage in women at risk: a randomized controlled trial. *Journal of Obstetrics and Gynaecology Research* 2012;**38**(11):1294-301.

Balki 2008 (published data only)

Balki M, Dhumne S, Kasodekar S, Kingdom J, Windrim R, Carvalho JC. Oxytocin-ergometrine co-administration does not reduce blood loss at caesarean delivery for labour arrest. *BJOG: an international journal of obstetrics and gynaecology* 2008;**115**(5):579-84.

Bamigboye 1998a {published data only}

Bamigboye AA, Hofmeyr GJ, Merrell DA. Rectal misoprostol in the prevention of postpartum haemorrhage: a placebo controlled trial. Proceedings of the 17th Conference on Priorities in Perinatal Care; 1998; South Africa. 1998:49-52.

* Bamigboye AA, Hofmeyr GJ, Merrell DA. Rectal misoprostol in the prevention of postpartum hemorrhage: a placebocontrolled trial. *American Journal of Obstetrics and Gynecology* 1998;**179**(4):1043-6.

Bamigboye 1998b {published data only}

Bamigboye AA, Merrell DA, Hofmeyr GJ, Mitchell R. Randomized comparison of rectal misoprostol with syntometrine for management of third stage of labor. *Acta Obstetricia et Gynecologica Scandinavica* 1998;**77**:178-81.

Barton 1996 (published data only)

Barton SR, Jackson A. The safety and efficiency of carbetocin to control uterine bleeding following caesarean section. *Prenatal and Neonatal Medicine* 1996;**1**(Suppl 1):185.

Baskett 2007 (published data only)

Baskett TF, Persad V, Clough H, Young D. Prophylactic use of misoprostol in the third stage of labor [abstract]. *Obstetrics & Gynecology* 2005;**105**(4 Suppl):39S.

* Baskett TF, Persad VL, Clough HJ, Young DC. Misoprostol versus oxytocin for the reduction of postpartum blood loss. *International Journal of Gynecology & Obstetrics* 2007;**97**(1):2-5.

Chandra S, Persad V, Young D, Baskett T. A preliminary study of cutaneous blood flow associated with postpartum use of oral misoprostol. *Journal of Obstetrics & Gynaecology Canada: JOGC* 2004;**26**(12):1073-6.

Begley 1990 {published data only}

Begley CM. Comparative studies in the third stage of labour [MSc thesis]. Dublin: Trinity College, University of Dublin, 1990.

* Begley CM. A comparison of 'active' and 'physiological' management of the third stage of labour. *Midwifery* 1990;**6**:3-17.

Begley CM. The effect of ergometrine on breast feeding. *Midwifery* 1990;**6**:60-72.

Begum 2015 (published data only)

Begum T, Yeasmin S, Chakma S. Sublingual misoprostol versus oxitocin infusion to reduce blood loss in caesarean section. *BJOG: an international journal of obstetrics and gynaecology* 2015;**122**(Suppl S1):258.

Bellad 2012 (published data only)

Bellad M, Ganachari TD, Mallapur M. Sublingual (SL) powdered misoprostol (400 mcg) vs IM oxytocin (10 IU) for prevention of postpartum blood loss - a randomized controlled trial. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S124-5.

* Bellad MB, Tara D, Ganachari MS, Mallapur MD, Goudar SS, Kodkany BS, et al. Prevention of postpartum haemorrhage with sublingual misoprostol or oxytocin: a double-blind randomised controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2012;**119**(8):975-86.

Benchimol 2001 (published data only)

Benchimol M, Gondry J, Mention JE, Gagneur O, Boulanger JC. Role of misoprostol in the delivery outcome. [French] [Place du misoprostol dans la direction de la delivrance.]. *Journal de Gynecologie, Obstetrique et Biologie de la Reproduction* 2001;**30**(6):576-83.

Bhatti 2014 (published data only)

Bhatti K, Mahar T, Hafeez R, Shoaib-u-Nisa. A randomized controlled trial on prevention of postpartum haemorrhage with sublingual misoprostol or oxytocin. *Medical Forum Monthly* 2014;**25**(1):10-2.

Bhullar 2004 (published data only)

Bhullar A, Carlan SJ, Hamm J, Lamberty N, White L, Richichi K. Buccal misoprostol to decrease blood loss after vaginal delivery: a randomized trial. *Obstetrics & Gynecology* 2004;**104**(6):1282-8.

Biswas 2007 (published data only)

Biswas A, Bal R, Kundu MK, Kyal A, Halder M. A study of prophylactic use of 15-methyl prostaglandin f2alpha in the active management of third stage of labour. *Journal of the Indian Medical Association* 2007;**105**(9):506-9.

Borruto 2009 (published data only)

Borruto F, Treisser A, Comparetto C. Utilization of carbetocin for prevention of postpartum hemorrhage after cesarean section: a



randomized clinical trial. *Archives of Gynecology and Obstetrics* 2009;**280**(5):707-12.

Boucher 1998 (published data only)

Boucher M, Horbay GL, Griffin P, Deschamps Y, Desjardins C, Schulz M, et al. Double-blind, randomized comparison of the effect of carbetocin and oxytocin on intraoperative blood loss and uterine tone of patients undergoing cesarean section. *Journal of Perinatology* 1998;**18**(3):202-7.

Boucher 2004 (published data only)

Boucher M, Nimrod C, Tawagi G. Carbetocin IM injection vs oxytocin IV infusion for prevention of postpartum hemorrhage in women at risk following vaginal delivery. *American Journal of Obstetrics and Gynecology* 2001;**185**(6 Pt 2):Abstract no: 494.

* Boucher M, Nimrod CA, Tawagi GF, Meeker TA, Rennicks White RE, Varin J. Comparison of carbetocin and oxytocin for the prevention of postpartum hemorrhage following vaginal delivery:a double-blind randomized trial. *Journal of Obstetrics & Gynaecology Canada: JOGC* 2004;**26**(5):481-8.

Bugalho 2001 {published data only}

Bugalho A, Daniel A, Foundes A, Cunha M. Misoprostol for the prevention of postpartum hemorrhage. *International Journal of Gynecology & Obstetrics* 2001;**73**:1-6.

Butwick 2010 {published data only}

Butwick AJ, Coleman L, Cohen SE, Riley ET, Carvalho B. A study of the minimum effective dose of oxytocin in patients undergoing elective cesarean delivery. American Society of Anaesthesiologists Annual Meeting; 2009 Oct 17-21; New Orleans, USA. 2009.

* Butwick AJ, Coleman L, Cohen SE, Riley ET, Carvalho B. Minimum effective bolus dose of oxytocin during elective caesarean delivery. *British Journal of Anaesthesia* 2010;**104**(3):338-43.

Caliskan 2002 (published data only)

Caliskan E, Meydanli M, Dilbaz B, Aykan B, Sonmezer M, Haberal A. Is rectal misoprostol really effective in the treatment of third stage of labor? A randomized controlled trial. *American Journal of Obstetrics and Gynecology* 2002;**187**:1038-45.

Caliskan 2003 {published data only}

Caliskan E, Dilbaz B, Meydanli MM, Ozturk N, Narin MA, Haberal A. Oral misoprostol for the third stage of labor: a randomized controlled trial. *Obstetrics & Gynecology* 2003;**101**(5 Pt 1):921-8.

Carbonell 2009 (published data only)

Carbonell i Esteve JL, Hernandez JM, Piloto M, Setien SA, Texido CS, Tomasi G, et al. Active management of the third phase of labour plus 400 mug of sublingual misoprostol and 200 mug of rectal misoprostol versus active management only in the prevention of post-partum haemorrhage. A randomised clinical trial [Manejo activo de la tercera fase del parto mas 400 mug de misoprostol sublingual y 200 mug de misoprostol rectal frente a manejo activo solo en la prevencion de la hemorragia posparto. Ensayo clinico aleatorizado]. *Progresos de Obstetricia y Ginecologia* 2009;**52**(10):543-51.

Carrillo-Gaucin 2016 (published data only)

Carrillo-Gaucin S, Torres-Gomez LG. Carbetocin and oxytocin: prevention of postpartum hemorrhage in patients with risk factors for uterine atony [Carbetocina y oxitocina: prevención de hemorragia posparto en pacientes con factores de riesgo para atonía uterina]. *Revista Médica del Instituto Mexicano del Seguro Social* 2016;**54 Suppl 3**:S284-90.

Cayan 2010 {published data only}

Cayan F, Doruk A, Sungur MA, Dilek S. Comparison of the different dosages of rectal misoprostol on I=intestinal motility and pain score in high risk cesarean delivery. *Turkiye Klinikleri Journal of Medical Sciences* 2010;**30**(4):1154-9.

Chalermpolprapa 2010 (published data only)

Chalermpolprapa V. Efficacy of sublingual misoprostol in prevention of postpartum hemorrhage in cesarean section: A randomized double-blinded, placebo-controlled trial. *Region 4-5 Medical Journal* 2010;**29**(3):325-35.

Chandhiok 2006 (published data only)

Chandhiok N, Dhillon BS, Datey S, Mathur A, Saxena NC. Oral misoprostol for prevention of postpartum hemorrhage by paramedical workers in India. *International Journal of Gynecology & Obstetrics* 2006;**92**(2):170-5.

Chaudhuri 2010 {published data only}

Chaudhuri P, Banerjee GB, Mandal A. Rectally administered misoprostol versus intravenous oxytocin infusion during cesarean delivery to reduce intraoperative and postoperative blood loss. *International Journal of Gynecology & Obstetrics* 2010;**109**(1):25-9.

Chaudhuri 2012 {published data only}

Chaudhuri P, Biswas J, Mandal A. Sublingual misoprostol versus intramuscular oxytocin for prevention of postpartum hemorrhage in low-risk women. *International Journal of Gynecology and Obstetrics* 2012;**116**(2):138-42.

Chaudhuri 2015 {published data only}

Chaudhuri P, Majumdar A. Sublingual misoprostol as an adjunct to oxytocin during cesarean delivery in women at risk of postpartum hemorrhage. *International Journal of Gynaecology & Obstetrics* 2015;**128**:48-52.

Chaudhuri 2016 {published data only}

Chaudhuri P, Majumdar A. A randomized trial of sublingual misoprostol to augment routine third-stage management among women at risk of postpartum hemorrhage. *International Journal of Gynecology and Obstetrics* 2016;**132**:191-5.

Chhabra 2008 (published data only)

Chhabra S, Tickoo C. Low-dose sublingual misoprostol versus methylergometrine for active management of the third stage of labor. *Journal of Obstetrics and Gynaecology Research* 2008;**34**(5):820-3.

Choy 2002 {published data only}

Choy CM, Lau WC, Tam WH, Yuen PM. A randomised controlled trial of intramuscular syntometrine and intravenous oxytocin in the management of the third stage of labour. *BJOG:*



an international journal of obstetrics and gynaecology 2002;**109**:173-7.

Chua 1995 {published data only}

Chua S, Chew SL, Yeoh CL, Roy AC, Ho LM, Selamat N, et al. A randomized controlled study of prostaglandin 15-Methyl F2 alpha compared with syntometrine for prophylactic use in the third stage of labour. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 1995;**35**:413-6.

Cook 1999 {published data only}

Cook CM, Spurrett B, Murray H. A randomized clinical trial comparing oral misoprostol with synthetic oxytocin or syntometrine in the third stage of labour. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 1999;**39**(4):414-9.

Dabbaghi Gale 2012 {published data only}

Dabbaghi Gale T, Elmizadeh KH, Moradi SD, Rashvand Melli E. Comparison of intravenous oxytocin and oral misoprostol in reduction of postpartum hemorrhage. *Journal of Zanjan University of Medical Sciences and Health Services* 2012;**20**(81):1-8.

Dansereau 1999 {published data only}

Dansereau J. Comparison of carbetocin vs. oxytocin in prevention of uterine atony post cesarean section. *Prenatal and Neonatal Medicine* 1996;**1**(Suppl 1):80.

Dansereau J, Gambling D, Joshi A, Helewa M, Doran T, Lange I, et al. Double-blind comparison of carbetocin vs oxytocin in preventing uterine atony post cesarean section. *International Journal of Gynecology & Obstetrics* 1994;**46 Suppl**:77.

* Dansereau J, Joshi AK, Helewa ME, Doran TA, Lange IR, Luther ER, et al. Double blind comparison of carbetocin versus oxytocin in prevention of uterine atony after cesarean section. American Journal of Obstetrics and Gynecology 1999;**18**(3 Pt 1):670-6.

Dansereau J, Joshi AK, Helewa ME, Doran TA, Lange IR, Luther ER, et al. Double-blind comparison of carbetocin vs oxytocin in preventing uterine atony post cesarean section. *European Journal of Obstetrics & Gynecology and Reproductive Biology* 1996;**69**:37.

Gambling D, Dansereau J, Schulz M, Horbay GL, Waasenaar W. Double-blind, randomized comparison of a single dose of carbetocin vs 8 hours oxytocin infusion after cesarean delivery: safety data. A Canadian multi-center trial [abstract]. *International Journal of Obstetric Anesthesia* 1994;**3**:113-4.

Gambling DR, Dansereau J, Wassenaar W, Schulz M, Horbay GLA. Double-blind randomized comparison of a single dose of carbetocin versus 8 hours oxytocin infusion after cesarean delivery: safety data. *Anesthesia & Analgesia* 1994;**78 Suppl**:S127.

Dasuki 2002 (published data only)

Dasuki D, Emilia O, Harini S. Randomized clinical trial: the effectiveness of oral misoprostol versus oxytocin in prevention of postpartum hemorrhage [abstract]. *Journal of Obstetrics and Gynaecology Research* 2002;**28**(1):46.

de Groot 1996 (published data only)

de Groot AN, Van Roosmalen J, Van Dongen PWJ, Borm GF. A placebo-controlled trial of oral ergometrine to reduce postpartum hemorrhage. *Acta Obstetricia et Gynecologica Scandinavica* 1996;**75**:464-8.

Del Angel-Garcia 2006 (published data only)

Del Angel-Garcia G, Garcia-Contreras F, Constantino-Casas P, Nevarez-Sida A, Lopez-Gonzalez N, Garcia-Constantino M, et al. Economic evaluation of carbetocin for the prevention of uterine atony in patients with risk factors in Mexico. *Value in Health* 2006;**9**(6):A254.

Derman 2006 {published data only}

* Derman RJ, Kodkany BS, Goudar SS, Geller SE, Naik VA, Bellad MB, et al. Oral misoprostol in preventing postpartum haemorrhage in resource-poor communities: a randomised controlled trial. *Lancet* 2006;**368**(9543):1248-53.

Geller SE, Goudar SS, Adams MG, Naik VA, Patel A, Bellad MB, et al. Factors associated with acute postpartum hemorrhage in low-risk women delivering in rural India. *International Journal of Gynecology & Obstetrics* 2008;**101**(1):94-9.

Geller SE, Patel A, Niak VA, Goudar SS, Edlavitch SA, Kodkany BS, et al. Conducting international collaborative research in developing nations. *International Journal of Gynecology & Obstetrics* 2004;**87**(3):267-71.

Goudar SS, Chakraborty H, Edlavitch SA, Naik VA, Bellad MB, Patted SS, et al. Variation in the postpartum hemorrhage rate in a clinical trial of oral misoprostol. *Journal of Maternal-Fetal & Neonatal Medicine* 2008;**21**(8):559-64.

Kodkany BS, Derman RJ, Goudar SS, Geller SE, Edlavitch SA, Naik VA, et al. Initiating a novel therapy in preventing postpartum hemorrhage in rural India: a joint collaboration between the United States and India. *International Journal of Fertility & Womens Medicine* 2004;**49**(2):91-6.

Kodkany BS, Goudar SS, Derman RJ. The efficacy of oral misoprostol in preventing postpartum hemorrhage in a community setting: a randomized double-blind placebocontrolled trial. *International Journal of Gynecology & Obstetrics* 2006;**94**(Suppl 2):S141-S142.

NCT00097123. RCT of misoprostol for postpartum hemorrhage in India. clinicaltrials.gov/ct2/show/NCT00097123 (first received 18 November 2004).

Patted SS, Goudar SS, Naik VA, Bellad MB, Edlavitch SA, Kodkany BS, et al. Side effects of oral misoprostol for the prevention of postpartum hemorrhage: results of a community-based randomised controlled trial in rural India. *Journal of Maternal-Fetal & Neonatal Medicine* 2009;**22**(1):24-8.

Dhananjaya 2014 (published data only)

Dhananjaya BS, Charishma S. Comparative study of efficacy and safety of intramuscular oxytocin with intramuscular methylergometrine in the active management of third stage of labour. Research Journal of Pharmaceutical, Biological and Chemical Sciences 2014;5(3):734-9.



Diallo 2017 (published data only)

* Diallo M, Sylla T, Diouf AA, Moreira PM, Gassama O, Gueye MD, et al. Active management of third stage of labour with low doses of oral misoprostol and oxytocin on low: risk parturient in a Sub-Saharan hospital, Dakar, Sénégal. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2017;**6**(2):516-22.

Vlassoff M, Diallo A, Philbin J, Kost K, Bankole A. Costeffectiveness of two interventions for the prevention of postpartum hemorrhage in Senegal. *International Journal of Gynecology and Obstetrics* 2016;**133**(3):307-11.

Diop 2016 {published data only}

* Diop A, Daff B, Sow M, Blum J, Diagne M, Sloan NL, et al. Oxytocin via Uniject (a prefilled single-use injection) versus oral misoprostol for prevention of postpartum haemorrhage at the community level: a cluster-randomised controlled trial. *Lancet. Global Health* 2016;**4**(1):e37-44.

NCT01713153. Comparing misoprostol and oxytocin in uniject for postpartum hemorrhage (PPH) prevention in Senegal. clinicaltrials.gov/ct2/show/NCT01713153 (first received: 22 October 2012).

Docherty 1981 {published data only}

Docherty PW, Hooper M. Choice of an oxytocic agent for routine use at delivery. *Journal of Obstetrics and Gynaecology* 1981;**2**:60.

Dutta 2016 {published data only}

Dutta BK, Gupta KR. A comparative study on rectal misoprostol versus intramuscular oxytocin to prevent postpartum haemorrhage. *New Indian Journal of OBGYN* 2016;**2**(2):98-103.

Eftekhari 2009 {published data only}

Eftekhari N, Doroodian M, Lashkarizadeh R. The effect of sublingual misoprostol versus intravenous oxytocin in reducing bleeding after caesarean section. *Journal of Obstetrics and Gynaecology* 2009;**29**(7):633-6.

El Behery 2015 {published data only}

El Behery MM, El Sayed GA, El Hameed AA, Soliman BS, Abdelsalam WA, Bahaa A. Carbetocin versus oxytocin for prevention of postpartum hemorrhage in obese nulliparous women undergoing emergency cesarean delivery. Journal of Maternal-Fetal & Neonatal Medicine 2015:1-4.

Elbohoty 2016 {published data only}

Elbohoty AE, Mohammed WE, Sweed M, Eldin AM, Nabhan A, Abd-El-Maeboud KH. Randomized controlled trial comparing carbetocin, misoprostol, and oxytocin for the prevention of postpartum hemorrhage following an elective cesarean delivery. *International Journal of Gynaecology and Obstetrics* 2016;**134**(3):324-8.

Elgafor El Sharkwy 2013 (published data only)

Elgafor El Sharkwy IA. Carbetocin versus sublingual misoprostol plus oxytocin infusion for prevention of postpartum hemorrhage at cesarean section in patients with risk factors: a randomized, open trail study. *Archives of Gynecology and Obstetrics* 2013;**288**(6):1231-6.

El-Refaey 2000 (published data only)

El-Refaey H, Nooh R, O'Brien P, Abdalla M, Geary M, Walder J, et al. The misoprostol third stage of labour study: a randomised controlled comparison between orally administered misoprostol and standard management. *BJOG: an international journal of obstetrics and gynaecology* 2000;**107**:1104-10.

Elsedeek 2012 {published data only}

Elsedeek MS. Impact of preoperative rectal misoprostol on blood loss during and after elective cesarean delivery. *International Journal of Gynecology & Obstetrics* 2012;**118**(2):149-52.

El Tahan 2012 (published data only)

El Tahan MR, Warda OM, Rashad A, Yasseen AM, Ramzy EA, Ahmady MS, et al. Effects of preoperative sublingual misoprostol on uterine tone during isoflurane anesthesia for cesarean section. *Revista Brasileira de Anestesiologia* 2012;**62**(5):625-35.

Enakpene 2007 (published data only)

Enakpene CA, Morhason-Bello IO, Enakpene EO, Arowojolu AO, Omigbodun AO. Oral misoprostol for the prevention of primary post-partum hemorrhage during third stage of labor. *Journal of Obstetrics and Gynaecology Research* 2007;**33**(6):810-7.

Ezeama 2014 (published data only)

Ezeama CO, Eleje GU, Ezeama NN, Igwegbe AO, Ikechebelu JI, Ugboaja JO, et al. A comparison of prophylactic intramuscular ergometrine and oxytocin for women in the third stage of labor. *International Journal of Gynecology & Obstetrics* 2014;**124**(1):67-71.

Fahmy 2015 (published data only)

Fahmy AA, Fawzy M. Oxytocin infusion after oxytocin bolus and carbetocin bolus to reduce blood loss during and after cesarean section - a randomized clinical trial. *Medical Journal of Cairo University* 2015;**83**(1):79-83.

Fahmy 2016 {published data only}

Fahmy NG, Yousef HM, Zaki HV. Comparative study between effect of carbetocin and oxytocin on isoflurane-induced uterine hypotonia in twin pregnancy patients undergoing cesarean section. *Egyptian Journal of Anaesthesia* 2016;**32**(1):117-21.

Fakour 2013 (published data only)

Fakour F, Mirzayi M, Reza Naghipour M, Ebrahimi H, Mahdavi M. Comparison between sublingual misoprostol and intravenous oxytocin in management of third stage of labor. *Iranian Journal of Obstetrics, Gynecology and Infertility* 2013;**15**(34):7-14.

Fararjeh 2003 {published data only}

Fararjeh C, Gezer A, Cepni I, Benian A, Ocal P, Kosebay D. The efficacy of misoprostol in preventing postpartum bleeding [Postpartum Kanama Profilaksisinde Rektal Mizoprostol Kulanımının Etkinliği.]. *Jinekoloji ve Obstetrik Dergisi* 2003;**17**(4):218-23.

Fawole 2011 {published data only}

Fawole AO, Sotiloye OS, Hunyinbo KI, Umezulike AC, Okunlola MA, Adekanle DA, et al. A double-blind, randomized,



placebo-controlled trial of misoprostol and routine uterotonics for the prevention of postpartum hemorrhage. *International Journal of Gynecology & Obstetrics* 2011;**112**(2):107-11.

Fawzy 2012 {published data only}

Fawzy AE, Swelem M, Abdelrehim AI, Titeli S, Elghazal ZS, El-Gahwagi MM, et al. Active management of third stage of labor by intravenous ergometrine and rectal versus sublingual misoprostol (a double-center study). *Alexandria Journal of Medicine* 2012;**48**(4):381-5.

Fazel 2013 {published data only}

Fazel MR, Mansoure-Samimi, Esmaeil-Fakharian. A comparison of rectal misoprostol and intravenous oxytocin on hemorrhage and homeostatic changes during cesarean section. *Middle East Journal of Anesthesiology* 2013;**22**(1):41-6.

Fekih 2009 (published data only)

Fekih M, Jnifene A, Fathallah K, Ben Regaya L, Memmi A, Bouguizene S, et al. Benefit of misoprostol for prevention of postpartum hemorrhage in cesarean section: a randomized controlled trial. *Journal de Gynecologie, Obstetrique et Biologie de la Reproduction* 2009;**38**(7):588-93.

Fenix 2012 (published data only)

Fenix AM. Double-blind randomized controlled trial comparing the effect of carbetocin with oxytocin for the prevention of postpartum hemorrhage among high risk women following vaginal delivery. *International Journal of Gynaecology & Obstetrics* 2012;**119**(Suppl 3):S347-S348.

Fu 2003 (published data only)

Fu YX, Ran KQ, Wang M. Prevention of early postpartum hemorrhage by way of oral misoprostol. *Journal of Nursing Science* 2003;**18**(12):910-1.

Fuks 2014 (published data only)

Fuks AM, Khanna P, Yusaf T, Aslian A, Kowalska D, Salafia CM. Use of prophylactic misoprostol in reduction of blood loss at vaginal delivery. *Obstetrics and Gynecology* 2014;**123**(5 Suppl):144S-5S.

Garg 2005 (published data only)

Garg P, Batra S, Gandhi G. Oral misoprostol versus injectable methylergometrine in management of the third stage of labor. *International Journal of Gynecology & Obstetrics* 2005;**91**(2):160-1.

Gavilanes 2016 {published data only}

Gavilanes P, Morales MF, Velasco S, Teran E. Sublingual misoprostol is as effective as intravenous oxytocin to reduce intra-operative blood loss during cesarean delivery in women living at high altitude. *Journal of Maternal-Fetal & Neonatal Medicine* 2016;**29**(4):559-61.

Gerstenfeld 2001 {published data only}

Gerstenfeld TS, Wing DA. Rectal misoprostol versus intravenous oxytocin for the prevention of postpartum hemorrhage after vaginal delivery. *American Journal of Obstetrics and Gynecology* 2001;**185**:878-82.

Gore 2017 (published data only)

Gore S, Padmawar A, Pathan SK. A prospective randomized controlled trial for comparison of oral misoprostol with methyl ergometrine in the third stage of labour for prevention of postpartum haemorrhage. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2017;**6**(7):2825-8.

Gulmezoglu 2001 (published data only)

* Gulmezoglu AM, Villar J, Ngoc NT, Piaggio G, Carroli G, Adetoro L, et al. WHO multicentre randomised trial of misoprostol in the management of the third stage of labour. *Lancet* 2001;**358**:689-95.

Lumbiganon P, Hofmeyr J, Gulmezoglu AM, Pinol A, Villar J. Misoprostol dose-related shivering and pyrexia in the third stage of labour Who collaborative trial of misoprostol in the management of the third stage of labour. *British Journal of Obstetrics and Gynaecology* 1999;**106**(4):304-8.

Lumbiganon P, Villar J, Piaggio G, Gulmezoglu AM, Adetoro L, Carroli G. Side effects of oral misoprostol during the first 24 hours after administration in the third stage of labour. *BJOG: an international journal of obstetrics and gynaecology* 2002;**109**:1222-6.

Gupta 2006 (published data only)

Gupta B, Jain V, Aggarwal N. Rectal misoprostol versus oxytocin in the prevention of postpartum hemorrhage - a pilot study. *International Journal of Gynecology & Obstetrics* 2006;**94**(Suppl 2):S139-S140.

Hamm 2005 {published data only}

Hamm J, Russell Z, Botha T, Carlan SJ, Richichi K. Buccal misoprostol to prevent hemorrhage at cesarean delivery: a randomized study. *American Journal of Obstetrics and Gynecology* 2005;**192**:1404-6.

Harriott 2009 (published data only)

Harriott J, Christie L, Wynter S, DaCosta V, Fletcher H, Reid M. A randomized comparison of rectal misoprostol with syntometrine on blood loss in the third stage of labour. *West Indian Medical Journal* 2009;**58**(3):201-6.

Hernandez-Castro 2016 *[published data only]*

* Hernandez-Castro F, Lopez-Serna N, Trevino-Salinas EM, Soria-Lopez JA, Sordia-Hernandez LH, Cardenas-Estrada E. Randomized double-blind placebo-controlled trial of buccal misoprostol to reduce the need for additional uterotonic drugs during cesarean delivery. *International Journal of Gynaecology and Obstetrics* 2016;**132**(2):184-7.

NCT01733329. Buccal misoprostol during cesarean section for preventing postpartum hemorrhage in women with risk factors for uterine atony. clinicaltrials.gov/ct2/show/NCT01733329 (first received 16 November 2012).

Hofmeyr 1998 {published data only}

Hofmeyr GJ, Nikodem C, de Jager M, Drakely A, Gilbart B. Oral misoprostol for labour third stage management: randomised assessment of side effects. Proceedings of the 17th Conference on Priorities in Perinatal Care; 1998; South Africa. 1998:53-4.



* Hofmeyr GJ, Nikodem VC, de Jager M, Gelbart BR. A randomised placebo controlled trial of oral misoprostol in the third stage of labour. *British Journal of Obstetrics and Gynaecology* 1998;**105**(9):971-5.

Hofmeyr GJ, de Jager M, Rose L, Nikodem VC, Lawrie T. Misoprostol for third stage of labour management: a double blind, placebo controlled clinical trial. Proceedings of the 16th Conference on Priorities in Perinatal Care; 1997; South Africa. 1997:29-31.

Hofmeyr 2001 {published data only}

Hofmeyr GJ, Nikodem VC, De Jager M, Drakely AJ. Side effects of oral misoprostol in the third stage of labour: a random allocation placebo controlled trial. *Journal of Obstetrics & Gynaecology* 2000;**20**(Suppl 1):S40-1.

* Hofmeyr GJ, Nikodem VC, de Jager M, Drakely A. Sideeffects of oral misoprostol in the third stage of labour--a randomised placebo-controlled trial. *South African Medical Journal* 2001;**91**(5):432-5.

Hofmeyr 2011 {published data only}

* Hofmeyr GJ, Fawole B, Mugerwa K, Godi NP, Blignaut Q, Mangesi L, et al. Administration of 400mug of misoprostol to augment routine active management of the third stage of labor. *International Journal of Gynecology and Obstetrics* 2011;**112**(2):98-102.

NCT00124540. Misoprostol for preventing postpartum hemorrhage. clinicaltrials.gov/ct2/show/NCT00124540 (first received: 26 July 2005).

Hoj 2005 {published data only}

* Hoj L, Cardoso P, Nielsen BB, Hvidman L, Nielsen J, Aaby P. Effect of sublingual misoprostol on severe postpartum haemorrhage in a primary health centre in Guinea-Bissau: randomised double blind clinical trial. *BMJ* 2005;**331**:723-7.

Nielsen BB, Hoj L, Hvidman LE, Nielsen J, Cardoso P, Aaby P. Reduced post-partum bleeding after treatment with sublingual misoprostol: a randomized doubleblind clinical study in a developing country - secondary publication [Reduceret post partum-blodning efter sublingval misoprostol: et randomiseret dobbeltblindt klinisk studie i et udviklingsland - sekundaerpublikation]. *Ugeskrift for Laeger* 2006;**168**(13):1341-3.

Hong 2007 (published data only)

Hong SC, Kim JW, Park HT, Seol HJ, Kim HJ, Kim SH, et al. Additional rectal misoprostol plus intravenous oxytocin versus intravenous oxytocin for the prevention of postpartum hemorrhage after cesarean section. *American Journal of Obstetrics and Gynecology* 2007;**197**(6 Suppl 1):S99, Abstract no: 321.

Humera 2016 {published data only}

Humera A, Kerure SB. Randomized comparative study of oral misoprostol with intravenous methyl ergometrine in prevention of postpartum haemorrhage in cases high risk for postpartum haemorrhage. *International Journal*

of Reproduction, Contraception, Obstetrics and Gynecology 2016:**5**(3):798-802.

Is 2012 {published data only}

Is S, Gr V, Keranahalli S. Comparison of intramuscular ergometrine and per rectal misoprostol for prophylaxis against atonic post partum haemorrhage. *International Journal of Gynecology & Obstetrics* 2012;**119**(Suppl 3):S797-S798.

Jago 2007 (published data only)

Jago AA, Ezechi OC, Achinge GI, Okunlola MA. Effect of oxytocics on the blood pressure of normotensive Nigerian parturients. Journal of Maternal-Fetal & Neonatal Medicine 2007;**20**(9):703-5.

Jangsten 2011 {published data only}

Jangsten E, Mattsson L, Lyckestam I, Hellstrom A, Berg M. A comparison of active management and expectant management of the third stage of labour: a Swedish randomised controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2011;**118**:362–9.

Jans 2017 {published data only}

Jans S, Herschderfer KC, van Diem MT, Aitink M, Rijnders M, van der Pal-Bruin K, et al. LENTE study: effectiveness of prophylactic intramuscular oxytocin during third stage of labour amongst low risk women. A randomized controlled trial. 31st International Confederation of Midwives Triennial Congress. Midwives - Making a Difference in the World; 2017 June 18-22; Toronto, Canada. 2017:Abstract no: P1.063.

Jerbi 2007 {published data only}

Jerbi M, Hidar S, Elmoueddeb S, Chaieb A, Khairi H. Oxytocin in the third stage of labor. *International Journal of Gynecology & Obstetrics* 2007;**96**(3):198-9.

Jirakulsawas 2000 {published data only}

Jirakulsawas J, Khooarmompattana S. Comparison of oral misoprostol and intramuscular methylergonovine for prevention of postpartum hemorrhage. *Thai Journal of Obstetrics and Gynaecology* 2000;**12**(4):332.

Kabir 2015 {published data only}

Kabir N, Akter D, Daisy TA, Jesmin S, Razzak M, Tasnim S, et al. Efficacy and safety of carbetocin in comparison to oxytocin in the active management of third stage of labour following vaginal delivery: an open label randomized control trial. Bangladesh Journal of Obstetrics and Gynecology 2015;**30**(1):3-9.

Karkanis 2002 (published data only)

Karkanis SG, Caloia D, Salenieks ME, Kingdom J, Walker M, Meffe F, et al. Randomized controlled trial of rectal misoprostol versus oxytocin in third stage management. *Journal of Obstetrics and Gynaecology Canada: JOGC* 2002;**24**(2):149-54.

Kerekes 1979 {published data only}

Kerekes L, Domokos N. The effect of prostaglandin F2alpha on third stage labour. *Prostaglandins* 1979;**18**:161-6.

Khan 1995 {published data only}

* Khan GQ, John IS, Chan T, Wani S, Hughes AO, Stirrat GM. Abu Dhabi third stage trial: oxytocin vs syntometrine in the



active management of the third stage of labour. *European Journal of Obstetrics & Gynecology and Reproductive Biology* 1995;**58**:147-51.

Khan GQ, Susheela JI, Chan T, Wani S, Hughes AO, Stirrat GM. Abu Dhabi third stage trial: oxytocin versus syntometrine in the active management of the third stage of labour. 27th British Congress of Obstetrics and Gynaecology; 1995 July 4-7; Dublin, Ireland. 1995:Abstract no: 212.

Khurshid 2010 (published data only)

Khurshid R, Fatima K, Parveen S, Ul Shamas I, Salman R. A comparison between intramuscular PGF2 a125 mG and intravenous methyl ergometrine 0.2 Mg in the active management of third stage labor. Internet Journal of Gynecology and Obstetrics 2010; Vol. 14, issue 1.

Koen 2016 (published data only)

Koen S, Snyman LC, Pattinson RC, Makin JA. A randomised controlled trial comparing oxytocin and oxytocin + ergometrine for prevention of postpartum haemorrhage at caesarean section. *South African Medical Journal = Suid-Afrikaanse Tydskrif Vir Geneeskunde* 2016;**106**(4):399-402.

Kumar 2016 (published data only)

Kumar SK, Shyam S, Batakurki P. Carboprost versus oxytocin for active management of third stage of labor: a prospective randomized control study. *Journal of Obstetrics and Gynecology of India* 2016;**66**(S1):S229-S234.

Kumru 2005 (published data only)

Kumru S, Gurates B, Parmaksiz C. Investigation of the usefulness of methyl ergonovine application in cesarean section cases [Sezaryen olgularinda metil ergonovin uygulamasinin yararliliginin arastirilmasi]. *Journal of the Turkish German Gynecology Association Artemis* 2005;**6**(1):42-5.

Kundodyiwa 2001 (published data only)

Kundodyiwa TW, Majoko F, Rusakaniko S. Misoprostol versus oxytocin in the third stage of labor. *International Journal of Gynecology & Obstetrics* 2001;**75**:235-41.

Kushtagi 2006 (published data only)

* Kushtagi P, Verghese LM. Evaluation of two uterotonic medications for the management of the third stage of labor. *International Journal of Gynecology & Obstetrics* 2006;**94**(1):47-8.

Verghese L, Kushtagi P. Evaluation of carboprost as a prophylactic oxytocic in the management of third stage of labour. *BJOG: an international journal of obstetrics and gynaecology* 2008;**115**(s1):74.

Lam 2004 {published data only}

Lam H, Tang OS, Lee CP, Ho PC. A pilot-randomized comparison of sublingual misoprostol with syntometrine on the blood loss in third stage of labor. *Acta Obstetricia et Gynecologica Scandinavica* 2004;**83**:647-50.

Lamont 2001 {published data only}

Lamont RF, Morgan DJ, Logue M, Gordon H. A prospective randomised trial to compare the efficacy and safety of hemabate and syntometrine for the prevention of primary

postpartum haemorrhage. *Prostaglandins & Other Lipid Mediators* 2001;**66**(3):203-10.

Lapaire 2006 {published data only}

Lapaire O, Schneider MC, Stotz M, Surbek DV, Holzgreve W, Hoesli IM. Oral misoprostol vs. intravenous oxytocin in reducing blood loss after emergency cesarean delivery. *International Journal of Gynecology & Obstetrics* 2006;**95**(1):2-7.

Leung 2006 (published data only)

Leung SW, Ng PS, Wong WY, Cheung TH. A randomised trial of carbetocin versus syntometrine in the management of the third stage of labour. *BJOG: an international journal of obstetrics and gynaecology* 2006;**113**:1459-64.

Lokugamage 2001 (published data only)

Lokugamage AU, Moodley J, Sullivan K, Rodeck CH, Niculescu L, Tigere P. The Durban primary postpartum haemorrhage study. Women's Health - into the new millennium. Proceedings of the 4th International Scientific Meeting of the Royal College of Obstetricians and Gynaecologists; 1999 October 3-6; Cape Town South Africa. RCOG, 1999:77-8.

Lokugamage AU, Paine M, Bassau-Balroop H, El-Refaey K, Sullivan K, Rodek C. Active management of the third stage at caesarean section: misoprostol vs syntocinon. XVI FIGO World Congress of Obstetrics & Gynecology; 2000 Sept 3-8; Washington DC, USA. 2000; Vol. Book 2:54.

* Lokugamage AU, Paine M, Bassaw-Balroop K, Sullivan KR, El-Refaey H, Rodeck CH. Active management of the third stage at caesarean section: a randomised controlled trial of misoprostol versus syntocinon. *Australian & New Zealand Journal of Obstetrics & Gynaecology* 2001;**41**(4):411-4.

Lumbiganon 1999 (published data only)

Lumbiganon P, Hofmeyr J, Gülmezoglu AM, Pinol A, Villar J. Misoprostol dose-related shivering and pyrexia in the third stage of labour. *British Journal of Obstetrics and Gynaecology* 1999;**106**:304-8.

Maged 2016 (published data only)

Maged AM, Hassan AM, Shehata NA. Carbetocin versus oxytocin for prevention of postpartum hemorrhage after vaginal delivery in high risk women. *Journal of Maternal-Fetal & Neonatal Medicine* 2016;**29**(4):532-6.

Maged 2017 {published data only}

Maged AM, Ragab AS, Elnassery N, Al Mostafa W, Dahab S, Kotb A. Carbetocin versus syntometrine for prevention of postpartum hemorrhage after cesarean section. *Journal of Maternal-Fetal & Neonatal Medicine* 2017;**30**(8):962-6.

Malik 2018 {published data only}

Sabha M Cimona LS Fida M. A comparative study between prostaglandin 2a and methyl ergometrine in the active management of third stage of labour. *Modern Approaches in Drug Designing* 2018;**1**(5):MADD.000522.

Mannaerts 2018 (published data only)

Mannaerts D, Van der Veeken L, Coppejans H, Jacquemyn Y. Adverse effects of carbetocin versus oxytocin in the



prevention of postpartum haemorrhage after caesarean section: a randomized controlled trial. *Journal of Pregnancy* 2018;**2018**:1374150.

McDonald 1993 {published data only}

McDonald S. The Perth third stage oxytocic trial. Proceedings of National Conference on Research in Midwifery; 1992; Reading, UK. 1992.

McDonald S, Prendiville WJ. A randomized controlled trial of syntocinon vs syntometrine as part of the active management of the third stage of labour. *Journal of Perinatal Medicine* 1992;**20**(Suppl 1):97.

McDonald S, Prendiville WJ, Blair E. A randomised controlled trial of syntocinon vs syntometrine as part of the active management of the third stage of labour. Proceedings of 26th British Congress of Obstetrics and Gynaecology; 1992 July 7-10; Manchester, UK. 1992:87.

McDonald SJ. Management in the Third Stage of Labour [thesis]. University of Western Australia, 1996.

* McDonald SJ, Prendiville WJ, Blair E. Randomised controlled trial of oxytocin alone vs oxytocin and ergometrine in active management of third stage of labour. *BMJ* 1993;**307**:1167-71.

Mitchell 1993 {published data only}

Mitchell GG, Elbourne DR. The Salford third stage trial: oxytocin plus ergometrine vs oxytocin alone in the active management of the third stage of labor. *Online Journal of Current Clinical Trials* 1993;**2**:Doc 83.

Mobeen 2011 {published data only}

* Mobeen N, Durocher J, Zuberi NF, Jahan N, Blum J, Wasim S, et al. Administration of misoprostol by trained traditional birth attendants to prevent postpartum haemorrhage in homebirths in Pakistan: a randomised placebo-controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2011;**118**:353-61.

Mobeen N, Durocher J, Zuberi NF, Jahan N, Blum J, Wasim S, et al. Use of misoprostol by trained traditional birth attendants to prevent postpartum haemorrhage during home deliveries in Pakistan: a randomised placebo-controlled trial. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S92.

NCT00120237. Misoprostol for the prevention of postpartum hemorrhage in rural Pakistan. clinicaltrials.gov/ct2/show/NCT00120237 (first received: 7 July 2005).

Modi 2014 {published data only}

Modi V, Goel JK, Kashyap A, Arya SB, Kar J, Goel R. Active management of third stage of labor: A comparison of various uterotonic. *Journal of South Asian Federation of Obstetrics and Gynaecology - SAFOG* 2014;**6**(3):151-5.

Moertl 2011 {published data only}

Moertl M, Kraschl J, Friedrich S, Pickel K, Ulrich D, Eder M, et al. Hemodynamic changes of carbetocin and oxytocin in women undergoing cesarean section. *American Journal of Obstetrics and Gynecology* 2008;**199**(6 Suppl 1):S112.

* Moertl MG, Friedrich S, Kraschl J, Wadsack C, Lang U, Schlembach D. Haemodynamic effects of carbetocin and oxytocin given as intravenous bolus on women undergoing caesarean delivery: a randomised trial. *BJOG: an international journal of obstetrics and gynaecology* 2011;**118**(11):1349-56.

Mortl M, Pickel K, Friedrich S, Ulrich D, Lang U, Schlembach D. Hemodynamic changes of carbetocin and oxytocin given as i.v. bolus on women undergoing cesarean section. *Geburtshilfe und Frauenheilkunde* 2008;**68**:S46.

Mohamed 2015 (published data only)

Mohamed HF, Mustafa GF, Ibrahim MA, Stefanos GE. Comparative study between intravenous bolus dose of carbetocin versus oxytocin during cesarean delivery in healthy parturients on blood loss: a randomized control trial. *Medical Journal of Cairo University* 2015;83(2):167-72.

Moir 1979 {published data only}

Moir DD, Amoa AB. Ergometrine or oxytocin? Blood loss and side-effects at spontaneous vertex delivery. *British Journal of Anaesthesia* 1979;**51**(2):113-7.

Moodie 1976 {published data only}

Moodie JE, Moir DD. Ergometrine, oxytocin and extradural analgesia. *British Journal of Anaesthesia* 1976;**48**:571-4.

Mukta 2013 (published data only)

Mukta M, Sahay PB. Role of misoprostol 600 mcg oral in active management of third stage of labor: a comparative study with oxytocin 10 IU i.m. *Journal of Obstetrics and Gynecology of India* 2013;**6**:325-7.

Musa 2015 (published data only)

Musa AO, Ijaiya MA, Saidu R, Aboyeji AP, Jimoh AA, Adesina KT, et al. Double-blind randomized controlled trial comparing misoprostol and oxytocin for management of the third stage of labor in a Nigerian hospital. International Journal of Gynecology & Obstetrics 2015; Vol. 129, issue 3:227-30.

Nankaly 2016 (published data only)

Nankaly A, Jalilian N, Eshghiali S, Rezaei M. The effects of sublingual misoprostol and intravenous oxytocin in reducing bleeding among cesarean deliveries. *Acta Medica Mediterranea* 2016;**32**:953-7.

Nasr 2009 {published data only}

Nasr A, Shahin AY, Elsamman AM, Zakherah MS, Shaaban OM. Rectal misoprostol versus intravenous oxytocin for prevention of postpartum hemorrhage. *International Journal of Gynecology & Obstetrics* 2009;**105**(3):244-7.

Nayak 2017 (published data only)

Nayak L Pradhan K Mishra S. Role of 400 mcg intraoperative sublingual misoprostol for reduction of caesarean blood loss. *Journal of Evidence Based Medicine and Healthcare* 2017;**4**(10):573-7.

Nellore 2006 {published data only}

Nellore V, Mittal S, Dadhwal V. Rectal misoprostol vs. 15-methyl prostaglandin F2alpha for the prevention of postpartum



hemorrhage. *International Journal of Gynecology & Obstetrics* 2006:**94**(1):45-6.

Ng 2001 {published data only}

Ng PS, Chan AS, Sin WK, Tang LC, Cheung KB, Yuen PM. Comparison of oral misoprostol and intramuscular syntometrine in the management of the third stage of labor - a multicenter randomised controlled trial. XVI FIGO World Congress of Obstetrics & Gynecology. Book 4; 2000 Sept 3-8; Washington DC, USA. 2000:29.

* Ng PS, Chan ASM, Sin WK, Tang LCH, Cheung KB, Yuen PM. A multicentre randomized controlled trial of oral misoprostol and im syntometrine in the management of the third stage of labour. *Human Reproduction* 2001;**16**(1):31-5.

Ng 2004 (published data only)

Ng PS, Yuen PM, Sahota DS. Comparison of oral misoprostol and intravascular syntocinon in the management of the third stage of labour - a double-blind randomised controlled trial. 30th British Congress of Obstetrics and Gynaecology; 2004 July 7-9; Glasgow, UK. 2004:69.

Ng 2007 {published data only}

* Ng PS, Lai CY, Sahota DS, Yuen PM. A double-blind randomized controlled trial of oral misoprostol and intramuscular syntometrine in the management of the third stage of labor. *Gynecologic and Obstetric Investigation* 2007;**63**(1):55-60.

Yuen PM, Ng PS, Sahota DS. A double-blind randomised controlled trial of oral misoprostol in addition to intra-muscular syntometrine in the management of the third stage of labour. 30th British Congress of Obstetrics and Gynaecology; 2004 July 7-9; Glasgow, UK. 2004:62.

Nirmala 2009 {published data only}

Nirmala K, Zainuddin AA, Ghani NA, Zulkifli S, Jamil MA. Carbetocin versus syntometrine in prevention of post-partum hemorrhage following vaginal delivery. *Journal of Obstetrics and Gynaecology Research* 2009;**35**(1):48-54.

Nordstrom 1997 {published data only}

Nordstrom L, Fogelstam K, Fridman G, Larsson A, Rydhstroem H. Routine oxytocin in the third stage of labour: a placebo controlled randomised trial. *British Journal of Obstetrics and Gynaecology* 1997;**104**(7):781-6.

Nuamsiri 2016 {published data only}

Nuamsiri T Kaewkiattikun K. Prevention of postpartum hemorrhage with oxytocin versus ergometrine plus oxytocin in the third stage of labor. *Thai Journal of Obstetrics and Gynaecology* 2016;**24**:97-103.

Oboro 2003 {published data only}

Oboro VO, Tabowei TO. A randomised controlled trial of misoprostol versus oxytocin in the active management of the third stage of labour. *Journal of Obstetrics & Gynaecology* 2003;**23**(1):13-6.

Ogunbode 1979 (published data only)

Ogunbode O, Obisesan K, Ayeni O. Methergin in the management of the third stage of labor: a comparative clinical trial with syntometrine and ergometrine. *Current Therapeutic Research, Clinical and Experimental* 1979;**26**:460-5.

Orji 2008 (published data only)

Orji E, Agwu F, Loto O, Olaleye O. A randomized comparative study of prophylactic oxytocin versus ergometrine in the third stage of labor. *International Journal of Gynecology & Obstetrics* 2008;**101**(2):129-32.

Ortiz-Gomez 2013 {published data only}

Ortiz-Gomez JR, Morillas-Ramirez F, Fornet-Ruiz I, Palacio-Abizanda FJ, Bermejo-Albares L. [Clinical and pharmacological study of the efficacy of carbetocin in elective caesareans compared to low and usual doses of oxytocin]. [Spanish]. *Revista Espanola de Anestesiologia y Reanimacion* 2013;**60**(1):7-15.

Othman 2016 (published data only)

Othman ER, Fayez MF, El Aal DEMA, El-Dine Mohamed HS, Abbas AM, Ali MK. Sublingual misoprostol versus intravenous oxytocin in reducing bleeding during and after cesarean delivery: a randomized clinical trial. *Taiwanese Journal of Obstetrics and Gynecology* 2016;**55**(6):791-5.

Owonikoko 2011 {published data only}

Owonikoko KM, Arowojolu AO, Okunlola MA. Effect of sublingual misoprostol versus intravenous oxytocin on reducing blood loss at cesarean section in Nigeria: A randomized controlled trial. *Journal of Obstetrics and Gynaecology Research* 2011;**37**(7):715-21.

Pakniat 2015 (published data only)

Pakniat H, Khezri MB. The effect of combined oxytocinmisoprostol versus oxytocin and misoprostol alone in reducing blood loss at cesarean delivery: a prospective randomized double-blind study. *Journal of Obstetrics and Gynaecology of India* 2015;**65**(6):376-81.

Parsons 2006 (published data only)

Parsons SM, Walley RL, Crane JM, Matthews K, Hutchens D. Oral misoprostol versus oxytocin in the management of the third stage of labour. *Journal of Obstetrics and Gynaecology Canada* 2006;**28**(1):20-6.

Parsons 2007 (published data only)

Parsons S, Ntumy YM, Walley RL, Wilson JB, Crane JM, Matthews K, et al. Rectal misoprostol vs intramuscular oxytocin in the routine management of the third stage of labour. 30th British Congress of Obstetrics and Gynaecology; 2004 July 7-9; Glasgow, UK. 2004:18.

* Parsons SM, Walley RL, Crane JM, Matthews K, Hutchens D. Rectal misoprostol versus oxytocin in the management of the third stage of labour. *Journal of Obstetrics and Gynaecology Canada* 2007;**29**(9):711-8.



Patil 2013 (published data only)

Patil NB, Patted SS. A randomised controlled trial of oral misoprostol vs injection methylergometrine for prevention of post partum hemorrhage. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2013;**2**(3):296-303.

Patil 2016 {published data only}

Patil AS, Dadavate V, Thobbi VA. Carboprost versus oxytocin for active management of third stage of labour. *Al Ameen Journal of Medical Sciences* 2016;**9**(3):196-201.

Penaranda 2002 (published data only)

Penaranda WA, Arrieta OB, Yances BR. Active management of childbirth with sublingual misoprostol: a controlled clinical trial in the Hospital de Maternidad Rafael Calvo [Manejo activo del alumbramiento con misoprostol sublingual: un estudio clinico controlado en al hospital de maternidad rafael calvo de cartagena]. Revista Colombiana de Obstetricia y Ginecologia 2002;**53**(1):87-92.

Perez-Rumbos 2017 (published data only)

Perez-Rumbos A, Reyna-Villasmil E Rondon-Tapia M, Reyna-Villasmil N. Rectal misoprostol or intramuscular oxytocin in the management of the third phase of labour. *Perinatología y Reproducción Humana* 2017;**31**(2):78-84.

Poeschmann 1991 {published data only}

* Poeschmann RP, Doesburg WH, Eskes TK. A randomized comparison of oxytocin, sulprostone and placebo in the management of the third stage of labour. *British Journal of Obstetrics and Gynaecology* 1991;**98**:528-30.

Poeschmann RP, Eskes TK, Doesburg WH. Oxytocin and sulprostone reduce post partum blood loss and shorten the third stage in low risk term women. *International Journal of Gynecology & Obstetrics* 1991;**36 Suppl**:312.

Poeschmann RP, Eskes TK, Doesburg WH, Lemmens WA, Benneker JC. Oxytocin and sulprostone reduce postpartum blood loss in low risk term women compared to saline. Proceedings of 1st European Congress on Prostaglandins in Reproduction; 1988 July 6-9; Vienna, Austria. 1988:176.

Prendiville 1988 {published data only}

Prendiville WJ, Harding JE, Elbourne DR, Stirrat GM. The Bristol third stage trial: active versus physiological management of third stage of labour. *BMJ* 1988;**297**(6659):1295-300.

Quibel 2016 {published data only}

* Quibel T, Ghout I, Goffinet F, Salomon LJ, Fort J, Javoise S, et al. Active management of the third stage of labor with a combination of oxytocin and misoprostol to prevent postpartum hemorrhage: a randomized controlled trial. *Obstetrics and Gynecology* 2016;**128**(4):805-11.

Rozenberg P, Quibel T, Ghout I, Salomon L, Bussiere L, Goffinet F. Active management of the third stage of labor with routine oxytocin and misoprostol for the prevention of postpartum hemorrhage: a randomized controlled trial. *American Journal of Obstetrics and Gynecology* 2015;**212**(1 Suppl 1):S18.

Rajaei 2014 (published data only)

NCT01863706. Misoprostol versus oxytocin for prevention of post partum hemorrhage. clinicaltrials.gov/ct2/show/NCT01863706 (first received: 23 May 2013).

* Rajaei M, Karimi S, Shahboodaghi Z, Mahboobi H, Khorgoei T, Rajaei F. Safety and efficacy of misoprostol versus oxytocin for the prevention of postpartum hemorrhage. *Journal of Pregnancy* 2014;**2014**:713879.

Ramirez 2001 (published data only)

Ramirez O, Benito V, Jimenez R, Valido C, Hernandez C, Garcia JA. Third stage of labour: active or expectant management? preliminary results [abstract]. *Journal of Perinatal Medicine* 2001;**Suppl 1**(Pt 2):364.

Rashid 2009 (published data only)

Rashid M, Clark A, Rashid MH. A randomised controlled trial comparing the efficacy of intramuscular syntometrine and intravenous syntocinon, in preventing postpartum haemorrhage. *Journal of Obstetrics and Gynaecology* 2009;**29**(5):396-401.

Ray 2001 (published data only)

Chatterjee A. Misoprostol and the 3rd stage. XVI FIGO World Congress of Obstetrics & Gynecology; 2000 Sept 3-8; Washington DC, USA. 2000; Vol. Book 4:29.

* Ray A, Mukherjee P, Basu G, Chatterjee A. Misoprostol and third stage of labour. *Journal of Obstetrics and Gynecology of India* 2001;**51**(6):53-4.

Reddy 2001 {published data only}

Reddy R, Shenoy JV. Active management of third stage of labour. A comparative study in high risk patients for atonic postpartum haemorrhage. *Journal of Obstetrics and Gynecology of India* 2001;**51**(2):44-7.

Reyes, Gonzalez 2011 (published data only)

Reyes OA, Gonzalez GM. Carbetocin versus oxytocin for prevention of postpartum hemorrhage in patients with severe preeclampsia: a double-blind randomized controlled trial. *Journal of Obstetrics and Gynaecology Canada: JOGC* 2011;**33**(11):1099-104.

Reyes 2011 (published data only)

Reyes OA. Carbetocin vs oxytocin for the prevention of postpartum hemorrhage in grand multiparous patients: A randomized controlled trial. [Spanish]. Clinica e investigacion en ginecologia y obstetricia 2011; Vol. 38, issue 1:2-7.

Rogers 1998 (published data only)

Rogers J, Wood J, McCandlish R, Ayers S, Truesdale A, Elbourne D. Active versus expectant management of third stage of labour: the Hinchingbrooke randomised controlled trial. *Lancet* 1998;**351**(9104):693-9.

Rosseland 2013 (published data only)

Gawecka E, Rosseland LA. A secondary analysis of a randomized placebo-controlled trial comparing the analgesic effects of



oxytocin with carbetocin: postcesarean delivery morphine equivalents. *Anesthesia and Analgesia* 2014;**119**(4):1004.

* Rosseland LA, Hauge TH, Grindheim G, Stubhaug A, Langesaeter E. Changes in blood pressure and cardiac output during cesarean delivery: the effects of oxytocin and carbetocin compared with placebo. *Anesthesiology* 2013;**119**(3):541-51.

Sadiq 2011 {published data only}

Sadiq UG, Kwanashie O, Mairiga G, Gamaniel S, Isa H, Abdu A, et al. A randomised clinical trial comparing the efficacy of oxytocin injection and oral misoprostol tablet in the prevention of postpartum haemorrhage in Maiduguri Nigeria. *International Research Journal of Pharmacy* 2011;**2**(8):76-81.

Samimi 2013 (published data only)

Samimi M, Imani-Harsini A, Abedzadeh-Kalahroudi M. Carbetocin vs. syntometrine in prevention of postpartum hemorrhage: a double blind randomized control trial. *Iranian Red Crescent Medical Journal* 2013;**15**(9):817-22.

Shady 2017 (published data only)

Shady NW, Sallam HF, Elsayed AH, Abdelkader AM, Ali SS, Alanwar A, et al. The effect of prophylactic oral tranexamic acid plus buccal misoprostol on blood loss after vaginal delivery: a randomized controlled trial. *Journal of Maternal-Fetal & Neonatal Medicine* 2017;27:1-7.

Shrestha 2011 (published data only)

Shrestha A, Dongol A, Chawla CD, Adhikari RK. Rectal misoprostol versus intramuscular oxytocin for prevention of post partum hemorrhage. *Kathmandu University Medical Journal* 2011;**33**(1):8-12.

Singh 2009 {published data only}

Singh G, Radhakrishnan G, Guleria K. Comparison of sublingual misoprostol, intravenous oxytocin, and intravenous methylergometrine in active management of the third stage of labor. *International Journal of Gynecology & Obstetrics* 2009;**107**(2):130-4.

Sitaula 2017 (published data only)

Sitaula S Uprety DK Thakur A Pradhan T. Impact of preoperative rectal misoprostol on blood loss during and after elective cesarean delivery: a randomized controlled trial. *Nepal Journal of Obstetrics and Gynaecology* 2017;**11**(2):37-41.

Soltan 2007 (published data only)

Soltan MH, El-Gendi E, Imam HH, Fathi O. Different doses of sublingual misoprostol versus methylergometrine for the prevention of atonic postpartum haemorrhage. *International Journal of Health Sciences* 2007;**1**(2):229-36.

Sood 2012 {published data only}

Sood AK, Singh S. Sublingual misoprostol to reduce blood loss at cesarean delivery. *Journal of Obstetrics and Gynaecology of India* 2012;**62**(2):162-7.

Stanton 2013 {published data only}

NCT01108302. Effectiveness, safety and feasibility of auxiliary nurse midwives' (ANM) use of oxytocin in uniject™ to prevent

postpartum hemorrhage in India. clinicaltrials.gov/ct2/show/NCT01108302 (first received: 2 April 2010).

Stanton CK, Newton S, Mullany LC, Cofie P, Agyemang CT, Adiibokah E, et al. Impact on postpartum hemorrhage of prophylactic administration of oxytocin 10 IU via Uniject by peripheral health care providers at home births: Design of a community-based cluster-randomized trial. *BMC Pregnancy and Childbirth* 2012;**12**:42.

* Stanton CK, Newton S, Mullany LC, Cofie P, Tawiah Agyemang C, Adiibokah E, et al. Effect on postpartum hemorrhage of prophylactic oxytocin (10 IU) by injection by community health officers in Ghana: a community-based, cluster-randomized trial. PLOS Medicine. NCT01108289 2013; Vol. 10, issue 10:e1001524.

Su 2009 {published data only}

Su LL, Rauff M, Chan YH, Mohamad Suphan N, Lau TP, Biswas A, et al. Carbetocin versus syntometrine for the third stage of labour following vaginal delivery--a double-blind randomised controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2009;**116**(11):1461-6.

Sultana 2007 (published data only)

Sultana N, Khatun M. Misoprostol versus oxytocin in the active management of the third stage of labour. *Journal of Bangladesh College of Physicians and Surgeons* 2007;**25**(2):73-6.

Supe 2016 (published data only)

Supe PA, Kore SJ, Nandanwar YS. A comparative study of efficacy of misoprostol with methyl ergometrine and carboprost in active management of third stage of labour. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2016;**5**(5):1525-31.

Surbek 1999 {published data only}

Surbek DV, Fehr PM, Hoesli I, Holzgreve W. Misoprostol for prevention of postpartum hemorrhage: a randomized controlled trial [abstract]. XVI FIGO World Congress of Obstetrics & Gynecology; 2000 Sept 3-8; Washington DC, USA 2000; Book 1:33.

* Surbek DV, Fehr PM, Hoesli I, Holzgreve W. Oral misoprostol vs placebo for third stage of labour [Orales misoprostol reduziert den postpartalen blutverlust]. *Gynakologisch Geburtshilfliche Rundschau* 1999;**39**:144.

Taheripanah 2018 (published data only)

Taheripanah R, Shoman A, Ali Karimzadeh M, Zamaniyan M, Malih N. Efficacy of oxytocin vs. Carbetocin in prevention of postpartum hemorrhage after cesarean section under general anesthesia: a prospective randomized clinical trial. *Journal of Maternal-Fetal & Neonatal Medicine* 2018;**31**(21):2807-12.

Tewatia 2014 {published data only}

Tewatia R, Rani S, Srivastav U, Makhija B. Sublingual misoprostol versus intravenous oxytocin in prevention of post-partum hemorrhage. *Archives of Gynecology and Obstetrics* 2014;**289**:739-42.



Thilaganathan 1993 (published data only)

Thilaganathan B, Cutner A, Latimer J, Beard R. Management of the third stage of labour in women at low risk of postpartum haemorrhage. *European Journal of Obstetrics & Gynecology and Reproductive Biology* 1993;**48**:19-22.

Tripti 2006 (published data only)

Tripti N, Manju E. Intramuscular PGF2 alpha 125 microg versus intravenous methyl ergometrine 0.2 mg in the active management of third stage of labor. *Journal of Obstetrics and Gynecology of India* 2006;**56**(5):396-8.

Ugwu 2014 {published data only}

Ugwu IA, Enabor OO, Adeyemi AB, Lawal OO, Oladokun A, Olayemi O. Sublingual misoprostol to decrease blood loss after caesarean delivery: a randomised controlled trial. *Journal of Obstetrics and Gynaecology* 2014;**34**(5):407-11.

Uncu 2015 {published data only}

Uncu Y, Karahasan M, Uyaniklar O, Uncu G. Prophylactic misoprostol for the prevention of postpartum hemorrhage: a randomized controlled trial. *European Review for Medical and Pharmacological Sciences* 2015;**19**(1):15-22.

Un Nisa 2012 {published data only}

Un Nisa S, Usmani SY. Role of intravenous syntocinon in prevention of primary postpartum haemorrhage. *Pakistan Journal of Medical and Health Sciences* 2012;**6**(4):1020-4.

Vagge 2014 (published data only)

Vagge DS, Mamatha KR, Rohatgi V. A comparative study to assess the efficacy and tolerability of per rectal misoprostol versus intravenous oxytocin in prevention of primary postpartum haemorrhage in a tertiary care hospital. *Indian Journal of Pharmacology* 2013;**45 Suppl**:S45.

* Vagge DS, Mamatha KR, Shivamurthy G, Rohatgi V. A comparative study to assess the efficacy and tolerability of per rectal misoprostol and intravenous oxytocin in prevention of primary postpartum haemorrhage in a tertiary care hospital. *Journal of Chemical and Pharmaceutical Research* 2014;**6**(3):1134-40.

Vaid 2009 {published data only}

Vaid A, Dadhwal V, Mittal S, Deka D, Misra R, Sharma JB, et al. A randomized controlled trial of prophylactic sublingual misoprostol versus intramuscular methyl-ergometrine versus intramuscular 15-methyl PGF2alpha in active management of third stage of labor. *Archives of Gynecology and Obstetrics* 2009;**280**(6):893-7.

Van Selm 1995 {published data only}

Van Selm M, Kanhai HH, Keirse MJ. Preventing the recurrence of atonic postpartum hemorrhage: a double-blind trial. *Acta Obstetricia et Gynecologica Scandinavica* 1995;**74**:270-4.

Verma 2006 (published data only)

Verma P, Aggarwal N, Jain V, Suri V. A double-blind randomized controlled trial to compare sublingual misoprostol with methylergometrine for prevention of postpartum hemorrhage.

International Journal of Gynecology & Obstetrics 2006;**94**(Suppl 2):S137-S138.

Vimala 2004 (published data only)

Vimala N, Mittal S, Kumar S, Dadhwal V, Mehta S. Sublingual misoprostol versus methylergometrine for active management of third stage of labor. *International Journal of Gynecology & Obstetrics* 2004;**87**:1-5.

Vimala 2006 (published data only)

Vimala N, Mittal S, Kumar S. Sublingual misoprostol versus oxytocin infusion to reduce blood loss at cesarean section. *International Journal of Gynecology & Obstetrics* 2006;**92**(2):106-10.

Walley 2000 {published data only}

Walley RL, Wilson JB, Crane JM, Matthews K, Sawyer E, Hutchens D. A double-blind placebo controlled randomised trial of misoprostol and oxytocin in the management of the third stage of labour. *BJOG: an international journal of obstetrics and gynaecology* 2000;**107**(9):1111-5.

Whigham 2016 {published data only}

Whigham CA, Gorelik A, Loughnan T, Trivedi A. Carbetocin versus oxytocin in active labour. *BJOG: an international journal of obstetrics and gynaecology* 2014;**121**(Suppl 2):88.

* Whigham CA, Gorelik A, Loughnan TE, Trivedi A. Carbetocin versus oxytocin to reduce additional uterotonic use at non-elective caesarean section: a double-blind, randomised trial. *Journal of Maternal-Fetal & Neonatal Medicine* 2016;**29**(23):3866-9.

Widmer 2018 (published data only)

Gulmezoglu M. The WHO Champion Trial. *International Journal of Gynecology and Obstetrics* 2015;**131**(Suppl 5):E29-30.

Widmer M, Piaggio G, Abdel-Aleem H, Carroli G, Chong Y, Coomarasamy A, et al. Room temperature stable carbetocin for the prevention of postpartum haemorrhage during the third stage of labour in women delivering vaginally: study protocol for a randomized controlled trial. *Trials* 2016;**17**:143.

* Widmer M, Piaggio G, Nguyen TM, et al. Heat-stable carbetocin versus oxytocin to prevent hemorrhage after vaginal birth. *New England Journal of Medicine* 2018;**379**(8):743-52.

Yuen 1995 {published data only}

Yuen PM, Chan NS, Yim SF, Chang AM. A randomised double blind comparison of syntometrine and syntocinon in the management of the third stage of labour. *British Journal of Obstetrics and Gynaecology* 1995;**102**:377-80.

Zachariah 2006 {published data only}

Zachariah ES, Naidu M, Seshadri L. Oral misoprostol in the third stage of labor. *International Journal of Gynecology & Obstetrics* 2006;**92**(1):23-6.



References to studies excluded from this review

Abdel-Aleem 2013 (published data only)

Abdel-Aleem H, Alhusaini TK, Abdel-Aleem MA, Menoufy M, Gulmezoglu AM. Effectiveness of tranexamic acid on blood loss in patients undergoing elective cesarean section: randomized clinical trial. *Journal of Maternal-Fetal & Neonatal Medicine* 2013;**26**(17):1705-9.

Abdel-Aleem 2018 {published data only}

Abdel-Aleem AA, Abbas AM, Thabet AL, Badran E, El-Nashar IH. The effect of initiating intravenous oxytocin infusion before uterine incision on the blood loss during elective cesarean section: a randomized clinical trial. Journal of Maternal-Fetal & Neonatal Medicine 2018 [Epub ahead of print].

Abdollahy 2000 {published data only}

Abdollahy F. Comparison effect of oxytocin and normal salin injection intra umbelical venuse [abstract]. *Gynecological Endocrinology* 2000;**14**(Suppl 2):49.

Adhikari 2007 (published data only)

Adhikari S, Rana A, Bista KD. Active management of third stage of labour: comparison between prophylactic intramuscular methylergometrine and intramuscular oxytocin. *Nepal Journal of Obstetrics and Gynaecology* 2007;**2**(2):24-8.

Adnan 2017 (published data only)

Adnan N, Boland F, Murphy DJ. Intramuscular oxytocin versus intravenous oxytocin to prevent postpartum haemorrhage at vaginal delivery (LabOR trial): study protocol for a randomised controlled trial. *Trials* 2017;**18**(1):541.

Ahmed 2015 {published data only}

Ahmed MR, Sayed Ahmed WA, Madny EH, Arafa AM, Said MM. Efficacy of tranexamic acid in decreasing blood loss in elective caesarean delivery. *Journal of Maternal-Fetal & Neonatal Medicine* 2015;**28**(9):1014-8.

Akinaga 2016 (published data only)

Akinaga C, Uchizaki S, Kurita T, Taniguchi M, Makino H, Suzuki A, et al. Randomized double-blind comparison of the effects of intramyometrial and intravenous oxytocin during elective cesarean section. *Journal of Obstetrics and Gynaecology Research* 2016;**42**(4):404-9.

Alam 2017 {published data only}

Alam A, Shyam P, Goswami S. A comparative study of efficacy of oxytocin, methylergometrine and misoprostol in prevention of post-partum haemorrhage. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2017;**6**(5):1960-4.

Al-Harazi 2009 (published data only)

Al-Harazi AH, Frass KA. Sublingual misoprostol for the prevention of postpartum hemorrhage. *Saudi Medical Journal* 2009;**30**(7):912-6.

Ali 2012 {published data only}

Ali R, Hina F. Postpartum hemorrhage; comparison of efficacy of ergometrine with misoprostol in prophylaxis in cesarean section. *Professional Medical Journal* 2012;**19**(3):360-4.

Ali 2018 (published data only)

Ali AA, Nasr AA, Ahmed HH, El- Rasheedy MI, Badawy M. Carbetocin versus oxytocin and misoprostol in prevention of atonic post-partum hemorrhage in high risk patients planed for cesarean delivery. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2018;**7**(1):10-4.

Anandakrishnan 2013 (published data only)

Anandakrishnan S, Balki M, Farine D, Seaward G, Carvalho JC. Carbetocin at elective cesarean delivery: a randomized controlled trial to determine the effective dose, part 2. *Canadian Journal of Anaesthesia* 2013;**60**(11):1054-60.

Anjaneyulu 1988 (published data only)

Anjaneyulu R, Devi PK, Kanthamani CR, Vijaya R, Raghavan KS. Prophylactic use of 15(S)15 methyl PGF2alpha by intramuscular route - a controlled clinical trial. *Acta Obstetricia et Gynecologica Scandinavica Supplementa* 1988;**145**:9-11.

Anvaripour 2013 {published data only}

Anvaripour A, Shahryari H, Ahmadi S, Ghasemi S, Mirzaei K. Comparison the effects of oxytocin and methylergonovine in elective caesarean section under spinal anesthesia. *Archives of Gynecology and Obstetrics* 2013;**287**(5):979-83.

Ashwal 2016 (published data only)

Ashwal E, Hiersch L, Wertheimer A, Krispin E, Aviram A, Dayan DB, et al. The effect of post-partum oxytocin regimen on hemoglobin decline–a randomized controlled trial. *American Journal of Obstetrics and Gynecology* 2016;**214**(1 Suppl):S197-S198, Abstract no: 351.

Athavale 1991 {published data only}

Athavale RD, Nerurkar NM, Dalvi SA, Bhattacharya MS. Umbilical vein oxytocin in the management of third stage of labour. Journal of Postgraduate Medicine 1991; **37**(4):219-20.

Ayedi 2011 {published data only}

Ayedi M, Zouche I, Smaoui L, Bouaziz I, Smaoui M, Kolsi K. Comparison of 2 versus 5 units of oxytocin in caesarean section. European Journal of Anaesthesiology 2011;**28 Suppl**:159-60.

Ayedi 2011b {published data only}

Ayedi M, Jarraya A, Smaoui M, Zouari J, Smaoui L, Kolsi K. Effect of tranexamic acid on post partum hemorrhage by uterine atony: a preliminary result of a randomized, placebo controlled trial. *European Journal of Anaesthesiology* 2011;**28 Suppl 1**:165.

Ayedi 2012 {published data only}

Ayedi M, NCT01599468. Effects of tranexamic acid on post partum hemorrhage by uterine atony after cesarean section delivery: a randomized, placebo controlled trial. clinicaltrials.gov/ct2/show/NCT01599468 (first received 14 May 2012).



Aziz 2014 (published data only)

Aziz S, Kazi S, Haq G, Soomro N. Oral misoprostol versus oxytocin in the management of third stage of labour. *JPMA - Journal of the Pakistan Medical Association* 2014;**64**(4):428-32.

Bader 2000 (published data only)

* Bader W, Ast S, Hatzmann W. The significance of acupuncture in the third stage of labour [Dit Bedeutung der Akupunktur in der Plazentarperiode]. *Deutsche Zeitschrift fur Akupunktur* 2000;**43**:264-8.

Bader W, Ast S, Reinehr J, Hackmann J, Hatzmann W. Oxytocin versus acupuncture in the third stage of labour - a prospective randomized study [Oxytocin versus Akupunktur in der Plazentarperiode - eine prospektiv randomisierte Studie]. *Geburtshilfe und Frauenheilkunde* 2000;**60 Suppl 1**:S73.

Badhwar 1991 {published data only}

Badhwar L, Singh K, Sethi N, Gupta I, Aggarwal N. The value of nipple stimulation in the management of third stage of labour. *International Journal of Gynecology & Obstetrics* 1991;**36 Suppl**:16.

Bai 2014 (published data only)

Bai J, Sun Q, Zhai H. A comparison of oxytocin and carboprost tromethamine in the prevention of postpartum hemorrhage in high-risk patients undergoing cesarean delivery. *Experimental and Therapeutic Medicine* 2014;**7**(1):46-50.

Baig 2015 {published data only}

Baig FS, Shahzad N, Khurshid HN, Malik A. Postpartum haemorrhage; comparison of intra umbilical and intra venous injection of oxytocin on blood loss in third stage of labour. *Professional Medical Journal* 2015;**22**(6):793-7.

Balki 2006 (published data only)

* Balki M, Ronayne M, Davies S, Fallah S, Kingdom J, Windrim R, et al. Minimum oxytocin dose requirement after cesarean delivery for labor arrest. *Obstetrics & Gynecology* 2006;**107**(1):45-50.

Balki M, Ronayne M, Davies S, Kingdom J, Windrim R, Carvalho J. Oxytocin requirements at cesarean section for failure to progress in labor: a dose-finding study [abstract]. *Anesthesiology* 2005;**102**(Suppl 1):10.

Banovska 2013 (published data only)

Banovska J, Goffard P, Suball M, Origer P, Delatte P, Kapessidou P. Efficiency of temporary balloon occlusion of iliac arteries in patients at high hemorrhagic risk undergoing cesarean section. *European Journal of Anaesthesiology* 2013;**30**:176.

Barbaro 1961 {published data only}

Barbaro CA, Smith GO. Clinical trial of SE505 - a new oxytocic mixture. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 1961;**1**:147-50.

Baumgarten 1983 {published data only}

Baumgarten K, Schmidt J, Horvat A, Neumann M, Cerwenka R, Gruber W, et al. Uterine motility after post-partum application of

sulprostone and other oxytocics. *European Journal of Obstetrics* & *Gynecology and Reproductive Biology* 1983;**16**:181-92.

Bhattacharya 1988 {published data only}

* Bhattacharya P, Devi PK, Jain S, Kanthamani CR, Raghavan KS. Prophylactic use of 15(S)15 methyl PGF2alpha by intramuscular route for control of postpartum bleeding - a comparative trial with methylergometrine. *Acta Obstetricia et Gynecologica Scandinavica Supplement* 1988;**145**:13-5.

Devi PK, Sutaria UD, Raghavan KS. Prophylactic use of 15(S)15 methyl PGF2alpha for control of postpartum bleeding. *Acta Obstetricia et Gynecologica Scandinavica Supplementa* 1988;**145**:7-8.

Bhavana 2013 (published data only)

Bhavana G, Mittal S. Evaluation of efficacy of prophylactic injection tranexamic acid in decreasing blood loss before and after caesarean section. *BJOG: an international journal of obstetrics and gynaecology* 2013;**120**(Suppl s1):32.

Bider 1991 (published data only)

Bider D, Menashe Y, Dulitzky M, Mashiach S, Ben-Rafael Z. Oxytocin or saline injected intra-umbilically did not influence the third stage of labor. *Acta Obstetricia et Gynecologica Scandinavica* 1991;**70**:321-3.

Bider 1992 {published data only}

Bider D, Ben-Rafael Z, Dulitzky M, Menashe Y, Mashiach S, Barkai G. Effect of intraumbilical prostaglandin F2alpha injection on the third stage of labor. *Journal of Reproductive Medicine* 1992;**37**(4):317-9.

Bisri 2011 {published data only}

Bisri Y, Redjeki IS, Himendra A. The comparative of effect of bolus-infusion oxytocine with infusion oxytocine on blood pressure, heart rate, and uterine contraction of women undergoing elective caesarean section with general anesthesia N2O-sevoflurane. *European Journal of Anaesthesiology* 2011;**28 Suppl**:159.

Bivins 1993 {published data only}

* Bivins HA, Cope DA, Newman RB, Eller DP. Randomized trial of intraumbilical vein oxytocin. *American Journal of Obstetrics and Gynecology* 1993;**168**:435.

Bivins HA, Cope DA, Newman RB, Eller DP. Randomized trial of intraumbilical vein oxytocin in midtrimester pregnancy losses. *American Journal of Obstetrics and Gynecology* 1993;**169**(4):1070-3.

Blum 2010 (published data only)

Blum J, Winikoff B, Raghavan S, Dabash R, Ramadan MC, Dilbaz B, et al. Treatment of post-partum haemorrhage with sublingual misoprostol versus oxytocin in women receiving prophylactic oxytocin: a double-blind, randomised, non-inferiority trial. *Lancet* 2010;**375**(9710):217-23.

Bonham 1963 {published data only}

Bonham DG. Intramuscular oxytocics and cord traction in third stage of labour. *BMJ* 1963;**2**:1620-3.



Bonis 2012 (published data only)

Bonis M, Torricelli M, Leoni L, Berti P, Ciani V, Puzzutiello R, et al. Carbetocin versus oxytocin after caesarean section: similar efficacy but reduced pain perception in women with high risk of postpartum haemorrhage. *Journal of Maternal-Fetal & Neonatal Medicine* 2012;**25**(6):732-5.

Boopathi 2014 (published data only)

Boopathi A, Nayak SR, Rao A, Rao B. Oxytocin versus methylergometrine in the active management of third stage of labour. *Open Journal of Obstetrics and Gynecology* 2014;**4**:666-71.

Bose 2017 (published data only)

Bose D, Beegum R. Sublingual misoprostol vs intravenous tranexamic acid in reducing blood loss during cesarean section: a prospective randomized study. *Journal of South Asian Federation of Obstetrics and Gynaecology - SAFOG* 2017;**9**(1):9-13.

Bulusu 2017 (published data only)

Bulusu R, Ray P, Rani A, Handa P. Comparison of side effects of misoprostol by oral and rectal routes in active management of third stage of labour. *Journal of Evidence Based Medicine and Healthcare* 2017;**4**(3):146-9.

Cappiello 2006 (published data only)

Cappiello E, Lugo L, Kodali B, Hepner D, Harnett M, Tsen LC. A double-blinded, randomized, placebo-controlled trial of calcium chloride for the augmentation of uterine tone following cesarean delivery [abstract]. *Anesthesiology* 2006;**104**(Suppl 1):32.

Carvalho 2004 (published data only)

Carvalho JC, Balki M, Kingdom J, Windrim R. Oxytocin requirements at elective cesarean delivery: A dose finding study. *Obstetrics & Gynecology* 2004;**104**:1005-10.

Catanzarite 1990 {published data only}

Catanzarite VA. Prophylactic intramyometrial carboprost tromethamine does not substantially reduce blood loss relative to intramyometrial oxytocin at routine cesarean section. *American Journal of Perinatology* 1990;**7**:39-42.

Chaplin 2009 {published data only}

Chaplin AC, George RB, McKeen D, McLeod LC. Up-down determination of the ED90 of oxytocin infusions for the prevention of postpartum uterine atony in parturients undergoing an elective caesarean delivery. *Canadian Journal of Anaesthesia* 2009;**56**(Suppl 1):S62.

Chatterjee 2016 {published data only}

Chatterjee S, Sarkar A, Rao KD. Using misoprostol for primary versus secondary prevention of postpartum haemorrhage - Do costs matter?. *PLOS One* 2016;**11**(10):e0164718.

Chaudhuri 2014 (published data only)

Chaudhuri P, Mandi S, Mazumdar A. Rectally administrated misoprostol as an alternative to intravenous oxytocin infusion for preventing post-partum hemorrhage after cesarean

delivery. *Journal of Obstetrics and Gynaecology Research* 2014:**40**(9):2023-30.

Chestnut 1987 {published data only}

Chestnut DH, Wilcox LL. Influence of umbilical vein administration of oxytocin on the third stage of labor. Proceedings of 19th Annual Meeting of Society for Obstetric Anesthesia and Perinatology; 1987 May 20-23; Halifax, Nova Scotia, Canada. 1987:49.

* Chestnut DH, Wilcox LL. Influence of umbilical vein administration of oxytocin on the third stage of labor: a randomized, double-blind, placebo-controlled study. *American Journal of Obstetrics and Gynecology* 1987;**157**:160-2.

Chou 1994 (published data only)

Chou MM, MacKenzie IZ. A prospective, double-blind, randomized comparison of prophylactic intramyometrial 15-methyl prostaglandin F2, 125 micrograms, and intravenous oxytocin, 20 units, for the control of blood loss at elective cesarean section. *American Journal of Obstetrics and Gynecology* 1994;**171**:1356-60.

Chou 2015 {published data only}

Chou LT, Da AW, Murizah MZ, Rushdan M, Rashid Z. A randomised controlled trial on low dose versus high dose oxytocin infusion in prevention of uterine atony at caesarean delivery. *Journal of Obstetrics and Gynaecology Research* 2015;**41**(Suppl S1):44-5, Abstract no: FC 8.11.

Chukudebelu 1963 {published data only}

Chukudebelu WO, Marshall AT, Chalmers JA. Use of 'syntometrine' in the third stage of labour. *BMJ* 1963;**1**:1390-1.

Cooper 2004 (published data only)

ISRCTN07452238. A study to determine the cardiovascular effects of different methods of administering the oxytocic drug syntocinon. isrctn.com/ISRCTN07452238 (first received: 12 September 2003).

Cordovani 2011 {published data only}

Cordovani D, Farine D, Balki M, Seaward G, Carvalho JC. Carbetocin at elective cesarean delivery: A dose-finding study. Canadian Journal of Anesthesia 2011;**58**(Suppl 1):S90.

Cordovani 2012 {published data only}

Cordovani D, Balki M, Farine D, Seaward G, Carvalho JC. Carbetocin at elective cesarean delivery: a randomized controlled trial to determine the effective dose. *Canadian Journal of Anesthesia* 2012;**59**(8):751-7.

Dagdeviren 2016 {published data only}

* Dagdeviren H, Cengiz H, Heydarova U, Caypinar SS, Kanawati A, Guven E, et al. Intramuscular versus intravenous prophylactic oxytocin for postpartum hemorrhage after vaginal delivery: a randomized controlled study. *Archives of Gynecology* and Obstetrics 2016;**294**(5):911-6.

NCT02080104. Intramuscular versus intravenous prophylactic oxytocin for hemorrhage after vaginal delivery (oxytocin). clinicaltrials.gov/ct2/show/NCT02080104 (first received: 1 March 2014).



Dahiya 1995 (published data only)

Dahiya P, Puri M, Rathee S. Influence of intraumbilical oxytocin on the third stage of labour. *Indian Journal of Medical Science* 1995;**49**:23-7.

Daley 1951 {published data only}

Daley D. The use of intramuscular ergometrine at the end of the second stage of normal labour. *Journal of Obstetrics and Gynaecology of the British Empire* 1951;**57**:388-97.

Daly 1999 (published data only)

Daly S, Andolina K, Tolosa JE, Roberts N, Wapner R. A randomized controlled trial of misoprostol versus oxytocin in preventing postpartum blood loss. *American Journal of Obstetrics and Gynecology* 1999;**180**(1 Pt 2):S68.

Dao 2009 {published data only}

Dao B, Blum J, Barrera G, Cherine Ramadan M, Dabash R, Darwish E, et al. Side effect profiles for misoprostol and oxytocin in the treatment of postpartum hemorrhage. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S150.

Davies 2005 (published data only)

Davies GA, Tessier JL, Woodman MC, Lipson A, Hahn PM. Maternal hemodynamics after oxytocin bolus compared with infusion in the third stage of labor: a randomized controlled trial. *Obstetrics & Gynecology* 2005;**105**:294-9.

De bonis 2012 {published data only}

* De Bonis M, Torricelli M, Leoni L, Berti P, Ciani V, Puzzutiello R, et al. Carbetocin versus oxytocin after caesarean section: similar efficacy but reduced pain perception in women with high risk of postpartum haemorrhage. *Journal of Maternal-Fetal & Neonatal Medicine* 2012;**25**(6):732-5.

Voltolini C, De Bonis M, Vellucci F, Regini C, Orlandini C, Vannuccini S, et al. Carbetocin versus oxytocin after caesarean section: Similar efficacy but reduced pain perception in women with high risk of postpartum haemorrhage. *International Journal of Gynecology and Obstetrics* 2012;**119**:S806-7.

Dell-Kuster 2017 {published data only}

Dell-Kuster S, Hoesli I, Lapaire O, Seeberger E, Steiner LA, Bucher HC, et al. Efficacy and safety of carbetocin applied as an intravenous bolus compared to as a short-infusion for caesarean section: study protocol for a randomised controlled trial. *Trials* 2016;**17**(1):155.

* Dell-Kuster S, Hoesli I, Lapaire O, Seeberger E, Steiner LA, Bucher HC, et al. Efficacy and safety of carbetocin given as an intravenous bolus compared with short infusion for caesarean section - double-blind, double-dummy, randomized controlled non-inferiority trial. *British Journal of Anaesthesia* 2017;**118**(5):772-80.

Dell-Kuster S, Hoesli I, Lapaire O, Seeberger E, Steiner LA, Bucher HC, et al. Efficacy and safety of intravenous carbetocin as a bolus compared to a short infusion for caesarean section. *Journal of Obstetric Anesthesia* 2016;**26**(Suppl 1):S7.

Dennehy 1998 {published data only}

Dennehy KC, Rosaeg OP, Cicutti NJ, Krepski B, Sylvain JP. Oxytocin injection after caesarean delivery: intravenous or intramyometrial?. *Canadian Journal of Anaesthesia* 1998;**45**(7):635-9.

Deshpande 2016 {published data only}

Deshpande HG, Madkar CS, Patel KK. Comparative study between intravenous and intraumbilical oxytocin as active management of third stage in elective and emergency caesarean section. *Indian Journal of Obstetrics and Gynaecology Research* 2016;**3**(1):55-8.

Diab 1999 {published data only}

Diab KM, Ramy AR, Yehia MA. The use of rectal misoprostol as active pharmacological management of the third stage of labor. *Journal of Obstetrics and Gynaecology Research* 1999;**25**(5):327-32.

Dickinson 2009 (published data only)

Dickinson JE, Doherty DA. Optimization of third-stage management after second-trimester medical pregnancy termination. *American Journal of Obstetrics & Gynecology* 2009;**201**(3):303.e1-303.e7.

Diop 2011 {published data only}

NCT01487278. Comparing misoprostol and oxytocin in UnijectTM postpartum hemorrhage (PPH) prevention in Mali. clinicaltrials.gov/ct2/show/NCT01487278 Date first received: 5 December 2011.

Dommisse 1980 {published data only}

Dommisse J. The routine use of oxytocic drugs in the third stage of labour [letter]. South African Medical Journal 1980;**46**:549.

Dong 2011 (published data only)

Dong Y. Effects of carboprost on prevention of hemorrhage after induced labor with scarred uterus. *Journal of Shanghai Jiaotong University (Medical Science)* 2011;**31**(8):1212-5.

Dumoulin 1981 {published data only}

Dumoulin JG. A reappraisal of the use of ergometrine. *Journal of Obstetrics and Gynaecology* 1981;**1**:178-81.

Durocher 2012 {published data only}

Durocher J, Blum J, Sheldon WR, Trussell J, Winikoff B. Does the effect of oxytocin prophylaxis on post-partum blood loss depend on route of administration?. *International Journal of Gynecology & Obstetrics* 2012;**119**(Suppl 3):S332.

Dutta 2000 {published data only}

Dutta DK, Saha KK. Comparative study on role of syntometrine and prostaglandin in the prevention of PPH. XVI FIGO World Congress of Obstetrics & Gynecology (Book 4); 2000 Sept 3-8; Washington DC, USA. 2000:29.

Dweck 2000 {published data only}

Dweck MF, Lynch CM, Spellacy WN. Use of methergine for the prevention of postoperative endometritis in non-elective cesarean section patients. *Infectious Diseases in Obstetrics & Gynecology* 2000;**8**:151-4.



Dzuba 2012 (published data only)

Dzuba I, Durocher J, Dilbaz B, Gelisen O, Ngoc NT, Montesinos R, et al. Route of administration of oxytocin in prevention of postpartum hemorrhage. *International Journal of Gynecology & Obstetrics* 2012;**119**(Suppl 3):S333.

Elati 2011 (published data only)

Elati A, Elmahaishi MS, Elmahaishi MO, Elsraiti OA, Weeks AD. The effect of misoprostol on postpartum contractions: a randomised comparison of three sublingual doses. *BJOG:* an international journal of obstetrics and gynaecology 2011;**118**(4):466-73.

Erkkola 1984 {published data only}

Erkkola R, Kero P, Kanto J, Korvenranta H, Nanto V, Peltonen T. Delayed cord clamping in cesarean section with general anesthesia. *American Journal of Perinatology* 1984;**1**(2):165-9.

Farber 2013 (published data only)

NCT02026297. Tranexamic acid and thromboelastography during cesarean delivery (TA TEG). clinicaltrials.gov/ct2/show/NCT02026297 (first received: 22 December 2013).

Farber 2015 (published data only)

Farber MK, Schultz R, Lugo L, Liu X, Huang C, Tsen LC. The effect of co-administration of intravenous calcium chloride and oxytocin on maternal hemodynamics and uterine tone following cesarean delivery: a double-blinded, randomized, placebo-controlled trial. *International Journal of Obstetric Anesthesia* 2015;**24**(3):217-24.

Fatemeh 2011 {published data only}

Fatemeh F, Zohreh S, Abbas MG, Layla H. Maternal haemodynamic effects of oxytocin bolus or infusion in the third stage of labour. *Pakistan Journal of Medical Sciences* 2011;**27**(3):656-9.

Forster 1957 {published data only}

Forster FM. A comparative study of ergometrine and 'methergin' used in the management of the third stage of labour. *Medical Journal of Australia* 1957;**2**:155-6.

Francis 1965 (published data only)

Francis HH, Miller JM, Porteous CR. Clinical trial of an oxytocinergometrine mixture (first of two trials). *Australian and New Zealand Journal of Obstetrics and Gynaecology* 1965;**5**:47-51.

Friedman 1957 {published data only}

Friedman EA. Comparative clinical evaluation of postpartum oxytocics. *American Journal of Obstetrics and Gynecology* 1957;**73**:1306-13.

Fugo 1958 (published data only)

Fugo NW, Dieckmann WJ. A comparison of oxytocic drugs in the management of the placental stage. *American Journal of Obstetrics and Gynecology* 1958;**76**:141-6.

Gai 2004 (published data only)

Gai MY, Wu LF, Su QF, Tatsumoto K. Clinical observation of blood loss reduced by tranexamic acid during and after caesarian section: a multi-center, randomized trial. *European* Journal of Obstetrics & Gynecology and Reproductive Biology 2004;**112**:154-7.

Gavhane 2017 {published data only}

Gavhane Satyajit P, Harshad T, Bangal VB, Verma P, Bhavsar Dhruval K. Comparative outcome of active management of third stage of labour with prophylactic use of oxytocin, methyl ergometrine and misoprostol. *Pravara Medical Review* 2017;**9**(4):4-9.

George 2010 (published data only)

George RB, McKeen D, Chaplin AC, McLeod L. Up-down determination of the ED(90) of oxytocin infusions for the prevention of postpartum uterine atony in parturients undergoing Cesarean delivery. *Canadian Journal of Anaesthesia* 2010;**57**(6):578-82.

Ghulmiyyah 2007 {published data only}

Ghulmiyyah LM, Wehbe SA, Saltzman SL, Ehleban C, Sibai BM. Effects of intraumbilical vein injection of saline versus oxytocin plus saline on duration of the third stage of labor: a randomized double-blind placebo trial [abstract]. *American Journal of Obstetrics and Gynecology* 2005;**193**(6 Suppl):S18.

* Ghulmiyyah LM, Wehbe SA, Saltzman SL, Ehleben C, Sibai BM. Intraumbilical vein injection of oxytocin and the third stage of labor: randomized double-blind placebo trial. *American Journal of Perinatology* 2007;**24**(6):347-52.

Ghulmiyyah 2017 {published data only}

Ghulmiyyah LM, Usta IM, Ghazeeri G, Taher N, Abu-Ghannam G, Nassar AH, et al. Intravenous oxytocin use to decrease blood loss during scheduled cesarean delivery: a randomized double-blinded controlled trial (oxytrial). *American Journal of Perinatology* 2017;**34**(4):379-87.

Gobbur 2011 {published data only}

Gobbur VR, Reddy SV, Bijapur UJ. Efficacy of tranexamic acid in reducing blood loss during lower segment caesarean section. 54th All India Congress of Obstetrics and Gynaecology; 2011 January 5-9; Hyderabad, Andhra Pradesh, India. 2011:92.

Gohel 2007 {published data only}

Gohel M, Patel P, Gupta A, Desai P. Efficacy of tranexamic acid in decreasing blood loss during and after cesarean section: a randomized case controlled prospective study. *Journal of Obstetrics and Gynaecology of India* 2007;**57**(3):228-30.

Goswami 2013 (published data only)

Goswami U, Sarangi S, Gupta S, Babbar S. Comparative evaluation of two doses of tranexamic acid used prophylactically in anemic parturients for lower segment cesarean section: a double-blind randomized case control prospective trial. *Saudi Journal of Anaesthesia* 2013;**7**(4):427-31.

Groeber 1960 {published data only}

Groeber WR, Bishop EH. Methergine and ergonovine in the third stage of labor. *Obstetrics & Gynecology* 1960;**15**:85-8.



Gungorduk 2010 (published data only)

Gungorduk K, Asicioglu O, Besimoglu B, Gungorduk OC, Yildirm G, Ark C, et al. Using intraumbilical vein injection of oxytocin in routine practice with active management of the third stage of labor: a randomized controlled trial. *Obstetrics & Gynecology* 2010;**116**(3):619-24.

Gungorduk 2010b {published data only}

Gungorduk K, Asicioglu O, Celikkol O, Olgac Y, Ark C. Use of additional oxytocin to reduce blood loss at elective caesarean section: a randomised control trial. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 2010;**50**(1):36-9.

Gungorduk 2011 {published data only}

Gungorduk K, Yildirim G, Asicioglu O, Gungorduk OC, Sudolmus S, Ark C. Efficacy of intravenous tranexamic acid in reducing blood loss after elective cesarean section: a prospective, randomized, double-blind, placebo-controlled study. *American Journal of Perinatology* 2011;**28**(3):233-40.

Gungorduk 2013 (published data only)

Gungorduk K, Asicioglu O, Yildirim G, Ark C, Tekirdag AI, Besimoglu B. Can intravenous injection of tranexamic acid be used in routine practice with active management of the third stage of labor in vaginal delivery? A randomized controlled study. *American Journal of Perinatology* 2013;**30**(5):407-13.

Gupta 2014 {published data only}

Gupta M, Bhosale U. Comparative study of methylergometrine and low dose carboprost (PGF2-x) in active management of 3rd stage labor. *BJOG: an international journal of obstetrics and gynaecology* 2014;**121**(Suppl 2):139.

Habek 2007 {published data only}

Habek D, Franicevic D. Intraumbilical injection of uterotonics for retained placenta. *International Journal of Gynecology & Obstetrics* 2007;**99**(2):105-9.

Hacker 1979 {published data only}

Hacker NF, Biggs JS. Blood pressure changes when uterine stimulants are used after normal delivery. *British Journal of Obstetrics and Gynaecology* 1979;**86**:633-6.

Häivä 1994 (published data only)

Häivä L, Hartikainen A. Pharmacological management of third stage of labor in primiparae and multiparae [Kohtua supistavan lääkkeen valinta ensi- ja uudelleensynnyttäjille]. Suomen Lääkärilehti 1994;**49**(33):3442-4.

Halder 2013 {published data only}

Halder S, Samanta B, Sardar R, Chattopadhyay S. Tranexamic acid used before caesarean section reduces blood loss based on pre- and postoperative hemoglobin level: A case-control study. *Journal of the Indian Medical Association* 2013;**111**(3):184-6.

Hoffman 2006 {published data only}

* Hoffman M, Castagnola D, Naqvi F. A randomized trial of active versus expectant management of the third stage of labor [abstract]. *American Journal of Obstetrics and Gynecology* 2006;**195**(6 Suppl 1):S107.

Hoffman M, Naqvi F, Sciscione A. A randomized trial of active versus expectant management of the third stage of labor [abstract]. *American Journal of Obstetrics and Gynecology* 2004;**191**(6 Suppl 1):S82.

Hofmeyr 2004 {published data only}

Hofmeyr GJ, Ferreira S, Nikodem VC, Mangesi L, Singata M, Jafta Z, et al. Misoprostol for treating postpartum haemorrhage: a randomized controlled trial. *BMC Pregnancy and Childbirth* 2004;**4**(1):16.

Howard 1964 (published data only)

Howard WF, McFadden PR, Keettel WC. Oxytocic drugs in fourth stage of labor. *Journal of the American Medical Association* 1964;**189**:411-3.

Huh 2004 {published data only}

Huh W, Chelmow D, Malone FD. A randomized, double-blinded, placebo controlled trial of oxytocin at the beginning versus the end of the third stage of labor for prevention of postpartum hemorrhage. *American Journal of Obstetrics and Gynecology* 2000;**182**(1 Pt 2):S130.

* Huh WK, Chelmow D, Malone FD. A double-blinded, randomized, controlled trial of oxytocin at the beginning versus the end of the third stage of labour for prevention of postpartum hemorrhage. *Gynecologic and Obstetric Investigation* 2004;**58**(2):72-6.

Hunt 2013 {published data only}

Hunt BJ. Tranexamic acid for the treatment of postpartum haemorrhage-preliminary results of the woman trial. *Transfusion Medicine* 2013;**23**(Suppl 1):7.

Ilancheran 1990 {published data only}

Ilancheran A, Ratnam SS. Effect of oxytocics on prostaglandin levels in the third stage of labour. *Gynecologic and Obstetric Investigation* 1990;**29**:177-80.

Irons 1994 {published data only}

Irons DW, Sriskandabalan P, Bullough CH. A simple alternative to parenteral oxytocics for the third stage of labor. *International Journal of Gynecology & Obstetrics* 1994;**46**:15-8.

Islam 2008 {published data only}

Islam A, Siraj A, Arif N. Post partum hemorrhage prophylaxis; comparison of the efficacy of misoprostol and ergometrine in cesarean delivery. *Professional Medical Journal* 2008;**15**(3):323-7.

Jackson 2001 {published data only}

Jackson KW, Allbert JR, Schemmer GK, Elliot M, Humphrey A, Taylor J. A randomized controlled trial comparing oxytocin administration before and after placental delivery in the prevention of postpartum hemorrhage. *American Journal of Obstetrics and Gynecology* 2001;**185**:873-7.

Jagielska 2015 (published data only)

Jagielska I, Kazdepka-Zieminska A, Kaczorowska A, Madej A, Kolossa T, Grabiec M. [Evaluation of carbetocin and oxytocin efficacy in prevention of postpartum hemorrhage in



women after cesarean section]. [Polish]. *Ginekologia Polska* 2015:**86**(9):689-93.

Javadi 2015 (published data only)

Javadi EH, Sadeghipour Z, Barikani A, Javadi M. Tranexamic acid in the control of uterine atony during labor. *Biotechnology and Health Sciences* 2015;**2**(2):e26898.

Jiang 2001 (published data only)

Jiang Q, Wang P, Cao W. Effect on different doses of misoprostol to prevent postpartum hemorrhage. *Chinese Nursing Research* 2001;**15**(6):313-4.

Jin 2000 {published data only}

Jin LJ, Zhou L. Application of anus misoprostol to decrease the volume of post partum hemorrhage. *Journal of Practical Nursing* 2000;**16**(2):9-10.

Jolivet 1978 (published data only)

Jolivet A, Robyn C, Huraux-Rendu C, Gautray JP. Effect of ergot alkaloid derivatives on milk secretion in the immediate postpartum period. *Journal de Gynecologie, Obstetrique et Biologie de la Reproduction* 1978;**7**:129-34.

Jonsson 2010 {published data only}

* Jonsson M, Hanson U, Lidell C, Norden-Lindeberg S. ST depression at caesarean section and the relation to oxytocin dose. A randomised controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2010;**117**(1):76-83.

Jonsson M, Norden Lindeberg S, Hanson U. ST depression at caesarean section and the relation to oxytocin dose. A randomised controlled trial. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S214.

Kashanian 2010 (published data only)

Kashanian M, Fekrat M, Masoomi Z, Sheikh Ansari N. Comparison of active and expectant management on the duration of the third stage of labour and the amount of blood loss during the third and fourth stages of labour:a randomised controlled trial. *Midwifery* 2010;**26**:241-5.

Kemp 1963 {published data only}

Kemp J. Clinical trial of "syntometrine" in the third stage of labour. *British Medical Journal* 1963;**1**(5342):1391-2.

Khan 1997 {published data only}

Khan GQ, John IS, Wani S, Doherty T, Sibai BM. Controlled cord traction versus minimal intervention techniques in delivery of the placenta: a randomized controlled trial. *American Journal of Obstetrics and Gynecology* 1997;**177**(4):770-4.

Khan 2003 {published data only}

Khan RU, El-Refaey H. Pharmacokinetics and adverse-effect profile of rectally administered misoprostol in the third stage of labor. *Obstetrics & Gynecology* 2003;**101**(5 Pt 1):968-74.

Khan 2012 (published data only)

Khan MS, Sinha SK, Sultana T, Singhal S. Comparison of two oxytocin infusions in patients undergoing emergency cesarean sections: A double blind study. *International Journal of Gynecology and Obstetrics* 2012;**119**(Suppl 3):S389.

Khan 2013 (published data only)

Khan M, Balki M, Ahmed I, Farine D, Searward G, Carvalho JC. Carbetocin at elective cesarean delivery: a randomized controlled trial to determine the effective dose, part 3 final. Society for Obstetric Anesthesia and Perinatology (SOAP) 45th Annual Meeting; 2013 April 24-28; San Juan, Puerto Rico. 2013:Abstract no: GM 3.

Khanun 2011 {published data only}

Khanun A, Khanum S. Oral versus rectal misoprostol in the prevention of primary postpartum hemorrage. *Pakistan Journal of Medical and Health Sciences* 2011;**5**(3):587-8.

Kikutani 2003a {published data only}

Kikutani T, Shimada Y. Effects of methylergometrine and oxytocin on thoracic epidural pressure during cesarean section. *Journal of Obstetrics and Gynaecology Research* 2003;**29**(3):180-5.

Kikutani 2003b {published data only}

Kikutani T, Oshima M, Sugimoto K, Shimada Y. Effects of intravenous infusion rate of oxytocin on thoracic epidural pressure in parturients undergoing elective cesarean section. Journal of Nippon Medical School = Nihon Ika Daigahu Zasshi 2003;**70**(6):475-9.

Kikutani 2006 (published data only)

Kikutani T, Kikutani M, Oshima M, Sugimoto K, Shimada Y. Effects of methylergometrine and oxytocin on blood loss and uterine contraction during cesarean section. *Masui - Japanese Journal of Anesthesiology* 2006;**55**(5):590-4.

King 2010 {published data only}

King KJ, Douglas J, Unger W, Wong AB. A randomized double-blind comparison of a 5 unit intravenous oxytocin bolus versus placebo as a strategy to prevent uterine atony at cesarean section in women who are at increased risk of post-partum hemorrhage [abstract]. *Anesthesiology* 2006;**104**(Suppl 1):41.

King KJ, Douglas J, Unger W, Wong AB, Espinosa V, King RA. 5U bolus oxytocin at cesarean section in women at risk of atony [abstract]. *Anesthesiology* 2007;**106**(Suppl 1):14.

* King KJ, Douglas MJ, Unger W, Wong A, King RA. Five unit bolus oxytocin at cesarean delivery in women at risk of atony: a randomized, double-blind, controlled trial. *Anesthesia & Analgesia* 2010;**111**(6):1460-6.

Kintu 2012 {published data only}

Kintu A, Nakubulwa S, Mijumbi C, Kwizera A, Tindimwebwa J. Uterotonic efficacy of oxytocin 2.5 versus 10 units during caesarean section at mulago hospital: a double blinded placebo controlled randomised clinical trial. *British Journal of Anaesthesia* 2012;**108**:ii197-8.

Kiran 2012 (published data only)

Kiran S, Anand A, Singh T, Gupta N. Effective dose of oxytocin in caesarean delivery. *British Journal of Anaesthesia* 2012;**108**:ii195.



Kore 2000 {published data only}

Kore S, Srikrishna S, Hegde A, Ambiye VR, Vaidya PR. Active management of third stage of labour with intraumbilical oxytocin injection. *Journal of Obstetrics and Gynecology of India* 2000;**50**(3):54-5.

Kovacheva 2015 (published data only)

Kovacheva VP, Soens MA, Tsen LC. A randomized, double-blinded trial of a "rule of threes" algorithm versus continuous infusion of oxytocin during elective cesarean delivery. Anesthesiology 2015;**123**(1):92-100.

Kovavisarach 1998 {published data only}

Kovavisarach E, Rojsangruang S. Effect of umbilical vein oxytocin injection on the third stage of labor: a randomized controlled study. 9th Congress of the Federation of the Asia and Oceania Perinatal Societies; 1996 November 10-14; Singapore. 1996:59.

* Kovavisarach E, Rojsangruang S. Effect of umbilical vein oxytocin injection on the third stage of labor: a randomized controlled study. *Journal of the Medical Association of Thailand* 1998;**81**:693-7.

Le 2000 {published data only}

Le J. Prevention of postpartum hemorrhage by carboprost and oxytocin in 90 cases analysis. *Acta Medicinae Sinica* 2000;**13**(2):140-1.

Leader 2002 {published data only}

Leader J, Bujnovsky M, Carlan SJ, Triana T, Richichi K. Effect of oral misoprostol after second-trimester delivery: a randomized, blinded study. *Obstetrics & Gynecology* 2002;**100**(4):689-94.

Li 2002 {published data only}

Li X, Wang H, Wang J, Cao X L, Ma Y. Prophylactic and therapeutic effect of misoprofil plus oxytocin on postpartum hemorrhage in patients with pregnancy-induced hypertension syndrome. *Journal of Postgraduates of Medicine* 2002;**25**(7):34-5.

Li 2003 {published data only}

Li DP, Bei HZ. Clinical study on reduction of postpartum bleeding in the risk factors by misoprostol. *Hainan Medical Journal* 2003;**14**(11):11-2.

Li 2011 {published data only}

* Li H, Afzal A, Lian Q, Kramer GC, Svenson C, Prough D. Restricted fluid therapy decreases surgical blood loss - a clinical study of two fluid regimens during cesarean section under spinal anesthesia. *Anesthesia & Analgesia* 2011;**112**:S-291.

Li H, Simon M, Lian Q, Afzal A, Christer Svenson C, Prough D. Restricted fluid therapy decreases surgical blood loss - A clinical study of two fluid regimens during cesarean section under spinal anesthesia. American Society of Anesthesiologists Annual Meeting; 2011, October 15-19; Chicago, Illinois. 2011.

Lin 2009 {published data only}

Lin JH, Lin QD, Liu XH, Yan JY, He J, Li L, et al. [Multi-center study of motherwort injection to prevent postpartum hemorrhage

after caesarian section]. [Chinese]. Chung-Hua Fu Chan Ko Tsa Chih [Chinese Journal of Obstetrics & Gynecology] 2009;**44**(3):175-8.

Liu 1997 {published data only}

Liu C, Wang D, Li X. Clinical study on reduction of postpartum bleeding by methyl carprost suppository. *Chung-Hua Fu Chan Ko Tsa Chih/Chinese Journal of Obstetrics and Gynaecology* 1997;**32**(1):22-4.

Liu 2002 (published data only)

Liu DY, Fan L, Huang XH. Clinical observation on treatment of postpartum hemorrhage by xuesaitong soft capsule. *Chinese Journal of Integrated Traditional and Western Medicine* 2002;**22**(3):182-4.

Liu 2015 {published data only}

Liu Y, Chen HX, Kang DL, Kuang XH, Liu WX, Ni J. Influence of dexmedetomidine on incidence of adverse reactions introduced by hemabate in postpartum hemorrhage during cesarean section. *International Journal of Clinical and Experimental Medicine* 2015;**8**(8):13776-82.

Liu 2016 {published data only}

Liu W, Ma S, Pan W, Tan W. Combination of motherwort injection and oxytocin for the prevention of postpartum hemorrhage after cesarean section. *Journal of Maternal-Fetal & Neonatal Medicine* 2016;**29**(51):2489-92.

Luamprapas 1994 {published data only}

Luamprapas A. A study of umbilical vein administration of oxytocin to shorten the third stage of labor. *Chon Buri Hospital Journal* 1994;**19**(2):14-25.

Maged 2015 {published data only}

Maged AM, Helal OM, Elsherbini MM, Eid MM, Elkomy RO, Dahab S, et al. A randomized placebo-controlled trial of preoperative tranexamic acid among women undergoing elective cesarean delivery. *International Journal of Gynecology and Obstetrics* 2015;**131**:265-8.

Makvandi 2013 (published data only)

Makvandi S, Shoushtari SZ, Hosseini VZ. Management of third stage of labour: A comparison of intraumbilical oxytocin and placental cord drainage. *Shiraz E Medical Journal* 2013;**14**(2):83-90.

Mangla 2012 (published data only)

Mangla D, Goel JK, Goel R. Prophylactic intramyometrial oxytocin before placenta delivery during cesarean section prevents postpartum hemorrhage: a prospective randomized study of 150 women. *Journal of South Asian Federation of Obstetrics and Gynaecology* 2012;**4**(2):93-6.

Mankuta 2006 (published data only)

NCT00405626. Double blind placebo controlled bellis perenis and arnica montana as a drug for PPH. clinicaltrials.gov/ct2/show/NCT00405626 (first received: 29 November 2006).



Mansouri 2011 (published data only)

Mansouri HA, Alsahly N. Rectal versus oral misoprostol for active management of third stage of labor: a randomized controlled trial. *Archives of Gynecology and Obstetrics* 2011;**283**(5):935-9.

Martinez 2006 (published data only)

Martinez MM, Lopez Farfan JA, Ramos Alvarez G, Lopez Colombo A. Oxitocin trough umbilical vein to shorten the third stage of labor [Oxitocina transvena umbilical para acortar el tercer periodo de trabajo de parto]. *Ginecologia y Obstetricia de Mexico* 2006;**74**(2):89-94.

McGinty 1956 (published data only)

McGinty LB. A study of the vasopressor effects of oxytocics when used intravenously in the third stage of labour. *Western Journal of Surgery* 1956;**64**:22-8.

Miller 2009 (published data only)

Miller S, Tudor C, Thorsten V, Nyima, Kalyang, Sonam, et al. Randomized double masked trial of Zhi Byed 11, a Tibetan traditional medicine, versus misoprostol to prevent postpartum hemorrhage in Lhasa, Tibet. *Journal of Midwifery & Women's Health* 2009;**54**(2):133-41.

Mirghafourvand 2015 (published data only)

Mirghafourvand M, Alizadeh SM, Abasalizadeh F, Shirdel M. The effect of intravenous tranexamic acid on hemoglobin and hematocrit levels after vaginal delivery: a randomized controlled trial. *Iranian Journal of Obstetrics, Gynecology and Infertility* 2013;**16**(60):1-8.

* Mirghafourvand M, Mohammad-Alizadeh S, Abbasalizadeh F, Shirdel M. The effect of prophylactic intravenous tranexamic acid on blood loss after vaginal delivery in women at low risk of postpartum haemorrhage: a double-blind randomised controlled trial. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 2015;**55**(1):53-8.

Mirteimouri 2013 (published data only)

Mirteimouri M, Tara F, Teimouri B, Sakhavar N, Vaezi A. Efficacy of rectal misoprostol for prevention of postpartum hemorrhage. Iranian Journal of Pharmaceutical Research 2013; Vol. 12, issue 2:469-74.

Mockler 2015 {published data only}

Mockler JC, Malkoutzis V, Davis-Tuck M, Wallace EM. Oxytocin infusion at elective caesarean section: a double blind, randomised controlled trial. *Journal of Paediatrics and Child Health* 2015;**51**(Suppl 1):54.

Mohamadian 2013 {published data only}

Mohamadian S, Shorab NJ, Mirzakhani K. The effect of the timing of intramuscular oxytocin injection on maternal bleeding during the third stage of labour. *Journal of Midwifery & Reproductive Health* 2013;**1**(2):66-70.

Mollitt 2009 {published data only}

Mollitt C, Ssenoga A, Grassman C, Barclay PM. Randomised controlled trial comparing the effects of oxytocin i.v. bolus vs. oxytocin i.v. infusion on cardiac output during caesarean

section. *International Journal of Obstetric Anesthesia* 2009;**18**(Suppl 1):S11.

Moore 1956 (published data only)

Moore JH. Is methylergonovine tartrate superior to ergonovine maleate. *American Journal of Obstetrics and Gynecology* 1956;**71**:908-11.

Movafegh 2011 (published data only)

Movafegh A, Eslamian L, Dorabadi A. Effect of intravenous tranexamic acid administration on blood loss during and after cesarean delivery. *International Journal of Gynecology & Obstetrics* 2011;**115**(3):224-6.

Munishankarappa 2009 (published data only)

Munishankarappa B, McLeod GA, MacGregor H, Murphy D. Maternal haemodynamic at elective caesarean section following oxytocin 5-unit bolus and placebo infusion compared to oxytocin 5-unit bolus and 30-unit infusion. *International Journal of Obstetric Anesthesia* 2009;**18**(Suppl 1):S48.

Munn 2001 {published data only}

Munn MB, Owen J, Hauth J. Oxytocin regimens for the prevention of uterine atony at cesarean delivery [abstract]. *American Journal of Obstetrics and Gynecology* 2001;**184**(1):S14.

* Munn MB, Owen J, Vincent R, Wakefield M, Chestnut DH, Hauth JC. Comparison of two oxytocin regimens to prevent uterine atony at cesarean delivery: a randomized controlled trial. *Obstetrics & Gynecology* 2001;**98**(3):386-90.

Murphy 2009 {published data only}

ISRCTN17813715. A randomised controlled trial of oxytocin bolus versus oxytocin bolus and infusion for the control of blood loss at elective caesarean section. isrctn.com/ISRCTN17813715 (first received: 25 March 2008).

Murphy DJ, Carey M, Montgomery AA, Sheehan SR, the ECSSITSG. Study protocol. ECSSIT - Elective Caesarean Section Syntocinon Infusion Trial. A multi-centre randomised controlled trial of oxytocin (Syntocinon) 5 IU bolus and placebo infusion versus oxytocin 5 IU bolus and 40 IU infusion for the control of blood loss at elective caesarean section. *BMC Pregnancy and Childbirth* 2009;**9**:36.

* Murphy DJ, MacGregor H, Munishankar B, McLeod G. A randomised controlled trial of oxytocin 5IU and placebo infusion versus oxytocin 5IU and 30IU infusion for the control of blood loss at elective caesarean section--pilot study. *European Journal of Obstetrics & Gynecology and Reproductive Biology* 2009;**142**(1):30-3.

Murphy 2015 {published data only}

Murphy D, ISRCTN14718882. A study to compare the effectiveness of intravenous oxytocin with intramuscular oxytocin given at the third stage of labour at preventing bleeding at vaginal birth. isrctn.com/ISRCTN14718882 (first received 14 December 2015).

Nankali 2013 (published data only)

Nankali A, Keshavarzi F, Fakheri T, Zare S, Rezaei M, Daeichin S. Effect of intraumbilical vein oxytocin injection on third



stage of labor. *Taiwanese Journal of Obstetrics & Gynecology* 2013:**52**(1):57-60.

Narenji 2012 {published data only}

IRCT201206099981N1. Comparison the effect of intramuscular injection of oxytocin and nipple stimulation on the third stage of delivery length and bleeding. en.irct.ir/trial/10487? revision=10487 (first received 25 June 2012).

Nelson 1983 (published data only)

Nelson GH. Use of 15-methyl prostaglandin F2alpha postpartum to contract the uterus in normal pregnant women. *Journal of the Medical Association of Georgia* 1983;**72**:703-6.

Neri-Mejia 2016 {published data only}

Neri-Mejia M, Pedraza-Aviles AG. Active management of the third stage of labor: three schemes of oxytocin: randomised clinical trial. *Ginecologia y Obstetricia De Mexico* 2016;**84**(5):306-13.

Newton 1961 {published data only}

Newton M, Mosey LM, Egli GE, Gifford WB, Hull CT. Blood loss during and immediately after delivery. *Obstetrics & Gynecology* 1961;**17**:9-18.

Nguyen-Lu 2015 (published data only)

Nguyen-Lu N, Carvalho J, Farine D, Seaward G, Downey K, Balki M. Carbetocin at cesarean delivery for labor arrest: A randomised controlled trial to determine ED90. *Canadian Journal of Anesthesia* 2013;**60**(Suppl 1):S112.

* Nguyen-Lu N, Carvalho JC, Farine D, Seaward G, Ye XY, Balki M. Carbetocin at cesarean delivery for labour arrest: a sequential allocation trial to determine the effective dose. *Canadian Journal of Anaesthesia//Journal Canadien D'anesthesie* 2015;**62**(8):866-74.

Nieminen 1964 {published data only}

Nieminen U, Jaervinen PA. A comparative study of different medical treatments of the third stage of labour. *Annales Chirurgiae et Gynaecologiae Fenniae* 1964;**53**:424-9.

Oberbaum 2005 {published data only}

NCT01156194. Effect of a homeopathic remedy on the third stage of delivery: a prospective, randomized, double-blind study. clinicaltrials.gov/show/NCT01156194 (first received: 1 July 2010).

* Oberbaum M, Galoyan N, Lerner-Geva L, Singer SR, Grisaru S, Shashar D, et al. The effect of the homeopathic remedies arnica montana and bellis perennis on mild postpartum bleeding-a randomized, double-blind, placebo-controlled study-preliminary results. *Complementary Therapies in Medicine* 2005;**13**(2):87-90.

Oguz 2014 {published data only}

Oguz Orhan E, Dilbaz B, Aksakal SE, Altinbas S, Erkaya S. Prospective randomized trial of oxytocin administration for active management of the third stage of labor. *International Journal of Gynecology & Obstetrics* 2014;**127**(2):175-9.

Ononge 2015 (published data only)

Ononge S, Campbell OM, Kaharuza F, Lewis JJ, Fielding K, Mirembe F. Effectiveness and safety of misoprostol distributed to antenatal women to prevent postpartum haemorrhage after child-births: a stepped-wedge cluster-randomized trial. *BMC Pregnancy and Childbirth* 2015;**15**:315.

Ozalp 2010 (published data only)

Ozalp E, Tanir HM, Sener T. Dinoprostone vaginal insert versus intravenous oxytocin to reduce postpartum blood loss following vaginal or cesarean delivery. *Clinical and Experimental Obstetrics and Gynecology* 2010;**37**(1):53-5.

Ozcan 1996 {published data only}

Ozcan T, Sahin G, Senoz S. The effect of intraumbilical oxytocin on the third stage of labour. *Australian and New Zealand Journal of Obstetrics and Gynaecology* 1996;**36**:9-11.

Ozkaya 2005 (published data only)

Ozkaya O, Sezik M, Kaya H, Desdicioglu R, Dittrich R. Placebocontrolled randomized comparison of vaginal with rectal misoprostol in the prevention of postpartum hemorrhage. *Journal of Obstetrics & Gynaecology Research* 2005;**31**(5):389-93.

Padhy 2006 {published data only}

Padhy AK, Panigrahi R, Mohapatra KR. Alternative method of active management of 3rd stage of labour with 10 units of intraumbilical oxytocin injection [abstract]. 49th All India Congress of Obstetrics and Gynaecology; 2006 January 6-9; Cochin, Kerala State, India. 2006:76.

Palacio 2011 {published data only}

Palacio FJ, Morillas F, Ortiz-Gomez JR, Fornet I, Bermejo L, Cantalejo F. [Efficacy of low-dose oxytocin during elective cesarean section]. [Spanish]. *Revista Espanola de Anestesiologia y Reanimacion* 2011;**58**(1):6-10.

Paull 1977 {published data only}

Paull JD, Ratten GJ. Ergometrine and third stage blood loss. *Medical Journal of Australia* 1977;**1**:178-9.

Pei 1996 {published data only}

Pei JL, Zhao DF. Study of the effects of using uterine stimulants on milk secretion during delivery. *Zhonghua Hu Li Za Zhi* 1996;**31**(7):384-5.

Perdiou 2009 (published data only)

Perdiou A. The effect of 3rd generation colloids on primary haemostasis in pregnant women. European Hematology Association 14th Annual Congress; 2009 June 4-7; Berlin, Germany. 2009.

* Perdiou A, Kousoulakou A, Papadopoulou G, Leveta G, Trigka A, Andromida M, et al. The effect of 3rd generation colloids on primary haemostasis in pregnant women. *Haematologica* 2009;**94**(s2):527, Abstract no. 1334.

Phromboot 2010 {published data only}

Phromboot T. Efficacy of intraumbilical vein methylergonovine maleate on duration of third stage of labor. *Thai Journal of Obstetrics and Gynaecology* 2010;**20**:29-33.



Pierre 1992 (published data only)

Pierre F, Mesnard L, Body G. For a systematic policy of iv oxytocin inducted placenta deliveries in a unit where a fairly active management of third stage of labour is yet applied: results of a controlled trial. *European Journal of Obstetrics & Gynecology and Reproductive Biology* 1992;**43**:131-5.

Pinder 2002 (published data only)

Pinder AJ, Dresner M, Calow C, ORiordan J, Johnson R. Haemodynamic changes caused by oxytocin during caesarean section under spinal anaesthesia. *International Journal of Obstetric Anesthesia* 2002;**11**:156-9.

Pisani 2012 {published data only}

Pisani I, Tiralongo GM, Gagliardi G, Scala RL, Todde C, Frigo MG, et al. The maternal cardiovascular effect of carbetocin compared to oxytocin in women undergoing caesarean section. *Pregnancy Hypertension* 2012;**2**(2):139-42.

Porter 1991 (published data only)

Porter KB, O'Brien WF, Bruskivage L, Collins MK, Knuppel RA, Givens P. Prospective randomized study on the effects of umbilical vein oxytocin on puerperal blood loss, length of the third stage of labor and on alpha-fetoprotein levels. *American Journal of Obstetrics and Gynecology* 1991;**164**:326.

* Porter KB, O'Brien WF, Collins MK, Givens P, Knuppel R, Bruskivage L. A randomized comparison of umbilical vein and intravenous oxytocin during the puerperium. *Obstetrics & Gynecology* 1991;**78**:254-6.

Priya 2015 {published data only}

Priya GP, Veena P, Chaturvedula L, Subitha L. A randomized controlled trial of sublingual misoprostol and intramuscular oxytocin for prevention of postpartum hemorrhage. *Archives of Gynecology and Obstetrics* 2015;**292**(6):1231-7.

Puri 2012 {published data only}

Puri M, Taneja P, Gami N, Rehan HS. Effects of different doses of intraumbilical oxytocin on the third stage of labor. *International Journal of Gynecology & Obstetrics* 2012;**118**(3):210-2.

Qiu 1999 {published data only}

Qiu H, Zhu H, Ouyang W. Clinical study on chanlibao in accelerating second stage of labor [Chinese]. *Chung-Kuo Chung Hsi i Chieh Ho Tsa Chih* 1998;**18**:214-6.

* Qiu H, Zhu H, Ouyang W, Wang Z, Sun H. Clinical effects and mechanism of chanlibao in accelerating second stage of labor. Journal of Tongji Medical University 1999;**19**(2):141-4.

Quiroga 2009 (published data only)

Quiroga Diaz R, Cantu Mata R, Tello Gutierrez HE, Puente Villalobos M, Montemayor Garza R, Martinez Mendoza A. Intrauterine misoprostol for the prevention of bleeding cesarean. *Ginecologia y Obstetricia de Mexico* 2009;**77**(10):469-74.

Ragab 2016 (published data only)

Ragab A, Barakat R, Alsammani MA. A randomized clinical trial of preoperative versus postoperative misoprostol in elective

cesarean delivery. *International Journal of Gynaecology and Obstetrics* 2016;**132**(1):82-4.

Raghavan 2016 (published data only)

Raghavan S, Geller S, Miller S, Goudar SS, Anger H, Yadavannavar MC, et al. Misoprostol for primary versus secondary prevention of postpartum haemorrhage: a cluster-randomised non-inferiority community trial. *BJOG: an international journal of obstetrics and gynaecology* 2016:**123**:120-7.

Rahbar 2018 (published data only)

Rahbar N, Mirjan N, Ghorbani R. Comparison of sublingual misoprostol and intravenous oxytocin in the management of hemorrhage after cesarean section. *Middle East Journal of Rehabilitation and Health Studies* 2018;**5**(1):e62025.

Rajwani 2000 {published data only}

Rajwani J, Survana K. Active management of third stage of labor - a comparative study [abstract]. XVI FIGO World Congress of Obstetrics & Gynecology (Book 3); 2000 Sept 3-8; Washington DC, USA. 2000:54.

Ray 2012 {published data only}

Ray D, Ghosh S, Bhattacharya S, Mandal RD, Basak A. Oxytocin administration during caesarian delivery: comparison between bolus versus infusion. Society for Obstetric Anesthesia and Perinatology (SOAP) 44th Annual Meeting; 2015 May 2-5; Monterey, USA. 2012:T-26.

Razali 2016 (published data only)

Razali N, Md Latar IL, Chan YK, Omar SZ, Tan PC. Carbetocin compared to oxytocin in emergency cesarean section: a randomized trial. *European Journal of Obstetrics, Gynecology, and Reproductive Biology* 2016;**198**:35-9.

Reddy 1989 (published data only)

Reddy VV, Carey JC. Effect of umbilical vein oxytocin on puerperal blood loss and length of the third stage of labor. *American Journal of Obstetrics and Gynecology* 1989;**160**(1):206-8.

Rezk 2018 (published data only)

Rezk M. A randomized clinical trial of sublingual versus rectal misoprostol for the prevention of postpartum hemorrhage in low resource settings. pactr.org/ATMWeb/appmanager/atm/atmregistry?dar=true&tNo=PACTR201802003154375 (first received 27 February 2018).

Rooney 1985 {published data only}

Rooney I, Hughes P, Calder AA. Is routine administration of syntometrine still justified in the management of the third stage of labour?. *Health Bulletin* 1985;**43**:99-101.

Rosales-Ortiz 2014 {published data only}

Rosales-Ortiz S. Prophylaxis of obstetric haemorrhage: Experience using carbetocin vs. oxytocin in patients with risk factors for postpartum haemorrhage. *Journal of Perinatal Medicine* 2013;**41**(Suppl 1):140.



* Rosales-Ortiz S, Aguado RP, Hernandez RS, Castorena M, Cristobal FL, Gonzalez MC, et al. Carbetocin versus oxytocin for prevention of postpartum haemorrhage: A randomised controlled trial. *Lancet* 2014;**383**:S51.

Rouse 2011 {published data only}

Rouse D, Abramovici A, Szychowski J, Seals S, Andrews W, Hauth J, et al. Oxytocin dose-regimens to prevent uterine atony after vaginal delivery: does treatment efficacy vary by risk status?. *American Journal of Obstetrics and Gynecology* 2011;**204**(1 Suppl):S50-S51.

Sadeghipour 2013 (published data only)

IRCT2013052613473N1. The role of tranexamic acid in management of uterine atony during delivery. en.search.irct.ir/view/13710 (first received: 14 July 2013).

Saito 2007 {published data only}

Saito K, Haruki A, Ishikawa H, Takahashi T, Nagase H, Koyama M, et al. Prospective study of intramuscular ergometrine compared with intramuscular oxytocin for prevention of postpartum hemorrhage. *Journal of Obstetrics and Gynaecology Research* 2007;**33**(3):254-8.

Sallam 2018 (published data only)

Sallam HF Shady NW. Adjunctive sublingual misoprostol for secondary prevention of post-partum hemorrhage during cesarean delivery: double blind placebo randomized controlled trial. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2018;**7**(2):495-502.

Samuels 2005 (published data only)

Samuels N, Oberbaum M. The effect of the homoeopathic remedies arnica montana and bellis perennis on postpartum bleeding - a randomised, double-blind, placebo-controlled study [abstract]. Focus on Alternative and Complementary Therapies 2005;10 Suppl 1:47.

Sangkhomkhamhang 2012 {published data only}

Sangkhomkhamhang U, ACTRN12612000624886. A randomised controlled trial of intravenous versus intramuscular oxytocin in the management of third stage of labor. anzctr.org.au/Trial/Registration/TrialReview.aspx?ACTRN=12612000624886 (first received 12 June 2012).

Sariganont 1999 (published data only)

Sariganont J. Comparative study between syntocinon and methergin in prevention of postpartum hemorrhage. *Thai Journal of Obstetrics and Gynaecology* 1999;**11**(4):248.

Sarna 1997 {published data only}

Sarna MC, Soni AK, Gomez M, Oriol NE. Intravenous oxytocin in patients undergoing elective cesarean section [see comments]. *Anesthesia & Analgesia* 1997;**84**(4):753-6.

Sartain 2008 {published data only}

Sartain JB, Barry JJ, Howat PW, McCormack DI, Bryant M. Intravenous oxytocin bolus of 2 units is superior to 5 units during elective caesarean section. *British Journal of Anaesthesia* 2008;**101**(6):822-6.

Savitha 2017 {published data only}

Savitha A Sarita H Kashinath G. Randomized controlled trial of rectal misoprostol and intramuscular oxytocin in the prevention of PPH. *International Journal of Basic & Clinical Pharmacology* 2017;**6**(5):1101-3.

Schaefer 2004 (published data only)

Schaefer A, Klein L, Wolfe P, Heindricks G, Downs L, Guinn D. Double blind rct of early versus traditional oxytocin management in the third stage to prevent blood loss [abstract]. *American Journal of Obstetrics and Gynecology* 2004;**191**(6 Suppl 1):S69.

Schemmer 2001 {published data only}

Schemmer G. A randomized controlled trial comparing prophylactic administration of oxytocin before and after placental delivery in the prevention of postpartum hemorrhage [abstract]. *American Journal of Obstetrics and Gynecology* 2001;**184**(1):S20.

Sekhavat 2009 (published data only)

Sekhavat L, Tabatabaii A, Dalili M, Farajkhoda T, Tafti AD. Efficacy of tranexamic acid in reducing blood loss after cesarean section. *Journal of Maternal-Fetal & Neonatal Medicine* 2009;**22**(1):72-5.

Sentilhes 2015 {published data only}

NCT02302456. Tranexamic acid for preventing postpartum haemorrhage following a vaginal delivery (TRAAP). clinicaltrials.gov/ct2/show/NCT02302456 (first received: 17 November 2014).

* Sentilhes L, Daniel V, Darsonval A, Deruelle P, Vardon D, Perrotin F, et al. Study protocol. TRAAP - TRAnexamic Acid for Preventing postpartum hemorrhage after vaginal delivery: a multicenter randomized, double-blind, placebo-controlled trial. BMC Pregnancy and Childbirth 2015;**15**:135.

Senturk 2013 (published data only)

Senturk MB, Cakmak Y, Yildiz G, Yildiz P. Tranexamic acid for cesarean section: a double-blind, placebo-controlled, randomized clinical trial. *Archives of Gynecology and Obstetrics* 2013;**287**(4):641-5.

Senturk 2016 (published data only)

Senturk S, Kagitci M, Balik G, Arslan H, Kir Sahin F. The effect of the combined use of methylergonovine and oxytocin during caesarean section in the prevention of post-partum haemorrhage. *Basic & Clinical Pharmacology & Toxicology* 2016;**118**(5):338-43.

Shahid 2013 {published data only}

Shahid A, Khan A. Tranexamic acid in decreasing blood loss during and after caesarean section. Journal of the College of Physicians and Surgeons--Pakistan: JCPSP 2013; Vol. 23, issue 7:459-62.

Sharma 2014 (published data only)

Sharma M, Kaur P, Kaur K, Kaur A, Kaur PK, Kaur MM. A comparative study of oxytocin/misoprostol/methylergometrine



for active management of the third stage of labor. *Journal of Obstetrics and Gynecology of India* 2014;**64**(3):175-9.

Sheehan 2011 {published data only}

Sheehan S, Carey M, Murphy D. A cohort study of 500 patients recruited to ECSSIT - Elective Caesarean Section Syntocinon Infusion Trial. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S492.

Sheehan S, Montgomery AA, Carey M, McAuliffe F, Eogan M, Gleeson R, et al. Ecssit-elective caesarean section Syntocinon infusion trial a multicentre randomized controlled trial of oxytocin (Syntocinon) 5 IU bolus and placebo infusion versus oxytocin 5 IU bolus and 40 IU infusion for the control of blood loss at elective caesarean section. *Irish Journal of Medical Science* 2011;**180**(Suppl 4):S119.

* Sheehan SR, Montgomery AA, Carey M, McAuliffe FM, Eogan M, Gleeson R, et al. Oxytocin bolus versus oxytocin bolus and infusion for control of blood loss at elective caesarean section: Double blind, placebo controlled, randomised trial. *BMJ* 2011;**343**(7819):d4661.

Shirazi 2013 (published data only)

IRCT201204079399N1. A placebo-controlled clinical trial to assess efficacy of tranexamic acid in reducing hemorrhage after vaginal delivery. en.search.irct.ir/view/9264 (first received: 15 November 2012).

Shrestha 2007 (published data only)

Shrestha P, Babu CS. Influence of umbilical vein oxytocin on blood loss and length of third stage of labour. *Nepal Medical College Journal* 2007;**9**(3):176-8.

Shrestha 2008 (published data only)

Shrestha A, Urala MS, Upreti D, Niraula S. Comparison of intramyometrial and intramuscular 15 methyl PGF2x against traditional prophylactic intramuscular methergin for the active management of third stage of labor. *Nepal Journal of Obstetrics and Gynaecology* 2008;**3**(2):35-9.

Singh 2005 (published data only)

Singh N, Singh U. Methylergometrine and carboprost tromethamine prophylaxis for postpartum hemorrhage. *Journal of Obstetrics and Gynaecology of India* 2005;**55**(4):325-8.

Siriwarakul 1991 {published data only}

Siriwarakul W. A study of umbilical vein administration of oxytocin to shorten the third stage of labor. *Chon Buri Hospital Journal* 1991;**16**(1):40-51.

Soiva 1964 (published data only)

Soiva K, Koistinen O. Clinical experience with simultaneous intramuscular injection of oxytocin and methylergometrine. *Annales Chirurgiae et Gynaecologiae Fenniae* 1964;**53**:173-8.

Soleimani 2014 {published data only}

Soleimani Z, Naini AA. The effectiveness of sublingual misoprostol in prevention of bleeding during cesarean delivery. [Persian]. Iranian Journal of Obstetrics, Gynecology and Infertility 2014; Vol. 17, issue 125:1-7.

Sorbe 1978 (published data only)

Sorbe B. Active pharmacologic management of the third stage of labor. A comparison of oxytocin and ergometrine. *Obstetrics & Gynecology* 1978;**52**:694-7.

Soriano 1995 (published data only)

Soriano D, Dulitzki M, Schiff E, Barkai G, Seidman DS. A randomized prospective trial of oxytocin plus ergometrin versus oxytocin alone for prevention of postpartum hemorrhage. *American Journal of Obstetrics and Gynecology* 1995;**172**(1 Pt 2):361.

Sreelatha 2017 (published data only)

Sreelatha S, Nethra HS, Nadagoudar S, Ambastha V, Rajeshwari. A comparative study of different route of administration of misoprostol in the management of third stage of labour. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2017;**6**(9):3865-71.

Stearn 1963 (published data only)

Stearn RH. Syntometrine in the management of the third stage of labour. *Journal of Obstetrics and Gynaecology of the British Commonwealth* 1963;**70**:593-6.

Svanstrom 2008 (published data only)

Svanstrom MC, Biber B, Hanes M, Johansson G, Naslund U, Balfors EM. Signs of myocardial ischaemia after injection of oxytocin: a randomized double-blind comparison of oxytocin and methylergometrine during caesarean section. *British Journal of Anaesthesia* 2008;**100**(5):683-9.

Swapnika 2018 (published data only)

Swapnika D, Prema Priya G, Senthil Priya S, Allirathinam AS. A comparative study between intramuscular oxytocin and intramuscular methyl ergometrine in the active management of third stage of labour. *International Journal of Reproduction, Contraception, Obstetrics and Gynecology* 2018;**7**(5):1943-8.

Symes 1984 {published data only}

Symes JB. A study on the effect of ergometrine on serum prolactin levels following delivery. *Journal of Obstetrics and Gynaecology* 1984;**5**:36-8.

Taj 2014 (published data only)

Taj N, Firdous A, Akhtar N, Chaudhary MH, Sarah, Bajwa Z, et al. Efficacy of tranexamic acid in reducing blood loss during and after cesarean section [Nergis Taj, Ahsan Firdaus, Nadeem Akhtar, Muhammad Hamid Chaudhary, Sarah, Zunaira Bajwa, Ehsan Ullah.]. Rawal Medical Journal 2014;39(3):311-3.

Takagi 1976 (published data only)

Takagi S, Yoshida T, Togo Y, Tochigi H, Abe M, Sakata H, et al. The effects of intramyometrial injection of prostaglandin F2alpha on severe post-partum hemorrhage. *Prostaglandins* 1976;**12**(4):565-79.

Tali 2016 {published data only}

Tali K, Ignacio Alensuela A. The effect of prophylactic intravenous tranexamic acid in reducing blood loss after vaginal delivery in women at low risk of postpartum haemorrhage: A prospective, randomised, double-blind, placebo-controlled



study. Australian and New Zealand Journal of Obstetrics and Gynaecology 2016;**56**(Suppl 1):61.

Tanir 2009 {published data only}

Tanir H, Sener T, Ozalp E. Dinoprostone vaginal insert versus intravenous oxytocin to reduce the postpartum blood loss following vaginal or cesarean delivery. *International Journal of Gynecology & Obstetrics* 2009;**107**(Suppl 2):S506-7.

Tarabrin 2012 (published data only)

Tarabrin O, Kaminskiy V, Galich S, Tkachenko R, Gulyaev A, Shcherbakov S, et al. Efficacy of tranexamic acid in decreasing blood loss during cesarean section. *Critical Care* 2012;**16 Suppl 1**:S157.

Tariq 2015b {published data only}

Tariq N, Khakwani M, Parveen R. Effectiveness of misoprostol in the prevention of postpartum hemorrhage. *Pakistan Journal of Medical and Health Sciences* 2015;**9**(1):268-70.

Tehseen 2008 (published data only)

Tehseen F, Anwar A, Arfat Y. Intraumbilical veinous injection oxytocin in the active management of third stage of labour. *Journal of the College of Physicians and Surgeons--Pakistan* 2008;**18**(9):551-4.

Terry 1970 {published data only}

Terry MF. A management of the third stage to reduce fetomaternal transfusion. *Journal of Obstetrics and Gynaecology of the British Commonwealth* 1970;**77**:129-32.

Tessier 2000 {published data only}

Tessier JL, Davies GA, Woodman MC, Lipson A. Maternal hemodynamics after oxytocin bolus versus infusion in the third stage of labor. *American Journal of Obstetrics and Gynecology* 2000;**182**(1 Pt 2):S128.

Tharakan 2008 {published data only}

* Tharakan T, Jha J. Randomized double blind prospective trial of active management of the third stage of labor. *Archives of Medical Science* 2008;**4**(1):79-82.

Tharakan T, Jha J. Randomized double-blind prospective trial of active management of the third stage of labor [abstract]. *Obstetrics & Gynecology* 2007;**109**(4 Suppl):1S.

Thomas 2007 {published data only}

Thomas JS, Koh SH, Cooper GM. Haemodynamic effects of intravenous bolus or infusion of oxytocin in women undergoing caesarean section [abstract]. *International Journal of Obstetric Anesthesia* 2006;**15 Suppl 1**:S13.

* Thomas JS, Koh SH, Cooper GM. Haemodynamic effects of oxytocin given as i.v. bolus or infusion on women undergoing caesarean section. *British Journal of Anaesthesia* 2007;**98**(1):116-9.

Thornton 1988 {published data only}

* Thornton S, Davison JM, Baylis PH. Plasma oxytocin during third stage of labour: comparison of natural and active management. *BMJ* 1988;**297**:167-9.

Thornton S, Davison JM, Baylis PH. Plasma oxytocin in the third stage of human labour with and without syntometrine [abstract]. *Clinical Science* 1987;**73 Suppl 17**:2P.

Tita 2012 {published data only}

Tita AT, Szychowski JM, Rouse DJ, Bean CM, Chapman V, Nothern A, et al. Higher-dose oxytocin and hemorrhage after vaginal delivery: A randomized controlled trial. *Obstetrics and Gynecology* 2012;**119**(2 Pt 1):293-300.

Tripti 2009 (published data only)

Tripti N, Balram S. 400 ug oral misoprostol versus 0.2mg intravenous methyl ergometrine for the active management of third stage of labor. *Journal of Obstetrics and Gynecology of India* 2009;**59**(3):228-34.

Tudor 2006 {published data only}

Tudor C, Miller S, Nyima, Sonam, Droyoung, Varner M. Preliminary progress report: randomized double-blind trial of Zhi Byed 11, a Tibetan traditional medicine, versus misoprostol to prevent postpartum hemorrhage in Lhasa, Tibet. *International Journal of Gynecology & Obstetrics* 2006;**94**(Suppl 2):S145-6.

Ugwu 2016 {published data only}

Ugwu IA, Oluwasola TA, Enabor OO, Anayochukwu-Ugwu NN, Adeyemi AB, Olayemi OO. Randomized controlled trial comparing 200mug and 400mug sublingual misoprostol for prevention of primary postpartum hemorrhage. *International Journal of Gynaecology and Obstetrics* 2016;**133**:173-7.

Van den Enden 2009 {published data only}

Van den Enden E, Lahousse J, Devlieger R, Vandermeersch E, Van de Velde M. Haemodynamic effects of a bolus or infusion of oxytocin: a randomised double-blind trial. *International Journal of Obstetric Anesthesia* 2009;**18**(Suppl 1):S45.

Vasegh 2005 {published data only}

Vasegh FR, Bahiraie A, Mahmoudi M, Salehi L. Comparison of active and physiologic management of third stage of labor. HAYAT: The Journal of Tehran Faculty of Nursing & Midwifery 2005;10(23):102.

Vaughan Williams 1974 (published data only)

Vaughan Williams CA, Johnson A, Ledward R. A comparison of central venous pressure changes in the third stage of labour following oxytocic drugs and diazepam. *Journal of Obstetrics and Gynaecology of the British Commonwealth* 1974;**81**:596-9.

Ventoskovskiy 1990 (published data only)

Ventoskovskiy BM, Popov AV. Homoeopathy as a practical alternative to traditional obstetrics methods. *British Homoeopathic Journal* 1990;**79**:201-5.

Vogel 2004 {published data only}

Vogel D, Burkhardt T, Rentsch K, Schweer H, Watzer B, Zimmerman R, et al. Misoprostol versus methylergometrine: pharmacokinetics in human milk. *American Journal of Obstetrics and Gynecology* 2004;**191**:2168-73.



Wallace 2007 (published data only)

Wallace EM. A double-blind randomised controlled trial of oxytocin bolus plus placebo infusion versus oxytocin bolus plus oxytocin infusion at elective caesarean section. anzctr.org.au/Trial/Registration/TrialReview.aspx?ACTRN=12607000631404 (first received 12 December 2007).

Walraven 2005 (published data only)

Walraven G, Blum J, Dampha Y, Sowe M, Morison L, Winikoff B, et al. Misoprostol in the management of the third stage of labour in the home delivery setting in rural Gambia: a randomised controlled trial. *BJOG: an international journal of obstetrics and gynaecology* 2005;**112**:1277-83.

Wang 2000 (published data only)

Wang BI, Du JM. Clinical study on reduction of postpartum bleeding using carprost suppository. *Henan Medical Research* 2000;**9**(2):155-6.

Wang 2018 (published data only)

Wang Y. Therapeutic efficacy and safety of carbetocin on postpartum hemorrhage. chictr.org.cn/showproj.aspx? proj=26431 (first received 11 April 2018).

Weeks 2015 (published data only)

Frye L, Durocher J, Weeks A, Ditai J, Ononge S, Faragher B, et al. On the trail of misoprostol in the community: A secondary analysis of self-administered misoprostol for the prevention of postpartum hemorrhage in Uganda. *International Journal of Gynaecology and Obstetrics* 2015;**131**(Suppl 5):E354-5.

Weeks A, Ditai J, Ononge S, Faragher B, Mirembe F, Byamugisha J, et al. Self-administered misoprostol to prevent bleeding after homebirths in Uganda: a placebo-controlled randomised trial. *BJOG: an international journal of obstetrics and gynaecology* 2013;**120**:76.

* Weeks AD, Ditai J, Ononge S, Faragher B, Frye LJ, Durocher J, et al. The MamaMiso study of self-administered misoprostol to prevent bleeding after childbirth in rural Uganda: a community-based, placebo-controlled randomised trial. *BMC Pregnancy and Childbirth* 2015;**15**(1):219.

Weihong 1998 {published data only}

Weihong H, Hanrong C, Hong L, Linan C. Preventing of postpartum haemorrhage by carboprost methylate suppository administered through vagina or sublingually. *Acta Academiae Medicinae Shanghai* 1998;**25**:137-9.

Weiss 1975 {published data only}

Weiss G, Klein S, Shenkman L, Kataoka K, Hollander CS. Effect of methylergonovine on puerperal prolactin secretion. *Obstetrics & Gynecology* 1975;**46**:209-10.

Wellmann 2016 (published data only)

Wellmann S, Koslowski A, Spanaus K, Zimmermann R, Burkhardt T. Fetal release of copeptin in response to maternal oxytocin administration: a randomized controlled trial. *Obstetrics and Gynecology* 2016;**128**(4):699-703.

Wetta 2013 (published data only)

Wetta L, Szychowski J, Seals S, Mancuso M, Hauth J, Tita A. Risk factors for uterine atony at vaginal delivery: a comprehensive evaluation. *American Journal of Obstetrics and Gynecology* 2011;**204**(1 Suppl):S71-2.

* Wetta LA, Szychowski JM, Seals S, Mancuso MS, Biggio JR, Tita AT. Risk factors for uterine atony/postpartum hemorrhage requiring treatment after vaginal delivery. *American Journal of Obstetrics and Gynecology* 2013;**209**(1):51e1-6.

Winikoff 2012 (published data only)

NCT01608958. Intravenous and intramuscular administration of oxytocin in the third stage of labor for prevention of postpartum hemorrhage. clinicaltrials.gov/ct2/show/NCT01608958 (first received: 29 May 2012).

Winikoff 2016 (published data only)

Winikoff B, Dzuba I, Carroli G, NCT02954068. Intravenous versus intramuscular administration of oxytocin and its relationship with postpartum bleeding and other clinical signs: a randomized placebo-controlled study. clinicaltrials.gov/show/NCT02954068 (first received 28 October 2016).

Wong 2005a {published data only}

NCT00257803. Does the rapid intravenous administration of oxytocin after delivery of the baby decrease the bleeding during cesarean section in women at risk of bleeding during cesarean section?. clinicaltrials.gov/ct2/show/NCT00257803 (first received: 21 November 2005).

Wong 2005b {published data only}

NCT01710566. Misoprostol and oxytocin in Uniject® for postpartum hemorrhage prevention in communities. clinicaltrials.gov/ct2/show/NCT01710566 (first received: 8 May 2012)

Wright 2005 (published data only)

NCT00147420. RCT of Zhi Byed 11 (ZB11) Versus Misoprostol in Tibet. clinicaltrials.gov/ct2/show/NCT00147420 (first received 7 September 2005).

Wu 2007 {published data only}

Wu LF, Liu Y, Ruan Y. Clinical study on prevention of postpartum hemorrhage of cesarean section using hemabat in high risk pregnant women. *Chinese Journal of Obstetrics & Gynecology* 2007;**42**(9):577-81.

Xu 2003 {published data only}

Xu H. Misoprostol on preventing postpartum bleeding in cesarean. *Hebei Medicine* 2003;**9**(9):806-7.

Xu 2013 {published data only}

Xu J, Gao W, Ju Y. Tranexamic acid for the prevention of postpartum hemorrhage after cesarean section: a double-blind randomization trial. *Archives of Gynecology and Obstetrics* 2013;**287**(3):463-8.

Yamaguchi 2011 {published data only}

Yamaguchi ET, Cardoso MM, Torres ML, Nascimento RC, Ribeiro MC, Frerichs E, et al. Serum oxytocin concentrations in



elective caesarean delivery: a randomized comparison of three infusion regimens. *International Journal of Obstetric Anesthesia* 2011;**20**(3):224-8.

Yan 2000 {published data only}

Yan WG, Ling MX, Mao HY. Clinical study on reduction of postpartum bleeding in cesarean operation by misoprostol. *Journal of Zhenjiang Medical College* 2000;**10**(3):440-1.

Yang 2001 (published data only)

Yang H, Zheng S, Shi C. [Clinical study on the efficacy of tranexamic acid in reducing postpartum blood lose: a randomized comparative, multicenter trial] [Chinese]. *Chung-Hua Fu Chan Ko Tsa Chih [Chinese Journal of Obstetrics and Gynaecology]* 2001;**36**(10):590-2.

Young 1988 {published data only}

Young SB, Martelly PD, Greb L, Considine G, Coustan DR. The effect of intraumbilical oxytocin on the third stage of labor. *Obstetrics & Gynecology* 1988;**71**:736-8.

Zamora 1999 {published data only}

Zamora LA. A randomized controlled trial of oxytocin administered at the end of the second stage of labor versus oxytocin administered at the end of the third stage of labor in the prevention of postpartum hemorrhage. *Philippine Journal of Obstetrics and Gynecology* 1999;**23**(4):125-33.

Zaporozhan 2013 {published data only}

Zaporozhan V, Tarabrin O, Gavrychenko D, Mazurenko G, Saleh O, Lyoshenko I. Effcacy of tranexamic acid in decreasing blood loss during cesarean section. *Critical Care* 2013;**17**(Suppl 2):S135-6.

Zhao 1998 {published data only}

Zhao Y, Li X, Peng Y. Clinical study on reduction of postpartum bleeding in cesarean section by misoprostol. *Chung-Hua Fu Chan Ko Tsa Chih [Chinese Journal of Obstetrics & Gynecology]* 1998;**33**:403-5.

Zhao 2003 {published data only}

Zhao SF, Sun XF. Clinical study on preventing and curing postpartum hemorrhage in the third stage of labor. *Journal of Practical Obstetrics and Gynecology* 2003;**19**(5):278-80.

Zhou 1994 {published data only}

Zhou HL, Zhang L. Study on the effect of third stage of labor through different channels of injection of oxytocin. *Chinese Journal of Nursing* 1994;**29**(8):453-5.

References to studies awaiting assessment

Abdel-Aleem 1997 (published data only)

Abdel-Aleem H. [Personal communication] Management of the third stage of labour with carboprost trometamol in high risk patients for postpartum haemorrhage. Letter to: Cochrane Pregnancy and Childbirth, Liverpool, UK 1997.

* Abdel-Aleem H, Mostafa SA, Makarem MH, Abol-Oyoun EM, Makhlouf A, Shoukry M. Management of the third stage of labour with carboprost trometamol in high risk patients for

postpartum hemorrhage. Research Activities on Reproductive Health: Annual Report of Assiut University Department of Obstetrics and Gynecology November 1997. Assiut University, Faculty of Medicine, 1997:75.

Alli 2013 (published data only)

Alli QO. Comparing effectiveness of sublingual misoprostol with oxytocin infusion to reduce blood loss at caesarean section: double blind, randomised study. *BJOG: an international journal of obstetrics and gynaecology* 2013;**120**:77-8.

Amornpetchakul 2017 {published data only}

* Amornpetchakul P, Lertbunnaphong T, Boriboonhiransarn D, Leetheeragul J, Sirisomboon R, Jiraprasertwong R. Efficacy of intravenous 100 mcg carbetocin versus intravenous 5 units oxytocin for prevention of immediate postpartum hemorrhage after normal vagina delivery among high risk pregnancies; a triple-blinded randomized controlled trial. *Journal of Obstetrics and Gynaecology Research* 2017;**43**:24-5.

Lertbunnaphong T. Efficacy of intravenous 100 mcg carbetocin versus intravenous 5 units oxytocin for prevention of immediate postpartum hemorrhage after normal vagina delivery among high risk pregnancies; a triple-blinded randomized controlled trial. clinicaltrials.in.th/index.php? tp=regtrials&menu=trialsearch&smenu=fulltext&task=search&task2=view1& (first received 15 July 2016).

Beigi 2009 (published data only)

Beigi A, Tabarestani H, Moini A, Zarrinkoub F, Kazempour M, Hadian Amree A. [Sublingual misoprostol versus intravenous oxytocin in the management of postpartum hemorrhage]. *Tehran University Medical Journal* 2009;**67**(8):556-61.

Muller 1996 (published data only)

Muller R, Beck G. Active management of the third stage of labour. 19th Swiss Congress of the Swiss Society of Gynecology and Obstetrics; 1996 June; Interlaken, Switzerland. 1996.

Norchi 1988 {published data only}

Norchi S, Beretta E, Zanini A, Bottino S. Prevention of primary post-partum haemorrhage (PPH). Controlled clinical trial: Sulprostone vs Metilergometrina. 12th FIGO World Congress of Gynecology and Obstetrics; 1988 October 23-28; Brazil. 1988.

Rabow 2017 {published data only}

Rabow S, Jonsson E, Jonsson H, Olofsson P. Cardiovascular effects of oxytocin and carbetocin at caesarean section, a prospective double-blind randomised study using non-invasive pulse wave analysis. Acta Anaesthesiologica Scandinavica 2017; Vol. 61, issue 8:1053.

Roy 2017 {published data only}

Roy R Vernekar M. Comparative study on effect of misoprostol and oxytocin in the active management of third stage of labor in a tertiary hospital in Manipur, India. *Indian Journal of Public Health Research and Development* 2017;**8**(4):376-81.

Said 2017 {published data only}

Said SK. A comparative study of rectal misoprostol to oxytocin infusion during cesarean delivery to reduce intra operative



& postoperative blood loss. *Al-Kufa Journal for Biology* 2017:**9**(3):55-62.

Shrivasatava 2012 (published data only)

Shrivasatava DD, Khamsara D. Critical evaluation of sublingual misoprostol and methyl ergometrine in active management of third stage of labour. *International Journal of Gynecology and Obstetrics* 2012;**119**(Suppl 3):S484.

Sunil 2016 (published data only)

Sunil Kumar KS, Shyam S, Batakurki P. Carboprost versus oxytocin for active management of third stage of labor: a prospective randomized control study. *Journal of Obstetrics and Gynaecology of India* 2016;**66**(Suppl 1):S229-34.

References to ongoing studies

Balki 2017 (published data only)

NCT03168698. Carbetocin vs. oxytocin at elective cesarean section: a double-blind, randomized controlled non-inferiority trial of high and low dose regimens. https://clinicaltrials.gov/ct2/show/NCT03168698 (first received 30 May 2017).

Draycott 2014 {published data only}

NCT02216383. Intramuscular oxytocics: a comparison study of intramuscular carbetocin, syntocinon and syntometrine for the third stage of labour following vaginal birth (IMox). clinicaltrials.gov/ct2/show/NCT02216383 (first received 15 August 2014).

Gomez 2011 {published data only}

ACTRN12610000550000. Comparison of the effectiveness of carbetocin vs oxytocin in managing the third stage of labor in a group of women with risk factors for postpartum hemorrhage. anzctr.org.au/Trial/Registration/TrialReview.aspx?id=335628 (first received 25 June 2010).

Goudar 2016 (published data only)

CTRI/2016/06/006996. Effect of carbetocin RTS vs oxytocin for preventing postpartum hemorrhage on post-delivery hemoglobin: a randomized controlled trial. ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid=10619 (first received 6 June 2016).

Kalahroudi 2010a {published data only}

IRCT138810212854N2. Comparison effect of carbetocine and syntometrin in prevention of post partum hemorrhage. en.search.irct.ir/view/2059 (first received 15 March 2010).

Kalahroudi 2010b {published data only}

IRCT201008212854N5. Comparison of the effect of rectal misoprostol and syntometrin in prevention of post partum hemorrhage. en.search.irct.ir/view/3932 (first received 9 September 2010).

Maged 2018 (published data only)

NCT03556852. Carbetocin versus rectal misoprostol for management of third stage of labor in women at low risk of postpartum hemorrhage. clinicaltrials.gov/ct2/show/NCT03556852 (first received 14 June 2018).

Moradi 2010 (published data only)

IRCT138812223548N1. Comparison of misoprostol and oxytocin in reduction of postpartum hemorrhage. en.search.irct.ir/view/2616 (first received 29 May 2010).

Sweed 2014 (published data only)

NCT02083107. Comparison between rectal and sublingual misoprostol before caesarian section to reduce intra & post-operative blood loss. clinicaltrials.gov/ct2/show/NCT02083107 (first received 7 March 2014).

Thakur 2015 (published data only)

CTRI/2015/06/005918. Efficacy of oxytocin, misoprostol, 15-methylprostaglandinF2alpha and methylergometrine in active management of third stage of labor. ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid=10133 (first received 16 June 2015).

Additional references

Alkema 2016

Alkema L, Chou D, Hogan D, Zhang S, Moller AB, Gemmill A, et al. Global, regional, and national levels and trends in maternal mortality between 1990 and 2015, with scenario-based projections to 2030: a systematic analysis by the UN Maternal Mortality Estimation Inter-Agency Group. *Lancet* 2016;**387**:462-74.

Begley 2015

Begley CM, Gyte GM, Devane D, McGuire W, Weeks A. Active versus expectant management for women in the third stage of labour. *Cochrane Database of Systematic Reviews* 2015, Issue 3. [DOI: 10.1002/14651858.CD007412.pub4]

Brignardello-Petersen 2018

Brignardello-Petersen R, Bonner A, Alexander PE, Siemieniuk RA, Furukawa TA, Rochwerg B, et al. Advances in the GRADE approach to rate the certainty in estimates from a network meta-analysis. *Journal of Clinical Epidemiology* 2018;**93**:36-44.

Caldwell 2005

Caldwell DM, Ades AE, Higgins JP. Simultaneous comparison of multiple treatments: combining direct and indirect evidence. *BMJ* 2005;**331**:897-900.

Combs 1991

Combs CA, Murphy EL, Laros RK Jr. Factors associated with postpartum hemorrhage with vaginal birth. *Obstetrics and Gynecology* 1991;**77**:69-76.

Davies 2001

Davies NM, Longstreth J, Jamali F. Misoprostol therapeutics revisited. *Pharmacotherapy* 2001;**21**:60-73.

de Groot 1996a

de Groot AN. The role of oral (methyl) ergometrine in the prevention of postpartum haemorrhage. *European Journal of Obstetrics, Gynecology, and Reproductive Biology* 1996;**69**:31-6.



de Groot 1998

de Groot AN, van Dongen PW, Vree TB, Hekster YA, van Roosmalen J. Ergot alkaloids. Current status and review of clinical pharmacology and therapeutic use compared with other oxytocics in obstetrics and gynaecology. *Drugs* 1998;**56**:523-35.

DerSimonian 1986

DerSimonian R, Laird N. Meta-analysis in clinical trials. *Controlled Clinical Trials* 1986;**7**(3):177-88.

Dias 2013

Dias S, Sutton AJ, Ades AE, Welton NJ. Evidence synthesis for decision making 2: a generalized linear modeling framework for pairwise and network meta-analysis of randomized controlled trials. Medical Decision Making 2013; Vol. 33, issue 5:607-17.

Higgins 2002

Higgins JP, Thompson SG. Quantifying heterogeneity in a meta-analysis. *Statistics in Medicine* 2002;**21**(11):1539-58.

Higgins 2011

Higgins JP, Green S, editor(s). Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 [updated March 2011]. The Cochrane Collaboration, 2011. Available from www.cochrane-handbook.org.

Higgins 2012

Higgins JP, Jackson D, Barrett JK, Lu G, Ades AE, White IR. Consistency and inconsistency innetwork meta-analysis: Concepts and models for multi-arm studies. *Research Synthesis Methods* 2012;**3**(2):98-110.

Hogerzeil 1993

Hogerzeil HV, Walker GJ, de Goeje MJ. Stability of Injectable Oxytocics in Tropical Climates. Results of Field Surveys and Simulation Studies on Ergometrine, Methylergometrine and Oxytocin. Geneva: World Health Organization (WHO), 1993.

Hunter 1992

Hunter DJ, Schulz P, Wassenaar W. Effect of carbetocin, a long-acting oxytocin analog on the postpartum uterus. *Clinical Pharmacology and Therapeutics* 1992;**52**:60-7.

Liabsuetrakul 2018

Liabsuetrakul T, Choobun T, Peeyananjarassri K, Islam QM. Prophylactic use of ergot alkaloids in the third stage of labour. *Cochrane Database of Systematic Reviews* 2018, Issue 6. [DOI: 10.1002/14651858.CD005456.pub2]

Lumley 2002

Lumley T. Network meta-analysis for indirect treatment comparisons. Statistics in Medicine 2002;21(16):2313-24.

McDonald 2004

McDonald SJ, Abbott JM, Higgins SP. Prophylactic ergometrineoxytocin versus oxytocin for the third stage of labour. *Cochrane Database of Systematic Reviews* 2004, Issue 1. [DOI: 10.1002/14651858.CD000201.pub2]

MEDICINES.ORG.UK

Medicines.org.uk. Syntocinon Ampoules 5 IU/ml - Summary of Product Characteristics (SPC) - (eMC). Available from: http://www.medicines.org.uk/emc/medicine/16423/SPC 2014; Vol. [cited 10 October 2014].

Nüesch 2010

Nüesch E, Trelle S, Reichenbach S, Rutjes AW, Tschannen B, Altman DG, et al. Small study effects in meta-analyses of osteoarthritis trials: meta-epidemiological study. *BMJ* 2010;**341**:c3515.

Penney 2007

Penney G, Brace V. Near miss audit in obstetrics. *Current Opinion in Obstetrics & Gynecology* 2007;**19**:145-50.

Puhan 2014

Puhan MA, Schünemann HJ, Murad MH, Li T, Brignardello-Petersen R, Singh JA, et al. A GRADE Working Group approach for rating the quality of treatment effect estimates from network meta-analysis. *BMJ* 2014;**349**:g5630.

RevMan 2014 [Computer program]

The Nordic Cochrane Centre, The Cochrane Collaboration. Review Manager (RevMan). Version 5.3. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014.

Salanti 2011

Salanti G, Ades AE, Ioannidis JP. Graphical methods and numerical summaries for presenting results from multiple-treatment meta-analysis: An overview and tutorial. *Journal of Clinical Epidemiology* 2011;**64**(2):163-71.

Say 2014

Say L, Chou D, Gemmill A, Tunçalp O, Moller A, Daniels J, et al. Global causes of maternal death: a WHO systematic analysis. *Lancet Global Health* 2014;**2**(6):e323-e333.

Schaff 2005

Schaff EA, DiCenzo R, Fielding SL. Comparison of misoprostol plasma concentrations following buccal and sublingual administration. *Contraception* 2005;**71**(1):22-5.

Souza 2013

Souza JP, Gülmezoglu AM, Vogel J, Carroli G, Lumbiganon P, Qureshi Z, et al. Moving beyond essential interventions for reduction of maternal mortality (the WHO Multicountry Survey on Maternal and Newborn Health): a cross-sectional study. *Lancet* 2013;**381**:1747-55.

Su 2012

Su LL, Chong YS, Samuel M. Carbetocin for preventing postpartum haemorrhage. *Cochrane Database of Systematic Reviews* 2012, Issue 4. [DOI: 10.1002/14651858.CD005457.pub4]

Tuncalp 2012

Tuncalp O, Hofmeyr GJ, Gulmezoglu AM. Prostaglandins for preventing postpartum haemorrhage. *Cochrane Database of Systematic Reviews* 2012, Issue 8. [DOI: 10.1002/14651858.CD000494.pub4]



Westhoff 2013

Westhoff G, Cotter AM, Tolosa JE. Prophylactic oxytocin for the third stage of labour to prevent postpartum haemorrhage. *Cochrane Database of Systematic Reviews* 2013, Issue 10. [DOI: 10.1002/14651858.CD001808.pub2]

White 2011

White I. Multivariate random-effects meta-regression:Updates to mvmeta. *The Stata Journal* 2011;**11**(2):255-270.

White 2012

White IR, Barrett JK, Jackson D, Higgins JP. Consistency and inconsistency in network metaanalysis:Model estimation using multivariate meta-regression. *Research Synthesis Methods* 2012;**3**(2):111-25.

White 2015

White IR. Network meta-analysis. *Stata Journal* 2015;**15**(4):951-85.

WHO 1993

WHO. Stability of Injectable Oxytocics in Tropical Climates: Results of Field Surveys and Simulation Studies on

Ergometrine, Methylergometrine, and Oxytocin. Geneva: World Health Organization, 1993.

WHO 2012

WHO. WHO Recommendations for the Prevention and Treatment of Postpartum Haemorrhage. Geneva: Department of Reproductive Health, World Health Organization, 2012.

References to other published versions of this review Gallos 2015

Gallos ID, Williams HM, Price MJ, Merriel A, Gee H, Lissauer D, et al. Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis. *Cochrane Database of Systematic Reviews* 2015, Issue 5. [DOI: 10.1002/14651858.CD011689]

Gallos 2018

Gallos ID, Williams HM, Price MJ, Merriel A, Gee H, Lissauer D, et al. Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis. *Cochrane Database of Systematic Reviews* 2018, Issue 4. [DOI: 10.1002/14651858.CD011689.pub2]

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Abdel-Aleem 1993

Methods	2-arm active-controlled randomised trial		
Participants	150 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with risk factors for PPH: duration of labour less than 2 hours or prolonged labour more than 24 hours, MgSO4 for pre-eclampsia, chorioamnionitis, multiple pregnancy, previous PPH, APH and episiotomy.		
Interventions	200 mcg of ergometrine administered by an IV bolus versus 250 mcg of carboprost administered IM		
Outcomes	The study recorded the following outcomes: blood loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; abdominal pain.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Table of random numbers was used.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias)	Unclear risk	Blinding (of study participants and caregivers) was unclear.	

^{*} Indicates the major publication for the study



Abdel-Aleem 1993 (Continued)

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Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Blood was collected in a tray and measured. Sterile pads were placed over the vulva before and after use for a period of 4 hours.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Carboprost kindly supplied by Prof. S. Bergstrom, Sweden but source(s) of funding for the study were not reported.

Abdel-Aleem 2010

Methods	3-arm controlled randomised trial
Participants	1964 women were randomised in a hospital setting in Egypt and South Africa. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with medical complications such as hypertension and diabetes, previous caesarean section, or an abdominal wall that was not thin enough to allow easy palpation of the uterus after delivery.
Interventions	10 IU of oxytocin administered IM versus no treatment
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta.
Notes	Contact with study authors for additional information: no. Additional data from authors: no

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocated to 1 of 3 groups by selecting the next number in a computer-generated random number sequence
Allocation concealment (selection bias)	Low risk	The allocated group was noted inside opaque sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.



Abdel-Aleem 2010 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	In Assiut, investigators appraised blood loss by collection with a calibrated plastic drape placed under the mother within 30 minutes of delivery. At the East London Hospital Complex, investigators appraised blood loss by collection with a low profile plastic "fracture" bedpan placed under the mother.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Investigators were unable to collect outcome data from 14 randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The study protocol was registered retrospectively (ACTRN: 12609000372280).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the institution of the authors, or conducted without external funding.

Acharya 2001

Participants 60 women were randomised in a hospital setting in UK. The population cor liparous and multiparous, either singleton or multiple pregnancy, at high relective caesarean section. Exclusion criteria were not specified.	
cteedive edesared. Section: Exclusion enteria were not specified.	
Interventions 10 IU of oxytocin administered by an IV bolus versus 400 mcg of misoprosto	l administered orally
Outcomes The study recorded the following outcomes: PPH at 1000; additional uteror loss (mL; change in Hb; vomiting; shivering.	onics; transfusion; blood
Notes Contact with study authors for additional information: no. Additional data	rom authors: no

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Low risk	Randomisation was performed using sealed opaque envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Acharya 2001 (Continued)		
Objective assessment of blood loss	High risk	Investigators appraised intra-operative blood loss by the estimation of attending physicians, and by measurement of preoperative and postoperative Hb concentration and hematocrit.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Adanikin 2012

Methods	2-arm active-controlled double-dummy randomised trial
Participants	218 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with altered serum electrolytes, peritonitis, sepsis, previous bowel surgery, thyroid disease, inflammatory bowel disease, or chronic constipation.
Interventions	25 IU of oxytocin administered by an IV bolus + infusion versus 600 mcg plus 5 IU of misoprostol plus oxytocin administered rectally plus by an IV bolus
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; nausea; vomiting; fever; shivering.
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes

KISK OI DIUS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocation sequence developed by 1 researcher (0.0.) using a computer-gener ated table of random numbers with varied permutated blocks.
Allocation concealment (selection bias)	Low risk	Used sealed opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The same researcher administered the drugs intra-operation and set up the infusions in the operating room; he was the only person who was not blind to the drug allocation and he did not take any further part in the active running of the study".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.



Adanikin 2012 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Adanikin 2013

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	50 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with asthma or with hypersensitivity to prostaglandins.		
Interventions	600 mcg of misoprostol administered rectally versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	1:1 computer-generated randomisation.
Allocation concealment (selection bias)	Low risk	The pharmacy department provided the study drugs and placebos in unidentifiable form but the resident doctor was responsible for the patient's allocation according to the randomisation table.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Outcome assessors were blinded.
Objective assessment of blood loss	Low risk	Investigators weighed the pads 4 hours postpartum for assessment of blood loss.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.



Adanikin 2013 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Afolabi 2010

Methods	2-arm active-controlled randomised trial		
Participants	200 women were randomised in a hospital setting in Nigeria. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction of labour or caesarean section, or those with hematocrit of less than 30%, pre-eclampsia/eclampsia, grand multiparity (5 or more), multiple pregnancy, coagulopathy, or medical disorders.		
Interventions	10 IU of oxytocin administered IM versus 400 mcg of misoprostol administered orally		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised into 2 groups, A and B, by blocked (restrictive) double-blind randomisation using random table generated numbers.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss at delivery by collection with a large kidney dish, for measurement in a graduated measuring jar.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.



Afolabi 2010 (Continued)		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ahmed 2014

Methods	2-arm active-controlled randomised trial.	
Participants	80 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with risk factors for excessive blood loss e.g. those with placenta praevia or placental abruption.	
Interventions	100 mcg of carbetocin administered by an IV bolus versus 10 IU of oxytocin administered by an IV bolus	
Outcomes	The study recorded the following outcomes: blood loss (mL).	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study was "single-blind" but the identity of those blinded and the method of blinding were not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.



Al-Sawaf 2013

Methods	3-arm controlled randomised trial		
Participants	120 women were randomised in a hospital setting in Egypt. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction of labour or instrumental delivery, or those with previous caesarean section, extensive perineal, vaginal or cervical lacerations, bleeding disorders, HB less than 100 g/L, uterine malformations, grand multiparity, multiple pregnancy, polyhydramnios, intrauterine fetal death, medical problems such as pre-eclampsia, diabetes, cardiopulmonary problems, bowel disease, or allergy to prostaglandins.		
Interventions	No treatment versus 20 tered IM	00 mcg of misoprostol administered sublingually versus 5 IU of oxytocin adminis-	
Outcomes	The study recorded the sion; blood loss (mL); c	e following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu- change in Hb.	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Used closed envelopes.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with sterile packs weighed beforehand and afterwards.	
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "Following randomisation, 16 study participants were excluded from our analysis. Of these, 14 patients received intrapartum oxytocin, 1 patient experienced extensive vaginal laceration and another experienced a cervical laceration".	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	



Alwani 2014		
Methods	2-arm active-controlle	d randomised trial
Participants	3 or less, either singlet	omised in a hospital setting in India. The population comprised women of parity on or multiple pregnancy, at high risk for PPH, who delivered by both elective or Exclusion criteria were not specified.
Interventions	600 mcg of misoprosto	ol administered rectally versus 10 IU of oxytocin administered IM
Outcomes		e following outcomes: additional uterotonics; transfusion; death; change in Hb; ertension; fever;.shivering.
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The patients were randomised in 2 groups using random number table generated online (http://www.graphpad.com/quickcalcs/randomize1/).
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	No funding was sought for this study.

Amant 1999

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	213 women were randomised in a hospital setting in Belgium. The population comprised women of both nulliparous and multiparous, either singleton or multiple pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with hypertensive disorders, gestational age less than 32 weeks, intrauterine fetal death, uterine	



Amant 1999 (Continued)	malformations, inflammatory bowel disease, obliterative vascular or coronary disease, sepsis, allergy to prostaglandins or alkaloids.		
Interventions	600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered by an IV bolus		
Outcomes		e following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfuor placenta; diarrhoea; nausea; vomiting; headache; fever; shivering.	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Allocation by a computer-generated list and randomisation in blocks.	
Allocation concealment (selection bias)	Low risk	The study box contained either 2 capsules of misoprostol and an ampoule containing placebo, or 2 capsules with placebo and an ampoule containing methylergometrine. The study boxes and capsules were indistinguishable in the 2 groups	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.	
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.	
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "213 women were enrolled in the study, but the data for 13 were excluded because a caesarean section was performed after randomisation (n = 3), or because no predelivery (n = 3) or postpartum (n = 7, short hospital stay) blood sample was taken".	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Amin 2014

Methods	2-arm active-controlled randomised trial
Participants	200 women were randomised in a hospital setting in Pakistan. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with traumatic PPH, bleeding disorders, prolonged labour, placenta praevia, placental abruption, multiple pregnancy, BMI more than 30, or previous PPH.



Amin 2014 (Continued)					
Interventions	5 IU of oxytocin administered by an IV bolus versus 800 mcg of misoprostol administered rectally				
Outcomes	The study recorded the following outcomes: PPH at 500; severe maternal morbidity: intensive care admissions; manual removal of placenta; death; blood loss (mL); third stage duration (minutes); diarrhoea; vomiting; fever; shivering.				
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.			
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.			
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.			
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with special drapes placed un der the mother until 1 hour postpartum, and weighed beforehand and afterwards. Blood was also collected in graduated plastic bags.			
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.			
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.			
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.			
Funding source	Unclear risk	Source(s) of funding for the study were not reported.			

Askar 2011

Methods	2-arm active-controlled double-blinded randomised trial
Participants	240 women were randomised in a hospital setting in Kuwait. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women less than 18 years old and those with known or suspected coagulopathy, grand multiparity (5 or more), uterine fibroids, polyhydramnios, multiple pregnancy, fetal macrosomia, severe anaemia, cervical tears or who required prophylactic oxytocin infusion. The presence of contraindications for the use of either syntometrine or carbetocin that include pre-existing hypertension,



Askar 2011 (Continued)	pre-eclampsia, asthma, cardiac, renal or liver diseases, epilepsy, or history of hypersensitivity to syntometrine or carbetocin.				
Interventions	100 mcg of carbetocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM				
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; nausea; vomiting; hypertension; headache; abdominal pain.				
Notes	Contact with study aut	chors for additional information: yes. Additional data from authors: yes			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	Allocation by a computer-generated code prepared before the recruitment.			
Allocation concealment (selection bias)	Low risk	Used sealed, consecutively-numbered, opaque envelopes			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.			
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.			
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a new plastic sheet placed under the mother following delivery of the placenta, and weighed (together with any gauzes, tampons and pads applied during the delivery) beforehand and 2 hours afterwards. A digital scale was used for weight measurement.			
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.			
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.			
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.			
Funding source	Unclear risk	Source(s) of funding for the study were not reported.			

Asmat 2017

Methods	2-arm active-controlled randomised trial
Participants	1678 women were randomised in a hospital setting in Pakistan. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with malpresentations such as breech, compound or transverse



Asmat 2017 (Continued)		pregnancy, placenta praevia type III, IV, placenta accreta, placental abruption, ectomy (uterine cavity opened), coagulation disorders, DIC, cardiac diseases, di-			
Interventions	800 mcg of misoprosto	l administered rectally versus 10 IU of oxytocin administered IM			
Outcomes	The study recorded the	e following outcomes: PPH at 500; blood loss (mL).			
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	A lottery method was used.			
Allocation concealment (selection bias)	High risk	Allocation concealment was not reported but unlikely to have been implemented with a lottery method of randomisation.			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.			
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.			
Objective assessment of blood loss	Unclear risk	Quote: "Pads soaked were used to asses the amount of blood loss." Methods of evaluating blood loss were not reported in sufficient detail.			
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.			
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.			
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.			
Funding source	Unclear risk	Source(s) of funding for the study were not reported.			

Attilakos 2010

Methods 2-arm active-controlled double-blinded randomised trial		
Participants	377 women were randomised in a hospital setting in UK. The population comprised women of both nulliparous and multiparous, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women undergoing caesarean section with general anaesthesia, gestational age less than 37 weeks performed for fetal or maternal distress where, due to time constraints, it was not possible to recruit or randomise, or those with multiple pregnancy, placenta praevia or placental abruption.	



Interventions	100 mcg of carbetocin administered by an IV bolus versus 5 IU of oxytocin administered by an IV bolus				
Outcomes	The study recorded the following outcomes: PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; death; blood loss (mL); change in Hb; nausea; vomiting; headache; tachycardia; hypotension; shivering; abdominal pain.				
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	The randomisation sequence (1:1 ratio—blocks of ten, no stratification) was generated by computer.			
Allocation concealment (selection bias)	Low risk	The preparation of the ampoules was undertaken by DHP Ltd. (Powys, UK) which provided sequentially numbered and labelled boxes each containing a 1-mL ampoule of the study drug. All boxes and ampoules were identically labelled, with the study number being the only differentiating feature between different drug packs. the random allocation sequence was not known to the investigators until the study had finished and the analysis was started.			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.			
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.			
Objective assessment of blood loss	High risk	Blood loss was estimated by the attending surgeon quote: "in the usual way (visual estimation, number of used swabs and amount of aspirated blood)".			
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.			
Selective reporting (re- porting bias)	Low risk	The study report matches the study protocol that was registered prospectively (EudraCT 2005-002812-94).			
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.			
Funding source	Low risk	Ferring Pharmaceuticals funded the cost of preparation of blinded medication ampoules. No other external funding was required for the study.			

Atukunda 2014

Methods	2-arm active-controlled double-dummy randomised trial
Participants	1140 women were randomised in a hospital setting in Uganda. The population comprised women of both nulliparous and multiparous, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or elective caesarean section, or those with intrauterine fetal death, heart disease, severe



Atukunda 2014 (Continued)	malaria or acute bacterial infection, multiple pregnancy, antepartum haemorrhage, altered cognitive status or reported hypersensitivity to prostaglandins.				
Interventions	10 IU of oxytocin administered IM versus 600 mcg of misoprostol administered sublingually				
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering; abdominal pain.				
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	A study biostatistician generated a randomisation list with a block size of 10.			
Allocation concealment (selection bias)	Low risk	The study clinical pharmacist prepared the study drugs and placebos. The midwife research assistants received opaque envelopes with affixed study codes, containing both an injection (1 mL of oxytocin 10 IU or its placebo) an 3 pills (misoprostol 600 mg or its placebo).			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "To achieve blinding of the participants and assessors, both inactive agents were manufactured and packaged to resemble actual study medicin in terms of shape, size, and colour".			
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.			
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a clean plastic sheet placed under the mother during and after the third stage of labour. The sheet was specifically designed and piloted for the purpose. Blood was then drained into a calibrated container to improve accuracy in blood loss measurement. Furthermore, quote: "mothers were given pre-weighed standard sanitary pads to place in the perineum at all times. These pads were changed and weighed hourly for the first 6 hours, and then every 6 hours until 24 hours postpartum. Blood loss was estimated as 1 mL per g of weight of the pad after subtracting the dry pad weight". Investigators added the estimated blood loss in pads, to the volume of blood already collected with the plastic sheet. To improve consistency in the estimation of blood loss, standardised electronic scales were used to weigh soiled sanitary pads.			
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.			
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov NCT01866241).			
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.			
Funding source	Low risk The study was supported by scholarship funding from the Father Bash Foundation (public funding).				



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Bauejoko 2012					
Methods	2-arm active-controlled double-dummy randomised trial				
Participants	264 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women in the second or third stage of labour, or those with cervical lacerations or coagulopathy.				
Interventions	30 IU of oxytocin administered by an IV bolus + infusion versus 600 mcg plus 20 IU of misoprostol plus oxytocin administered rectally plus by an IV infusion				
Outcomes		e following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu- (mL); vomiting; fever; shivering.			
Notes	Contact with study aut	chors for additional information: no. Additional data from authors: no			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	The randomisation code produced by an independent statistician using a computer-generated random number sequence			
Allocation concealment (selection bias)	Low risk	Used sequentially numbered sealed packets made of identical opaque brown-paper envelopes prepared by the hospital pharmacy			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.			
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.			
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a BRASS-V calibrated drape, quote: "which is a sterile intrapartum blood collection mat with a calibrated receptacle" placed under the mother after the delivery of the baby and immediate clamping of the umbilical cord. The drape included ribbons tied around the abdomen of the mother to optimise blood collection."			
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "6 women from the misoprostol group and 3 from the oxytocin group were excluded from statistical analysis. 5 of these women in the misoprostol group and all 3 in the oxytocin group were excluded because of the occurrence of cervical lacerations in them.			
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.			
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.			
Funding source	Low risk The study was conducted without external funding.				



Methods	2-arm active-controlled	d double-blinded randomised trial
Participants	48 women were randomised in a hospital setting in Canada. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women requiring general anaesthesia, or those with cardiac disease, hypertension or any condition predisposing to uterine atony and PPH, such as placenta praevia, multiple pregnancy, pre-eclampsia, macrosomia, polyhydramnios, uterine fibroids, bleeding disorders, chorioamnionitis, previous uterine atony, previous PPH or allergy/hypersensitivity to oxytocin or ergot derivatives.	
Interventions	250 mcg plus 20 IU of ergometrine plus oxytocin administered by an IV bolus versus 20 IU of oxytocin administered by an IV bolus + infusion	
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL); nausea; vomiting; hypertension; tachycardia; hypotension.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated list of numbers.
Allocation concealment (selection bias)	Low risk	Used consecutively-numbered opaque sealed packets or envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by measurement of hematocrit preoperatively and 48 hours postoperatively.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the institution of the authors.

Bamigboye 1998a

Methods	2-arm placebo-controlled randomised trial	



Bamigboye 1998a (Continued)	
Participants	550 women were randomised in a hospital setting in South Africa. The population comprised women of
	unspecified parity, unspecified whether singleton or multiple pregnancy, at low risk for PPH, who deliv-

o delivered by vaginal delivery. Exclusion criteria were not specified.

Interventions 400 mcg of misoprostol administered rectally versus placebo

> The study recorded the following outcomes: PPH at 1000; additional uterotonics; manual removal of placenta; third stage duration (minutes); diarrhoea; vomiting; shivering; abdominal pain.

Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Outcomes

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random sequence.
Allocation concealment (selection bias)	Low risk	Allocation concealment was by means of sealed, opaque containers containing 400 mg misoprostol or placebo tablets.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "The placebo tablets were similar in size and colour but were not identical in shape to the misoprostol tablets. Blinding of the midwife administering the tablets was therefore not possible".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with an absorbent plastic-backed linen saver and a low-profile plastic "fracture" bedpan placed under the mother. Blood collection in the plastic bedpan continued until 1 hour after delivery of the baby. At 1 hour after delivery, all the blood on the linen saver was scooped into the bedpan with the blood already collected there, and quote: "the total blood was carefully measured". All the used linen savers and vaginal pads were weighed, and the known dry weights of these materials were subtracted from the measured total weight.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Records of 4 of the 550 allocations (all from the placebo group) could not be traced".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Bamigboye 1998b

Methods 2-arm active-controlled randomised trial
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Bamigboye 1998b (Continued)			
Participants	491 women were randomised in a hospital setting in South Africa. The population comprised women of any parity, unspecified whether singleton or multiple pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	400 mcg of misoprostol administered rectally versus 500 mcg and 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random sequence.	
Allocation concealment (selection bias)	Low risk	Allocation concealment was by means of sealed opaque envelopes.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "About halfway through enrolment it was discovered that a small number of women had been excluded from the syntometrine [ergometrine plus oxytocin] group because of hypertension detected after enrolment (thus contraindicating the use of syntometrine [ergo	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were includ-	

Barton 1996

Funding source

Methods	2-arm placebo-controlled randomised trial
Participants	119 women were randomised in a hospital setting in USA. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria were not specified.

Council (public funding).

ed in the analysis, in the groups to which they were randomised.

The study was supported by funding from the South African Medical Research

Low risk



Barton 1996 (Continued)		
Interventions	100 mcg of carbetocin administered by an IV bolus versus placebo	
Outcomes	The study recorded the following outcomes: additional uterotonics.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Baskett 2007

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	622 women were randomised in a hospital setting in Canada. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with placenta praevia, placental abruption, coagulopathy or unstable asthma.	
Interventions	5 IU of oxytocin administered by an IV bolus versus 400 mcg of misoprostol administered orally	
Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; fever; shivering.	



Baskett 2007 (Continued)

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation cards.
Allocation concealment (selection bias)	Low risk	Used sealed, opaque, sequentially numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The packages were prepared by the hospital pharmacy and their active drug unknown to the physicians and nurses".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by a combination of the visual estimation of attending physicians and measurement of blood volume in a kidney dish placed under the mother during the third stage of labour.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Nova Scotia Health Research Foundation (public funding).

Begley 1990

Methods	2-arm controlled randomised trial	
Participants	1429 women were randomised in a hospital setting in Ireland. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, vaginal breech or instrumental delivery, or those with hypertension, epidural anaesthesia, antepartum haemorrhage, placenta praevia, placental abruption, first stage of labour more than 15 hours, "quick" delivery or needing resuscitation.	
Interventions	500 mcg of ergometrine administered IV bolus versus No treatment	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; trar sion; manual removal of placenta; blood loss (mL); third stage duration (minutes); nausea; vomiti Hypertension. Headache. Abdominal pain.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	



Begley 1990 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random number tables were used. The first number was selected from the table and the numbers were then allocated in blocks of 100, following in sequence.
Allocation concealment (selection bias)	Low risk	Used numbered, sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	Low risk	A sterile receiver was placed against the perineum to collect the blood lost and was measured.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No losses but dropouts for change in Hb.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by public funding, or conducted without external funding.

Begum 2015

Methods	2-arm active-controlled randomised trial		
Participants	100 women were randomised in a hospital setting in Bangladesh. The population comprised women o unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by caesarean. Exclusion criteria were not specified.		
Interventions	400 mcg of misoprostol administered sublingually versus 20 IU of oxytocin administered by an IVinfusion		
Outcomes	The study recorded the following outcomes: (No outcome data found)		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Begum 2015 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Bellad 2012

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	652 women were randomised in a hospital setting in India. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or instrumental delivery, or those with medical disorders, in active labour with more than 4 cm dilatation or stillbirths.	
Interventions	400 mcg of misoprostol administered sublingually versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death blood loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Paticipants were assigned to treatment with a 1:1 ratio using computer-generated simple randomisation



Low risk	The study medications and placebos were packaged in appropriately coded envelopes by administrative staff from the department of clinical pharmacy
Low risk	Study participants and caregivers were blinded to treatment allocations.
Low risk	Assessors were blinded to treatment allocations.
Low risk	Investigators appraised blood loss by collection with a BRASS-V calibrated drape placed under the mother before delivery of the baby. Quote: "The calibrated blood collection receptacle was opened after delivery and drainage of amniotic fluid. The blood collected in the drape was transferred to a measuring jar with 10-mL calibrations for accuracy. Blood-soaked swabs were weighed in g, and the known dry weight of the swabs was subtracted; this volume was added to the measured blood volume from the drape (assuming an equivalence of 1 g and 1 mL)". Blood loss was measured at 1 and 2 hours after delivery of the baby.
Low risk	Data were collected completely from all randomised study participants.
Unclear risk	The study protocol was registered retrospectively (ClinicalTrials.gov NCT01373359).
Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Low risk	The study was supported by funding from Jawaharlal Nehru Medical College (the institution of the authors). Study medications were donated by Cipla (misoprostol) and AstraZeneca (oxytocin).
	Low risk Low risk Low risk Unclear risk Low risk

Benchimol 2001

Methods	3-arm controlled randomised trial	
Participants	602 women were randomised in a hospital setting in France. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with gestational age less than 32 weeks, previous PPH, intrauterine fetal death, previous uterine scar, multiple pregnancy or preeclampsia.	
Interventions	No treatment versus 2.5 IU of oxytocin administered by an IV bolus versus 600 mcg of misoprostol administered orally	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; blood loss (mL); change in Hb; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		



Benchimol 2001 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Slips with the words "control," "Syntocinon," and "Cytotec" were placed into envelopes which were then drawn at random upon admission into the delivery room to determine to which group the woman would belong.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by weighing (methods of collecting blood were not reported).
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Bhatti 2014

Bias

Random sequence genera-

tion (selection bias)

2-arm active-controlled randomised trial	
120 women were randomised in a hospital setting in Pakistan. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with medical disorders, multiple pregnancy, instrumental births, stillbirths and over 42 weeks.	
400 mcg of Misoprostol administered sublingually versus 10 IU of Oxytocin administered IM	
The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL); nausea; vomiting; fever; shivering.	
Contact with study authors for additional information: no. Additional data from authors: no	

Support for judgement

1:1 simple randomisation but the sequence generation was not reported in

Authors' judgement

Unclear risk

sufficient detail.



Bhatti 2014 (Continued) Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Visual assessment of blood loss.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Bhullar 2004

Random sequence genera-

Allocation concealment

tion (selection bias)

(selection bias)

Methods	2-arm placebo-controlled randomised trial	
Metrious	z-arm placebo-controlled randomised that	
Participants	756 women were randomised in a hospital setting in USA. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with a bleeding disorder.	
Interventions	200 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 20 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes; vomiting shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	
Risk of bias		
Bias	Authors' judgement Support for judgement	

Agent vials were coded with a number, which had been assigned using a ran-

Used opaque vials containing either a 200 mcg misoprostol tablet or a place-

Low risk

Low risk

dom number table.



Bhullar 2004 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "The placebo tablets were similar in size and colour, but not identical in shape to the misoprostol tablet".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The placebo tablets were similar in size and colour, but not identical in shape to the misoprostol tablet".
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Biswas 2007

Methods	2-arm active-controlled randomised trial	
Participants	100 women were randomised in a hospital setting in India. The population comprised women of gravida 3 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with heart, renal or liver disease, previous caesarean and severe hypertension.	
Interventions	125 mcg of carboprost administered IM versus 200 mcg of ergometrine administered IM	
Outcomes	The study recorded the following outcomes:transfusion; manual removal of placenta; nausea; vomiting; hypertension; fever.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.



Biswas 2007 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Weighed blood clots and vaginal pads before and after use.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Borruto 2009

Methods	2-arm active-controlled randomised trial		
Participants	104 women were randomised in a hospital setting in Italy. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with toxemia, eclampsia or epilepsy.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 10 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; blood loss (mL); vom ing; headache; hypotension; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "The patients were divided in two groups with blinding to the study medication". Blinding of caregivers was not confirmed.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Borruto 2009 (Continued)		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by quote:" a sensitive colorimetric method".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	High risk	The authors, quote: "do not have a financial relationship with the organisation that sponsored the research". No other source(s) of funding for the study were reported.

Boucher 1998

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	60 women were randomised in a hospital setting in Canada. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with heart disease or cardiac arrhythmia, hypertension or liver/renal/endocrine disease.	
Interventions	100 mcg of carbetocin administered by an IV bolus versus 32.5 IU of oxytocin administered by an IV bolus + infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL); nausea; vomiting; headache; fever; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Pandom seguence genera-	Unclear risk	Sequence generation was not reported

NISK OF DIGS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Double-blinded.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by a sensitive colorimetric measurement of the Hb concentration of blood loss collected, quote: "by means of aspira-



Boucher 1998 (Continued)

tion from the operative field [that] began immediately after administration of the study drug and ceased at the time of skin closure. All gauzes used during this timeframe were placed in 15% Lyse solution. All aspirated blood, gauzes, and the reference blood sample were sent to the laboratory for quantification of total blood volume. Blood on gauzes was extracted with Lyse solution, and haemoglobin content was determined with a sensitive colorimetric method adapted to the Cobas FARA analyser. Haemoglobin concentration is proportional to the absorbance of a hydrogen peroxide-activated aminophenazonephenol mixture measured at a wavelength of 500 nm. The inter-assay coefficient of variation averaged 3.3%, and the limit of detection of the assay was 14 mg/dL. The amount of blood collected in gauzes was calculated with the following formula: blood loss in dL = amount of haemoglobin in surgical gauzes in mg/haemoglobin concentration in mg/dL before caesarean section. Total blood loss was calculated by means of summing the volumes of blood aspirated and collected with gauzes". Ouote: "3 patients who received general instead of epidural anaesthesia were

Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "3 patients who received general instead of epidural anaesthesia were excluded from the study and did not receive the study medication" but the study report did not specify whether these exclusions occurred before or after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	High risk	The study was supported by funding from Ferring Pharmaceuticals.

Boucher 2004

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	164 women were randomised in a hospital setting in Canada. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women younger than 18 years old, or those without known PPH risk, known or suspected coagulopathy, heart disease or cardiac arrhythmia, chronic liver/renal/endocrine disease or hypersensitivity to study drugs.	
Interventions	100 mcg of carbetocin administered IM versus 10 IU of oxytocin administered IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; blood loss (mL); change in Hb; nausea; vomiting; headache; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generaterd randomisation codes using a block size of 4.
Allocation concealment (selection bias)	Unclear risk	Used consecutively-numbered sealed envelopes.



Boucher 2004 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was 'double-blind': Quote: "for each study subject, kits containing both the study medication and a placebo were prepared in the hospital pharmacy according to the randomisation schedule, to assure blinding of the clinical staff".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	164 women were randomised in the study, but 4 were excluded because they did not receive the study medication (3 oxytocin and 1 carbetocin) after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	High risk	The study was supported by funding from Ferring Pharmaceuticals.

Bugalho 2001

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	700 women were randomised in a hospital setting in Mozambique. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour.	
Interventions	400 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); third stage duration (minutes); diarrhoea; vomiting; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Neither the investigators nor the nurses participating in the study had access to the codes until the completion of the study".



Bugalho 2001 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss with a metallic collector placed under the mother, from immediately after delivery of the baby until the mother was removed from the delivery room.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "A few subjects were excluded after randomisation for emergency caesarean section or incomplete data collection".
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of retained placenta were omitted).
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Low risk	This study was financed by the Maputo Central Hospital (the institution of the authors) and the Special Program on Research and Research Training in Human Reproduction of the WHO (public funding).

Butwick 2010

Methods	5-arm placebo-controlled randomised trial		
Participants	75 women were randomised in a hospital setting in the USA. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with active labour, ruptured membranes, drug allergy, multiple pregnancy, significant obstetric disease, risk factors for PPH (abnormal placentation, fibroids, previous PPH, previous classical uterine incision), coagulopathy or thrombocytopenia.		
Interventions	Placebo versus 5, 3, 1, or 0.5 IU of oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL); nausea; vomiting; tachycardia; hypotension.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised using Microsoft Excel-generated random number allocations
Allocation concealment (selection bias)	Unclear risk	Used opaque envelopes containing group assignments
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The obstetrician and anaesthetist involved in each case were blinded to the oxytocin dose assignments".



Butwick 2010 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss quote: "by estimating blood collected by suction and by calculating the weight of blood on surgical swabs".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "75 patients were enrolled, and 74 patients completed the study; 1 patient was excluded due to protocol violation (obstetrician request for supplemental oxytocin despite adequate uterine tone)".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Low risk	The study was supported by funding from the Department of Anesthesia of the Stanford University School of Medicine (the institution of the authors).

Caliskan 2002

Methods	4-arm active-controlled double-dummy randomised trial		
Participants	1633 women were randomised in a hospital setting in Turkey. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with gestational age less than 32 weeks or hypersensitivity to prostaglandins.		
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered rectally plus by an IV infusion versus 400 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered by an IV infusion versus 200 mcg plus 10 IU of ergometrine plus oxytocin administered IM plus by an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; change in Hb; third stage duration (minutes); diarrhoea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The randomisation was based on a table of computer-generated blocks of random numbers.
Allocation concealment (selection bias)	Low risk	Used sealed consecutively-numbered opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "To overcome the limitation of the shape of the placebo, all medications were applied by midwives, but residents who treat the birth and the third stage of labour were blinded to the identity of medication. Only the midwife who applied the medication opened the envelope once to read the code and then transferred the randomisation code into another identical envelope. The identities of the placebo and active medication were also concealed from care-



Caliskan 2002 (Continued)		givers and residents who followed up the patient for the next 24 hours. The randomisation code was not broken until study completion."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "To overcome the limitation of the shape of the placebo, all medications were applied by midwives, but residents who treat the birth and the third stage of labour were blinded to the identity of medication. Only the midwife who applied the medication opened the envelope once to read the code and then transferred the randomisation code into another identical envelope. The identities of the placebo and active medication were also concealed from caregivers and residents who followed the patient for the next 24 hours. The randomisation code was not broken until study completion."
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a sterile steel bedpan and plastic bed linen. Gauzes and pads were also collected and weighed until 1 hour after delivery of the placenta.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "The study enrolled 1633 women, but the data for 27 women were excluded because of lack of predelivery (n = 13) or postpartum (n = 14, short hospital stay) haemoglobin concentrations".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Caliskan 2003

Methods	4-arm active-controlled double-dummy randomised trial
Participants	1800 women were randomised in a hospital setting in Turkey. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with gestational age less than 32 weeks or hypersensitivity to prostaglandins.
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered orally plus by an IV infusion versus 400 mcg of misoprostol administered orally versus 10 IU of oxytocin administered by an IV versus 200 mcg plus 10 IU of ergometrine plus oxytocin administered IM plus by an IV infusion
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; vomiting; fever; shivering.
Notes	Contact with study authors for additional information: no. Additional data from authors: no
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated without any blocking or stratification.
Allocation concealment (selection bias)	Low risk	Used sealed, consecutively-numbered opaque envelopes.



Caliskan 2003 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The placebo tablets were similar in size and colour but were not identical in shape to the misoprostol tablets. To minimise this limitation, the preparation and administration of the medication were carried out by a midwife who had not been involved in the management of the patient except for drug administration."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The placebo tablets were similar in size and colour but were not identical in shape to the misoprostol tablets. To minimise this limitation, the preparation and administration of the medication were carried out by a midwife who had not been involved in the management of the patient except for drug administration."
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a sterile steel bedpan and plastic bed linen from immediately after delivery. Gauzes and pads were also collected 1 hour after delivery of the placenta and weighed.
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "The data for 226 patients were excluded because of caesarean deliveries performed after randomisation ($n = 206$) and the lack of predelivery ($n = 6$) or postpartum ($n = 14$, short hospital stay) haemoglobin concentrations."
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Carbonell 2009

Methods	2-arm active-controlled randomised trial	
Participants	1410 women were randomised in a hospital setting in Spain. The population comprised women of parity 4 or less, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or instrumental delivery, or those with gestational age less than 32 weeks, coagulopathy, Hb less than 80 g/L, liver or kidney disorder, grand multiparity (5 or more), hypersensitivity or any contraindication for use of prostaglandins.	
Interventions	400 mcg and 200 mcg plus 10 IU of misoprostol plus oxytocin administered sublingually and rectally plus IM versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity; intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); NNU admissions; diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random assignments generated by computer.



Low risk	Used sequentially-numbered, opaque, sealed envelopes prepared by people not related to the study. This process was supervised by an analyst. Every morning a secretary received the sealed envelopes for distribution and this process was monitored by someone working on the study.
Unclear risk	Blinding (of study participants and caregivers) was not reported.
Unclear risk	Assessor blinding was not reported.
Low risk	After delivery of the baby, investigators appraised blood loss by collection with a sterile waterproof cloth placed under the mother, to channel blood into a bottle with capacity of 2 L: the volume reading was collected once beyond the third stage of labour.
Low risk	1410 women were randomised in the study, but 10 were excluded because they did not receive the allocated agents (3 in the misoprostol plus oxytocin group and 7 in the oxytocin group) after randomisation.
Unclear risk	The protocol of the study was unavailable for verification.
High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Low risk	The study was supported by the Science and Ethics Committee of the Hospital Eusebio Hernandez in Habana, Cuba in conjunction with the Clinica Mediterranea Medica in Valencia, Spain (the institutions of the authors).
	Unclear risk Unclear risk Low risk Unclear risk High risk

Carrillo-Gaucin 2016

Methods	2-arm active-controlled randomised trial		
Participants	120 women were randomised in a hospital setting in Mexico. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women with allergies to oxytocin or carbetocin or previous coagulation disorder.		
Interventions	unspecified dose of carbetocin administered by an unspecified route versus unspecified dose of oxytocin administered by an unspecified route		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Simple randomisation but sequence generation was not reported in sufficient detail.	



Carrillo-Gaucin 2016 (Continue	ed)	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	It is mentioned that the study was double blinded but blinding methods (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	There were 3 losses to follow-up.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Cayan 2010

Methods	4-arm controlled randomised trial	
Participants	160 women were randomised in a hospital setting in Turkey. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with thyroid disorder, inflammatory bowel disease or other bowel diseases, previous bariatric surgery or hypersensitivity to prostaglandins.	
Interventions	200, 400, or 600 mcg of misoprostol plus oxytocin administered rectally plus by an IV infusion versus 10 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.



Cayan 2010 (Continued)		
Blinding of participants U and personnel (perfor- mance bias) All outcomes	Jnclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome as- sessment (detection bias) All outcomes	Jnclear risk	Assessor blinding was not reported.
Objective assessment of Ublood loss	Jnclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data L (attrition bias) All outcomes	ow risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Jnclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis L	ow risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source U	Jnclear risk	Source(s) of funding for the study were not reported.

Chalermpolprapa 2010

Methods	2-arm placebo-controlled randomised trial	
Participants	120 women were randomised in a hospital setting in Thailand. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by caesareans. Exclusion criteria were not specified.	
Interventions	Unspecified dose of misoprostol plus oxytocin administered by an unspecified route versus unspecified dose of oxytocin administered by an unspecified route	
Outcomes	The study recorded the following outcomes: (No outcome data found)	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias)	Unclear risk	Assessor blinding was not reported.



Chalermpolprapa 2010 (Continued)

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Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chandhiok 2006

Methods	2-arm cluster-controlled randomised trial	
Participants	1200 women were randomised in a community setting in India. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancy, known systemic disease or previous uterine surgery, or who were designated as high risk and scheduled for transfer to an advanced care facility at the time of labour.	
Interventions	600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); third stage duration (minutes; nausea; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Randomisation process not explained in sufficient detail.
Allocation concealment (selection bias)	Low risk	Randomisation process not explained in sufficient detail but lack of allocation concealment usually not an issue in cluster trials.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not applicable.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not applicable.



Chandhiok 2006 (Continued)		
Objective assessment of blood loss	Low risk	Immediately after the cord was clamped and cut, the paramedical worker in both groups placed a calibrated blood collection drape (BRASS-V drape) under the women's buttocks for quantification of blood loss. This consists of a plastic sheet to which a funnelled pouch is attached. The volume of blood collected in the first hour was recorded. In the event of persistent bleeding, another measurement was made at the end of 2 hours.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	This ICMR Task Force study was funded in part by the WHO Country Office, New Delhi; Cipla Pharmaceuticals provided the misoprostol tablets.

Chaudhuri 2010

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women undergoing caesarean section for cord prolapse or bradycardia, or those with cardiovascular, respiratory, liver or haematological disorders or known hypersensitivity to prostaglandins.	
Interventions	800 mcg of misoprostol administered rectally versus 40 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity; intensive care admissions; additional uterotonics; transfusion; death; blood loss (mL); change in Hb; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised using computer-generated random numbers in a 1:1 ratio
Allocation concealment (selection bias)	Low risk	The packets containing the 2 drugs were sealed and opaque, and could not be identified by the surgeons and anaesthetists
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The packets containing the 2 types of drug were sealed and opaque, and could not be identified by the surgeons and anaesthetist".
Blinding of outcome assessment (detection bias)	Low risk	Assessors were blinded to treatment allocations.



Chaudhuri 2010 (Continued)

All outcomes

Objective assessment of blood loss	Low risk	Investigators appraised intraoperative blood loss by collection with a suction bottle for volumetric measurement, combined with linen savers and mops weighed before and after delivery. They added the approximate volume of the contents of the suction bottle (a) to the difference in weight between dry (b) and soaked (c) linen savers and mops (1 g equivalent to 1 mL). Amniotic fluid volume (d) was calculated by multiplying amniotic fluid index by 30 mL. Finally, intraoperative blood loss was determined by subtracting amniotic fluid volume from approximate blood loss ((a + (c - b)) - d). Furthermore, investigators appraised postoperative bleeding over the next 8 hours by weighing soaked pads and subtracting the dry weight.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "4 women in group 1 [misoprostol] and 6 women in group 2 [oxytocin] were excluded from the analysis: 4 women required conversion to general anaesthesia, 5 women had traumatic intraoperative bleeding (extension of lower segment incision or broad ligament"
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (CTRI 2009/091/000075).
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chaudhuri 2012

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	530 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing augmentation of labour, caesarean section or instrumental delivery, or those with risk factors for PPH, including BMI more than 30, grand multiparity (5 or more), polyhydramnios, fetal macrosomia, antepartum haemorrhage, prolonged labour, previous PPH, Hb less than 80 g/L, severe pre-eclampsia, asthma or coagulopathy.		
Interventions	400 mcg of misoprosto	400 mcg of misoprostol administered sublingually versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death. Blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random number sequence.	
Allocation concealment (selection bias)	Low risk	Used pre-prepared sealed and opaque packet.	



Chaudhuri 2012 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The misoprostol and placebo tablets were similar in size, shape, and colour. The ampoules of oxytocin and placebo were also similar. Selection, enrolment, and randomisation were done by the resident doctors, whereas preparation of packets and confidential record maintenance was done by the labour room nursing staff in charge."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Quote: "Investigators appraised blood loss by collection with specially designed, pre-weighed absorbent thick cotton pads with plastic lining, placed under the mother. Blood clots, if any, were expressed from the vagina into a polythene bag. Any episiotomy wound was repaired immediately, and the swabs used for the purpose of episiotomy were not included in blood loss assessment. If necessary, pads were replaced during the observational hour after delivery. Then the soaked pad(s) and the blood clots were weighed. "The specific gravity of blood being 1.08, the amount of blood lost in mL was approximately equal to the weight in g".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "2 women in the study group and 1 woman in the control group refused sublingual administration of the drug".
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (CTRI 2009/091/000672).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chaudhuri 2015

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	396 women were randomised in a hospital setting in India. The population comprised women of parity 5 or less, either singleton or multiple pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women requiring conversion to general anaesthesia, or those with cardiovascular, hepatic, or haematologic disorders or any contraindication for the use of misoprostol or oxytocin.		
Interventions	400 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IM bolus and IV infusion versus 20 IU of oxytocin administered IM bolus plus an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL); change in Hb; diarrhoea; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Chaudhuri 2015 (Continued)		
Random sequence generation (selection bias)	Low risk	Randomisation was done using a computer-generated random number sequence and blocks of size 8.
Allocation concealment (selection bias)	Low risk	Assignments were contained in sealed, opaque and sequentially-numbered packets.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Randomisation and confidential record maintenance were performed by residents who were not involved in the trial, and the operation theatre midwife prepared the sealed packets and allocated and administered the drugs. Thus, clinicians, investigators, data analysts, and participants were masked to the treatment allocation."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised intraoperative blood loss from after delivery of the placenta. Blood was collected with a suction bottle, linen savers and mops: the dry weights of these materials were subtracted from the soaked weights, and the total volume of intraoperative blood loss calculated on the basis that 1 g is equivalent to 1 mL. Investigators also appraised postoperative blood loss by weighing soaked pads.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (CTRI 2013/05/003645).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chaudhuri 2016

Methods	2-arm placebo-controlled randomised trial
Participants	288 women were randomised in a hospital setting in India. The population comprised women of parity 5 or less, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women who had caesareans or instrumental birth, known hypersensitivity to misoprostol and/or oxytocin, major cardiovascular, hepatic, or haematological disorders or intrauterine fetal death or stillbirth.
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered sublingually plus IM versus 10 IU of oxytocin administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; diarrhoea; fever; shivering.
Notes	Contact with study authors for additional information: no. Additional data from authors: no
Risk of bias	



Chaudhuri 2016 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Used a computer-generated random number sequence and block randomisation (blocks of 6–8).
Allocation concealment (selection bias)	Low risk	Used sealed, opaque, and sequentially numbered packets.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants, investigators, and data analysts were masked to group assignment.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Participants, investigators, and data analysts were masked to group assignment.
Objective assessment of blood loss	Low risk	Linens soaked with amniotic fluid were removed soon after delivery of the newborn, and a pre-weighed thick cotton pad with plastic lining was placed under the buttocks. All blood clots were removed from the vagina and kept in a plastic bag. The pad was replaced if completely soaked during the 1-hour observation period. Episiotomies were repaired immediately after complete delivery of the placenta, and cotton swabs used during this procedure were not included in the blood loss assessment. The difference in weight between the soaked and dry pad was added to the weight of blood clots to calculate the total blood loss (1mL was considered equal to 1 g given the specific gravity of blood of 1.08).
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	Registered with Clinical Trial Registry India (Registration No. CTRI/2014/03/004491).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chhabra 2008

Methods	3-arm active-controlled randomised trial
Participants	300 women were randomised in a hospital setting in India. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing augmentation of labour, caesarean section or instrumental delivery, or those with grand multiparity (more than 5), multiple pregnancy, pregnancy-induced hypertension, antepartum haemorrhage, previous caesarean, Hb less than 80 g/L, other obstetric problems or known hypersensitivity to prostaglandins.
Interventions	100 or 200 mcg of misoprostol administered sublingually versus 200 mcg of ergometrine administered by an IV bolus



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Outcomes The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu-

sion; death; blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; headache;

fever; shivering.

Notes Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Used random number tables.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss by quote: "measuring blood and blood clots collected in sponges".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Choy 2002

Methods	2-arm active-controlled randomised trial		
Participants	991 women were randomised in a hospital setting in Hong Kong. The population comprised women of parity 3 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with medical conditions that precluded the use of ergometrine, such as pre-eclampsia, cardiac disease or conditions that required prophylactic oxytocin infusion after delivery such as grand multiparity (4 or more) or presence of uterine fibroids.		
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 10 IU of oxytocin administered by an IV bolus		



CI	hον	2002	(Continued)

Outcomes The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; trans-

fusion; manual removal of placenta; blood loss (mL); change in Hb; nausea; vomiting; hyperten-

sion; headache.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number.
Allocation concealment (selection bias)	Low risk	Used sealed consecutively-numbered opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The preparation and administration of the medication was carried out by a second midwife who was not involved in the management of the patient except for the drug administration. The medical attendant who delivered the baby was not informed of the type of oxytocics used."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss quote: "by measuring the amount of blood clots and weighing the towels and swabs used".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Chua 1995

Methods	2-arm active-controlled randomised trial	
Participants	115 women were randomised in a hospital setting in Singapore. The population comprised women of unspecified parity, a singleton pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	
Interventions	125 mcg of carboprost administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM	
Outcomes	The study recorded the following outcomes: additional uterotonics; manual removal of placenta; diarrhoea.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	



Chua 1995 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised by a random number table.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	All blood and blood clots lost in the first 2 hours after delivery were collected by mopping the blood and clots with absorbent paper, and collect the paper in a plastic bag. The bags were sent to the laboratory for processing within 2 hours of completion of blood collection.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	115 women were randomised in the study, but 3 were excluded because they gave birth precipitously before preparing the bed for accurate collection of blood after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Cook 1999

Methods	3-arm active-controlled randomised trial		
Participants	930 women were randomised in a hospital setting in Australia, Papua and China. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section, or those with coagulopathy, asthma, heart disease, severe renal disease, epilepsy or hypertension.		
Interventions	400 mcg of misoprostol administered orally versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 10 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			



Cook 1999 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was by random number list in blocks of 20 with a separate randomisation for each centre.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered sealed security (opaque) envelopes containing the appropriate drug label for each centre.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss by combining "estimated" and "measured" values according to the standard clinical practice of each study centre. The "estimated" blood loss was judged by the attending senior midwives and/or clinicians. The "measured" blood loss was calculated as the actual volume of blood collected in a calibrated measuring jug, combined with the difference in weight between dry and blood-stained undersheets and sanitary pads.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were not collected completely from 67 study participants: quote: "the main reasons for exclusion prior to randomisation, and following randomisation but before treatment, were the need for caesarean section and development of hypertension, either before or during labour."
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Dabbaghi Gale 2012

Methods	2-arm active-controlled randomised trial		
Participants	269 women were randomised in a hospital setting in Iran. The population comprised women of parity less than 3, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with previous PPH, asthma, clotting disorders, placental abruption, PPH due to lacerations, or those requiring instrumental delivery or caesarean section.		
Interventions	400 mcg of misoprostol administered orally versus 10 IU of oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: (No outcome data found)		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Random sequence genera- Un tion (selection bias)	clear risk	Sequence generation was not reported.
Allocation concealment Un (selection bias)	clear risk	Allocation concealment was not reported.
Blinding of participants Un and personnel (perfor- mance bias) All outcomes	clear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	clear risk	Assessor blinding was not reported.
Objective assessment of Un blood loss	clear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data Un (attrition bias) All outcomes	clear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	clear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis Un		The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source Un	clear risk	Source(s) of funding for the study were not reported.

Dansereau 1999

Methods	2-arm active-controlled double-blinded randomised trial		
Participants	694 women were randomised in a hospital setting in Canada. The population comprised women of parity 5 or less, either singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women undergoing general anaesthesia or requiring a classical uterine incision, or those with heart disease, chronic hypertension requiring treatment, liver, renal, or endocrine disorders, coagulopathy, placenta praevia or placental abruption.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 25 IU of oxytocin administered by an IV bolus + infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; change in Hb; nausea; vomiting; headache; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		
Risk of bias			

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation code, stratified by centre and with use of random blocks of 2.



Dansereau 1999 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "All physicians and nurses involved, all investigators and their staff, and all sponsor representatives were kept blinded to the treatment codes at all times".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	694 women were enrolled in the study, but 59 were excluded because of withdrawals ($n = 5$) or protocol violations ($n = 54$) after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	High risk	The study was supported by funding from Ferring Pharmaceuticals.

Dasuki 2002

Methods	2-arm active-controlled randomised trial		
Participants	196 women were randomised in a hospital setting in Indonesia. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	600 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: blood loss (mL); third stage duration (minutes); shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.



Dasuki 2002 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

de Groot 1996

Methods	3-arm placebo-controlled randomised trial		
Participants	371 women were randomised in a hospital and community setting in the Netherlands. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or instrumental delivery, requiring tocolysis or those who refuse to take part or with cardiac disease, multiple pregnancy, non-cephalic presentation, polyhydramnios, coagulopathy, stillbirth, antepartum haemorrhage, Hb less than 4.8 mmol/L or previous complication in third stage.		
Interventions	Placebo versus 5 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation list.
Allocation concealment (selection bias)	Low risk	Used identical study boxes. Care was taken that no difference could be seen or heard between the packages of the ergometrine/placebo tablets and the oxytocin ampoules.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study made use of placebo tablets to minimise detection bias between the placebo and the oral ergometrine arm but also included an unblinded oxytocin arm and the comparison of oxytocin versus placebo was unblinded.
Blinding of outcome assessment (detection bias)	Unclear risk	Assessor blinding was not reported.



de Groot 1996 (Continued)

All outcomes

Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a "fresh" perineal pad placed under the mother from immediately after birth until 1 hour after the delivery of the placenta. The difference in the weight of the pad before and after delivery was calculated on the basis that 1 g is equivalent to 1 mL of blood. "During delivery some blood was usually spattered on the drapes and gowns of the attendants, although attempts were made to minimise such losses. This gave a constant error of approximately 10%. In addition, the placental interstices contain maternal blood (about 9% of placental weight). As systematic overestimations (amniotic fluid) and underestimations (blood loss) are likely to be equally distributed among the groups, no corrections have been made for them".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "4 women with exclusion criteria were entered erroneously (3 forceps, 1 augmentation). They are considered as non-participants".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Del Angel-Garcia 2006

Methods	2-arm active-controlled randomised trial
Participants	152 women were randomised in a hospital setting in Mexico. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by an unspecified method. Exclusion criteria were not specified.
Interventions	unspecified dose of oxytocin administered by an unspecified route versus unspecified dose of carbetocin administered by an unspecified route
Outcomes	The study recorded the following outcomes: (No outcome data found)
Notes	Contact with study authors for additional information: no. Additional data from authors: no (Abstract only)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.



Del Angel-Garcia 2006 (Continu	ued)	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Derman 2006

Methods	2-arm placebo-control	lled randomised trial
Participants	any parity, a singleton teria comprised wome dia's ministry of health vaginal delivery, or tho hypertension, multiple	domised in a community setting in India. The population comprised women of pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion crien at high risk and inappropriate for home or community births according to Inguidelines including those undergoing elective caesarean section or breech ose previous caesarean section, Hb less than 80 g/L, antepartum haemorrhage, a pregnancy, history of previous antepartum or PPH, retained placenta, uterine eart disease, seizures, placenta praevia, asthma or contraindications to misopros-
Interventions	600 mcg of misoprostol administered orally versus placebo	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study aut	thors for additional information: yes. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Generated randomisation list with a random block size by the data co-ordinating centre and was stratified by the midwife.

(selection bias)

The envelopes were numbered and each envelope had a 5-digit code number assigned to it. The first 2 digits were the auxiliary nurse midwife number, followed by a sequence number beginning with 001 and ending with 100, assigned to the individual participant. Non-distinguishable envelopes in batch-

es of 100 were distributed to each of the midwifes affiliated with the 4 selected primary-health centres.



Derman 2006 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The identical placebo was specifically manufactured for the study".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a polyurethane blood collection drape placed under the mother from immediately after birth until 1 hour after delivery of the baby. The blood collection drape included a calibrated receptacle specifically developed for the study. In the event of persistent bleeding beyond 1 hour, the drape was removed at 1 hour, blood loss measured, and a new drape used with a second measurement made at 2 hours.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov NCT00097123).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the National Institute of Child Health and Human Development (public funding) and the Bill and Melinda Gates Foundation (public funding).

Dhananjaya 2014

Methods	2-arm active-controlled	d randomised trial
Participants	100 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with grand multiparity (not defined), rhesus negative blood group, cardiac disease, diabetes, bleeding disorder, precipitated labour, overdistended uterus, traumatic PPH, PROM/chorioamnionitis, intrauterine death, previous caesarean section/scar on uterus or inability to obtain the informed consent.	
Interventions	10 IU of oxytocin admir	nistered IM versus 200 mcg of ergometrine administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; blood loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; headache.	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Systematic random sampling method.



Allocation concealment (selection bias) Blinding of participants and personnel (performance bias) All outcomes Blinding of outcome assessment (detection bias) All outcomes Unclear risk Assessor blinding was not reported. Assessor blinding was not reported. Assessor blinding was not reported. Dijective assessment of blood loss Unclear risk Investigators appraised blood loss by collection with drapes that were weighed together with mops and clots, and by measurement of Hb concentration and haematocrit of a sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours a	Dhananjaya 2014 (Continued)		
and personnel (performance bias) All outcomes Blinding of outcome assessment (detection bias) All outcomes Unclear risk Assessor blinding was not reported. Dipective assessment of blood loss Unclear risk Investigators appraised blood loss by collection with drapes that were weighed together with mops and clots, and by measurement of Hb concentration and haematocrit of a sample of venous blood before delivery and 24 hours after the birth. A sample of venous blood before delivery and 24 hours after the birth was also collected, for Hb and haematocrit measurement quote: "as an objective index of blood loss". Incomplete outcome data (attrition bias) All outcomes Selective reporting (reporting (reporting bias) Unclear risk The protocol of the study was unavailable for verification. The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		Unclear risk	Allocation concealment was not reported.
Objective assessment of blood loss Low risk Investigators appraised blood loss by collection with drapes that were weighed together with mops and clots, and by measurement of Hb concentration and haematocrit of a sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after birth was also collected, for Hb and haematocrit measurement quote: "as an objective index of blood loss". Incomplete outcome data (attrition bias) All outcomes Selective reporting (reporting bias) Unclear risk The protocol of the study was unavailable for verification. The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	and personnel (perfor- mance bias)	Unclear risk	Blinding (of study participants and caregivers) was not reported.
blood loss weighed together with mops and clots, and by measurement of Hb concentration and haematocrit of a sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after the birth was also collected, for Hb and haematocrit measurement quote: "as an objective index of blood loss". Incomplete outcome data (attrition bias) All outcomes Selective reporting (reporting (reporting bias) Unclear risk The protocol of the study was unavailable for verification. The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	sessment (detection bias)	Unclear risk	Assessor blinding was not reported.
(attrition bias) All outcomes Selective reporting (reporting bias) Intention to treat analysis Unclear risk The protocol of the study was unavailable for verification. The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		Low risk	weighed together with mops and clots, and by measurement of Hb concentration and haematocrit of a sample of venous blood before delivery and 24 hours after birth. A sample of venous blood before delivery and 24 hours after the birth was also collected, for Hb and haematocrit measurement quote: "as
Intention to treat analysis Unclear risk The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	(attrition bias)	Unclear risk	The study authors did not mention any incomplete outcome data.
ly allocated to treatment were included in the analysis, in the groups to which they were randomised.		Unclear risk	The protocol of the study was unavailable for verification.
Funding source Unclear risk Source(s) of funding for the study were not reported.	Intention to treat analysis	Unclear risk	ly allocated to treatment were included in the analysis, in the groups to which
	Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Diallo 2017

Methods	2-arm active-controlled randomised trial	
Participants	304 women were randomised in a hospital setting in Senegal. The population comprised women of un specified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal de livery. Exclusion criteria comprised women who could not give their consent, those requiring a caesard an delivery and those with asthma allergy to misoprostol, pregnancies of less than 36 weeks, tempera ture above 38°C, chorioamnionitis, multiple pregnancy, severe cardiopathy, severe anaemia, clotting disorders, or complex perineal tear.	
Interventions	400 mcg of misoprostol administered orally versus 5 IU of oxytocin administered by an IV bolus	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion' blood loss (mL); change in Hb; diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement Support for judgement	



Diallo 2017 (Continued)		
Random sequence generation (selection bias)	Low risk	A computer-generated randomised sequence.
Allocation concealment (selection bias)	Low risk	Cards assigning patients into groups were placed in envelopes which were then sealed and numbered as and when patients were included.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	If an oxytocin drip was used during labour, it was continued for patients in the "oxytocin" group and replaced by a bottle of 5% glucose solution in the "misoprostol" group. The patient was then attended by the midwife who was not informed of the type of uterotonic administered.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The patient was then attended by the midwife who was not informed of the type of uterotonic administered."
Objective assessment of blood loss	Low risk	The blood lost was collected in a basin placed after the clamping of the umbilical cord and the removal of the amniotic fluid. Episiotomies were repaired immediately after delivery. Blood loss was collected for up to 2 hours after delivery. This blood was transferred into a graduated jar to measure its exact volume.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	No funding sought for this study.

Diop 2016

Methods	2-arm active-controlled randomised trial
Participants	1820 women were randomised in a community setting in Senegal. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with known allergies to prostaglandins or pregnancy complications.
Interventions	600 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM
Outcomes	The study recorded the following outcomes: death; change in Hb; diarrhoea; nausea; vomiting; fever; shivering; maternal satisfaction.;
Notes	Contact with study authors for additional information: yes. Additional data from authors: no
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The computer-generated random allocation was overseen by Gynuity Health Projects, which also assigned clusters. Maternity huts with auxiliary midwives



Diop 2016 (Continued)		located 3–21 km from the closest referral centre were randomly assigned (1:1) by staff at Gynuity Health Projects to either oral misoprostol or oxytocin in Uniject, stratified by reported previous year clinic volume (deliveries) and geographical location (inland or coastal).
Allocation concealment (selection bias)	Low risk	Study drugs were packed into individually numbered single-dose envelopes by staff at Gynuity Health Projects and supplied to maternity huts by ChildFund Senegal.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Objective assessment of blood loss	High risk	The perceived amount of blood loss was documented as "normal", "moderate", or "significant".
Incomplete outcome data (attrition bias) All outcomes	High risk	There were 1820 recruited initially through the clusters but 1412 were included in the analysis and 1049 had data available for the study's primary outcome.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (ClinicalTrials.gov, number NCT01713153).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	This study was funded by the Bill & Melinda Gates Foundation.

Docherty 1981

Methods	2-arm active-controlled randomised trial
Participants	50 women were randomised in a hospital setting in UK. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.
Interventions	10 IU of oxytocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM
Outcomes	The study recorded the following outcomes: blood loss (mL).
Notes	Contact with study authors for additional information: no. Additional data from authors: no
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.



Docherty 1981 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Dutta 2016

Methods	2-arm active-controlled	d randomised trial
Participants	400 women were randomised in a hospital setting in India. The population comprised women of parity 2 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring caesarean section or instrumental delivery, Hb less than 8 g/dL, APH, severe pregnancy-induced hypertension, pre-eclampsia or eclampsia, prolonged labour or precipitate labour, fetal weight > 3.5 kg, polyhydramnios, and medical disorders (cardiovascular disease, diabetes mellitus, thyroid disorders and other coagulation abnormalities.	
Interventions	600 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; transfusion; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering; abdominal pain.	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.



Dutta 2016 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Study is stated to be double-blinded but blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Any blood clot which expressed from the uterus was measured in the calibrated glass container.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Eftekhari 2009

ITERIIAI I 2003			
Methods	2-arm active-controlled	2-arm active-controlled randomised trial	
Participants	100 women were randomised in a hospital setting in Iran. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with multiple pregnancy, prolonged labour more than 12 hours, 2 or more previous caesarean sections, previous uterine rupture, Hb less than 80 g/L,who had a history of heart, renal or liver disorders or had a coagulopathy did not enter the study.		
Interventions	400 mcg of misoprostol administered sublingually versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; blood loss (mL); change in Hb		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	By a simple randomisation method, patients were allocated into 2 equal groups.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	



Eftekhari 2009 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection in a suction bottle, and with drapes and pads beneath the mother. Amniotic fluid was suctioned and measured, and then subtracted from the total volume of the suction bottle. Meanwhile the known dry weight(s) of drapes and pads were subtracted from the soaked weights of these materials. Measurements of blood collected in the suction bottle and on drapes and pads were added together.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of transfusion were omitted).
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

El Behery 2015

<u> </u>		
Methods	2-arm active-controlled double-dummy randomised trial	
Participants	180 women were randomised in a hospital setting in Egypt. The population comprised women of nulliparous, a singleton pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women undergoing elective caesarean section, vaginal delivery or general anaesthesia, or those who are multigravida, or with malpresentation, fetal anomalies, placenta praevia, diabetes, hypertension, pre-eclampsia or cardiac disease.	
Interventions	100 mcg of carbetocin a	administered by an IV bolus versus 20 IU of oxytocin administered by an IV infu-
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL;. change in Hb; headache; fever.	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated code.



El Behery 2015 (Continued)		
Allocation concealment (selection bias)	Low risk	Used sealed, opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote: "double-blinded": "a double dummy system for administration was used".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss quote: "in the usual way (visual estimation, number of used swabs and amount of aspirated blood)".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	180 women were included in the study, but 100 were excluded because 4 had congenital fetal anomalies, 7 cases had placenta praevia, 5 cases were diabetic, 8 had hypertension, 9 had pre-eclampsia, 3 cases were cardiac, 28 cases needs general anaesthesia, 17 cases delivered vaginally and 19 cases delivered by elective caesarean section). It was unclear if these were excluded before or after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

El Tahan 2012

Methods	2-arm placebo-controlled randomised trial		
Participants	382 women were randomised in a hospital setting in Egypt. The population comprised women of parity 3 or less, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with asthma, anaemia, bleeding disorders, cardiac disease, inflammatory disease, bowel disease, multiple pregnancy, pre-eclampsia, placenta praevia, placental abruption, previous APH, previous PPH, grand multiparity (not defined), fibroids, growth restriction, fetal malformations or allergy to prostaglandins.		
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered sublingually plus by an IV bolus versus 10 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; death; blood loss (mL); diarrhoea; vomiting; fever; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		
Risk of bias			
Bias	Authors' judgement Support for judgement		



El Tahan 2012 (Continued)		
Random sequence generation (selection bias)	Low risk	Used a computer-generated randomisation code.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered sealed opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo and misoprostol tables quote: "looked identical in size, colour, and packing".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised intraoperative blood loss by collection in a suction bottle minus sonographically estimated amniotic fluid volume, together with visual estimates of the volume of blood on the floor and the weight differences between dry and used towels, linens, and swabs. Visual estimates were performed by obstetricians blinded to treatment allocation. Towels, linen and swabs were weighed with an electronic scale. Weights were added to volumetric values on the basis that 1 g is equivalent to 1 mL. Investigators appraised postoperative blood loss by weighing bed linen, gowns and perineal pads. Furthermore, blinded investigators estimated blood loss by multiplying maternal blood volume in mL by the difference between preoperative and postoperative hematocrit measurements, all divided by preoperative haematocrit measurements.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "4 patients in the placebo group and 12 patients in the misoprostol group were excluded from the study due to loss to follow-up or missed preoperative haematocrit data".
Selective reporting (reporting bias)	Unclear risk	The study report matches the study protocol that was registered retrospectively (ClinicalTrials.gov NCT01466530).
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Low risk	The study was supported by funding from Mansoura University (the institution of the authors).

El-Refaey 2000

Methods	2-arm active-controlled randomised trial
Participants	1000 women were randomised in a hospital setting in UK. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or water birth, or those with severe asthma.
Interventions	500 mcg of misoprostol administered orally versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes). Diarrhoea. Nausea. Vomiting. Headache. Fever. Shivering. Abdominal pain.



El-Refaey 2000 (Continued)

Notes Contact with study authors for additional information: no. Additional data from authors: no

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Statistician using computer-generated block randomisation with varying block size
Allocation concealment (selection bias)	Low risk	Used opaque, sequentially-numbered sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Elbohoty 2016

Methods	3-arm active-controlled triple-dummy randomised trial
Participants	270 women were randomised in a hospital setting in Egypt. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with hypersensitivity to oxytocin, carbetocin, or prostaglandins; contraindication to treatment with prostaglandins (e.g. glaucoma); history of significant heart disease; severe asthma; epilepsy; history or evidence of liver, renal, or vascular disease; history of coagulopathy, thrombocytopenia, or anticoagulant therapy; HELLP syndrome or eclampsia; placental abruption; or contraindication to spinal anaesthesia.
Interventions	100 mcg of carbetocin administered by an IV bolus versus 400 mcg of misoprostol administered sublingually versus 30 IU of oxytocin administered by an IV bolus + infusion
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; nausea; vomiting; headache; fever; shivering; abdominal pain.
Notes	Contact with study authors for additional information: no. Additional data from authors: no



Elbohoty 2016 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was performed in a 1:1:1 ratio using a computer-generated sequence.
Allocation concealment (selection bias)	Low risk	Numbered, sealed envelopes were prepared, with each envelope containing 1 of the 3 study drugs and placebos for the other 2 drugs. The randomisation protocol was concealed from the research team and the primary investigator contacted a central co-ordinating investigator to identify the envelope to be distributed to each patient.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Tablet placebos, containing hydrogenated castor oil, hypromellose, microcrystalline cellulose, and sodium starch glycolate were prepared to be identical in size, colour, shape, and packing to the tablet study drug. Intravenous placebo ampoules containing normal saline were prepared and were identical in shape and packing to the IV study drugs used. All envelopes were prepared by Sigma Pharmaceuticals and were already sealed when received by the research team.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Consequently, patients, investigators, and data analysts were masked to group assignments and unmasking only occurred after data analysis was completed.
Objective assessment of blood loss	Low risk	Surgical towels were weighed with their wrapping before and after delivery using a highly accurate digital balance. The difference in mass between the dry and soaked towels was calculated. Operative blood loss was calculated using 3 parameters: (A) the volume of the suction bottle contents (mL), (B) the difference in towel mass (g), and © the amniotic fluid volume (mL). Intraoperative blood loss (mL) was calculated as: Intraoperative blood loss = (A + B) –C.
Incomplete outcome data (attrition bias) All outcomes	Low risk	270 women were randomised in the study, but 7 were excluded because they had general anaesthesia (n = 4) or the drug ampoules were damaged after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov: NCT02053922).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Elgafor El Sharkwy 2013

Methods	2-arm active-controlled double-dummy randomised trial
Participants	380 women were randomised in a hospital setting in Egypt. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women undergoing general anaesthesia, or those with coagulopathy, coronary artery disease, hypertension, PPH due to causes other than uterine atony or hypersensitivity to carbetocin.
Interventions	400 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 100 mcg of carbetocin administered by an IV bolus



Elgafor El Sharkwy 2013 (Continued)

Outcomes	The study recorded the following outcomes: severe maternal morbidity: additional uterotonics; trans-
	fusion; death; change in Hb; nausea; vomiting; headache; hypotension; fever; shivering.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number sequence.
Allocation concealment (selection bias)	Low risk	Drugs were in pre-prepared sealed and opaque packets.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Caesarean delivery was performed by 4 senior obstetricians who were blinded to the allocation.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss quote: "in the usual way (visual estimation, number of used swabs and amount of aspirated blood)".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Elsedeek 2012

Methods	2-arm placebo-controlled randomised trial	
Participants	400 women were randomised in a hospital setting in Egypt. The population comprised women of parity 4 or less, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women undergoing their first elective caesarean section, or those unsure of gestation or with hypertension, diabetes, oligohydramnios, abnormal placenta or abnormal laboratory investigations.	
Interventions	400 mcg plus 10 IU of Misoprostol plus Oxytocin administered rectally plus by an intravenous infusion versus 10 IU of Oxytocin administered by an intravenous infusion	
Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb. NNU admissions; fever; shivering.	
	loss (mL); change in Hb. NNU admissions; fever; shivering.	



Elsedeek 2012 (Continued)

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated tables.
Allocation concealment (selection bias)	Unclear risk	Allocation was placed in sealed envelopes until the time of operation.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Attending obstetricians and other caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss from after uterine incision, by collection in 2 separate suction sets administered by a nurse, and by weighing surgical towels before and after each operation.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The study protocol was registered retrospectively (ACTRN 12611000638932).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the institution of the authors, or conducted without external funding.

Enakpene 2007

Methods	2-arm active-controlled randomised trial
Participants	864 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, a singleton pregnancy, at Low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with pre-eclampsia, hypertension, cardiac disease, severe anaemia, asthma, renal/hepatic disorders, grand multiparity (not defined), multiple pregnancy, polyhydramnios, previous PPH, fibroids or contraindications to misoprostol or ergometrine.
Interventions	400 mcg of misoprostol administered orally versus 500 mcg of ergometrine administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering.
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes



Enakpene 2007 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was by simple random selection. An independent statistician generated sets of 4 random letters, which were in boxes, and each box contained 4 separate random allocations which was equivalent to an opaque sealed envelope stratified in a block of 4.
Allocation concealment (selection bias)	Low risk	Used opaque sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study was "single-blinded". The identity of those blinded was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by a combination of careful collection in a receptacle after the delivery of the baby, by visual estimation of blood loss, and by extrapolation of blood loss using the weight difference of the total perineal pad used up to 24 hours postpartum.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of transfusion, chest pain and abdominal pain were omitted).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the National Postgraduate Medical College and Faculty of Obstetrics and Gynecology of the University College Hospital in Ibadan, Nigeria (the institution of the authors).

Ezeama 2014

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	300 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with premature labour (less than 28 weeks), multiple pregnancy, APH, hypertension in pregnancy, severe anaemia or haemoglobinopathy.	
Interventions	10 IU of oxytocin administered IM versus 500 mcg of ergometrine administered IM	



Ezeama 2014 (Co	ontinued)
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Outcomes The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; manual

removal of placenta; blood loss (mL); third stage duration (minutes); nausea; vomiting; hypertension;

headache.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation numbers.
Allocation concealment (selection bias)	Low risk	A person uninvolved with the study prepared the study drugs. The labels on the ampoules (which were similar in size and colour) were removed and the ampoules were placed in opaque sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "A person uninvolved with the study prepared the study drugs: 1-mL ampoules containing either 10 IU of oxytocin (Labtocin; Laborate Pharmaceutical India, Panipat, India) or 0.5 mg of ergometrine (Ergosav; Savorite Pharmaceuticals, Vadodara, India). The labels on the ampoules (which were similar in size and colour) were removed and the ampoules were placed in opaque sealed envelopes, such that only the computer generated randomisation numbers on the envelopes were available to identify the study drug. Both drugs were purchased from a public pharmacy."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with "a fresh large perineal pad with plastic backing". They placed all the gauzes and perineal pads used to absorb the blood into a polythene bag, and subtracted the dry weight from the wet weight. Volume of blood loss was calculated on the basis that 1 g is equivalent to 1 mL.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study protocol was registered (PACTR 201105000292708).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the institution of the authors.

Fahmy 2015

Methods	4-arm active-controlled double-dummy randomised trial	
Participants	200 women were randomised in a hospital setting in Egypt. The population comprised women of parity 4 or less, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with coagulopathy, thrombocytope-	



Fahmy 2015 (Continued)		praevia, history of previous obstetric haemorrhage more than 1 litre, and women ulant and antiplatelets therapy.	
Interventions	10 IU of oxytocin administered by an IV bolus versus 100 mcg of carbetocin administered by an IV bolus		
Outcomes	The study recorded the	e following outcomes: additional uterotonics; transfusion; blood loss (mL).	
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	An online randomisation program (http://ww.randomizer.org) was used to generate random list and to allocate patients into the 4 study groups.	
Allocation concealment (selection bias)	Low risk	Random allocation numbers were concealed in opaque closed envelops but there is no mention of the envelopes being sequentially numbered.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear as a placebo saline infusion is mentioned but no sufficient details of how blinding was achieved.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	The calculated estimated blood loss = Estimated blood volume X (preoperative PCV – postoperative PCV)/preoperative PCV. (Where estimated blood volume = Booking weight (kg) X 85 mL)	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Fahmy 2016

Methods	2-arm active-controlled randomised trial
Participants	60 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a twin pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with hypertension, pre-eclampsia, cardiac, respiratory, renal or liver disease, pre-existing bleeding disorder such as haemophilia and women taking therapeutic anticoagulants, hypersensitivity to carbetocin or oxytocin. Patients with Hb less than 9.5 g% and those who are pregnant with more than 2 babies.



Fahmy 2016 (Continued)			
Interventions	100 mcg of carbetocin	administered by an IV bolus versus 20 IU of oxytocin administered by an IV bolus	
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL).		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Randomisation was performed by using computer-generated program.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Both drugs were prepared preoperatively and coded so that the working investigator and the obstetrician were blinded to the type of drug injected.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Fakour 2013

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	200 women were randomised in a hospital setting in Iran. The population comprised women of nulliparous, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	400 mcg of misoprostol administered sublingually versus 20 IU of oxytocin administered IV		
Outcomes	The study recorded the following outcomes: (No outcome data found)		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		



Fakour 2013 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study used double-dummy.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The study used double-dummy.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Fararjeh 2003

Methods	2-arm active-controlled randomised trial
Participants	97 women were randomised in a hospital setting in Turkey. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section or instrumental delivery, or those with premature labour (less than 37 weeks), postmaturity (more than 43 weeks), grand multiparity (more than 4), twin pregnancy, growth restriction, macrosomia, Hb less than 100 g/L, systemic disorder, prolonged third stage, manual removal of placenta or additional lacerations due to episiotomy or where it took longer than 30 minutes to repair lacerations after episiotomy.
Interventions	400 mcg of misoprostol administered rectally versus 200 mcg plus 10 IU of ergometrine plus oxytocin administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; blood loss (mL); change in Hb.
Notes	Contact with study authors for additional information: no. Additional data from authors: no
Risk of bias	



Fararjeh 2003 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Used urn block randomisation.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with scale vessels, and by subtraction of the dry weight(s) of cloths and pads from the soaked weight(s) of these items.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Fawole 2011

Bias	Authors' judgement Support for judgement		
Risk of bias			
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes, but data not provided separate for each drug used and could not be included in the meta-analysis.		
Outcomes	Could not include in the analysis as could not separate out the patients who received oxytocin from those who received ergometrine.		
Interventions	400 mcg of misoprostol administered sublingually plus 10 IU of oxytocin or 250 mcg to 500 mcg of ergometrine administered IM or by an intravenous bolus (n = 658) or IV bolus versus 10 IU of oxytocin or 250 mcg to 500 mcg of ergometrine administered IM or intravenously (n = 660).		
Participants	1345 parturients were randomised in a hospital setting in Nigeria. The population comprised multiparous women, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered vaginally. Exclusion criteria comprised severe allergic conditions or asthma, agbelow 18 years, pyrexia above 38°C, or abortion of the pregnancy.		
Methods	2-arm placebo-controlled randomised trial.		



Fawole 2011 (Continued)		
Random sequence generation (selection bias)	Low risk	Treatment was allocated in blocks of 6-8 women by the research nurse, who used a computer-generated randomisation sequence.
Allocation concealment (selection bias)	Low risk	The trial drugs were concealed in sealed, sequentially numbered opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Placebo was identical in shape, colour, size, and design.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinded.
Objective assessment of blood loss	Low risk	Blood collection was initiated as soon as possible after administration of the trial medication. A low-profile plastic fracture bedpan was placed below the woman's perineum to collect all subsequent blood loss for a period of 1 hour. Blood collected in the bedpan and all blood-soaked small gauze swabs were emptied into a plastic measuring jar and the volume was measured.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No losses stated by authors but 27 women randomised were not included in the analysis for the primary outcome.
Selective reporting (reporting bias)	Unclear risk	No available protocol.
Intention to treat analysis	Unclear risk	27 women randomised were not included in the analysis for the primary outcome.
Funding source	Low risk	The trial was funded by theMedical Research Council of South Africa.

Fawzy 2012

Methods	3-arm active-controlled randomised trial		
Participants	300 women were randomised in a hospital setting in Egypt, Libya. The population comprised women of nulliparous, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women at high risk for PPH such as multiple pregnancy, polyhydramnios, placenta praevia, diabetes mellitus, renal disorders.		
Interventions	500 mcg of ergometrine administered by an IV bolus versus 200 mcg of misoprostol administered sub- lingually or rectally		
Outcomes	The study recorded the following outcomes: death; blood loss (mL); third stage duration (minutes); shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Fawzy 2012 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Randomly allocated but no further details were reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	No blinding.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Objective assessment of blood loss	High risk	All patients were closely observed for time of placental delivery, amount of blood loss by Hb and haematocrit value pre and immediately post delivery (within 1 hour), {then calculation of estimated blood loss using the following equation EBL = (BV)X(HCTO-HCTf)/HCT where: EBL = estimated blood loss, BV: blood volume= body weight X600 cc KG&HCTO = initial haematocrit HCTf = final haematocrit HCTave = (HCTO + HCTF)/2}
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Fazel 2013

Methods	2-arm active-controlled randomised trial		
Participants	100 women were randomised in a hospital setting in Iran. The population comprised women of parity 3 or less, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with twin pregnancy, fetal distress, pregnancy-induced hypertension, oligohydramnios, polyhydramnios, macrosomia, grand multiparity (4 or more), HELLP syndrome, coagulopathy, asthma, heart/lung/liver disease, previous more than 1 caesarean section, previous myomectomy, previous other abdominal operations, febrile diseases or sensitivity to prostaglandins.		
Interventions	400 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: transfusion; blood loss (mL); nausea; vomiting; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Fazel 2013 (Continued)		
Random sequence generation (selection bias)	Low risk	Using a table of random numbers
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised intraoperative blood loss by collection with an isolated suction. The volume of blood collected in suction was combined with the volume of blood collected in gauzes and gowns: every small gauze soaked with blood was considered to contain 20 mL, and every large gauze soaked with blood 50 mL, and every g increase in the weight of a gown was considered as equivalent to 1 mL of blood.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Kashan University of Medical Sciences (the institution of the authors).

Fekih 2009

Methods	2-arm active-controlled randomised trial
Participants	250 women were randomised in a hospital setting in Tunisia. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women undergoing caesarean section with general anaesthesia, or those with placenta praevia, retroplacental clot, multiple pregnancy, premature labour (less than 32 weeks), intrauterine death, Hb less than 80 g/L, coagulopathy, HELLP syndrome, antepartum haemorrhage, ruptured uterus, previous more than 2 caesareans or other uterine scar, prolonged labour (more than 12 hours) or pyrexia.
Interventions	200 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV bolus and infusion versus 20 IU of oxytocin administered by an IV bolus + infusion
Outcomes	The study recorded the following outcomes: PPH at 1000; transfusion; blood loss (mL); change in Hb; nausea; vomiting; headache; fever; shivering.
Notes	Contact with study authors for additional information: no. Additional data from authors: no



Fekih 2009 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The randomisation was computer-generated.
Allocation concealment (selection bias)	Low risk	A slip of paper was placed inside an opaque, sealed envelope.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised perioperative blood loss as a combination of the volume of liquid in the suction collection jar, and the weight of swabs and pads.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Fenix 2012

2-arm active-controlled double-dummy randomised trial		
75 women were randomised in a hospital setting in Phillipines. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with pre-existing hypertension, pre-eclampsia, diabetes, asth ma, cardiac/renal diseases, coagulopathy, abnormal laboratory tests or allergy to the study medication.		
100 mcg of carbetocin administered by an IV bolus versus 10 IU of oxytocin administered by an IV infusion		
The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb; nausea; vomiting; headache; tachycardia; abdominal pain.		
Contact with study authors for additional information: yes. Additional data from authors: yes		
Authors' judgement Support for judgement		



Fenix 2012 (Continued)		
Random sequence generation (selection bias)	Low risk	Computer-generated code.
Allocation concealment (selection bias)	Unclear risk	Used sealed, consecutively-numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The patient and the principal investigator attending the delivery were blinded to the type of medication administered" [additional information from the authors].
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The patient and the principal investigator attending the delivery were blinded to the type of medication administered" [additional information from the authors].
Objective assessment of blood loss	High risk	Investigators appraised blood loss by visual estimation, not including blood loss considered to result from repair of lacerations.
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "9 women in the carbetocin group and 6 women in the oxytocin group failed to have a paired haemoglobin test to measure the change in haemoglobin 24 hours after delivery because they refused further blood extraction. These 15 women were excluded".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Fu 2003

Fu 2003			
Methods	2-arm controlled randomised trial		
Participants	156 women were randomised in a hospital setting in China. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	400 mcg of misoprostol administered orally versus no treatment		
Outcomes	The study recorded the following outcomes: PPH at 500; blood loss (mL).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	



Fu 2003 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss in the 2 hours after delivery and after all amniotic fluids had been drained, by collection in a small tray and absorption into disposable, sterile, water-resistant gauze. The contents were weighed and volume was determined on the basis that 1.05 g is equivalent to 1 mL of blood. A measuring cup was used to estimate the blood in the tray; blood that soaked into the gauze was measured on the basis that material measuring 10 cm by 10 cm holds 10 mL of blood. These 3 measurements were combined to ascertain total blood loss.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis (Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source U	Unclear risk	Source(s) of funding for the study were not reported.

Fuks 2014

1 UK3 2014		
Methods	2-arm active-controlled double-blinded randomised trial	
Participants	143 women were randomised in a hospital setting in Jamaica. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancy, grand multiparous, intrauterine fetal demise, pre-eclampsia, polyhydramnios, third- or fourth-degree laceration, and caesarean delivery.	
Interventions	600 mcg plus 10 IU of misoprostol plus oxytocin administered rectally plus IM versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: (No outcome data found)	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.



Fuks 2014 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.
Garg 2005 Methods	2-arm active-controller	d randomicod trial
Metrious	Z-arm active-controlled	a randomised triat
Participants		omised in a hospital setting in India. The population comprised women of nulligrancy, at both high and low risk for PPH, who delivered by vaginal delivery. Expectified.
Interventions	600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered by an IV bolus	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; manual removal of placenta; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Randomised in 1:1 ratio by random number sequence.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.

Unclear risk

Blinding of participants

and personnel (perfor-

mance bias)

Blinding (of study participants and caregivers) was not reported.



Garg 200	(Continued)
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ΛI	l outcome	_
Αl	courcome	S

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Gavilanes 2016

Methods	2-arm active-controlled randomised trial
Participants	100 women were randomised in a hospital setting in Equador. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with Hb less than 80 g/L, multiple pregnancy, polyhydramnios, previous uterine rupture, bleeding disorders, intrauterine death or hyperthermia (more than 38.5C).
Interventions	400 mcg of misoprostol administered sublingually versus 10 IU of oxytocin administered by an IV infusion
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; blood loss (mL); nausea; vomiting; headache; shivering.
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.



Gavilanes 2016 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised postoperative blood loss by collection with quote: "suction apparatus and sterile drapes before irrigation" and by weighing the blood collected in abdominal swabs and gauzes with a calibrated scale (Zhongshan Camry Electronic Co Ltd, model EK 4052-E, Guangdong, China). Investigators estimated the volume of blood loss quote: "by subtraction of amniotic fluid at 30 cc per each centimetre reported by amniotic fluid index".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Gerstenfeld 2001

Methods	2-arm placebo-controlled randomised trial
Participants	400 women were randomised in a hospital setting in the USA. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancy, coagulopathy, Hb less than 70 g/L, indication for caesarean section or contraindication to prostaglandin or oxytocin use.
Interventions	400 mcg of misoprostol administered rectally versus 20 IU of oxytocin administered by an IV infusion
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; diarrhoea; nausea; vomiting; shivering.
Notes	Contact with study authors for additional information: Yes. Additional data from authors: No

Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	The randomisation was carried out by an uninvolved party and was determined by a random number sequence.			
Allocation concealment Low risk (selection bias)		The random number sequence was prepared by a third party and was concealed until the patient was enrolled. Packets were prepared in advance of randomisation.			
Blinding of participants Low risk and personnel (performance bias) All outcomes		The random number sequence was quote: "concealed until the patient was enrolled" and "packets were prepared in advance of randomisation".			



Gerstenfeld 2001 (Continued)				
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss (a) by collection with drapes placed under the mother. Each drape included a plastic pouch and measured volume in mL. Meanwhile the dry weights of delivery linen and sponges were subtracted from bloodied weights to determine the volume of blood collected with these materials, on the basis that 1 g is equivalent to 1 mL. The volumes of blood in drapes and linen were added together. Furthermore quote: "if amniotic fluid loss [after placement of the drape] was significant the approximate percentage was recorded on the data sheet and blood loss was adjusted accordingly". Investigators appraised blood loss (b) by estimation of the delivery attendant(s). Investigators appraised blood loss (c) by measurement of Hb and haematocrit values were obtained on admission and on postpartum day 1. The differences between these 2 values were recorded.		
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "Of the 75 women who were excluded from analysis, 73 underwent cesarean deliveries, one woman was discharged to home before delivery, and one had an initial haemoglobin of 6.8 mg/dL".		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.		
Funding source	Unclear risk	Source(s) of funding for the study were not reported.		

Gore 2017

Methods	2-arm active-controlled	2-arm active-controlled randomised trial			
Participants	364 women were randomised in a hospital setting in India. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women of gestational age less than 37 years, polyhydramnios, APH, pre-eclampsia, multiple pregnancy, intrauterine fetal distress, coagulation disorders, asthma, epilepsy, heart disease, kidney disease, severe anaemia with Hb less than 7 g/dL, complicated or eventful first and second stage of labour.				
Interventions	400 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered by an IV bolus				
Outcomes	The study recorded the following outcomes: change in Hb; third stage duration (minutes).				
Notes	Contact with study authors for additional information: no. Additional data from authors: no				
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.			



Gore 2017 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	The evaluation of blood loss was assessed by placing cotton pads under the buttocks prior to the delivery of baby. After the delivery of the placenta the total pads and linen used were weighed in grams. The weight of 1 g of cotton pad or linen was equal to 1 mL (Langford 2000). From this the known dry weight subtracted and the calculated volume added.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The authors report no funding sources.

Gulmezoglu 2001

Methods	2-arm active-controlled double-blinded randomised trial			
Participants	18,530 women were randomised in a hospital setting in Argentina, China, Egypt, Ireland, Nigeria, South Africa, Switzerland, Thailand and Vietnam Nigeria, South Africa, Switzerland, Thailand, and Vietnam. The population comprised women of both nulliparous and multiparous, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective or emergency caesarean section after randomisation, or those with asthma, severe chronic allergic conditions, abortion, pyrexia (more than 38°C) or inability to give consent.			
Interventions	600 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM or by an IV bolus			
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000. Severe maternal morbidity; intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.			
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes			
Risk of bias				
Bias	Authors' judgement Support for judgement			



Low risk	The random allocation schedule was generated centrally at WHO, Geneva, Switzerland, by computer-generated random numbers and was stratified by country. Within the strata, women were individually randomised into 1 of 2 intervention groups with randomly varying block sizes of 4–6 women.
Low risk	The treatment packs were sealed, numbered sequentially, and could only be taken from the dispenser consecutively.
Low risk	Quote: "The treatment packs and their contents were identical in shape, colour, weight, and feel."
Low risk	Assessors were blinded to treatment allocations.
Low risk	Investigators appraised blood loss from the time of delivery of the baby until the third stage of the labour was completed, when the mother was transferred to postnatal care (usually up to 1 hour postpartum). Immediately after the cord was clamped and cut, they passed a flat bedpan or an unsoiled receiver under the mother. The collected blood was poured into a standard measuring jar provided by WHO for volumetric measurement. Quote: "To simplify the procedure small gauze swabs soaked with blood were put into the measuring jar and included in the measurement together with the blood and clots".
Low risk	Investigators excluded quote: "37 and 34 women with emergency caesarean section, and 13 and 4 women lost to follow-up in misoprostol and oxytocin groups, respectively, for blood loss ≥ 1000 mL, and 2 and 4 women without information on the need for additional uterotonics".
Low risk	The study report matches the study protocol that was published in advance.
High risk	Not all study participants were included in the analysis.
Low risk	The study was supported by funding from the UNDP/UNFPA/WHO/World Bank (public funding). Special Programme of Research, Development and Research Training Human Reproduction of WHO. Searle (Skokie, IL, USA) and Novartis (Basel, Switzerland) donated the active and placebo medications used in the trial.
	Low risk Low risk Low risk Low risk Low risk High risk

Gupta 2006

Methods	2-arm active-controlled double-blinded randomised trial
Participants	200 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.
Interventions	600 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); nausea; fever; shivering.



Gupta 2006 (Continued)

Notes Contact with study authors for additional information: no. Additional data from authors: no

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was achieved using computer-generated random tables.
Allocation concealment (selection bias)	Unclear risk	A sealed envelope with a code number was opened when vaginal delivery was imminent. The code was not broken till the end of the study.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was "double-blind". "Each envelope contained either 3 tablets of 200 mcg misoprostol and an ampoule of normal saline or 3 identical looking placebo tablets and an ampoule of 10 IU oxytocin".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a BRASS-V calibrated drape placed under the mother. Pre-weighed gauzes were used to clean any perineal tears or episiotomy. After 1 hour the dry weight of the sponges was subtracted from the soiled weight, and added to the volume of blood collected in the drape on the basis that 1 g is equivalent to 1 mL.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Hamm 2005

Methods	2-arm placebo-controlled randomised trial	
Participants	352 women were randomised in a hospital setting in the USA. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria were not specified.	
Interventions	200 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 20 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	



Hamm 2005 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Low risk	Quote: "The group assignments were available only to the pharmacy. The nurse selected an opaque vial from the drug cabinet that contained either a 200-mg misoprostol tablet or placebo. The vial number (which had been assigned in the pharmacy) and patient identification were sent to the pharmacy."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were unclear.

Harriott 2009

Methods	2-arm active-controlled randomised trial	
Participants	140 women were randomised in a hospital setting in the West Indies. The population comprised women of both nulliparous and multiparous, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with previous PPH, hypertension, previous caesarean, intrauterine death, sepsis/pyrexia (more than 38°C), APH or Hb less than 80 g/L.	
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 400 mcg of misoprostol administered rectally	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb.;third stage duration (minutes); diarrhoea; nausea; vomiting; hypertension; fever; shivering; maternal satisfaction.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	



Harriott 2009 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated block randomisation was used to randomly assign participants.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Both the patient and the midwife conducting the delivery were aware of the drug administered".
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a modified plastic drape placed under the mother from the commencement of the third stage of labour, until 1 hour after delivery. The collection drape measured 168 cm by 84 cm, and contained folded over side-wings (to act as a chute) and a 34-cm collection pouch made by folding the distal end of the drape. Standard sterile drapes were placed above the blood collection drape. Every effort was made to avoid soiling the sterile drapes before delivery of the baby, because they were not weighed. After delivery, overlying sterile drapes were removed to facilitate the use of the collection drape.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Mona Campus and Research Publication Committee of the University of the West Indies (the institution of the authors).

Hernandez-Castro 2016

Methods	2-arm placebo-controlled randomised trial	
Participants	123 women were randomised in a hospital setting in Mexico. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by either elective or emergency caesarean delivery. Exclusion criteria comprised women with hypersensitivity to prostaglandins, hyperthermia, coagulation defects, or history of vaginal bleeding (placental abruption or placenta praevia) and those who required general anaesthesia.	
Interventions	400 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 20 IU of oxytocin administered by an IV infusion	



Hernandez-Castro 2016 (Continued)

Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; transfusion.
Notes	Contact with study authors for additional information: no. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was based on a computer-generated sequence in blocks of 6.
Allocation concealment (selection bias)	Low risk	The drugs were kept in opaque containers, prepared by the hospital's pharmacy department, marked with the number assigned to the patient.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Patients, clinicians, investigators, and data analysts were masked to group assignment is stated but the placebo used was folic acid tablets which are different shape than misoprostol.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Patients, clinicians, investigators, and data analysts were masked to group assignment is stated but the placebo used was folic acid tablets which are different shape than misoprostol.
Objective assessment of blood loss	High risk	Visual estimation of blood loss was performed by the anaesthesiologist.
Incomplete outcome data (attrition bias) All outcomes	Low risk	123 women were randomised in the study, but 3 were excluded because of inadequate drug administration ($n=1$), uterine artery injury ($n=1$) and incorrect fetal weight calculation ($n=1$) after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (ClinicalTrials.gov:NCT01733329).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Hofmeyr 1998

Methods	2-arm placebo-controlled randomised trial	
Participants	500 women were randomised in a hospital setting in South Africa. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing augmentation of labour, or those with hypertension, diabetes or previous caesarean.	
Interventions	400 mcg of misoprostol administered orally versus placebo	
Outcomes	The study recorded the following outcomes:PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; shivering; abdominal pain	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	
Risk of bias		



Hofmeyr 1998 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random sequence, in balanced blocks of 8.
Allocation concealment (selection bias)	Low risk	The containers were ordered according to a computer-generated random sequence, in balanced blocks of 8.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The tablets were either misoprostol 2 x 200 mcg or 2 placebo tablets similar in size and colour but not shape. Efforts to obtain identical placebo tablets were unsuccessful. This method of blinding proved to be effective. In only 1 case did the attending midwife inadvertently catch sight of the tablets.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinded.
Objective assessment of blood loss	Low risk	Within a minute of delivery, investigators removed any linen soiled with amniotic fluid, and placed a fresh, disposable absorbent linen-saver sheet with plastic backing, and a low wedge-shaped plastic "fracture" bedpan under the mother. Quote: "This was found to be a comfortable and efficient way of collecting the great majority of blood lost after delivery, and could be left in place without discomfort even during perineal suturing. When active bleeding had stopped, any blood clots were expressed from the uterus, the bedpan was removed and a sanitary towel was applied. The [volume of] blood in the bedpan was measured in a measuring jug. An hour after delivery, any bloodstained linen-savers and sanitary towels were placed in a plastic bag and weighed in g".
		After subtracting the known dry weights of these materials, the bloodstained weights were added to the volume of blood collected in the bedpan to ascertain the total blood loss in the first hour after delivery.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the South African Medical Research Council (public funding).

Hofmeyr 2001

Methods	2-arm placebo-controlled randomised trial	
Participants	600 women were randomised in a hospital setting in South Africa. The population comprised women of any parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	
Interventions	600 mcg of misoprostol administered orally versus placebo	



Outcomes	The study recorded the following outcomes:PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; diarrhoea; nausea; vomiting; fever; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Random assignments generated by computer in blocks of 18.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque test tubes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Misoprostol and placebo were similar in size and colour but not shape. Efforts to obtain identical placebo tablets were unsuccessful. This method of blinding proved to be effective.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Blinded.
Objective assessment of blood loss	Low risk	Within a minute of delivery, investigators removed any linen soiled with amniotic fluid, and placed a fresh, disposable absorbent linen-saver sheet with plastic backing, and a low wedge-shaped plastic "fracture" bedpan under the mother. Quote: "This was found to be a comfortable and efficient way of collecting the great majority of blood lost after delivery, and could be left in place without discomfort even during perineal suturing. When active bleeding had stopped, any blood clots were expressed from the uterus, the bedpan was removed and a sanitary towel was applied. The [volume of] blood in the bedpan was measured in a measuring jug. An hour after delivery, any bloodstained linen-savers and sanitary towels were placed in a plastic bag and weighed in g".
		After subtracting the known dry weights of these materials, the bloodstained weights were added to the volume of blood collected in the bedpan to ascertain the total blood loss in the first hour after delivery.
ncomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "There were no withdrawals after randomisation and all outcomes were analysed in the allocated group". However the primary outcome data of 1 study participant in the placebo group were unavailable.
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.
ntention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the South African Medical Research Council (public funding) and University of the Witwatersrand (the institution of the authors).



Hofmeyr 2011			
Methods	2-arm placebo-control	led randomised trial	
Participants	1103 women were randomised in a hospital setting in South Africa, Uganda, and Nigeria. The population comprised women of any parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or instrumental delivery, or those who declined participation or were unable to consent, were too ill or distressed to participate or with a not viable pregnancy.		
Interventions	400 mcg plus 10 IU of r tocin administered IM	nisoprostol plus oxytocin administered sublingually plus IM versus 10 IU of oxy-	
Outcomes		The study recorded the following outcomes :PPH at 500; PPH at 1000; manual removal of placenta; death; blood loss (mL); fever; shivering.	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random numbers and was stratified by country in blocks of 6–8.	
Allocation concealment (selection bias)	Low risk	Quote: "The trial medication was provided, and the study drug packs were prepared, by Gynuity Health Projects. When a participant enrolled, the researcher took the next study drug pack from the dispenser and immediately wrote the woman's name both on the pack and in the participant number list, which was kept separate from the case record forms. Enrolment took place when the pack was removed from the pack dispenser. The pack could not be used for another woman or returned to the dispenser."	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was "double-blind". Quote: "The packs were identical in shape, colour, weight, and feel, and contained either 2 tablets of 200 mcg of misoprostol (HRA Pharma, Paris, France) or 2 matching placebo tablets".	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.	
Objective assessment of blood loss	Low risk	Similarly to the study team of Gulmezoglu 2001, investigators appraised blood loss by collection with a fresh non-absorbent sheet and low plastic "fracture" bedpan placed under the mother from as soon as possible after delivery until 1 hour postpartum. Investigators considered that quote: "longer-term blood loss measurement is more difficult to standardise". They transferred the blood collected in the sheet and the bedpan (together with any soaked small gauze swabs) to a measuring jar to ascertain the volume. Alternatively, they collected blood with a plastic sheet placed under the mother immediately after delivery. If bleeding continued beyond 1 hour, investigators restarted collection and measurement until bleeding subsided. Attempts were made to minimise any losses on the drapes and gowns of delivery attendants. In addition, quote: "the placental interstices also contain maternal blood (about 9% of placental weight)."	
		Because overestimations (amniotic fluid) and underestimations (blood loss) were likely to be distributed equally between the 2 study groups, and most would have occurred before the onset of measurement, the data were not corrected.	



Hofmeyr 2011 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Data for the primary outcome were not available for 4 of the 1103 women".
Selective reporting (reporting bias)	High risk	The prospectively registered protocol of the study (ClinicalTrials.gov NCT 00124540) lists some secondary outcomes different to those included the study report (≥ 1000 mL within the first hour only, transfusion, Hb < 8 g/dL 24 hours after delivery).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from Gynuity Health Projects through a grant from the Bill and Melinda Gates Foundation (public funding).

Hoj 2005

Methods	2-arm placebo-controlled randomised trial		
Participants	661 women were randomised in a community setting in Guinea-Bissau. The population comprised women of parity 3 or less, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	600 mcg of misoprostol administered sublingually versus placebo		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Using a list of random numbers.
Allocation concealment (selection bias)	Low risk	Used opaque envelopes that were consecutively-numbered and filled with the study drugs.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Misoprostol and placebo tablets of identical form, size, colour, and packing were produced".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	After delivery of the baby and drainage of the amniotic fluid, investigators placed a clean plastic-lined absorbent drape under the mother. They changed the drape as many times as needed. The mother stayed on the drape or was asked to wear a pad over the next 60 minutes. All drapes and pads were weighed with an electronic scale and the known dry weights were subtracted



Hoj 2005 (Continued)		in order to ascertain the volume of blood loss on the basis that 1 g is equivalent to 1 mL.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Danish Society of Obstetrics and Gynaecology, the Illum Foundation, and the Danish International Development Agency (public funding).

Hong 2007

Methods	2-arm placebo-controlled randomised trial		
Participants	214 women were randomised in a hospital setting in Korea. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at high risk for PPH, who delivered by caesarean (unspecified whether elective or emergency). Exclusion criteria were not specified.		
Interventions	20 IU of oxytocin administered by an IV infusion versus 400 mcg plus 20 IU of misoprostol plus oxytocin administered rectally plus by an IV infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; change in HB; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Placebo is mentioned but insufficient detail is reported to decide on blinding (of study participants and caregivers).
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Placebo is mentioned but insufficient detail is reported to decide on blinding of outcome assessors.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.



Hong 2007 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Humera 2016

Methods	2-arm active-controlled randomised trial
Participants	100 women were randomised in a hospital setting in India. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with pre-eclampsia or eclampsia, previous caesarean, previous retained placenta, APH, coagulation disorder, cardiac diseases, diabetes, hypertension and epilepsy.
Interventions	600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered by an IV bolus
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); third stage duration (minutes); nausea.; vomiting; hypertension; headache; fever; shivering.
Notes	Contact with study authors for additional information: no. Additional data from authors: no

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	After delivery of the baby amniotic fluid was allowed to drain away (if present) and amniotic fluid soaked bed linen covered with dry disposable linen saver, corrugated rubber sheet placed under buttocks, sterile kidney tray placed at the vulva was used to collect blood loss over next 1 hour. Collected blood was measured using a measuring jar, blood clots weighed separately (1 g = 1 mL).



Humera 2016 (Continued)		Blood soaked swabs were weighed, the known dry weight subtracted and the calculated volume added to that of the blood volume of measuring jar.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	No funding sought for this study.

Is 2012

Methods	2-arm active-controlled randomised trial		
Participants	200 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	400 mcg of misoprostol administered rectally versus unspecified of ergometrine administered IM		
Outcomes	The study recorded the following outcomes: third stage duration (minutes); nausea; vomiting; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.



Is 2012 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Jago 2007

Methods	2-arm active-controlled randomised trial
Participants	510 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or instrumental delivery, or those requiring epidural analgesia or with hypertension in pregnancy, existing hypertension, chronic renal disease, diabetes, vascular diseases, cardiac disease, anticoagulation therapy or allergy to ergometrine or oxytocin.
Interventions	500 mcg of ergometrine administered IM versus 10 IU of oxytocin administered by an IV bolus
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; blood loss (mL); hypertension.
Notes	Contact with study authors for additional information: no. Additional data from authors: no

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated list of random numbers.
Allocation concealment (selection bias)	Unclear risk	Used numbers that were labelled on envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.



Jago 2007 (Continued)		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Jangsten 2011

Methods	2-arm controlled randomised trial		
Participants	1802 women were randomised in a hospital setting in Sweden. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section, or those who were non-Swedish speaking or with previous PPH, pre-eclampsia, grand multiparity (more than 4) or intrauterine death.		
Interventions	10 IU of oxytocin administered by an IV bolus versus no treatment		
Outcomes	The study recorded the following outcomes: PPH at 1000; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); maternal satisfaction.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated sequence.
Allocation concealment (selection bias)	Low risk	Used sealed envelopes containing the randomisation group prepared in consecutive order and kept in another unit. At randomisation, midwives phoned the staff at the other unit who opened the envelopes and disclosed the assigned intervention and trial number.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Because of the nature of the study, blinding was not possible for the midwives, but the women were not informed of which management was to be used for them".
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by removing pads soaked with amniotic fluid and placing a dry sanitary pad under the mother, immediately after the birth of the baby. They weighed all sanitary towels and pads before and after use. Blood loss was recorded (a) between the birth of the baby and the expulsion of the placenta, and (b) from expulsion of the placenta up to 2 hours postpartum.
Incomplete outcome data (attrition bias) All outcomes	Low risk	171 randomised women were not included in the study analysis. Among those randomised to receive oxytocin, 4 withdrew consent, 75 had caesareans, and 14 were lost to follow up. In the control group, 2 withdrew consent, 56 had caesareans, and 20 were lost to follow up.



Jangsten 2011 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	The authors excluded 131 randomised study participants from the analysis because they experienced caesarean deliveries.
Funding source	Low risk	The study was supported by funding from the Research and Development Board in Göteborg and Bohuslän, Baby Bag and the SU Foundation in Sweden (public funding).

Jans 2017

Methods	2-arm controlled randomised trial		
Participants	1704 women were randomised in a community setting in the Netherlands. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with indications for a prophylactic approach to the third stage management in primary midwifery care and women with poor command of the Dutch language.		
Interventions	5 IU of oxytocin administered IM versus no treatment		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000.;additional uterotonics; transfusion; third stage duration (minutes); breastfeeding; nausea; vomiting; headache; abdominal pain; maternal well-being.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was carried out by a lottery method Quote: "Randomization was achieved using two numbered and sealed opaque envelopes. Each envelope contained a sticker indicating one of the allotted treatments. When the midwife was confident that the birth would be completed in her care (defined for primigravid women when a large part of the baby's head was presenting and for multiparous women at the beginning of the second stage of labor), the woman herself or someone else designated by her would choose one of the two envelopes."
Allocation concealment (selection bias)	High risk	Allocation concealment was not reported but unlikely to have been implemented with a lottery method of randomisation.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Objective assessment of blood loss	Low risk	Used digital scales, 10 disposable pre-weighed incontinence pads (a small impermeable multilayered sheet with high absorbency) and graduated measuring cups.



Jans 2017 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	1704 women were randomised in the study, but 18 were excluded because of referral to hospital ($n=16$) and were lost to follow-up or withdrew from the study ($n=2$) after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The trial was funded by the Prevention Fund of the Netherlands.

Jerbi 2007

Methods	2-arm controlled randomised trial		
Participants	130 women were randomised in a hospital setting in Tunisia. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with placenta praevia, APH, non-cephalic presentation, intrauterine death, grand multiparity, (more than 5), fibroids, anticoagulation therapy, previous PPH or previous caesarean.		
Interventions	5 IU of oxytocin administered by an IV bolus versus no treatment		
Outcomes	The study recorded the following outcomes: PPH at 1000; transfusion; manual removal of placenta; death; change in Hb; third stage duration (minutes).		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.



Jerbi 2007 (Continued)		
Selective reporting (re- Unclear risk The protocol of th porting bias)		The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were unclear.

Jirakulsawas 2000

Methods	2-arm active-controlled randomised trial		
Participants	140 women were randomised in a hospital setting in Thailand. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; blood loss (mL).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.



Jirakulsawas 2000 (Continued)

Funding source Unclear risk Source(s) of funding for the study were not reported.

Kabir 2015

Methods	2-arm active-controlled randomised trial		
Participants	110 women were randomised in a hospital setting in Bangladesh. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with placenta praevia, multiple pregnancy, placental abruption, hypertensive disorders, pre-eclampsia, cardiac/renal/liver disorders, epilepsy, moderate anaemia (Hb < 9 g/dLl), intrauterine fetal death and unwilling to participate in the study.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 10 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
	A discussion of the state of th		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Used a computer generated randomisation.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Used pre-weighted standardised delivery mat (Quaiyum's mat) and pre- weighted sanitary pads for blood collection after delivery to each of the preg- nant woman to measure blood loss and measured the amount of blood loss in g by digital postal scale.
Incomplete outcome data (attrition bias) All outcomes	High risk	110 women were randomised in the study, but 16 were excluded because of pre-eclampsia ($n = 5$), eclampsia ($n = 5$), placenta praevia ($n = 2$), placental abruption ($n = 2$) and multiple pregnancy ($n = 2$) after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.



Karkanis 2002

(al Kallis 2002			
Methods	2-arm active-controlled randomised trial		
Participants	238 women were randomised in a hospital setting in Canada. The population comprised women of parity 5 or less, unspecified whether singleton or multiple pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with coagulopathy, anticoagulation therapy, previous PPH or previous caesarean.		
Interventions	400 mcg of misoprosto	ol administered rectally versus 5 IU of oxytocin administered by an IV bolus or IM	
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; manual removal of placenta; change in Hb; third stage duration (minutes); nausea; vomiting; headache; fever; shivering; abdominal pain.		
Notes	Contact with study aut	thors for additional information: yes. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	A statistician developed blocked randomisation tables for each centre.	
Allocation concealment (selection bias)	Low risk	Pharmacy assembled consecutively-numbered opaque, sealed packets that contained the group allocation.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.	
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.	
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "13 women randomised subsequently delivered by caesarean and wer excluded from analysis. 2 women were lost to follow-up early in the trial when their packets were opened but the manoeuvre was not completed and no dat were recorded".	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	High risk	Not all study participants were included in the analysis.	
Funding source	Low risk	The study was supported by funding from the physicians of Ontario, through	

Kerekes 1979

Methods	3-arm controlled randomised trial	

the Physician Services Incorporated Foundation (public funding).



Kerekes 1979 (Continued)			
Participants	140 women were randomised in a hospital setting in Hungary. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
Interventions	200 mcg of ergometrin	e administered IV bolus versus no treatment	
Outcomes	The study recorded the	e following outcomes: third stage duration (minutes).	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss by collection in a container placed under the mother during the third stage of labour until 2 hours postpartum. The contents of the container were transferred to a measuring cylinder. However, blood loss data were not reported in a format that could be extracted for the purpose of this review.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Khan 1995

Methods	2-arm active-controlled double-blinded randomised trial
Participants	2040 women were randomised in a hospital setting in United Arab Emirates. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour, caesarean section or instrumental delivery, or requiring general anaesthesia, epidural or diazepam, or



Khan 1995 (Continued)		ypertension (160/100 mmHg or more), hypertension on antihypertensive drugs, ordiac disease or Hb of 90 g/L or less.		
Interventions	10 IU of oxytocin admii IM	10 IU of oxytocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the placenta; vomiting; he	e following outcomes: PPH at 500; PPH at 1000; transfusion; manual removal of adache.		
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Number code by the hospital pharmacist who alone was aware of the content of the ampoules.		
Allocation concealment (selection bias)	Low risk	Participants were assigned an opaque sealed envelope. Each envelope carried the instruction to use a numbered vial of the study drug.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss "in the standard way" by measurement of blood and clots in a graduated jug, and by weighing swabs and linen.		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "12 patients had to be excluded from the trial (oxytocin 5; ergometrine plus oxytocin 7) after randomisation because they no longer fulfilled the inclusion criteria (2 who required caesarean section and 10 who were delivered by forceps or ventouse (oxytocin, 4; Ergometrine plus oxytocin 6)."		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.		
Funding source	Unclear risk	Source(s) of funding for the study were not reported.		

Khurshid 2010

Methods	2-arm active-controlled randomised trial
Participants	200 women were randomised in a hospital setting in India. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with hypertension, cardiac disease, renal disease, gastro-intestinal disorders, respiratory disease, endocrinal problems, coagulation disorder and sensitivity to prostaglandin or methergin.



(hurshid 2010 (Continued) Interventions	125 mcg of carboprost	administered IM versus 200 mcg of ergometrine administered by an IV bolus
Outcomes	The study recorded the following outcomes: additional uterotonics; manual removal of placenta; blooloss (mL;,third stage duration (minutes).	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Randomisation was done using random tables.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Blood loss was estimated by collecting blood and blood clots in the kidney tray and adding the difference in the weight of the drapes before use and after birth.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Koen 2016

Methods	2-arm active-controlled double-dummy randomised trial
Participants	540 women were randomised in a hospital setting in South Africa. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women not willing or not able to provide consent, previous classic CS, < 18 years of age, pre-eclampsia, eclampsia, uncontrolled hypertension, cardiac/liver/renal disorders, hypersensitivity to oxytocin or oxytocin + ergometrine, occlusive vascular disease, autoimmune vasculitis.



Koen 2016 (Continued)		
Interventions		ninistered by an IV bolus + infusion versus 500 mcg plus 15 IU of ergometrine plus IM plus by an IV infusion
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL); nausea; vomiting; headache.	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomisation was carried out by a lottery method "Randomisation was by means of sealed not transparent envelopes. Each had a label inside with a letter A (oxytocin group) or B (oxytocin + ergometrine), which corresponded to a pair of pre-packed colour-coded ampoules that were used for the two different groups."
Allocation concealment (selection bias)	High risk	Quote: "Randomisation was carried out by a lottery method "Randomisation was by means of sealed not transparent envelopes. Each had a label inside with a letter A (oxytocin group) or B (oxytocin + ergometrine), which corresponded to a pair of pre-packed colour-coded ampoules that were used for the two different groups."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Double-blinded.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Blinded.
Objective assessment of blood loss	Low risk	Calculation of blood loss was done using calculated pregnancy preoperative blood volume (0.75 \times [{height inches \times 50} + {weight pounds \times 25}) \times percentage of blood volume lost ([pre-delivery haematocrits – post-delivery haematocrits]/pre-delivery haematocrits).
Incomplete outcome data (attrition bias) All outcomes	High risk	540 women were randomised in the study, but 124 were excluded because of giving birth vaginally (n = 80), incomplete data or protocol violations (n = 44) after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (ClinicalTrials.gov NCT02046499).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Kumar 2016

Methods	2-arm active-controlled randomised trial
Participants	201 women were randomised in a hospital setting in India. The population comprised women of any parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by



Kumar 2016 (Continued)		sion criteria comprised women undergoing caesareans, with hypersensitivity to diseases, epilepsy, psychiatric disorders, liver and renal diseases.
Interventions	125 mcg of carboprost	administered IM versus 10 IU of oxytocin administered IM
Outcomes		e following outcomes: PPH at 500; additional uterotonics; transfusion; death; stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Perineal drapes were replaced by calibrated Brasss V obstetric drape after the delivery of the baby. The average time taken for episiotomy suturing was around 10 minutes in both the groups and did not have any significant impact on the blood loss and duration of bleeding. Brasss V drape was removed 10 minutes after the episiotomy suturing in all patients unless the patient continued to have significant PPH.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	1 women was excluded because of a fourth degree tear after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Kumru 2005

Methods	2-arm active-controlled randomised trial		
Participants	55 women were randomised in a hospital setting in Turkey. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with multiple pregnancy, hypertension or vascular diseases.		



(umru 2005 (Continued)		
Interventions		nistered by an IV bolus + infusion versus 200 mcg plus 10 IU of ergometrine plus by an IV bolus plus by IV bolus plus infusion
Outcomes	The study recorded the following outcomes: blood loss (mL).	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised intraoperative blood loss by weighing compresses and rolls before and after the birth of the baby, and calculating the difference between these measurements. Pre-weighted pads were distributed in advance to each mother, and collected at intervals of 3-6 hours hour intervals after the aspiration of amniotic fluid.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Kundodyiwa 2001

Methods	2-arm placebo-controlled randomised trial	
Participants	500 women were randomised in a hospital setting in Zimbabwe. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing instrumental delivery, or those with previous PPH, antepartum haemorrhage, coagulopathy, multiple pregnancy, asthma or allergies to prostaglandins or oxytocin.	



Interventions	400 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; bloc loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Computer-generated using a random sequence.
Allocation concealment (selection bias)	Low risk	The participant was asked to randomly pick a numbered sealed opaque envelope from the study cooler-box.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Identical placebo tablets could not be obtained from the manufacturers. The tablets were similar in size and colour but not in shape. However, most reviewed trials on misoprostol had this similar problem although this method of blinding proved to be effective."
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Quote: "The data sheet was completed by the midwife supervising the deliver and collected and checked by the research assistant".
Objective assessment of blood loss	Low risk	After delivery, investigators appraised blood loss by removing linen soiled with amniotic fluid, and then placing a fresh disposable incontinence pad with a plastic backing under the mother. Blood expressed from the uterus was measured with a calibrated measuring jug. The volume of blood soiling linen savers and sanitary pads was determined as the difference between dry weights and soiled weights: these measurements were added to the volume recorded by the calibrated jug.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Data for 1 woman were excluded because she delivered undiagnosed twins after randomisation".
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Kushtagi 2006

Methods	2-arm active-controlled randomised trial	
Participants	215 women were randomised in a hospital setting in India. The population comprised women of any parity, unspecified whether singleton or multiple pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	



(ushtagi 2006 (Continued)		
Interventions	200 mcg of ergometrine administered by an IV bolus versus 125 mcg of carboprost administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; blood loss (mL); third stage duration (minutes); hypertension.	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Amount of blood loss was quantified by noting the increment in weight of standardised tampons which were placed high up in the vagina immediately after placental delivery.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and random ly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Lam 2004

Methods	2-arm active-controlled randomised trial		
Participants	60 women were randomised in a hospital setting in China (Hong Kong SAR). The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour, or those with antepartum haemorrhage, anaemia, 2 or more surgical terminations, previous manual removal of placenta, previous PPH or previous third stage complications.		
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered by an IV bolus versus 600 mcg of miso- prostol administered sublingually		



Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; manual removal of placenta; death; fever.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Allocated using a random number-generated table.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	Investigators appraised blood loss during the third stage by visual estimation, and by objective measurement on the basis of a method previously described by Newton and colleagues. Whilst any blood clots were collected and measured with a jug, white linen was placed under the mother during delivery and subsequently processed for 15 minutes with sodium hydroxide solution in an automatic stomacher (laboratory blender), to achieve the formation of alkaline hematin. Quote: "The optical density at 550 nm of the alkaline hematin was measured by spectrophotometry and compared with that of a known volume of a sample of the patient's venous blood" to calculate the volume of blood loss.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Unclear risk	It was unclear from the study report whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Lamont 2001

Methods	2-arm active-controlled randomised trial	
Participants	529 women were randomised in a hospital setting in the UK. The population comprised women of both nulliparous and multiparous, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by both caesarean and vaginal delivery. Exclusion criteria comprised women with known sensitivity to either prostaglandins, ergometrine or oxytocin, had a history of asthma, glauco-	



Lamont 2001 (Continued)		pressure or were known to have cardiac, pulmonary, renal or hepatic disease,	
		r obliterative vascular disorders. Women were excluded if they were currently reatment or participating in other clinical trials.	
Interventions	250 mcg of carboprost administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; manual removal of placenta; blood loss (mL); diarrhoea; nausea; vomiting.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The randomisation slips were contained in envelopes which were opened by a person not involved in the postpartum assessments who resealed the envelope and drew 1 mL of the appropriate medication into a syringe. The nature of the medication was not revealed and the resealed envelope was retained in the woman's notes. The medication was administered by a competent person other than the one who had opened the envelope and filled the syringe.	
Objective assessment of blood loss	Unclear risk	Blood loss was measured as accurately as possible, taking into consideration the liquor amnii and soiling of the surgical drapes.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	530 women were randomised in the study, but 1 was excluded because did not receive the allocated agent (carboprost) after randomisation.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Lapaire 2006

Methods	2-arm active-controlled double-blinded randomised trial
Participants	56 women were randomised in a hospital setting in Switzerland. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women undergoing emergency caesarean section,



Lapaire 2006 (Continued)	vere systemic disorder	ress, fetal malformations, pre-eclampsia, HELLP syndrome, coagulopathy, ses, an American Society of Anesthesiologists physical status of 3 or greater, severe mectomy, pyrexia (more than 38.5C) or hypersensitivity to prostaglandins.		
Interventions	25 IU of oxytocin administered by an IV bolus + infusion versus 800 mcg plus 5 IU of misoprostol plus oxytocin administered orally plus by an IV bolus			
Outcomes		The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL); nausea; headache; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	The hospital pharmacy performed the 1:1 computer-generated randomisation that assigned the participants to their group.		
Allocation concealment (selection bias)	Low risk	Used identical study boxes from pharmacy.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was "double-blind": Quote: "the study drugs and placebos [were provided by the pharmacy] in unidentifiable form".		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.		
Objective assessment of blood loss	Low risk	When the membranes ruptured before delivery, investigators appraised intraoperative and postoperative blood loss by determining the difference in weight of cloths and pads used to absorb blood during surgery and in the intermediate care unit. When membranes did not rupture preoperatively, investigators appraised blood loss by collection in suction bottles and subtracting estimated amniotic fluid volume. Investigators considered that 1 g is equivalent to 1 mL of blood or amniotic fluid.		
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "3 patients in the oxytocin group were excluded from statistical analysis because of errors in drug administration". Moreover calculated blood loss data were unavailable in 13 cases and for these women the primary outcome was estimated clinically."		
Selective reporting (reporting bias)	High risk	The study protocol that was registered retrospectively (ClinicalTrials.gov) lists PPH as the primary outcome of the study, but the study report lists the primary outcomes as intraoperative and postoperative blood loss and drug-related adverse effects (these items are listed only as secondary outcomes in the registration file). The study does not report the incidence of PPH ≥ 500 mL, nor PPH ≥ 1000 mL.		
Intention to treat analysis	High risk	The authors excluded 3 study participants in the oxytocin group from the analysis because they incurred errors in drug administration.		
Funding source	Low risk	The study was supported by funding from the Scientific Pool of Basel University Hospital (the institution of the authors).		



Leung		

eung 2006			
Methods	2-arm active-controlled double-dummy randomised trial		
Participants	329 women were randomised in a hospital setting in Hong Kong. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring prophylactic oxytocin infusion, or those with pre-existing hypertension, pre-eclampsia, asthma, cardiac/renal/liver diseases, grand multiparity or fibroids.		
Interventions	100 mcg of carbetocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hbl third stage duration (minutes); nausea; vomiting; hypertension; headache; tachycardia; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated code before the recruitment.	
Allocation concealment (selection bias)	Low risk	This was performed by opening a sealed, consecutively-numbered, opaque envelope.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.	
Objective assessment of blood loss	High risk	Investigators appraised blood loss by visual estimation.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "15 women in the carbetocin group and 14 women in the ergometrine plus oxytocin group failed to have a paired haemoglobin test to measure the change in haemoglobin 48 hours after delivery either because they had requested early home or refused further blood taking. These 29 women were excluded."	
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of fever were omitted).	
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.	
Funding source	High risk	The study was supported by funding from Ferring Pharmaceuticals.	



Methods	2-arm active-controlled randomised trial			
Participants	40 women were randomised in a hospital setting in UK. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with two or more previous caesarean sections or previous uterine rupture.			
Interventions	10 IU of oxytocin admir	nistered by an IV bolus versus 500 mcg of misoprostol administered orally		
Outcomes		The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb; fever;.shivering.		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	The randomisation was undertaken by means of computer-generated random numbers.		
Allocation concealment (selection bias)	Low risk	Used sealed opaque envelopes.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "The obstetrician, surgical assistant, scrub nurse and recovery midwife were blinded to the treatment. The anaesthetist and the anaesthetic assistan were not blinded as it was important for patient safety that a record was kept of all drugs administered."		
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.		
Objective assessment of blood loss	High risk	Investigators appraised intraoperative and postoperative (up to 1 hour) blood loss by visual estimation, quote: "in a standard manner (volume of blood in suction bottle plus soiling of swabs and bed sheets)".		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.		
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		
Funding source	Low risk	The study was supported by "assistance" from the Department of Anaesthesia at University College London Hospitals NHS Trust (the institution of the authors).		

Methods

 $\hbox{3-arm active-controlled double-dummy randomised trial}\\$



Funding source

Lumbiganon 1999 (Continued)				
Participants	597 women were randomised in a hospital setting in South Africa and Thailand. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section or abortion, or those with asthma, other severe chronic allergic conditions a contraindication to use of misoprostol or if they were not willing or able to give informed consent.			
Interventions	600 mcg or 400 mcg of	misoprostol administered orally versus 10 IU of oxytocin administered IM		
Outcomes		The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); diarrhoea; nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Random allocation sequence, generated centrally.		
Allocation concealment (selection bias)	Low risk	The treatment packs were consecutively-numbered and sealed.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The packs were identical in shape, colour, weight and feel. Each woman received an injection and 3 tablets. Thus, the trial was double-blinded using double placebos".		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss from the delivery of the baby until the mother was transferred to postnatal care. The collected blood was poured into a standard measuring jar provided by WHO for the purpose of volumetric measurement. Linen was not weighed but clots and small gauze swabs soaked with blood were included in the measurement.		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Exclusion after randomisation: 8 women in the oxytocin group did not comp with treatment (6 had an emergency caesarean section, 1 was HIV positive a mistakenly excluded, 1 whose ampoule was not located). 1 woman in the 60 mcg group was excluded.		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		

Low risk

The study was supported by funding from the WHO (public funding). Active $\,$ and placebo medications, syringes and swabs were donated by Searle, Novar-

tis Pharma AG and Becton Dickinson International.



Methods	2-arm active-controlled	d double-blinded randomised trial	
Metrious			
Participants	200 women were randomised in a hospital setting in Egypt. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with placenta praevia, coagulopathy, pre-eclampsia, cardiac/renal/liver disorders, epilepsy or known hypersensitivity to oxytocin or carbetocin.		
Interventions	100 mcg of carbetocin	administered IM versus 5 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion.; blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; headache; tachycardia; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Participants were equally randomised using automated web-based randomisation system.	
Allocation concealment (selection bias)	Unclear risk	Only states that ensured allocation concealment with no further details.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported in sufficient detail even though the authors state it was double-blinded.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	High risk	Investigators appraised blood loss by weighing swabs and using pictorial charts.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Maged 2017

Methods	2-arm active-controlled double-blinded randomised trial
Participants	300 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by both



Maged 2017 (Continued)		caesarean. Exclusion criteria comprised women with placenta previa, coaguor known sensitivity to oxytocin or methergine.	
Interventions	100 mcg of carbetocin administered by an IV bolus versus 200 mcg plus 5 IU of ergometrine plus oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; blood loss (mL; change in Hb; nausea; vomiting; headache; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Randomised using automated web based randomisation system.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported in sufficient detail.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The authors state the study was double-blinded but blinding (of study participants and caregivers) was not described in sufficient detail.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	Calculated estimated blood loss.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Malik 2018

Methods	2-arm active-controlled randomised trial
Participants	200 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with anaemia, pregnancy-induced hypertension, placental abruption/placenta praevia, multiple pregnancy, grand multiparous, malpresentation, polyhydramnios, previous uterine scar, chorioamnionitis, prolonged labour, intrauterine fetal death, coagulation disorder, asthma/epilepsy/heart/renal disorder.



Malik 2018 (Continued)			
Interventions	125 mcg of carboprost administered IM versus 200 mcg of ergometrine administered by an IV bolus		
Outcomes	The study recorded the following outcomes: blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.	
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	Amount of blood loss was calculated by weighing the gauzes/sponges before delivery followed by again weighing them after delivery.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.	
Selective reporting (re- porting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Mannaerts 2018

Methods	2-arm active-controlled double-blinded randomised trial
Participants	68 women were randomised in a hospital setting in Belgium. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with medical conditions potentially influencing outcome measures (nausea, vomitus, and hypotension): diabetes, preexisting hypertension, pre-eclampsia, gestational hypertension, and known gastrointestinal diseases.
Interventions	15 IU of oxytocin administered by an IV bolus + infusion versus 100 mcg of carbetocin administered by an IV bolus



Mannae	rts 2018	(Continued)
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Outcomes	The study recorded the following outcomes: additional uterotonics; change in Hb; nausea.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Participants are randomly assigned following simple randomisation procedure in 1:1 ratio to 1 of the 2 treatment groups. A computer-generated randomisation list was generated using SPSS21.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Medication was prepared by a midwife not treating the patient to make sure that patient, gynaecologist, anaesthesiologist, and midwife clinically in charge of the patient are blinded for the medication.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Medication was prepared by a midwife not treating the patient to make sure that patient, gynaecologist, anaesthesiologist, and midwife clinically in charge of the patient are blinded for the medication.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	68 women were randomised in the study, but 10 were excluded because of incomplete data after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (ISRCTN 95504420).
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

McDonald 1993

Methods	2-arm active-controlled double-blinded randomised trial
Participants	3497 women were randomised in a hospital setting in Australia. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing emergency or elective caesarean section, or requiring general anaesthetic for instrumental delivery, or those with hypertension in labour (more than 150/100 mm Hg), antenatal hypertension, maternal distress, advanced stage in labour, language barrier, fetal abnormality, intrauterine death or medical disorder.
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 10 IU of oxytocin administered IM



McDona	lc	1993	(Continued)
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Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu-
	sion; manual removal of placenta; NNU admissions; breastfeeding; nausea; vomiting.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The ampoules were numbered by Sandoz by using simple randomisation. There was no blocking or prognostic stratification.
Allocation concealment (selection bias)	Low risk	The ampoules were numbered by third party (Sandoz).
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Delivery attendants were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending obstetricians and midwives.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "All women allocated to receive a drug were included in that group, excluding only the 14 women for whom drug allocation was not recorded".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	High risk	The study was supported by funding from Sandoz.

Mitchell 1993

Methods	2-arm active-controlled double-blinded randomised trial	
Participants	461 women were randomised in a hospital setting in the UK. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section, or those with significant hypertension or cardiac disease.	
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 5 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; manual removal of placenta; blood loss (mL); third stage duration (minutes).	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	



Mitchell 1993 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Unclear sequence: described as without any blocking or stratification.
Allocation concealment (selection bias)	Low risk	Used identical study boxes prepared by third party (Sandoz).
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss quote: "in the standard way by graduated jug measurement plus an allowance for spillage".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Perinatal Trials Service (public funding), for the Department of Health for England and Wales, and for Birthright (the charitable arm of the RCOG). Coded medication ampoules were provided by Sandoz.

Mobeen 2011

Methods	2-arm placebo-controlled randomised trial		
Participants	1119 women were randomised in a community setting in Pakistan. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with hypertension, non-cephalic presentation, polyhydramnios, previous caesarean, multiple pregnancy, intrauterine death, antepartum haemorrhage or Hb less than 80 g/L.		
Interventions	600 mcg of misoprostol administered orally versus placebo		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		



Mobeen 2011 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A computer-generated random code in blocks of 6 was maintained by Gynuity Health Projects in New York and not revealed until data collection and cleaning were completed.
Allocation concealment (selection bias)	Low risk	Study medication was packed in numbered colour-coded boxes by Gynuity Health Projects in New York.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Both women and TBAs were blinded to study assignment".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	To appraise postpartum blood loss, blood was collected with a perineal sheet and bedpan placed under the mother for a minimum of 1 hour or until active bleeding stopped (whichever occurred last). Quote: "Blood collected in the bedpan was transferred to a measuring jar, which was then closed, and the perineal sheet and cotton roll were placed in a sealed plastic bag. The closed measuring jar and sealed plastic bag were then placed inside a plastic cooler which was tightly closed and stored in a secure place in the woman's home until the local health visitor or community health nurse arrived for weighing, 1–2 days after delivery".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Invalid blood loss measures, which mainly occurred when monitoring visits were not possible because of poor weather conditions, were excluded from our analysis".
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov NCT00120237).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Bill and Melinda Gates Foundation (public funding).

Modi 2014

Methods	4-arm active-controlled randomised trial
Participants	100 women were randomised in a hospital setting in India. The population comprised women of parity 3 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with gestations less than 37 or more than 42 weeks, intrauterine death, fetal growth restriction, hypertensive or cardiac or renal disorders, multiple pregnancies, placenta praevia, placenta abruption, grand multiparous, coagulation disorders, anaemia (< 8 g/dL), tachycardia or hypotension, malpresentations, chorioamnionitis, or known allergy to prostaglandins.
Interventions	10 IU of oxytocin administered IM versus 200 mcg of ergometrine administered by an IV bolus versus 125 mcg of carboprost administered IM versus 600 mcg of misoprostol administered rectally



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The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; Blood loss (mL); third stage duration (minutes).	
Contact with study authors for additional information: no. Additional data from authors: no	
Authors' judgement	Support for judgement
Unclear risk	Sequence generation was not reported.
Unclear risk	Allocation concealment was not reported.
Unclear risk	Blinding (of study participants and caregivers) was unclear.
Unclear risk	Assessor blinding was not reported.
Low risk	Used BRASS-V drapes to measure the blood loss.
Unclear risk	The study authors did not mention any incomplete outcome data.
Unclear risk	The protocol of the study was unavailable for verification.
Unclear risk	The authors did not specify whether all those who were enrolled and random ly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Low risk	No funding sought for this study.
	Authors' judgement Unclear risk

Methods	2-arm active-controlled double-blinded randomised trial		
Participants	84 women were randomised in a hospital setting in Austria. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women requiring general anaesthesia, or those with placenta praevia, placental abruption, multiple pregnancy, pre-eclampsia, gestational diabetes, pre-existing insulin-dependent diabetes, cardiovascular/renal disorders, hypo-/hyperthyroidism or women on cardiovascular system medications.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 5 IU of oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: additional uterotonics change in Hb; nausea; headache.		



Moertl 2011 (Continued)

Notes

Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was performed by a computer-generated randomisation sequence 1:1 ratio—blocks of 10 without stratification.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Study medication was double-blinded to the clinical staff (obstetricians as well as anaesthesiologists) and the technicians performing the measurements".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators did not appraise blood loss.
Incomplete outcome data (attrition bias) All outcomes	High risk	After randomisation, investigators excluded 28 women from analysis for technical problems (n = 15), change to general anaesthesia (n = 9), recording artefacts (n = 3) and patient withdrawal (n = 1).
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (EudraCT 2007-005498-78).
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Low risk	CNSystems Medizintechnik AG in Graz, Austria provided the Task Force® Monitor 3040i system used to measure haemodynamic parameters. No other external funding was required for the study.

Mohamed 2015

Methods	2-arm active-controlled randomised trial	
Participants	172 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with medical disorder as hypertension, diabetes or on an anticoagulant, severe polyhydramnios, multiple pregnancy, placenta praevia or placental abruption, previous uterine scar other than lower segment caesarean section or who had more than 1 previous section.	
Interventions	5 IU of oxytocin administered by an IV bolus versus 100 mcg of carbetocin administered by an IV bolus	
Outcomes	The study recorded the following outcomes: blood loss (mL).	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		



Mohamed 2015 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was performed by computer generated randomisation system.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	After delivery of the placenta, the volume of blood loss was assessed by weight or saturation assessment techniques by subtracting the dry weight of absorbing materials (pads, sponges, etc) from the weight of blood-containing materials and using the conversion 1 g weight = 1 mL to quantify the blood volume contained in the materials.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Moir 1979

2-arm active-controlled randomised trial		
88 women were randomised in a hospital setting in the UK. The population comprised women of primigravidas, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.		
500 mcg of ergometrine administered by an IV bolus versus 10 IU of oxytocin administered by an IV bolus		
The study recorded the following outcomes: PPH at 500; PPH at 1000; blood loss (mL); nausea.		
Contact with study authors for additional information: yes. Additional data from authors: no.		
Authors' judgement Support for judgement		



Moir 1979 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by quote: "the haemoglobin extraction-dilution technique, which is acceptably accurate (Roe, Gardiner and Dudley, 1962; Thornton et al, 1963) and particularly suited to obstetric use (Moir and Wallace, 1967; Wallace, 1967). The perdometer apparatus was used and all blood and blood-stained linen were collected".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Moodie 1976

Into date 2010		
Methods	2-arm active-controlled randomised trial	
Participants	148 women were randomised in a hospital setting in the UK. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	
Interventions	500 mcg of ergometrine administered by an IV bolus versus 5 IU of oxytocin administered by an IV bolus	
Outcomes	The study recorded the following outcomes: PPH at 500; blood loss (mL); nausea.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.



Moodie 1976 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with the placenta bowl and soiled linen and swabs. Quote: "The principles of the haemoglobin extraction-dilution technique employed have been discussed by Roe, Gardiner and Dudley (1962) and Thornton and colleagues (1963).
Incomplete outcome data (attrition bias) All outcomes	High risk	There were 148 study participants but blood loss data were available in only 80 cases.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Mukta 2013

(selection bias)

Mukta 2013		
Methods	2-arm active-controlled randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of any parity, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing emergency or elective caesarean section, or those with eclampsia, asthma, epilepsy, cardiac/kidney disorder or coagulopathy.	
Interventions	600 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; diarrhoea; nausea; vomiting; fever; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Randomly divided into 2 equal groups.
Allocation concealment	Unclear risk	Allocation concealment was not reported.



Mukta 2013 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss in mL, by collection with a calibrated plastic drape, after the drainage of amniotic fluid and delivery of the baby until the third stage of labour was completed.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Musa 2015

Methods	2-arm active-controlle	d double-dummy randomised trial
Participants	235 women were randomised in a hospital setting in Nigeria. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing planned instrumental, or those who received oxytocin and/or misoprostol other than in the third stage of labour, or those with grand multiparity (more than 4), multiple pregnancy, fibroids, polyhydramnios, pre-eclampsia, eclampsia, hypertension, cardiac disorder, asthma,APH, previous PPH, prolonged rupture of membranes or Hb less than 100 g/L).	
Interventions	600 mcg of misoprosto	ol administered orally versus 10 IU of oxytocin administered IM
Outcomes	The study recorded the following outcomes: PPH at 500; severe maternal morbidity: intensive care admissions; additional uterotonics; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: yes
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocation was done by blocked (restrictive), using computer-generated random numbers prepared by an independent statistician.
Allocation concealment (selection bias)	Unclear risk	Used opaque envelopes but no other details provided.



Musa 2015 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participants, caregivers, and outcome assessors (researchers or research assistants) were masked to group allocation. Investigators were not masked for data analysis".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Participants, caregivers, and outcome assessors (researchers or research assistants) were masked to group allocation. Investigators were not masked for data analysis".
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by quote: "the gravimetric method" (Ambardekar 2009) until 1 hour after delivery.
Incomplete outcome data (attrition bias) All outcomes	High risk	235 study participants were randomised but only 200 were analysed due to protocol deviations and missing data.
Selective reporting (reporting bias)	Unclear risk	The study protocol was registered retrospectively (PACTR 201407000825227).
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Low risk	The study was supported by funding from the University of Ilorin Teaching Hospital (the institution of the authors).

Nankaly 2016

Methods	3-arm active-controlled randomised trial
Participants	185 women were randomised in a hospital setting in Iran. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with anaemia, multiple pregnancy, polyhydramnios, prolonged labour, PROM, placenta praevia, placental abruption, vaginal bleeding, diabetes, blood pressure, kidney disease, cardiovascular disease and coagulation disorders or other underlying disease.
Interventions	20 IU of oxytocin administered by an IV infusion versus 400 mcg or 200 mcg of misoprostol administered sublingually
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; fever; shivering.
Notes	Contact with study authors for additional information: no. Additional data from authors: no

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported in Quote: "The randomisation was done via block randomisation and in the form of four blocks".
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.



Nankaly 2016 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Objective assessment of blood loss	Unclear risk	Lost blood volume gained from calculating the total collected blood in suction container and counting the number of blood gases.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Nasr 2009

2-arm active-controlled double-dummy randomised trial		
514 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with antepartum haemorrhage, coagulopathy, hypertension in pregnancy or the need for anticoagulants.		
800 mcg of misoprostol administered rectally versus 5 IU of oxytocin administered by an IV infusion		
The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.		
Contact with study authors for additional information: yes. Additional data from authors: yes		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocated by a computer-generated random allocation system created at the Statistics Unit of Assiut University Hospital.
Allocation concealment (selection bias)	Low risk	Allocation codes were placed in sealed, opaque, consecutively-numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote: "double-blind": active treatments and placebo treatments were "identical-looking".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.



Nasr 2009 (Continued)		
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were unclear.

Nayak 2017

Nayak 2017		
Methods	2-arm placebo-control	led randomised trial
Participants	200 women were randomised in a hospital setting in India. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women having severe medical and surgical complications including the heart, liver, kidney, brain disease and blood disorders, any contraindication to misoprostol including mitral stenosis, glaucoma and diastolic blood pressure over 100 mmHg and known allergic to prostaglandins, history of thromboembolic disorders, abnormal placentation such as placenta praevia, placental abruption and placental adhesions caused by repeated artificial abortions, pregnancy complications such as severe pre-eclampsia, multiple pregnancies, macrosomia and polyhydramnios, complication with myoma and with any blood dyscrasia.	
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 10 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; blood loss (mL); change in Hb.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Nayak 2017 (Continued)		
Objective assessment of blood loss	Low risk	The quantity of blood (mL) = (weight of (used material + unused material) after surgery-weight of all materials prior to surgery)/1.05 plus the volume included in the suction container after placental delivery. In addition, pads used after completion of caesarean section to 2 hours postpartum weighed.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Nellore 2006

Methods	2-arm active-controlled randomised trial		
Participants	120 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring oxytocin induction or augmentation of labor, consarean delivery, or those with gestational age less than 37 weeks, multiple pregnancy, Hb concentration less than 8 g/dL, and known allergy to prostaglandins.		
Interventions	400 mcg of misoprostol administered rectally versus 125 mcg of carboprost administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Nellore 2006 (Continued)		
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ng 2001

Methods	2-arm active-controlled randomised trial		
Participants	2058 women were randomised in a hospital setting in Hong Kong. The population comprised women of parity 3 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring oxytocin infusion in the third stage, or those with pre-eclampsia, cardiac disorder, asthma, grand multiparity (more than 3), fibroids or contraindications for the use of either misoprostol or syntometrine.		
Interventions	600 mcg of misoprostol administered orally versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death. Blood loss (mL); change in Hb; nausea; vomiting; hypertension; headache; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation based on a table of computer-generated blocks of random numbers.
Allocation concealment (selection bias)	Low risk	Consecutively-numbered opaque sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "This was not a double-blinded study".
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.



Ng 2001 (Continued)		
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ng 2004

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	298 women were randomised in an unspecified setting in Hong Kong. The population comprised women of unspecified parity, a singleton pregnancy, at unspecified risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancy or non-vaginal delivery.		
Interventions	400 mcg of misoprostol administered orally versus 1 mL of oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: (No outcome data found)		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Double-blinding of personnel and participants (placebo use) but insufficient details from abstract only.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.



Ng 2004 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ng 2007

Methods	2-arm active-controlled double-dummy randomised trial 360 women were randomised in a hospital setting in Hong Kong. The population comprised women of parity 3 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring oxytocin infusion in the third stage, or those with pre-eclampsia, cardiac disorder, asthma, grand multiparity (more than 3), fibroids or contraindications for the use of either misoprostol or syntometrine.		
Participants			
Interventions	400 mcg of misoprostol administered orally versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu sion; manual removal of placenta; blood loss (mL); change in Hb; diarrhoea; nausea; vomiting; hypertension; headache; fever; shivering; maternal satisfaction.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The randomisation was based on a table of computer-generated random numbers.
Allocation concealment (selection bias)	Low risk	Used consecutively-numbered and sealed opaque packages.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The placebo was identical in size and colour but had a different shape to the misoprostol tablet. All women were asked to swallow the tablets directly from the opaque cup without looking at them. The identity of the active medication and placebo were concealed from the caregivers and the parturient."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "5 women were excluded from the analysis because of missing post-de- livery haemoglobin level".



Ng 2007 (Continued)		
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of tachycardia and dizziness were omitted).
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Nirmala 2009

Methods	2-arm active-controlled randomised trial	
Participants	120 women were randomised in a hospital setting in Malaysia. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women younger than 18 years old, or those with cardiac disorder, hypertension requiring treatment, liver/renal/vascular/endocrine disorder (excluding gestational diabetes) or hypersensitivity to oxytocin or carbetocin.	
Interventions	100 mcg of carbetocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; nausea; vomiting; hypertension; headache; shivering; abdominal pain.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation.
Allocation concealment (selection bias)	Unclear risk	Used sealed, sequentially-numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "The preparation and administration of the medication was carried out by midwives who were not involved in the management of the patient except for the drug administration".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by "the gravimetric method" from immediately after drug administration. They used a digital scale (Soehnle, Venezia) for weight measurement. In order to minimise confounding by fluid absorbed into drapes, they collected blood with a new plastic sheet placed under the mother after delivery of the baby. They also weighed any gauzes, tampons and pads used in the first hour after delivery of the placenta, and subtracted the dry weights of these materials to calculate blood loss on the basis that 1 g is equivalent to 1 mL.



Nirmala 2009 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Nordstrom 1997

Methods	2-arm placebo-controlled randomised trial	
Participants	1000 women were randomised in a hospital setting in Sweden. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	
Interventions	10 IU of oxytocin administered by an IV bolus versus placebo	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL).	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation.
Allocation concealment (selection bias)	Low risk	Ampoules were prepared at the hospital pharmacy and consecutively-numbered.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The content of the ampoules was unknown to mothers, midwives and doctors until the study was completed".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss quote: "by measuring collected blood and adding what was estimated to have been absorbed by surgical cloths and tissues".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.



Nordstrom 1997 (Continued)			
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Low risk	The study was supported by funding from the County Council and County Health Authority Research and Development Foundation in the County of Jämtland, Sweden (public funding).	

Nuamsiri 2016

Methods	2-arm placebo-controlled randomised trial	
Participants	323 women were randomised in a hospital setting in Thailand. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancy, polyhydramnios, uterine fibroids, previous PPH, APH, parity greater than 4, previous caesarean section, severe anaemia (Hb level of ≤ 8 g/dL), coagulopathy, contraindications to the use of ergometrine, estimated fetal birthweight > 4000 g. and inability to obtain written informed consent. Women who ended up having a caesarean section or instrumental delivery were also excluded from this study.	
Interventions	200 mcg plus 20 IU of ergometrine plus oxytocin administered by an IV bolus + infusion versus 20 IU of oxytocin administered by an IV infusion	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfession; death; blood loss (mL); change in Hb; third stage duration (minutes); nausea; hypertension.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

RISK OT DIAS		
Authors' judgement	Support for judgement	
Low risk	Random allocation scheme using a computer-generated list of numbers.	
Low risk	Used sealed and consecutively numbered opaque envelopes were prepared by a research assistant not involved in the study. The women were randomly allocated to 1 of the 2 study groups by opening the next available envelope just before delivery.	
Low risk	The study drug and placebo were prepared by the research assistant not involved in the study. The obstetrician and nursing staff were all blinded to the type of injectable substance.	
Low risk	The study drug and placebo were prepared by the research assistant not involved in the study. The obstetrician and nursing staff were all blinded to the type of injectable substance.	
Low risk	Used the blood collection drape, which was placed under the buttocks after placental delivery. Blood-soaked swabs were weighed in grams, and the known dry weight of the swabs was subtracted, this volume was added to the	
	Low risk Low risk Low risk	



Nuamsiri 2016 (Continued)		measured blood volume from the drape (assuming an equivalence of 1 g to 1 mL).
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The study report matches the study protocol that was registered retrospectively (TCTR20150820001).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding were not reported.

Oboro 2003

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	496 women were randomised in a hospital setting in Nigeria. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour, or those with previous caesarean, Hb less than 80 g/L, previous PPH, grand multiparity (not defined), multiple pregnancy, polyhydramnios, fibroids or precipitate labour.		
Interventions	10 IU of oxytocin administered IM versus 600 mcg of misoprostol administered orally		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Generated by using random tables.
Allocation concealment (selection bias)	Low risk	Pharmacy prepared opaque sealed sequentially-numbered packets.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The identity of the active medication and placebo were concealed from the caregivers and women".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending obstetricians.



Oboro 2003 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ogunbode 1979

Methods	3-arm active-controlled randomised trial		
Participants	144 women were randomised in a hospital setting in Nigeria. The population comprised women of specified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal livery. Exclusion criteria comprised women undergoing instrumental delivery, or those with previous PPH, multiple pregnancy, polyhydramnios or vaginal lacerations.		
Interventions	200 mcg or 500 mcg of ergometrine administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; manual removal of placenta; blood loss (mL).		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

RISK OT DIAS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Restricted random allocation.
Allocation concealment (selection bias)	Unclear risk	Used sealed sequentially-numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "The identity of the various drugs was not known to the investigators until after completion of the trial".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by collection in a dish pressed against the vulva for 3 minutes: the contents were carefully measured.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.



Ogunbode 1979 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	High risk	The study was supported by funding from Sandoz.

Orji 2008

Methods	2-arm active-controlled randomised trial	
Participants	600 women were randomised in a hospital setting in Nigeria. The population comprised women of parity 6 or less, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or those with hypertension in pregnancy, packed cell volume less than 30%, previous PPH, haemoglobinopathy or cardiac disorder.	
Interventions	10 IU of oxytocin administered by an IV bolus versus 250 mcg of ergometrine administered by an IV bolus	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; manual removal of placenta. Blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; hypertension; headache.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

11.51. 67 2.46		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation was done by sealed sequentially-numbered envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss by "using a pre-weighed gauze that was weighed again after delivery".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.



Orji 2008 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of transfusion and PPH ≥ 1000 mL were omitted).
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ortiz-Gomez 2013

Methods	3-arm active-controlled randomised trial	
Participants	156 women were randomised in a hospital setting in Spain. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with comorbidities, refractory hypotension due to neuraxial blockage, vasoactive drugs needed to control haemodynamic issues or multiple pregnancy.	
Interventions	100 mcg of carbetocin administered by an IV bolus versus 61 IU of oxytocin administered by an IV bolus + infusion	
Outcomes	The study recorded the following outcomes: additional uterotonics; nausea; vomiting; headache; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated sequence.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss by the estimation of delivery attendants, but blood loss data were not reported in a format that could be extracted for the purpose of this review.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.



Ortiz-Gomez 2013 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Othman 2016

Methods	2-arm active-controlled randomised trial		
Participants	120 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with anaemia (Hb < 8 g/dL), multiple pregnancy, placental abnormality (e.g. placenta praevia, placenta abruption), polyhydramnios, 2 or more previous caesar an deliveries, current or previous history of heart disease, liver, renal disorders or known coagulopates.		
Interventions	400 mcg of misoprostol administered sublingually versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; blood loss (mL); vomiting; headache; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was done using a computer-generated random table.
Allocation concealment (selection bias)	Low risk	Quote: "Allocation concealment was done using serially numbered closed opaque envelopes. Each envelope was labelled with a serial number and had a card noting the intervention type inside. Allocation was never changed after opening the envelopes."
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Quote: "The volume of blood loss during caesarean delivery and 2 hours post- operatively was assessed. Total blood loss during caesarean delivery was measured by adding the volume of the suction bottle with the blood soaked sponges (know dry weight). Blood loss 2 hours after caesarean delivery was measured by using blood collection drape. The whole blood loss was estimat- ed by adding the blood in the suction bottle, blood soaked sponges and blood collection drape."



Othman 2016 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	120 women were randomised in the study, but 10 were excluded from the analysis from the oxytocin group after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (NCT02562300).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Owonikoko 2011

Methods	2-arm active-controlled randomised trial	
Participants	100 women were randomised in a hospital setting in Nigeria. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women requiring general anaesthesia, or those with multiple pregnancy, placenta praevia, antepartum haemorrhage, cardiac/renal/liver disorders, coagulopathy, asthma, glaucoma, pre-eclampsia, eclampsia, prolonged labour or contraindications to administration of prostaglandins.	
Interventions	20 IU of oxytocin administered by an IV infusion versus 400 mcg of misoprostol administered sublingually	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); nausea; vomiting; headache; hypotension; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	

NISK OF DIAG		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocation sequence was developed by a statistician who was not otherwise involved with the study using computer-generated table of random numbers and varied permutated blocks.
Allocation concealment (selection bias)	Low risk	Used sealed, opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "The anaesthetist was blind to the allocation until he opened each participant's envelope at surgery. The obstetricians were unaware of what oxytocic was given as the faces of the patients were screened off during the surgery".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The obstetricians were unaware of what oxytocic was given as the faces of the patients were screened off during the surgery".
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection in a suction bottle, and by weighing delivery drapes and gauzes on the basis that 1 g is equivalent to 1 mL of blood. Quote: "Both the surgeon and anaesthetist estimated blood loss independently. The scrub nurse weighed the drapes and gauze before and after the operation, noted the amount of blood in the suction bottle, and recorded



Owonikoko 2011 (Continued)		
		these. The postoperative care nurse also recorded the blood loss during the first 4 hours after surgery". Finally a research assistant (not part of the medical team) calculated the mean estimated blood loss from all these values.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Pakniat 2015

Methods	3-arm active-controlled double-dummy randomised trial		
Participants	150 women were randomised in a hospital setting in Iran. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women with any risk factor of postpartum haemorrhage i.e. anaemia (Hb < 8 g/dL), multiple pregnancy, antepartum haemorrhage, polyhydramnios, 2 or more previous caesarean sections and/or a history of previous uterine rupture, cardiac/liver/renal disorders, or known coagulopathy.		
Interventions	400 mcg of misoprostol administered sublingually versus 200 mcg plus 5 IU of misoprostol plus oxytocin administered sublingually plus by an IV bolus versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; change in Hb; nausea; vomiting; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study is stated to be double-blinded but blinding (of study participants and caregivers) was unclear. The study used dummy infusion and tablets but there was no mention of a dummy for the IV bolus that 1 of the groups received. There is insufficient detail reported to decide on the adequacy of the blinding.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Pakniat 2015 (Continued)		
Objective assessment of blood loss	Low risk	The volume of blood in the suction bottle and blood-soaked sponges was measured.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (NCT01571323).
	Low risk Low risk	

Parsons 2006

Methods	2-arm active-controlled randomised trial		
Participants	450 women were randomised in a hospital setting in Ghana. The population comprised women of both nulliparous and multiparous, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with asthma, epilepsy or contraindications to prostaglandins.		
Interventions	10 IU of oxytocin administered IM versus 800 mcg of misoprostol administered orally		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; hypertension; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

RISK Of DIAS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated allocation.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "We acknowledge that unblinding for some participants was possible because the envelopes for women who were initially randomised but who subsequently underwent caesarean section were returned and used for the next women enrolled".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians and midwives.
Incomplete outcome data (attrition bias)	Low risk	Data were collected completely from all randomised study participants.



Parsons 2006	(Continued)
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All outcomes

Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from Matercare International and the Society of Obstetricians and Gynaecologists of Canada (public funding).

Parsons 2007

Methods	2-arm active-controlled randomised trial		
Participants	450 women were randomised in a hospital setting in Ghana. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with asthma, epilepsy or contraindications to prostaglandins.		
Interventions	10 IU of oxytocin administered IM versus 800 mcg of misoprostol administered rectally		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; hypertension; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Unblinding for some participants was possible because the envelopes for women who were initially randomised but who subsequently underwent caesarean section were returned and used for the next women enrolled".
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians and midwives.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Estimated blood loss data were unavailable in 9 cases (misoprostol 7; oxytocin 2) and haemoglobin measurements (misoprostol 4; oxytocin 6) were unavailable in 10 cases.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.



Parsons 2007 (Continued)		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from Matercare International and the Society of Obstetricians and Gynaecologists of Canada (public funding).

Patil 2013

2-arm active-controlled randomised trial	
200 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with Hb level less than 7 g/dL, APH, multiple pregnancy, non-cephalic presentations, pregnancy induced hypertension, previous LSCS, induced labour, instrumental delivery, cervical tear and third-degree perineal tear, body temperature > 38° C on admission, cardiac disease, hepatic disorders and known hypersensitivity to prostaglandins.	
600 mcg of misoprostol administered orally versus 200 mcg of ergometrine administered by an IV b	
The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); diarrhoea; nausea; vomiting; headache; fever; shivering.	
Contact with study authors for additional information: no. Additional data from authors: no	
Authors' judgement Support for judgement	

Kisk of blus		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A computer-generated randomisation table was used to randomise participants.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Once the active bleeding stopped, collected blood was weighed. Swabs and pads used during 3rd stage were not counted for blood loss, but were kept to minimum of < 3.
Incomplete outcome data (attrition bias) All outcomes	Low risk	200 women were randomised in the study, but 2 were excluded because of third degree perineal tear (n = 1) and adherent placenta (n = 1) after randomisation.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.

Unclear risk



Patil 2013 (Continued)		
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.

Source(s) of funding for the study were not reported.

Patil 2016

Funding source

Patit 2010			
Methods	2-arm active-controlle	d randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with hypersensitivity to drugs, respiratory diseases, cardiac disease, renal, liver disorder, epilepsy, psychiatric disorders, pre-eclampsia, severe anaemia, multiple pregnancy, poly/oligohydramnios, previous PPH, grand multiparous.		
Interventions	10 IU of oxytocin admi	nistered IM versus 125 mcg of carboprost administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; blood loss (mL); third stage duration (minutes); diarrhoea; nausea; vomiting; headache; shivering.		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.	
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	The blood loss during third stage of labour and the immediate postpartum period (1 hour after delivery) was estimated quantitatively using Brasss V Drape.	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and random-	

they were randomised.

ly allocated to treatment were included in the analysis, in the groups to which



Patil 2016 (Continued)

Funding source Unclear risk Source(s) of funding for the study were not reported.

Penaranda 2002

Methods	3-arm active-controlled randomised trial		
Participants	78 women were randomised in a hospital setting in Colombia. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with asthma, multiple pregnancy, intrauterine death, coagulopathy, cervical tear or water in the blood collector.		
Interventions	50 mcg of misoprostol administered sublingually versus 16 mIU/min of oxytocin administered by an IV versus 200 mcg of ergometrine administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; blood loss (mL; third stage duration (minutes); vomiting; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss from cord clamping until 1 hour after delivery.
Incomplete outcome data (attrition bias) All outcomes	Low risk	3 women were excluded from the analysis after entering the study because of liquor contamination during blood collection.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.



Perez-Rumbos 2017			
Methods	2-arm active-controlled randomised trial		
Participants	500 women were randomised in a hospital setting in Venezuela. The population comprised women of parity 4 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesareans, grand multiparous (>= 5), multiple pregnancy, previous caesareans, precipitate labour, anaemia (< 6 g/dL), chorioamnionitis, previous PPH, polyhydramnios, intrauterine fetal death, APH, asthma and hypersensitivity in any of the agents, clotting disorders, renal/liver disorders, epilepsy, hypertension, or those who did not consent to the study.		
Interventions	600 mcg of misoprosto	l administered rectally versus 20 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; headache; fever; shivering.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	The numbers for the assignment to each treatment group were generated with a table of random numbers.	
Allocation concealment (selection bias)	Unclear risk	A sealed system was used that contained the location of each patient to the treatment groups. The envelopes were opened at the beginning of each treatment.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.	
Objective assessment of blood loss	Low risk	The blood lost was collected in a calibrated and all the gauzes used were weighed.	
Incomplete outcome data (attrition bias) All outcomes	High risk	500 women were randomised in the study, but 108 were excluded because of missing data after randomisation.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.	
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	



Poeschmann 1991				
Methods	3-arm controlled randomised trial			
Participants	77 women were randomised in a hospital setting in the Netherlands. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women if they had a Hobel Score of more than 10.			
Interventions	5 IU of oxytocin admin	istered IM versus 500 mcg of sulprostone administered IM versus placebo		
Outcomes		The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; manual removal of placenta; blood loss (mL); third stage duration (minutes); nausea.		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Randomisation was within blocks of 10 but the sequence generation method was not reported.		
Allocation concealment (selection bias)	Low risk	Allocated identical numbered boxes containing trial medications.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	A nurse not involved with the delivery room prepared the injections.		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinded.		
Objective assessment of blood loss	Low risk	Blood loss was calculated by measuring the amount of blood and clots collected in the bedpan and by weighing the blood-stained swabs and linen obtained for 1 hour postpartum.		
Incomplete outcome data (attrition bias) All outcomes	Low risk	77 women were randomised in the study, but 3 were excluded because of induction of labour ($n=2$) and instrumental delivery ($n=1$) after randomisation.		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.		
Funding source	Unclear risk	Sulprostone was supplied by Schering without charge but no other funding sources are reported.		

Prendiville 1988

Methods	2-arm controlled randomised trial	
Participants	1695 women were randomised in a hospital setting in the UK. The population comprised women of both nulliparous and multiparous, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with cardiac disorder, antepartum haem-	



Prendiville 1988 (Continued)	orrhage, non-cephalic presentation, multiple pregnancy, intrauterine death but after change in the protocol multiple other exclusion criteria were introduced.			
Interventions	500 mcg plus 5 IU of er	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus no treatment		
Outcomes		The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; change in Hb; NNU admissions; breastfeeding; vomiting; headache.		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.		
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.		
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		
Funding source	Low risk	The study was supported by funding from the South Western Regional Health Authority of the UK (public funding).		

Quibel 2016

Methods	2-arm placebo-controlled randomised trial
Participants	1721 women were randomised in a hospital setting in France. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with multiple pregnancies, known hypersensitivity to prostaglandins, caesarean delivery, or participation in any other treatment trial.
Interventions	400 mcg plus 10 IU of misoprostol plus oxytocin administered orally plus by an IV bolus versus 10 IU of oxytocin administered by an IV bolus



Quibel 2016 (Continued)

Outcomes

The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL;.change in Hb; diarrhoea; nausea; vomiting; fever; shivering.

Notes

Contact with study authors for additional information: no. Additional data from authors: no

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	An independent, centralised, computer-generated randomisation sequence (Clean-Web; Télémedecine Technologies, Boulogne, France) was used for this allocation based on a randomisation list established by an independent statistician according to a permuted block method balanced and stratified by centre.
Allocation concealment (selection bias)	Low risk	To conceal allocation, treatment boxes were sealed and numbered sequentially according to the randomisation sequence and were stored in the predelivery unit of each maternity ward.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The placebo tablets were provided by the pharmacy of the Assistance Publique-Hôpitaux de Paris. They were identical to misoprostol tablets in colour but their shape was slightly different. Therefore, the treatment was administered by a research midwife who did not otherwise participate in this trial, to maintain the treatment blind of patients and staff, before or after randomisation."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The placebo tablets were provided by the pharmacy of the Assistance Publique-Hôpitaux de Paris. They were identical to misoprostol tablets in colour but their shape was slightly different. Therefore, the treatment was administered by a research midwife who did not otherwise participate in this trial, to maintain the treatment blind of patients and staff, before or after randomisation."
Objective assessment of blood loss	Low risk	Quote: "Blood loss was collected into a calibrated plastic bag placed under the mother's pelvis. The transparent, graduated bag allowed continuous monitoring of blood loss and was maintained in place for at least 2 hours after the neonate's delivery. It did not require sterilization and could be used in a dorsal, lateral, or lithotomy position. Blood from blood-soaked gauze swabs was also transferred into the plastic bag."
Incomplete outcome data (attrition bias) All outcomes	Low risk	1721 women were randomised in the study, but 118 were excluded because of caesarean during labour (n = 113) and withdrawals from the study (n = 5) after randomisation.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (NCT01113229).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Low risk	Supported by a grant from Programme Hospitalier de Recherche Clinique Clinique—PHRC 2009 (Ministère de la Santé N° AOR 09010).



Methods	2-arm active-controlled	d double-dummy randomised trial		
Methods		z-ann active-controlled double-duminy randomised that		
Participants	ified parity, a singleton Exclusion criteria comp	400 women were randomised in a hospital setting in Iran. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with placenta praevia, placental abruption, coagulopathy, previous caesarean, macrosomia (more than 4 kg), polyhydramnios or uncontrolled asthma.		
Interventions	20 IU of oxytocin admi	nistered by an IV infusion versus 400 mcg of misoprostol administered orally		
Outcomes		The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL); change in Hb; hypotension; fever; shivering.		
Notes	Contact with study aut	hors for additional information: yes. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Allocation using simple randomisation with computer-generated numbers in 1:1 ratio.		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote: "double-blind": "for blinding the study, identical-appearing solutions and tablets corresponding to the 2 pharmacological groups were prepared by the pharmacy and kept in the fridge until required".		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss during the first hour after delivery, by collection with pads weighed before and after absorbance of blood.		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.		
Selective reporting (reporting bias)	High risk	The study protocol was registered (ClinicalTrials.gov NCT01863706) but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of diarrhoea, nausea and vomiting were not completely reported). Moreover, the study publication reports outcomes (hypotension, nausea, transfusion) not listed in the registered protocol.		
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		
Funding source	Low risk	The study was supported by funding from the Hormozgan University of Medical Sciences (the institution of the authors).		



Ramirez 2001				
Methods	3-arm active-controlled	3-arm active-controlled randomised trial		
Participants	comprised women of n	An unspecified number of parturients were randomised in a hospital setting in Spain. The population comprised women of nulliparous, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised multiparous women, severe anaemia, hypertensive disorders.		
Interventions	5 IU of oxytocin adminiversus no treatment	istered by an IV bolus versus 200 mcg of ergometrine administered by an IV bolus		
Outcomes	The study recorded the	e following outcomes: (No outcome data found)		
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.		
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		
Funding source	Unclear risk	Source(s) of funding for the study were not reported.		

Rashid 2009

Methods	2-arm active-controlled randomised trial
Participants	686 women were randomised in a hospital setting in Saudi Arabia. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section or requiring oxytocin



tashid 2009 (Continued)		age, or those with pre-eclampsia, cardiac disorder, hypertension on treatment, less than 37 weeks), postmaturity (more than 42 weeks) or Hb less or equal to 90	
Interventions	500 mcg plus 5 IU of er by an IV infusion	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 10 IU of oxytocin administered by an IV infusion	
Outcomes		e following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfuor placenta; blood loss (mL); third stage duration (minutes); nausea; vomiting;	
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Computer-generated random sequence of numbers.	
Allocation concealment (selection bias)	Unclear risk	Used sequentially-numbered, sealed envelopes.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.	
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.	
Objective assessment of blood loss	Low risk	Investigators appraised blood loss quote: "clinically in a standard way" by collection with a plastic sheet that was subsequently drained (with clots) into a graduated measuring jug, and by weighing swabs and towels. Quote "Any delayed haemorrhage within 24 hours after delivery was calculated".	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Outcome data were collected completely from all randomised study participants.	
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of requirement for additional syntometrine [ergometrine plus oxytocin] were omitted).	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Methods

2-arm active-controlled randomised trial



Ray 2001	(Continued)
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Participants	200 women were randomised in a hospital setting in India. The population comprised women of un-
	specified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal
	delivery. Exclusion criteria comprised women undergoing elective caesarean section, or those with
	pre-term labour (less than 32 weeks), prolonged labour, antepartum haemorrhage, pre-eclampsia,
	intrauterine death, multiple pregnancy, epilepsy, asthma, cardiac/kidney disorder, coagulopathy or
	anaemia

Interventions	400 mcg of misoprostol administered orally versus unspecified dose of ergometrine administered by an unspecified route

Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; manual removal of placenta; hypertension.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss in the first 2 hours after delivery of the placenta, by "clinical estimation". However, blood loss data were not reported in a format that could be extracted for the purpose of this review.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Reddy 2001

Methods	3-arm active-controlled randomised trial



Reddy 2001 (Continued)				
Participants	120 women were randomised in a hospital setting in India. The population comprised women of any parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with heart, liver or renal disease, asthma, epilepsy, Rhesus-negative, traumatic PPH, severe anaemia (< 6 g/dL) or hypertension.			
Interventions	200 mcg of ergometrin	e administered by an IV bolus versus 250 mcg of carboprost administered IM		
Outcomes	The study recorded the headache.	The study recorded the following outcomes: blood loss (mL); third stage duration (minutes); diarrhoea; headache.		
Notes	Contact with study aut	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.		
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.		
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.		
Funding source	Unclear risk	Source(s) of funding for the study were not reported.		

Reyes 2011

Methods	2-arm active-controlled randomised trial
Participants	144 women were randomised in a hospital setting in Panama. The population comprised women of parity 5 or more, a singleton pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing emergency caesarean section, or those with coagulopathy, unknown parity or known allergy to carbetocin.



Reyes 2011 (Continued)					
Interventions	$100\ mcg$ of carbetocin administered by an IV bolus versus 20 IU of oxytocin administered by an IV infusion				
Outcomes		The study recorded the following outcomes: additional uterotonics; transfusion; manual removal of placenta; breastfeeding; nausea; vomiting; headache; shivering; abdominal pain.			
Notes	Contact with study aut	Contact with study authors for additional information: yes. Additional data from authors: no			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.			
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.			
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.			
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.			
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.			
Selective reporting (reporting bias)	High risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of PPH were omitted).			
Intention to treat analysis	High risk	Not all study participants were included in the analysis.			
Funding source	Low risk	Ferring Pharmaceuticals donated carbetocin. No other external funding was required for the study.			

Reyes, Gonzalez 2011

Methods	2-arm active-controlled double-dummy randomised trial
Participants	57 women were randomised in a hospital setting in Panama. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by both caesarean and vaginal delivery. Exclusion criteria comprised women with HELLP syndrome, blood dyscrasia or multiple pregnancy.
Interventions	100 mcg of carbetocin administered by an IV bolus versus 10 IU of oxytocin administered by an IV infusion



Reyes, Gonzalez 2011 (Continued)

Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; change in Hb; third
	stage duration (minutes); breastfeeding; vomiting; headache; fever.

Notes Contact with study authors for additional information: yes. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated code.
Allocation concealment (selection bias)	Low risk	Used opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote "double-blind": "because the two drugs are administered differently, a double dummy system for administration was used".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	2 women were excluded from the study analysis after randomisation (quote "1 given drug before expulsion of placenta; 1 ampoule of the drug broken before use").
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who withdrew from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Rogers 1998

Methods	2-arm controlled randomised trial
Participants	1512 women were randomised in a hospital setting in the UK. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing augmentation of labour or instrumental delivery or requiring epidural analgesia, or those with placenta praevia, previous PPH, antepartum haemorrhage, Hb less than 100 g/L or mean corpuscular volume less than 75 fL, non-cephalic presentation, multiple pregnancy, intrauterine death, grand multiparity (more than 5), fibroids, anticoagulation therapy, pre-term labour (less than 32 weeks) or contraindications to any of the drugs.
Interventions	Unspecified of ergometrine plus oxytocin administered IM versus no treatment



Rogers 1998	(Continued)
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Outcomes The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; trans-

fusion; manual removal of placenta; blood loss (mL); third stage duration (minute); NNU admissions;

breastfeeding; nausea; vomiting; headache; maternal satisfaction.

Notes Contact with study authors for additional information: no. Additional data from authors: no

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The randomisation schedule used variably sized balanced blocks, and the randomisation envelopes were prepared in advance in the National Perinatal Epidemiology Unit (NEPU).
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending midwives.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Blood loss data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Public Health and Operational Research Committee of the Anglia and Oxford Regional Health Authority, UK (public funding).

Rosseland 2013

Methods	3-arm placebo-controlled randomised trial
Participants	76 women were randomised in a hospital setting in Norway. The population comprised women of unspecified parity, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Exclusion criteria comprised women with pre-eclampsia, placenta praevia, placenta accreta, von Willebrand disease or other bleeding disorder or preoperative systolic arterial pressure less than 90 mmHg.
Interventions	5 IU of oxytocin administered by an IV bolus versus 100 mcg of carbetocin administered by an IV bolus versus placebo



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Rosse	land	201	3 (Co	ntinuad)

Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; blood
	loss (mL); change in Hb; headache.

Notes Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A computer-generated list of random numbers was used. The block size varied between 6 and 9. Stratified randomisation with 2 strata, BMI less than 30 and BMI of 30 or more.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote "double-blinded": "to maintain blinding of the participants and investigators, the test medicine was delivered to the Department of Anaesthesiology in 10 mL syringes containing 5 mL of solution marked only with trial identification and randomisation numbers. The 10-mL syringes with the test medicines were prepared by a staff anaesthesiologist, who was otherwise uninvolved in the study.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss with the following formula: $(0.75 \times 1.075 \times 1.075$
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov NCT00977769).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	High risk	The study was supported by funding from Ferring Pharmaceuticals.

Sadiq 2011

Methods	2-arm active-controlled randomised trial
Participants	1865 women were randomised in a hospital setting in Nigeria. The population comprised women of parity 6 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing instrumental delivery, or those with diabetes, non-cephalic presentation, anaemia, APH, multiple pregnancy, grand multiparity (more than 6) or known allergy.
Interventions	10 IU of oxytocin administered by an IV bolus versus 600 mcg of misoprostol administered orally



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Sad	ia 2	011	(Continued)

Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfu-
	sion; blood loss (mL); change in Hb.

Notes Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random assignments generated by dice-box.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss at delivery by collection with pre-calibrated kidney dishes.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote "46 of the administered questionnaires were invalidated leaving a total of 1819 valid questionnaires (912 for oxytocin and 907 for misoprostol)." The data were further reduced through a process of computer randomisation so as to have equal study populations.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Low risk	The study was supported by funding from the University of Maiduguri Teaching Hospital. Study medications were donated by Emzor Pharmaceutical Industries.

Samimi 2013

Methods	2-arm active-controlled double-blinded randomised trial	
Participants	216 women were randomised in a hospital setting in Iran. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with hypertension, pre-eclampsia, uterine rupture, cervical tear, asthma, cardiovas-cular/renal/liver disorders, grand multiparity (not defined), fibroids or previous PPH.	
Interventions	100 mcg of carbetocin administered IM versus 200 mcg plus 5 IU of ergometrine plus oxytocin administered IM	



Samimi 2013 (Continued)

Outcomes The study recorded the following outcomes: severe maternal morbidity: intensive care admissions; ad-

ditional uterotonics; death; change in Hb; nausea; vomiting; tachycardia; hypotension; shivering ab-

dominal pain.

Notes Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was performed using a random number table.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote "Patients and medical personnel were blinded to the type of drug".
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Blood loss was not measured.
Incomplete outcome data (attrition bias) All outcomes	Low risk	At 24 hours postpartum, blood samples could not be collected from 16 women (9 in the carbetocin group and 7 in the ergometrine plus oxytocin group).
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (Iranian registry of clinical trials number 138810212854N2).
Intention to treat analysis	High risk	The authors excluded 16 study participants from the analysis because postpartum haemoglobin measurements were not available.
Funding source	Unclear risk	The study was supported by funding from the Kashan University of Medical Sciences (the institution of the authors).

Shady 2017

Methods	3-arm active-controlled randomised trial
Participants	360 women were randomised in a hospital setting in Egypt. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with medical disorders as cardiac, hepatic, renal, neurologic disorders, thromboembolic disease, blood disorders, diabetes, gestational hypertension and pre-eclampsia, grand multiparous (> 5), multiple pregnancy, polyhydramnios, macrosomia, APH, prolonged and obstructed labour, scarred uterus or previous instrumental delivery and those suffering from hypersensitivity to tranexamic acid.
Interventions	10 IU of oxytocin administered by an IV bolus versus 600 mcg of misoprostol administered sublingually



Shady 2017 (Continued)		
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; diarrhoea; nausea; vomiting.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A statistician prepared computer-generated randomisation tables.
Allocation concealment (selection bias)	Low risk	Investigators placed the allocation data in serially numbered closed opaque envelopes. Each envelope had a card noting the intervention type inside. The envelopes were opened only by the principal investigator administering the study medications according to the order of attendance of women.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Immediately after delivery of the baby, and after liquor drainage, the patient was placed over a blood drape of known weight and a graduated container was placed under the delivery bed to collect blood. The amount of blood collected in the blood drape was measured. Then the patient was given preweighed pads, which were weighed 4 hours postpartum.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Shrestha 2011

Methods	2-arm active-controlled randomised trial
Participants	200 women were randomised in a hospital setting in Nepal. The population comprised women of any parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with polyhydramnios, chorioamnionitis, preterm labour, previous caesarean, asthma, cardiac disorder or contraindication/hypersensitivity to the use of prostaglandin and uterotonics.
Interventions	1000 mcg of misoprostol administered rectally versus 10 IU of oxytocin administered IM



Shrestha 2011 (Continued)	
Outcomes	The study recorded the following outcomes: PPH at 500: severe maternal

The study recorded the following outcomes: PPH at 500; severe maternal morbidity: intensive care admissions; death; blood loss (mL); change in Hb; third stage duration (minutes); fever; abdominal pain.

Notes Contact with study authors for additional information: yes. Additional data from authors: yes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomly allocated as per the lottery technique.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss in the 48 hours postpartum, by collection with pre-weighed sterile pads and a calibrated bucket. All the soaked drapes and pads were weighed and the dry weights of these materials were subtracted to calculate blood loss on the basis that 1 g is equivalent to 1 mL.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Singh 2009

Methods	4-arm active-controlled double-dummy randomised trial		
Participants	300 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing augmentation of labour, or those with intrauterine death, antepartum haemorrhage, multiple pregnancy, malpresentation, cardiac disorder, Rhesus-negative mother, hypertension, Hb less than 70 g/L or hypersensitivity/contraindication to prostaglandins.		
Interventions	400 or 600 mcg of misoprostol administered sublingually versus 5 IU of oxytocin administered by an IV bolus versus 200 mcg of ergometrine administered by an IV bolus		



Singh 2009 (Continued)			
Outcomes		e following outcomes: PPH at 500; additional uterotonics; transfusion; manual lood loss (mL); third stage duration (minutes); fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	The drug packets were sealed and coded using a computer-generated random number chart by the same individual.	
Allocation concealment (selection bias)	Unclear risk	Used sealed drug packets.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote "double-blind": active treatments and placebo treatments were "identical" and investigators were "thus blinded".	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.	
Objective assessment of blood loss	Low risk	Investigators removed any linen soiled with amniotic fluid, and placed a disposable and absorbent pre-weighed linen saver sheet with a pre-weighed polythene bag under the mother to collect blood from the uterine cavity. Any blood clots were expressed from the vagina into the polythene bag, which was then removed and weighed. A fresh pre-weighed sanitary napkin was applied. Separate swabs were not included in the final calculation (addition of the various gravimetric measurements), that was performed 1 hour after delivery. Quote "The specific gravity of blood being 1.08, the amount of blood lost in mL was equal to the weight in grams".	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.	
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (changes in Hb measurements were unspecified beyond textual summary that quote "all groups showed a slight decrease in mean haemoglobin concentration 24 hours postpartum [maximum decrease of 0.6 g/dL]; however, the difference was not significant [ANOVA, $P > 0.05$]").	
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.	
Funding source	Unclear risk	Source(s) of funding for the study were not reported.	

Sitaula 2017

Methods	2-arm active-controlled randomised trial
Participants	200 women were randomised in a hospital setting in Nepal. The population comprised women of parity 4 or less, a singleton pregnancy, at high risk for PPH, who delivered by elective caesarean section. Ex-



Sitaula 2017 (Continued)	more caesarean delive	sed women with polyhydramnios, uncontrolled diabetes mellitus, previous 2 or ries, severe pre-eclampsia, multiple pregnancy, grand multipara, known coaguean delivery under GA, previous myomectomy, previous uterine rupture, abnoritivity to misoprostol.
Interventions		nisoprostol plus oxytocin administered rectally plus by an IV infusion versus 20 tered by an Quote infusion
Outcomes	The study recorded the	e following outcomes: PPH at 1000; transfusion; blood loss (mL); change in Hb.
Notes	Contact with study aut	hors for additional information: no. Additional data from authors: no
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generated random table.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were objective involved weighing the swabs but also visual estimation quote "fist full of clot was 500 ml".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Soltan 2007

Methods	4-arm active-controlled randomised trial
Participants	1228 women were randomised in a hospital setting in Egypt. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with traumatic PPH, blood disorders, chorioamnionitis, placenta praevia or placental abruption.



Soltan 2007 (Continued)

Interventions	200 mcg of ergometrine administered IM versus 600-1000 mcg of misoprostol administered sublingually			
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; vomiting; fever; shivering.			
Notes	Contact with study authors for additional information: yes. Additional data from authors: no			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Randomisation was computer-generated.		
Allocation concealment (selection bias)	Low risk	Used opaque, closed envelopes.		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.		
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessor blinding was not reported.		
Objective assessment of blood loss	Low risk	Investigators appraised blood loss by collection with a graduated plastic bag, and by weighing towels, linen and gauzes.		
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote "144 women were excluded from analysis because they were exposed to trauma to the perineum, vagina or cervix during labour and had traumatic excessive bleeding".		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.		
Intention to treat analysis	High risk	Not all study participants were included in the analysis.		
Funding source	Unclear risk	Source(s) of funding for the study were not reported.		

Sood 2012

Methods	2-arm placebo-controlled randomised trial		
Participants	174 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria were not specified.		
Interventions	400 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb; nausea; vomiting; fever; shivering.		



Sood 2012 (Continued)

Notes Contact with study authors for additional information: no. Additional data from authors: no

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was by computer-generated random numbers.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes made at pharmacy.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Study participants and caregivers were blinded to treatment allocations.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised intraoperative blood loss by collection with suction apparatus and sterile drapes before irrigation, and by evaluating the blood in abdominal swabs and gauzes.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Stanton 2013

Stalitoli 2013				
Methods	2-arm cluster-controlled randomised trial			
Participants	1586 women were randomised in a community setting in Ghana. The population comprised women of unspecified parity, either singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.			
Interventions	10 IU of oxytocin administered IM versus no treatment			
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; death.			
Notes	Contact with study authors for additional information: no. Additional data from authors: no			
Risk of bias				
Bias	Authors' judgement Support for judgement			



Stanton 2013 (Continued)		
Random sequence generation (selection bias)	Low risk	The 52 CHOs were randomly allocated equally to either the intervention or the control group; this allocation was stratified by both district and distance (#10 km or .10 km) to emergency obstetric care. The randomisation sequence was determined using Stata (version 12)
Allocation concealment (selection bias)	Low risk	Allocation concealment was not reported but less of an issue in cluster-randomised trials.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote "The random allocation was not masked".
Blinding of outcome assessment (detection bias) All outcomes	High risk	Assessors were not blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised postpartum blood loss by collection with a BRASS-V calibrated plastic drape placed under the mother, who was asked to remain recumbent for 1 hour following delivery of the baby, or for 2 hours if active bleeding persisted. Quote "Fluids, urine, and faeces were excluded from the blood loss measure by sweeping them to the side and into a receptacle".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote "7 and 9 enrolled women in the oxytocin and control arms, respectively, lacked a blood-loss measure".
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered (ClinicalTrials.gov NCT01108289).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Bill and Melinda Gates Foundation (public funding).

Su 2009

Methods	2-arm active-controlled double-blinded randomised trial		
Participants	370 women were randomised in a hospital setting in Singapore. The population comprised women of unspecified parity, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing elective caesarean section, or those with multiple pregnancy, previous PPH, coagulopathy, coronary artery disease, hypertension or hypersensitivity/contraindications for the use of syntometrine or carbetocin.		
Interventions	100 mcg of carbetocin administered IM versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); third stage duration (minutes); nausea; vomiting; headache; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		



Su 2009 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was blocked and stratified by parity. The randomisation list with the allocation of the mode of intervention was forwarded from the Biostatistics Unit to the Department of Pharmacy at National University Hospital, where the purchased medications were kept.
Allocation concealment (selection bias)	Low risk	Used opaque packages made at pharmacy.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote "The identities of the medications were not known to the midwives, obstetricians and the participants. The medication codes were only broken following completion of the trial".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the visual estimation of attending obstetricians and midwives.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study protocol was registered 2 years after beginning recruitment (ClinicalTrial.gov NCT00499005).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the National Healthcare Group of Singapore (public funding).

Sultana 2007



Sultana 2007 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians after collection in a plastic bowl.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Supe 2016

Random sequence genera-

tion (selection bias)

Supe 2016			
Methods	4-arm controlled randomised trial		
Participants	200 women were randomised in a hospital setting in India. The population comprised women of any parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with medical disorders like pregnancy-induced hypertension, cardiac disease, sensitivity to prostaglandins, and history of previous caesarean section.		
Interventions	800 mcg of misoprostol administered rectally versus 200 mcg of ergometrine administered IM versus 125 mcg of carboprost administered IM versus no treatment		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement Support for judgement		

Randomisation was carried out by using a randomisation table.

Low risk



Supe 2016 (Continued) Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	The blood and blood clots in the kidney tray were weighed. A plastic pouch was placed under the buttocks prior to the delivery. The blood lost was collected in this pouch. After the delivery of the placenta, the content of the pouch was transferred to a graduated jar.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	Funding was not required.

Surbek 1999

2-arm placebo-controlled randomised trial		
65 women were randomised in a Hospital setting in Switzerland. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with multiple pregnancy, pre-eclampsia, previous PPH or antepartum haemorrhage.		
600 mcg of misoprostol administered orally versus placebo		
The study recorded the following outcomes: PPH at 500; additional uterotonics; blood loss (mL); third stage duration (minute); NNU admissions; shivering.		
Contact with study authors for additional information: yes. Additional data from authors: no		
Authors' judgement	Support for judgement	
Low risk	Generated by random tables.	
Low risk	Randomisation performed by pharmacy.	
	65 women were randor unspecified parity, a sidelivery. Exclusion crit pregnancy, pre-eclamp 600 mcg of misoprosto. The study recorded the stage duration (minute.) Contact with study aut. Authors' judgement. Low risk	



Surbek 1999 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was quote "double-masked": "for proper masking, the study drugs were prepared by the hospital pharmacy as three identical gelatine capsules".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Taheripanah 2018

Methods	2-arm active-controlled randomised trial		
Participants	220 women were randomised in a hospital setting in Iran. The population comprised women of nulliparous and multiparous, a singleton pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women refusing to co-operate, major therapeutic side effects, history of cardiac and renal diseases, pre-eclampsia, and twin pregnancy.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 30 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; blood loss (mL); change in Hb; nausea; vomiting; headache.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Described as block randomisation.
Allocation concealment (selection bias)	Low risk	Selection and randomisation of the patients were performed by a coordinating nurse, using a series of sequentially numbered sealed envelopes; therefore, the sequence of allocation was hidden.
Blinding of participants and personnel (perfor- mance bias)	Unclear risk	The authors state Quote "The women and practitioners were not aware of the type of intervention" but blinding (of study participants and caregivers) was unclear as it is not described in sufficient detail.



Taheripanah 2018 (Continued)

Αl	l outcomes
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Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of evaluating blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was registered retrospectively (NCT02079558).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Tewatia 2014

Methods	2-arm active-controlled randomised trial	
Participants	100 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with grand multiparity (more than 4), anaemia, malpresentation, polyhydramnios, antepartum haemorrhage, liver/renal disorder, previous caesarean, previous PPH, uterine anomaly, traumatic PPH or contraindications to use misoprostol or oxytocin.	
Interventions	10 IU of oxytocin administered by an IV infusion versus 600 mcg of misoprostol administered sublingually	
Outcomes	The study recorded the following outcomes: PPH at 500. PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; death; blood loss (mL); change in Hb; thir stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number sequence.
Allocation concealment (selection bias)	Unclear risk	Used sequentially-numbered, opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote "Due to [the] nature of administration of the drugs, [the] patient or clinical care team could not be blinded. However, [the] statistician was unaware of the group allocation".



Tewatia 2014 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators removed any linen soiled with amniotic fluid, and placed a calibrated plastic bag under the mother to collect blood from the uterine cavity. After delivery of the placenta, a pre-weighed pad was placed high up in vagina until 1 hour afterwards. In cases of episiotomy, a separate pad was applied to the episiotomy site, and the fluid collected by this pad was not included in blood loss measurements.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Thilaganathan 1993

Methods	2-arm controlled randomised trial	
Participants	193 women were randomised in a hospital setting in UK. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or instrumental delivery, or those with grand multiparity (not defined), malpresentation, multiple pregnancy, previous caesarean, previous PPH, APH, hypertension in pregnancy, intrauterine death, PROM, cervical lacerations or third degree perineal tears.	
Interventions	No treatment versus 500 mcg plus 5 IU of ergometrine plus oxytocin administered IM	
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes).	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	
Risk of bias		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomly allocated using standard randomisation tables.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.



d)	
Unclear risk	Assessor blinding was not reported.
High risk	Investigators appraised blood loss by the estimation of attending physicians.
Low risk	Data were collected completely from all randomised study participants.
Unclear risk	The protocol of the study was unavailable for verification.
Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Low risk	The study was conducted without external funding.
	Unclear risk High risk Low risk Unclear risk Low risk

Tripti 2006

Methods	2-arm active-controlled randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of nulliparous and multiparous, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with hypertension, cardiac disease, renal disease, gastrointestinal disorders, respiratory disease, endocrinal problems, coagulation disorder, and sensitivity to prostaglandin or methergin.	
Interventions	125 mcg of carboprost administered IM versus 200 mcg of ergometrine administered by an IV bolus	
Outcomes	The study recorded the following outcomes: additional uterotonics; manual removal of placenta; blood loss (mL); third stage duration (minutes).	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was done using random number tables.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Tripti 2006 (Continued)		
Objective assessment of blood loss	Unclear risk	Blood loss was estimated by blood and blood clots collected in the kidney tray and adding the difference in the weight of the drapes before use and after delivery.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Ugwu 2014

Methods	2-arm active-controlled randomised trial		
Participants	120 women were randomised in a hospital setting in Nigeria. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emergency caesarean. Exclusion criteria comprised women requiring general anaesthesia, or those with multiple pregnancy, placenta praevia, pre-eclampsia, eclampsia, undiagnosed vaginal bleeding, prolonged labour, prolonged obstructed labour, cardiac/renal/liver disorders or fever.		
Interventions	400 mcg plus 20 IU of misoprostol plus oxytocin administered sublingually plus by an IV infusion versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additionaluUterotonics; transfusion; death; blood loss (mL); fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		
Risk of bias			
Bias	Authors' judgement Support for judgement		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Generated by random tables.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote There were no look-alike placebo tablets for women who had oxytocin alone but the fact that no member of the obstetric team had knowledge of which agent the patient received, is expected to ensure allocation concealment. In addition, there was incomplete blinding of the anaesthetist, although this was not likely to affect the study outcome, since the anaesthetist's estimated blood loss was not used."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote "There were no look-alike placebo tablets for women who had oxytocin alone but the fact that no member of the obstetric team had knowledge of which agent the patient received, is expected to ensure allocation conceal-



Ugwu 2014 (Continued)		ment. In addition, there was incomplete blinding of the anaesthetist, although this was not likely to affect the study outcome, since the anaesthetist's estimated blood loss was not used."
Objective assessment of blood loss	Low risk	Investigators appraised intraoperative and postoperative blood loss by collection in a suction bottle. Furthermore, soiled drapes, abdominal packs and pieces of gauze were weighed and the known dry weights subtracted. Finally, vulva pads applied during the 4 hours post-operation, were also weighed and the known dry weights subtracted. Measurements obtained by these 3 methods were added together. Weight measurements were performed with a weighing scale made in China, of total weighing capacity of 5 kg and graduations of 0.25 g. Investigators considered that 1 g is equivalent to 1 mL of blood.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification, but not all of the outcomes projected by methodological descriptions were reported as results in the study report (cases of nausea, vomiting, diarrhoea, headaches, fatigue, dizziness, chills, flatulence and abdominal pain were omitted).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Un Nisa 2012

Methods	2-arm active-controlled randomised trial 100 women were randomised in a hospital setting in India. The population comprised women of parity 2 to 4, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with previous PPH, multiple pregnancy, previous caesarean, macrosomia, preeclampsia, diabetes, cardiac/lung/bleeding/clotting disorders or taking anticoagulants.	
Participants		
Interventions	10 IU of oxytocin administered by an IV bolus versus 500 mcg plus 5 IU of ergometrine plus oxytoci ministered IM	
Outcomes	The study recorded the following outcomes: PPH at 500.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Study participants (patients) were divided by lottery system in the 2 groups, each group comprising of 50 patients.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias)	High risk	Study participants and caregivers were not blinded to treatment allocations.



Un Nisa 2012 (Continued)

ΛI	l outcome	_
Αl	courcome	S

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss after the delivery of baby quote "by squeezing the soaked pads and quantifying the amount of blood clots in a kidney tray of standard size to be equal to 500 mL".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Uncu 2015

Methods	5-arm controlled randomised trial		
Participants	248 women were randomised in a hospital setting in Turkey. The population comprised women of parity 5 or less, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with placenta praevia, previous PPH, APH, non-cephalic presentation, multiple pregnancy, intrauterine death, grand multiparity (more than 5), fibroids, pre-eclampsia or anticoagulation therapy.		
Interventions	No treatment versus 400 mcg to 800 mcg of misoprostol administered orally, vaginally or rectally		
Outcomes	The study recorded the following outcomes: additional uterotonics; transfusion; third stage duration (minutes); diarrhoea; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Generated by random tables.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.



Uncu 2015 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Vagge 2014

Methods	2-arm active-controlled randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with cardiac disorder in pregnancy, uterine tumour in pregnancy, secondary PPH, grand multiparity (not defined), multiple pregnancy, polyhydramnios, anaemia, coagulopathy, antepartum haemorrhage, previous PPH, prolonged labour, precipitate labour or known allergic or hypersensitivity reaction to prostaglandins.	
Interventions	10 IU of oxytocin administered by an IV infusion versus 800 mcg of misoprostol administered rectally	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; blood loss (mL); diarrhoea; nausea; fever; shivering.	
Notes	Contact with study authors for additional information: no. Additional data from authors: no	
Risk of bias		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Used simple random sampling.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Vagge 2014 (Continued)		
Objective assessment of blood loss	Unclear risk	Methods of appraising blood loss were not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Vaid 2009

Methods	3-arm active-controlled randomised trial		
Participants	200 women were randomised in a hospital setting in India. The population comprised women of parity 4 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with grand multiparity (more than 4), multiple pregnancy, preterm labour (less than 32 weeks), HELLP syndrome, polyhydramnios, coagulopathy, asthma, cardiac/renal disorce epilepsy, hypertension, Hb less than 80 g/L or known drug allergy.		
Interventions	400 mcg of misoprostol administered sublingually versus 200 mcg of ergometrine administered IM ve sus 125 mcg of carboprost administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; transfusion; manual removal of placenta; diarrhoea; nausea; vomiting; fever; shivering; abdominal pain.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Allocation by a computer-generated random number.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was unclear.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	After the drainage of amniotic fluid, investigators appraised blood loss by collection with a sterile calibrated BRASS-V drape placed under the mother. The drape remained in placed for 1 hour. Furthermore, quote "blood loss in gauze



Vaid 2009 (Continued)		pieces was calculated by subtracting the weight of dry gauze from the weight of blood-soaked gauze pieces".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Van Selm 1995

Methods	2-arm active-controlled double-dummy randomised trial		
Participants	81 women were randomised in a hospital setting in the Netherlands. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women with coagulation disorder, anticoagulant medication, multiple pregnancy, fibroids, hypertension, induction of labour.		
Interventions	200 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 500 mcg of sulprostone administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; transfusion; manual removal of placenta; blood loss (mL); third stage duration (minutes).		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Low risk	Assignment to pharmacy coded boxes occurred, after informed consent, in first stage labour.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Double-blinding of personnel and participants (placebo use).
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Double-blinding of personnel and participants (placebo use).
Objective assessment of blood loss	Low risk	Measured the blood and clots by collecting and weighing the blood stained linen and pads.
Incomplete outcome data (attrition bias)	High risk	81 women were randomised in the study, but 12 were excluded because of exclusion criteria all in the ergometrine plus oxytocin group after randomisation.



Van Selm	1995	(Continued)
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All outcomes

Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Verma 2006

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	200 women were randomised in a hospital setting in India. The population comprised women of unspecified parity, unspecified whether singleton or multiple pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria were not specified.	
Interventions	400 mcg of misoprostol administered sublingually versus 200 mcg of ergometrine administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; additional uterotonics; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); nausea; fever; Shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: no	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation was not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The study was "double-blind": active treatments and placebo treatments were "identical-looking".
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss quote: "accurately with a specially designed calibrated blood collection drape (BRASS-V drape)".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.



Verma 2006 (Continued)		
Intention to treat analysis	Unclear risk	It was unclear from the study report whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were unclear.

Vimala 2004

Methods	2-arm active-controlled randomised trial		
Participants	120 women were randomised in a hospital setting in India. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or caesarean section, or those with preterm labour (less than 37 weeks), grand multiparity (more than 5), multiple pregnancy, hypertension in pregnancy, Hb less than 80 g/L or known hypersensitivity to prostaglandins.		
Interventions	400 mcg of misoprostol administered sublingually versus 200 mcg of ergometrine administered by an IV bolus		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; blood loss (mL); change in Hb; third stage duration (minutes); nausea; vomiting; headache; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			

RISK OT DIAS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Generated by random tables.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Treatments were administered via different routes and the authors did not report any double dummy.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	0 11 ,	
Incomplete outcome data (attrition bias)	Unclear risk	The study authors did not mention any incomplete outcome data.



Vimala 2004 (Continued) All outcomes		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Vimala 2006

Methods	2-arm active-controlled randomised trial		
Participants	100 women were randomised in a hospital setting in India. The population comprised women of u specified parity, a singleton pregnancy, at high risk for PPH, who delivered by both elective or emegency caesarean. Exclusion criteria comprised women with multiple pregnancy, APH, polyhydram prolonged labour (more than 12 hours), previous more than 1 caesarean, previous uterine rupture diac/liver/renal disorder, coagulopathy or Hb less than 80 g/L.		
Interventions	400 mcg of misoprostol administered sublingually versus 20 IU of oxytocin administered by an IV infusion		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; blood loss (mL); change in Hb; vomiting; headache; fever; shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random number.
Allocation concealment (selection bias)	Low risk	Used opaque, sealed envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Study participants and caregivers were not blinded to treatment allocations.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.
Objective assessment of blood loss	Low risk	Investigators appraised blood loss intraoperatively and in the first hour post- operatively "in a standard manner". They measured the volume of blood in the suction bottle, and weighed blood-soaked sponges and linen savers. Then they added the difference between dry and blood-soaked weights of sponges and linen savers, to the volume measured in the suction bottle.
Incomplete outcome data (attrition bias)	Low risk	Data were collected completely from all randomised study participants.



Vima	la 2006	(Continued)

All outcome	25
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Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from the Division of Reproductive Health and Nutrition, Indian Council of Medical Research (public funding).

Walley 2000

Methods	2-arm active-controlled double-dummy randomised trial	
Participants	401 women were randomised in a hospital setting in Ghana. The population comprised women of parity 5 or less, a singleton pregnancy, at low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing induction or augmentation of labour or caesarean section, or those with grand multiparity (more than 5), multiple pregnancy, preterm labour (less than 32 weeks), hypertension in pregnancy, HELLP syndrome, polyhydramnios, previous PPH, coagulopathy, precipitate labour, chorioamnionitis, Hb less than 80 g/L or a known hypersensitivity to prostaglandins.	
Interventions	400 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM	
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes); diarrhoea; nausea; vomiting; fever; shivering.	
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes	

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated random numbers.
Allocation concealment (selection bias)	Low risk	Used sequentially-numbered, opaque packets made by administrative staff.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The identity of the placebo and active medications were concealed from caregivers and participants".
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	High risk	Investigators appraised blood loss by the estimation of attending physicians.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Of those women randomised, blood loss measurements were unavailable in 3 cases, and postpartum Hb samples were unavailable in 9 cases.



Walley 2000 (Continued)		
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The study was supported by funding from MaterCare International and the Canadian International Development Agency (public funding).

Whigham 2016

Methods	2-arm active-controlled double-blinded randomised trial		
Participants	122 women were randomised in a hospital setting in Australia. The population comprised women of any parity, a singleton pregnancy, at high risk for PPH, who delivered by emergency caesarean section. Exclusion criteria comprised women undergoing elective caesarean section or requiring general anaesthesia, or those with vascular/liver/renal disorders, preterm labour (less than 37 weeks), multiple pregnancy, placenta praevia, placental abruption, previous more than 2 caesareans or an adverse reaction to carbetocin/oxytocin.		
Interventions	100 mcg of carbetocin administered by an IV bolus versus 5 IU of oxytocin administered by an IV bolus		
Outcomes	The study recorded the following outcomes: PPH at 1000; additional uterotonics; transfusion; blood loss (mL); change in Hb.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated randomisation at pharmacy level and none of the operating or anaesthetic doctors will have access to this.
Allocation concealment (selection bias)	Low risk	Randomisation performed by pharmacy.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Pharmacy used a study label, which included study title, number and expiry date to cover the trade label. Patients, anaesthetists and operating obstetricians were blinded to the intervention drug. These ampoules were stocked in the emergency theatre fridge in boxes labelled only with the matching study label.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Low risk	Investigators appraised intra-operative blood loss by the estimation of attending physicians. Excess blood was collected in measuring container by suction, and weighed together with any swabs soaked in blood.
Incomplete outcome data (attrition bias) All outcomes	Low risk	114 women were randomised in the study, but 10 were excluded because they had a general anaesthetic (n = 2) or ampoules discarded (n = 8) after randomisation.



Whigham 2016 (Continued)		
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (ACTRN 12612000466842).
Intention to treat analysis	High risk	Those who were excluded from the study after randomisation were not included in the analysis.
Funding source	Low risk	This project was awarded the Peninsula Health Grant for Health Research.

Widmer 2018

Methods	2-arm active-controlled double-blinded randomised trial		
Participants	29645 women were randomised in a hospital setting in Argentina, Egypt, India, Kenya, Nigeria, Singapore, South Africa, Thailand, Uganda and the UK. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women in an advanced stage of labour (cervical dilatation > 6 cm) or who were too distressed to give informed consent, who had known allergies to carbetocin, oxytocin homologues or excipients, who had serious cardiovascular disorders, serious hepatic or renal disease, or who had epilepsy.		
Interventions	100 mcg of carbetocin administered IM versus 10 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; vomiting; abdominal pain.		
Notes	Contact with study authors for additional information: no. Additional data from authors: no		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	The random allocation sequence was generated at WHO using computer-generated random numbers. Randomisation was stratified by country using permuted blocks of size ten, with an allocation ratio 1:1.
Allocation concealment (selection bias)	Low risk	Both HS carbetocin and oxytocin were in 1 mL ampoules in consecutively numbered treatment packs arranged in dispensers. Allocation was by opening the consecutively numbered treatment pack in the dispenser.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The ampoules, trial packs and dispensers were identical in shape, size and weight ensuring that investigators were blinded to individual treatment allocation. Although carbetocin was heat stable and did not require cold storage we kept the dispensers in cold storage (2°C to 8°C) to give oxytocin maximum efficacy and maintain double-blinding.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinded.
Objective assessment of blood loss	Low risk	Once the cord was clamped and cut, a blood collection plastic drape (BRASSS-V™) was placed under the woman's buttocks. The blood was collected for 1 hour, or 2 hours if the bleeding continued beyond 1 hour. The drape with the blood was then weighed by a digital scale, the weight recorded in grams and then converted to volume (mL) at the analysis stage.



Widmer 2018 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Data were collected completely from all randomised study participants.
Selective reporting (reporting bias)	Low risk	The study report matches the study protocol that was registered prospectively (Trial registration: HRP Trial A65870; UTN U1111-1162-8519; ACTRN12614000870651; CTRI/2016/05/006969, EUDRACT 2014-004445-26).
Intention to treat analysis	Low risk	All those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Low risk	The research in this publication was supported by funding from MSD, through its MSD for Mothers Program. MSD for Mothers is an initiative of Merck & Co., Inc., Kenilworth, N.J., USA The funder had no commercial interest in the investigational drug, no influence on the protocol, the statistical analysis plan and the final manuscript; the funder could provide comments, but there was no obligation on the trial team to accept any. The HS carbetocin was provided by Ferring International Center S.A. (Saint Prex, Switzerland) and oxytocin by Novartis (Basel, Switzerland) free of charge. Neither company had any influence on any of the trial documents or processes.

Yuen 1995

Methods	2-arm active-controlled	d double-blinded randomised trial	
Participants	1000 women were randomised in a hospital setting in Hong Kong. The population comprised women of unspecified parity, a singleton pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women requiring oxytocin infusion in the third stage, or those with pre-eclampsia or cardiac disorder.		
Interventions	500 mcg plus 5 IU of ergometrine plus oxytocin administered IM versus 10 IU of oxytocin administered IM		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; severe maternal morbidity: intensive care admissions; additional uterotonics; transfusion; manual removal of placenta; death; change in Hb; nausea; vomiting; headache.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: no		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Used computer-generated random numbers.	
Allocation concealment (selection bias)	Unclear risk	Used sequentially-numbered, opaque envelopes.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "When a patient entered the study, a nursing officer who was not involved in the management of the patient drew up the indicated medication and handed this to the patient's attendants". Study participants and caregivers were thus blinded to treatment allocations until the codes were revealed after all data were collected in the study.	



Yuen 1995 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessors were blinded to treatment allocations.
Objective assessment of blood loss	Unclear risk	Investigators appraised blood loss during delivery quote: "by measuring the amount of blood clots and weighing the towels used".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "9 [randomised participants] were excluded: 3 had a twin pregnancy, 1 had blood transfusion during labour, and the other 5 had unavailable records".
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	High risk	Not all study participants were included in the analysis.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

Zachariah 2006

Methods	3-arm active-controlled randomised trial		
Participants	2023 women were randomised in a hospital setting in India. The population comprised women of both nulliparous and multiparous, unspecified whether singleton or multiple pregnancy, at both high and low risk for PPH, who delivered by vaginal delivery. Exclusion criteria comprised women undergoing caesarean section, or those with asthma, cardiac disorder, rhesus factor incompatibility or hypertension.		
Interventions	400 mcg of misoprostol administered orally versus 10 IU of oxytocin administered IM versus 200 mcg of ergometrine administered by an IV bolus		
Outcomes	The study recorded the following outcomes: PPH at 500; PPH at 1000; additional uterotonics; transfusion; manual removal of placenta; death; blood loss (mL); change in Hb; third stage duration (minutes). Diarrhea. Nausea. Vomiting. Headache. Fever. Shivering.		
Notes	Contact with study authors for additional information: yes. Additional data from authors: yes		
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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Used computer-generated random numbers.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Blinding (of study participants and caregivers) was not reported.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Assessor blinding was not reported.



Zachariah 2006 (Continued)		
Objective assessment of blood loss	Low risk	After the drainage of amniotic fluid, investigators appraised blood loss by collection with a large sterile plastic bag placed under the mother until she was transferred to the postnatal department. The blood collected in the plastic bag was then transferred to a measuring jar. Mops were not used in the labour room, and gauze pieces were counted.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study authors did not mention any incomplete outcome data.
Selective reporting (reporting bias)	Unclear risk	The protocol of the study was unavailable for verification.
Intention to treat analysis	Unclear risk	The authors did not specify whether all those who were enrolled and randomly allocated to treatment were included in the analysis, in the groups to which they were randomised.
Funding source	Unclear risk	Source(s) of funding for the study were not reported.

ACTRN: Australian Clinical Trials Registration Number; ANOVA: one-way Analysis of variance; APH: antepartum haemorrhage; ASA I or II: ASA Physical Status Classification System: ASA I represents a normal healthy patient, ASA II represents a patient with mild systemic disease; BMI: Body Mass Index; cc: cubic centimetres; CHOs: community health officers; cm: centimetres; CS: caesarean section; CTRI: Clinical Trials Registry of India; DIC: disseminated intravascular coagulopathy; dL: decilitres; EudraCT: European Clinical Trials database; fL: femtolitres (measurement of mean corpuscular volume); g: gram; Hb: haemoglobin; HELLP syndrome: Hemolysis (destruction of red blood cells), Elevated Liver enzymes (which indicate liver damage), and Low Platelet count; HIV: human Immunodeficiency virus; Hong Kong SAR: Hong Kong Special Adminstrative Region; IM: intramuscularly; IU: International Units; IV: intravenous; kg: kilograms; km: kilometres; L, litres; mcg: micrograms; MCV: mean cell volume; mg: milligrams; mgSO4: magnesium sulphate; min: minutes; mL: millilitres; mmHG: millimetres of mercury (unit of pressure); mmol: millimoles; NCT: National Clinical Trial (number); NEPU: National Perinatal Epidemiology Unit; NHS: National Health Service; nm: nanometres; NNU: Neonatal Unit; PACTR: Pan African Clinical Trials Registry; PCV: packed cell volume; PPH: postpartum haemorrhage; PROM: premature rupture of membranes; RCOG: Royal College of Obstetricians and Gynaecologists; UK: United Kingdom; UNDP/UNFPA: United Nations Development Programme/United Nations Population Fund; USA: United States of America; WHO: World Health Organization.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abdel-Aleem 2013	Not eligible intervention
Abdel-Aleem 2018	Same drug intervention both arms and only different timing of oxytocin administration
Abdollahy 2000	Not eligible intervention
Adhikari 2007	Quasi-randomised
Adnan 2017	Same drug intervention both arms only different route of oxytocin administration
Ahmed 2015	Not eligible intervention
Akinaga 2016	Not eligible intervention
Al-Harazi 2009	Same drug intervention both arms and only different route of misoprostol administration
Alam 2017	Not randomised
Ali 2012	Quasi-randomised



Study	Reason for exclusion
Ali 2018	Not randomised
Anandakrishnan 2013	Same drug intervention both arms and only different dose of carbetocin administration
Anjaneyulu 1988	Not eligible intervention
Anvaripour 2013	Intervention given after the third stage of labour
Ashwal 2016	Same drug intervention both arms only different regimen of oxytocin administration
Athavale 1991	Not eligible intervention
Ayedi 2011	Same drug intervention both arms and only different dose of oxytocin administration
Ayedi 2011b	Not eligible intervention
Ayedi 2012	Not eligible intervention
Aziz 2014	Quasi-randomised
Bader 2000	Not eligible intervention
Badhwar 1991	Not eligible intervention
Bai 2014	Not eligible intervention
Baig 2015	Not eligible intervention
Balki 2006	Same drug intervention both arms and only different dose of oxytocin administration
Banovska 2013	Not eligible intervention
Barbaro 1961	Not eligible intervention
Baumgarten 1983	Intervention given after the third stage of labour
Bhattacharya 1988	Not eligible intervention
Bhavana 2013	Not eligible intervention
Bider 1991	Not eligible intervention
Bider 1992	Not eligible intervention
Bisri 2011	Same drug intervention both arms and only different regimen of oxytocin administration
Bivins 1993	Not eligible intervention
Blum 2010	Intervention for treatment of PPH
Bonham 1963	Quasi-randomised
Bonis 2012	Quasi-randomised
Boopathi 2014	Quasi-randomised



Study	Reason for exclusion
Bose 2017	Not eligible intervention
Bulusu 2017	Same drug intervention both arms only different route of misoprostol administration
Cappiello 2006	Not eligible intervention
Carvalho 2004	Same drug intervention both arms and only different dose of oxytocin administration
Catanzarite 1990	Not eligible intervention
Chaplin 2009	Same drug intervention both arms and only different dose of oxytocin administration
Chatterjee 2016	Intervention given after the third stage of labour
Chaudhuri 2014	Intervention given after the third stage of labour
Chestnut 1987	Not eligible intervention
Chou 1994	Not eligible intervention
Chou 2015	Same drug intervention both arms only different dose of oxytocin administration
Chukudebelu 1963	Quasi-randomised
Cooper 2004	Same drug intervention both arms and only different regimen of oxytocin administration
Cordovani 2011	Same drug intervention all arms only different dose of carbetocin administration
Cordovani 2012	Same drug intervention both arms and only different dose of carbetocin administration
Dagdeviren 2016	Same drug intervention both arms only different route of oxytocin administration
Dahiya 1995	Not eligible intervention
Daley 1951	Quasi-randomised
Daly 1999	Inappropriate population
Dao 2009	Intervention for treatment of PPH
Davies 2005	Same drug intervention both arms and only different regimen of oxytocin administration
De bonis 2012	Quasi-randomised
Dell-Kuster 2017	Same drug intervention both arms only different infusion rate of carbetocin administration
Dennehy 1998	Not eligible intervention
Deshpande 2016	Not eligible intervention
Diab 1999	Quasi-randomised
Dickinson 2009	Not eligible population (terminations 2nd trimester)
Diop 2011	Study withdrawn



Study	Reason for exclusion
Dommisse 1980	Not randomised
Dong 2011	Not eligible intervention
Dumoulin 1981	Not randomised
Durocher 2012	Not randomised
Dutta 2000	Quasi-randomised
Dweck 2000	Not eligible intervention
Dzuba 2012	Same drug intervention both arms and only different route of oxytocin administration
Elati 2011	Same drug intervention both arms and only different dose of misoprostol administration
Erkkola 1984	Not eligible intervention
Farber 2013	Not eligible intervention
Farber 2015	Not eligible intervention
Fatemeh 2011	Same drug intervention both arms and only different regimen of oxytocin administration
Forster 1957	Quasi-randomised
Francis 1965	Quasi-randomised
Friedman 1957	Quasi-randomised
Fugo 1958	Quasi-randomised
Gai 2004	Not eligible intervention
Gavhane 2017	Not randomised
George 2010	Same drug intervention both arms and only different dose of oxytocin administration
Ghulmiyyah 2007	Not eligible intervention
Ghulmiyyah 2017	Same drug intervention all arms only different dose of oxytocin administration
Gobbur 2011	Not eligible intervention
Gohel 2007	Not eligible intervention
Goswami 2013	Not eligible intervention
Groeber 1960	Quasi-randomised
Gungorduk 2010	Not eligible intervention
Gungorduk 2010b	Same drug intervention both arms and only different regimen of oxytocin administration
Gungorduk 2011	Not eligible intervention



Study	Reason for exclusion
Gungorduk 2013	Not eligible intervention
Gupta 2014	Not eligible intervention
Habek 2007	Not eligible intervention
Hacker 1979	Not randomised
Halder 2013	Not eligible intervention
Hoffman 2006	Same drug intervention both arms and only different timing of oxytocin administration
Hofmeyr 2004	Intervention for treating PPH
Howard 1964	Not eligible intervention
Huh 2004	Same drug intervention both arms and only different timing of oxytocin administration
Hunt 2013	Not eligible intervention
Häivä 1994	Quasi-randomised
Ilancheran 1990	Not randomised
Irons 1994	Inappropriate population (excluded women who had PPH)
Islam 2008	Not randomised
Jackson 2001	Same drug intervention both arms and only different timing of oxytocin administration
Jagielska 2015	Not randomised
Javadi 2015	Not eligible intervention
Jiang 2001	Same drug intervention both arms and only different dose of oxytocin administration
Jin 2000	Not eligible intervention
Jolivet 1978	Not eligible intervention
Jonsson 2010	Same drug intervention both arms and only different dose of oxytocin administration
Kashanian 2010	Intervention administered after the third stage of labour
Kemp 1963	Quasi-randomised
Khan 1997	Not eligible intervention
Khan 2003	Same drug intervention both arms and only different route of misoprostol administration
Khan 2012	Same drug intervention both arms and only different regimen of oxytocin administration
Khan 2013	Same drug intervention all arms only different dose of carbetocin administration
Khanun 2011	Same drug intervention both arms and only different route of misoprostol administration



Study	Reason for exclusion
Kikutani 2003a	Innapropriate population
Kikutani 2003b	Innapropriate population
Kikutani 2006	Not randomised
King 2010	Same drug intervention both arms and only different regimen of oxytocin administration
Kintu 2012	Same drug intervention both arms and only different dose of oxytocin administration
Kiran 2012	Same drug intervention both arms and only different dose of oxytocin administration
Kore 2000	Not eligible intervention
Kovacheva 2015	Same drug intervention both arms and only different regimen of oxytocin administration
Kovavisarach 1998	Not eligible intervention
Le 2000	Not randomised
Leader 2002	Not eligible population (2nd trimester)
Li 2002	Not eligible intervention
Li 2003	Not eligible intervention
Li 2011	Not eligible intervention
Lin 2009	Not eligible intervention
Liu 1997	Not eligible intervention
Liu 2002	Not eligible intervention
Liu 2015	Not eligible intervention
Liu 2016	Not eligible intervention
Luamprapas 1994	Not eligible intervention
Maged 2015	Not eligible intervention
Makvandi 2013	Not eligible intervention
Mangla 2012	Not eligible intervention
Mankuta 2006	Not eligible intervention
Mansouri 2011	Same drug intervention both arms and only different route of misoprostol administration
Martinez 2006	Not eligible intervention
McGinty 1956	Quasi-randomised
Miller 2009	Not eligible intervention



Study	Reason for exclusion
Mirghafourvand 2015	Not eligible intervention
Mirteimouri 2013	Not randomised
Mockler 2015	Same drug intervention both arms only different route of oxytocin administration
Mohamadian 2013	Same drug intervention both arms only different timing of oxytocin administration
Mollitt 2009	Same drug intervention both arms and only different regimen of oxytocin administration
Moore 1956	Same drug intervention both arms and only different type of the same drug
Movafegh 2011	Not eligible intervention
Munishankarappa 2009	Same drug intervention both arms and only different regimen of oxytocin administration
Munn 2001	Same drug intervention both arms and only different regimen of oxytocin administration
Murphy 2009	Same drug intervention both arms and only different regimen of oxytocin administration
Murphy 2015	Same drug intervention both arms only different route of oxytocin administration
Nankali 2013	Not eligible intervention
Narenji 2012	Not randomised
Nelson 1983	Not eligible intervention
Neri-Mejia 2016	Same drug intervention all arms only different route and regimen of oxytocin administration
Newton 1961	Quasi-randomised
Nguyen-Lu 2015	Same drug intervention all arms only different dose of carbetocin administration
Nieminen 1964	Not randomised
Oberbaum 2005	Not eligible intervention
Oguz 2014	Same drug intervention both arms and only different route and timing of oxytocin administration
Ononge 2015	Self-administration of uterotonic agent
Ozalp 2010	Not eligible intervention
Ozcan 1996	Not eligible intervention
Ozkaya 2005	Inappropriate population (excluded women who had PPH)
Padhy 2006	Not eligible intervention
Palacio 2011	Same drug intervention both arms and only different dose of oxytocin administration
Paull 1977	Same drug intervention both arms and only different doses of drug administration
Pei 1996	Not randomised



Study	Reason for exclusion
Perdiou 2009	Not eligible intervention
Phromboot 2010	Not eligible intervention
Pierre 1992	Quasi-randomised
Pinder 2002	Same drug intervention both arms and only different doses of drug administration
Pisani 2012	Quasi-randomised
Porter 1991	Not eligible intervention
Priya 2015	Inappropriate population (measured blood loss after the delivery of the placenta)
Puri 2012	Not eligible intervention
Qiu 1999	Not eligible population (second stage of labour)
Quiroga 2009	Not eligible intervention
Ragab 2016	Same drug intervention both arms only different timing of misoprostol administration
Raghavan 2016	Intervention given for treatment of PPH
Rahbar 2018	Quasi-randomised
Rajwani 2000	Not eligible intervention
Ray 2012	Same drug intervention both arms only different regimen of oxytocin administration
Razali 2016	Quasi-randomised
Reddy 1989	Not eligible intervention
Rezk 2018	Same drug intervention both arms only different route of misoprostol administration
Rooney 1985	Quasi-randomised
Rosales-Ortiz 2014	Quasi-randomised
Rouse 2011	Same drug intervention both arms and only different doses of drug administration
Sadeghipour 2013	Not eligible intervention
Saito 2007	Quasi-randomised
Sallam 2018	Intervention administered for treatment of PPH
Samuels 2005	Not eligible intervention
Sangkhomkhamhang 2012	Same drug intervention both arms only different route of oxytocin administration
Sariganont 1999	Not randomised
Sarna 1997	Same drug intervention both arms and only different doses of drug administration



Study	Reason for exclusion
Sartain 2008	Same drug intervention both arms and only different doses of drug administration
Savitha 2017	Quasi-randomised
Schaefer 2004	Same drug intervention both arms and only different timings of drug administration
Schemmer 2001	Same drug intervention both arms and only different timings of drug administration
Sekhavat 2009	Not eligible intervention
Sentilhes 2015	Not eligible intervention
Senturk 2013	Not eligible intervention
Senturk 2016	Not randomised
Shahid 2013	Not eligible intervention
Sharma 2014	Not randomised
Sheehan 2011	Same drug intervention both arms and only different doses of drug administration
Shirazi 2013	Not eligible intervention
Shrestha 2007	Not eligible intervention
Shrestha 2008	Not eligible intervention
Singh 2005	Quasi-randomised
Siriwarakul 1991	Not eligible intervention
Soiva 1964	Not randomised
Soleimani 2014	Quasi-randomised
Sorbe 1978	Quasi-randomised
Soriano 1995	Quasi-randomised
Sreelatha 2017	Same drug intervention all arms only different route of misoprostol administration
Stearn 1963	Not randomised
Svanstrom 2008	Innapropriate population
Swapnika 2018	Not randomised
Symes 1984	Inapropriate population
Taj 2014	Not eligible intervention
Takagi 1976	Not eligible intervention
Tali 2016	Not eligible intervention



Study	Reason for exclusion
Tanir 2009	Not eligible intervention
Tarabrin 2012	Not eligible intervention
Tariq 2015b	Administered for treatment of PPH
Tehseen 2008	Not eligible intervention
Terry 1970	Not eligible intervention
Tessier 2000	Same drug intervention both arms and only different regimen of drug administration
Tharakan 2008	Same drug intervention both arms and only different timings of drug administration
Thomas 2007	Same drug intervention both arms and only different regimen of drug administration
Thornton 1988	Quasi-randomised
Tita 2012	Same drug intervention both arms and only different doses of drug administration
Tripti 2009	Not randomised
Tudor 2006	Not eligible intervention
Ugwu 2016	Same drug intervention both arms and only different doses of drug administration
Van den Enden 2009	Same drug intervention both arms and only different regimen of drug administration
Vasegh 2005	Quasi-randomised
Vaughan Williams 1974	Innapropriate population
Ventoskovskiy 1990	Not eligible intervention
Vogel 2004	Not eligible outcomes
Wallace 2007	Same drug intervention both arms and only different regimen of oxytocin administration
Walraven 2005	Not eligible uterotonic (oral ergometrine)
Wang 2000	Not eligible intervention
Wang 2018	Not randomised
Weeks 2015	Self-administration of uterotonic agent
Weihong 1998	Same drug intervention both arms and only different routes of drug administration
Weiss 1975	Not eligible outcomes
Wellmann 2016	Intervention administered before the third stage of labour
Wetta 2013	Same drug intervention both arms and only different doses of drug administration
Winikoff 2012	Same drug intervention both arms and only different routes of drug administration



Study	Reason for exclusion
Winikoff 2016	Same drug intervention all arms only different route of oxytocin administration
Wong 2005a	Same drug intervention both arms and only different doses of drug administration
Wong 2005b	Study withdrawn
Wright 2005	Not eligible intervention
Wu 2007	Not eligible intervention
Xu 2003	Not eligible intervention
Xu 2013	Not eligible intervention
Yamaguchi 2011	Same drug intervention both arms and only different regimens of drug administration
Yan 2000	Not eligible intervention
Yang 2001	Not eligible intervention
Young 1988	Not eligible intervention
Zamora 1999	Same drug intervention both arms and only different timings of drug administration
Zaporozhan 2013	Not eligible intervention
Zhao 1998	Not eligible intervention
Zhao 2003	Not eligible intervention
Zhou 1994	Same drug intervention both arms and only different routes of drug administration

PPH: postpartum haemorrhage

Characteristics of studies awaiting assessment [ordered by study ID]

Abdel-Aleem 1997

Methods	Randomised trial
Participants	High-risk women after vaginal delivery in Assiut, Egypt.
Interventions	Carboprost 250 mcg IM versus methylergonovine maleate 0.4 mg IV versus oxytocin 10 IU IV
Outcomes	Blood loss
Notes	Abstract only and awaiting reply from authors for additional information or full text

Alli 2013

Methods	Randomised double-blinded trial.
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Alli 2013 (Continue

Participants	Women undergoing caesareans.
Interventions	Sublingual misoprostol 600 mcg or 10 IU bolus intravenous oxytocin
Outcomes	Blood loss, need for additional uterotonics, and side effects.
Notes	Abstract only and unable to contact authors for additional information or full text

Amornpetchakul 2017

Methods	Randomised controlled trial
Participants	Women undergoing vaginal delivery in high-risk singleton pregnancies.
Interventions	5 IU of oxytocin or 100 mcg of carbetocin intravenously.
Outcomes	Blood loss, PPH, additional uterotonics.
Notes	Abstract only and awaiting reply from authors for additional information or full text

Beigi 2009

Methods	Randomised trial.
Participants	542 nulliparous pregnant women
Interventions	20 IU of oxytocin administered intravenously or 400 mcg of misoprostol administered sublingually
Outcomes	PPH (not defined), third-stage duration (minutes), headache, shivering.
Notes	Method of randomisation not clear and 'Risk of bias' assessment is uncertain. Written in Persian and awaiting translation.

Muller 1996

Methods	Randomised trial
Participants	Women with singleton pregnancies in hospital setting
Interventions	Oxytocin 5 IU IV versus no treatment
Outcomes	Change in Hb level
Notes	Abstract only and unable to contact authors for additional information or full text

Norchi 1988

ed clinical trial		
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Norchi 1988	(Continued)
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Participants	No details available
Interventions	Sulprostone versus methylergometrine
Outcomes	No details available
Notes	Abstract only and unable to contact authors for additional information or full text

Rabow 2017

Methods	Randomised trial
Participants	Healthy, singleton pregnant women undergoing elective caesarean section in spinal anaesthesia
Interventions	Carbetocin 100 mcg IV versus oxytocin 5 IU IV
Outcomes	Cardiovascular parameters and need for additional uterotonics
Notes	Abstract only and awaiting reply from authors for additional information or full text

Roy 2017

Methods	Randomised trial
Participants	Women in the third stage of labour
Interventions	Misoprostol 400 mcg PO versus oxytocin 10 IU IM
Outcomes	Blood loss, postpartum Hb and side effects
Notes	Abstract only and unable to contact authors for additional information or full text

Said 2017

Methods	Randomised trial
Participants	Women undergoing elective caesareans
Interventions	Misoprostol 600 mcg PR versus oxytocin unspecified dose IV
Outcomes	Blood loss and postpartum Hb
Notes	Abstract only and unable to contact authors for additional information or full text

Shrivasatava 2012



Shrivasatava 2012 (Continued)

Participants	Not known how many women randomised
Interventions	200 mcg of methylergometrine of unknown route or 400 mcg of misoprostol administered sublingually.
Outcomes	PPH (not defined), additional uterotonics, change in Hb level, third-stage duration (minutes), blood

Notes Method of randomisation not clear and 'Risk of bias' assessment is uncertain. Abstract only. Unable to contact authors.

Sunil 2016

Methods	Randomised trial.
Participants	Women at term with spontaneous onset of labour
Interventions	Oxytocin 10 IU IM versus carboprost tromethamine 125 mcg IM
Outcomes	Blood loss, PPH, diarrhoea
Notes	Method of randomisation not clear and 'Risk of bias' assessment is uncertain. Abstract only. Unable to contact authors.

Hb: haemoglobin; **IM:** intramuscular; **IU:** international unit; **IV:** intravascular; **mcg:** microgram; **mL:**: millilitre; **P0:** by mouth; **PPH:** postpartum haemorrhage; **PR:** rectally.

Characteristics of ongoing studies [ordered by study ID]

Balki 2017

Trial name or title	Carbetocin versus oxytocin at elective cesarean section: a double-blind, randomized controlled non-inferiority trial of high and low dose regimens
Methods	Randomised double-blinded
Participants	Women undergoing elective caesareans.
Interventions	Carbetocin 20 mcg IV versus carbetocin 100 mcg IV versus oxytocin 0.5 IU IV versus oxytocin 5 IU IV
Outcomes	Additional uterotonics, side effects
Starting date	May 25, 2017
Contact information	Mrinalini Balki, Samuel Lunenfeld Research Institute, Mount Sinai Hospital
Notes	Recruiting

Draycott 2014

Trial name or title	Intramuscular oxytocics: a comparison study of intramuscular carbetocin, syntocinon and syn-
	tometrine for the third stage of labour following vaginal birth (IMox)



Dray	ycott 2	014	(Continued)
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Methods	Randomised trial
Participants	Women delivering vaginally, singleton pregnancy
Interventions	1 dose of 100 mcg intramuscular carbetocin given for active management of the third stage of labour, immediately after the birth of the baby.
	1dose of $10IU$ intramuscular syntocinon given for active management of the third stage of labour, immediately after the birth of the baby.
	1 dose of 500 mcg/5 IU intramuscular Ssyntometrine® given for active management of the third stage of labour, immediately after the birth of the baby.
Outcomes	Use of additional uterotonic agents
Starting date	February 2015
Contact information	Tim Draycott, North Bristol NHS Trust/University of Bristol
Notes	Study Chair:

Gomez 2011

Trial name or title	Efficiency of carbetocin in the prevention of the postpartum haemorrhage: a clinical double-blinded randomised study
Methods	Open-label randomised trial.
Participants	Women undergoing a vaginal birth at home with a trained study provider.
Interventions	600 mcg of misoprostol administered orally or 10 IU of oxytocin administered intramuscularly.
Outcomes	PPH at 1000, additional uterotonics, transfusion, nausea, headache, abdominal pain.
Starting date	15/07/2010
Contact information	Milton Cesar Gomez
Notes	This study is shown as not yet recruiting.

Goudar 2016

Trial name or title	Room temperature stable carbetocin for preventing blood loss after delivery
Methods	Randomised, parallel group, active controlled trial
Participants	Pregnant women and women who have had a vaginal birth
Interventions	Carbetocin RTS 100 mcg IM versus oxytocin 10 IU IM
Outcomes	Primary: post-delivery (48-72 hours) Hb level adjusted for pre-delivery haemoglobin. Secondary:



Goudar 2016 (Continued)	
, ,	1 Blood loss of 500 mL or more within 1 hour
	2 Blood loss of 1000 mL or more within 1 hour
	3 Additional uterotonics
	4 Blood transfusion
	5 Manual removal of placenta
	6 Additional surgical procedures
	7 Maternal death
	8 Composite outcome of maternal death or severe morbidity up to time of discharge.
	9 Incidence and severity of adverse or serious adverse events up to time of discharge.

Starting date	01/07/2016
Contact information	Dr Shivaprasad S Goudar
	Womens and Childrens Health Research Unit KLE Universitys J N Medical College Nehru Nagar
	Belgaum
	KARNATAKA
	590010
	India
Notes	Completed but results not available to date

Kalahroudi 2010a

Trial name or title	Comparison of the effect of rectal misoprostol and syntometrin in prevention of postpartum hemorrhage
Methods	Double-blinded randomised trial.
Participants	200 women with a singleton pregnancy undergoing a vaginal birth.
Interventions	500 mcg of ergometrine plus 5 IU of oxytocin administered intramuscularly or 600 mcg of misoprostol administered rectally.
Outcomes	Additional uterotonics, change in Hb level.
Starting date	21/4/2010
Contact information	Dr Mansoureh Samimi
Notes	This study is shown as recruitment complete.

Kalahroudi 2010b

Trial name or title	Comparison effect of carbetocine and syntometrin in prevention of postpartum hemorrhage	
THAT HATTIC OF LILL	comparison check of carbetocine and syntometrin in prevention of postpartain nemorinage	



Kalahroudi 2010b (Continued)	
Methods	Double-blinded randomised trial.
Participants	200 women with a singleton pregnancy undergoing a vaginal birth.
Interventions	500 mcg of ergometrine plus 5 IU of oxytocin administered intramuscularly or 100 mcg of carbetocin administered intramuscularly.
Outcomes	Additional uterotonics, change in Hb level.
Starting date	21/1/2010
Contact information	Dr Mansoureh Samimi
Notes	This study is shown as recruitment complete.

Maged 2018

Trial name or title	Carbetocin versus rectal misoprostol for management of third stage of labor in women at low risk of postpartum hemorrhage
Methods	Interventional (clinical trial)
Participants	Women admitted for spontaneous, induced or augmented vaginal delivery and categorized as low risk for PPH
Interventions	Carbetocin 100 mcg IV versus misoprostol 800 mcg PR
Outcomes	Prevention of PPH after vaginal delivery and side effects
Starting date	July 2, 2017
Contact information	Ahmed Maged, Cairo University
Notes	Completed but no results posted

Moradi 2010

Trial name or title	Comparison of misoprostol and oxytocin in reduction of postpartum hemorrhage
Methods	Randomised trial.
Participants	300 women with singleton, term pregnancies.
Interventions	10 IU of oxytocin administered intravenously or 400 mcg of misoprostol administered orally.
Outcomes	Change in Hb.
Starting date	22/12/2009
Contact information	Simindokht Moradi
Notes	This study is shown as recruitment complete.



Sweed 2014

Trial name or title	Comparison between rectal and sublingual misoprostol before caesarian section to reduce intra and post-operative blood loss
Methods	Placebo-controlled randomised trial.
Participants	635 women undergoing elective caesarean with a singleton term pregnancy and only 1 previous caesarean.
Interventions	400 mcg of misoprostol administered rectally or 400 mcg of misoprostol administered sublingually or placebo.
Outcomes	Change in Hb level, blood loss.
Starting date	February 2013
Contact information	Mohamed S Sweed
Notes	Completed but no report available.

Thakur 2015

Trial name or title	A clinical trial to compare the effects of 4 drugs, oxytocin, misoprostol, 15-methylprostaglandin-F2alpha and methylergometrine in active management of third stage of labor.
Methods	Randomised, parallel group, multiple-arm trial
Participants	All non high-risk women at term pregnancy (37 to 40 weeks of gestation) who delivered vaginally
Interventions	Oxytocin 10 IU IM versus misoprostol 600 mcg PR versus 15-methyl prostaglandin F2alpha 125 mcg IM versus methylergometrine 200 mcg IM
Outcomes	Blood loss, PPH, blood transfusion, need for additional uterotonics, side effects
Starting date	01/01/2013
Contact information	Dr Priyanka Thakur
Notes	Completed but no report available

Hb: haemoglobin; **IM:** intramuscular; **IU:** international unit; **IV:** intravenous; **mcg:** microgram; **PPH:** postpartum haemorrhage; **PR:** rectally; **RTS:** room temperature stable

DATA AND ANALYSES



Comparison 1. Oxytocin vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
1 Death	3	1920	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
2 PPH >= 1000 mL	11	8782	Risk Ratio (IV, Random, 95% CI)	0.61 [0.52, 0.73]	
3 Blood transfusion	8	6717	Risk Ratio (IV, Random, 95% CI)	0.75 [0.51, 1.12]	
4 Severe maternal morbidity: intensive care admissions	1	1950	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	9	7021	Risk Ratio (IV, Random, 95% CI)	0.61 [0.52, 0.71]	
7 Additional uterotonics	8	5047	Risk Ratio (IV, Random, 95% CI)	0.43 [0.32, 0.58]	
8 Blood loss	8	3521	Mean Difference (IV, Random, 95% CI)	-118.52 [-141.40, -95.64]	
9 Change in haemoglobin	5	2304	Mean Difference (IV, Random, 95% CI)	-2.68 [-4.47, -0.89]	
10 Breastfeeding	1	1540	Risk Ratio (IV, Random, 95% CI)	1.00 [0.95, 1.05]	
11 Nausea	3	1788	Risk Ratio (IV, Random, 95% CI)	0.82 [0.47, 1.42]	
12 Vomiting	3	2150	Risk Ratio (IV, Random, 95% CI)	1.40 [0.44, 4.41]	
13 Headache	2	1704	Risk Ratio (IV, Random, 95% CI)	1.56 [0.52, 4.74]	
14 Abdominal pain	1	1665	Risk Ratio (IV, Random, 95% CI)	0.89 [0.80, 1.00]	
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16 Shivering	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
17 Fever	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
19 Maternal sense of well-being	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected	
19.1 Women's perceptions of well-being at 3 months postpar- tum: less energy than before birth	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
19.2 Women's perceptions of well-being at 3 months postpartum: experiencing (some) fatigue	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20.1 Did management influence positively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Did management influence negatively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Did management make no difference in the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 1.1. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 1 Death.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, F	Random, 95	5% CI			IV, Random, 95% CI	
de Groot 1996	0/78	0/143							Not estimable	
Jerbi 2007	0/65	0/65							Not estimable	
Stanton 2013	0/682	0/887							Not estimable	
Total (95% CI)	825	1095							Not estimable	
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)									
Heterogeneity: Not applicable										
Test for overall effect: Not applicable										
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo o	r no treatment	

Analysis 1.2. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Abdel-Aleem 2010	4/1291	0/659	-	0.36%	4.6[0.25,85.26]
Al-Sawaf 2013	1/37	6/39		0.72%	0.18[0.02,1.39]
Benchimol 2001	12/196	13/220		5.28%	1.04[0.48,2.22]
de Groot 1996	7/78	16/143		4.29%	0.8[0.34,1.87]
Jangsten 2011	82/810	138/821	=	43.27%	0.6[0.47,0.78]
Jans 2017	54/851	99/835	-	28.87%	0.54[0.39,0.74]
Jerbi 2007	0/65	0/65			Not estimable
Nordstrom 1997	32/513	43/487	-+ 	15.43%	0.71[0.45,1.1]
Poeschmann 1991	2/28	3/24		1.06%	0.57[0.1,3.14]
Rosseland 2013	0/26	0/25			Not estimable
Stanton 2013	1/682	8/887		0.71%	0.16[0.02,1.3]
Total (95% CI)	4577	4205	•	100%	0.61[0.52,0.73]
Total events: 195 (Oxytocin), 32	26 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =8.	14, df=8(P=0.42); I ² =1.77%				
		Favours Oxytocin 0	0.005 0.1 1 10 20	Favours Placebo or	no treatment



Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio			Weight Risk Ratio	
	n/N	n/N		IV, Ra	ndom, 9	5% CI		IV, Random, 95% CI
Test for overall effect: Z=5.43(P<0.0001)							
		Favours Oxytocin	0.005	0.1	1	10	200	Favours Placeho or no treatment

Analysis 1.3. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 3 Blood transfusion.

Study or subgroup	Oxytocin	tocin Placebo or no treatment			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
Abdel-Aleem 2010	8/1257	7/642		_	+		15.56%	0.58[0.21,1.6]
Al-Sawaf 2013	0/37	1/39					1.58%	0.35[0.01,8.35]
Butwick 2010	0/59	0/15						Not estimable
de Groot 1996	2/78	3/143			+		5.08%	1.22[0.21,7.16]
Jangsten 2011	18/810	23/821			-		42.77%	0.79[0.43,1.46]
Jans 2017	10/851	12/835					22.83%	0.82[0.36,1.88]
Jerbi 2007	0/65	0/65						Not estimable
Nordstrom 1997	5/513	7/487		_	+		12.19%	0.68[0.22,2.12]
Total (95% CI)	3670	3047			•		100%	0.75[0.51,1.12]
Total events: 43 (Oxytocin), 53 (Place	bo or no treatment)							
Heterogeneity: Tau ² =0; Chi ² =0.85, df=	=5(P=0.97); I ² =0%							
Test for overall effect: Z=1.39(P=0.16)								
		Favours Oxytocin	0.01	0.1	1 1	0 100	Favours Placebo or i	no treatment

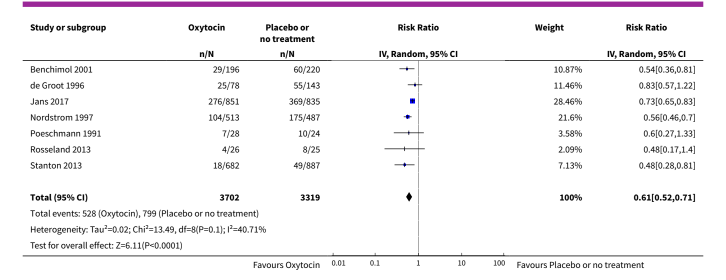
Analysis 1.4. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Abdel-Aleem 2010	0/1291	0/659							Not estimable
Total (95% CI)	1291	659							Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	oo or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	e								
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or n	o treatment

Analysis 1.6. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Ran	ndom, 9	5% CI			IV, Random, 95% CI
Abdel-Aleem 2010	63/1291	65/659		-	-			13.7%	0.49[0.35,0.69]
Al-Sawaf 2013	2/37	8/39			\pm	1		1.12%	0.26[0.06,1.16]
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or n	no treatment





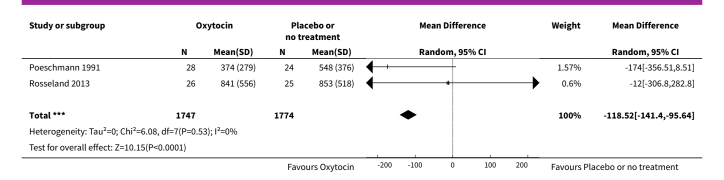
Analysis 1.7. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 7 Additional uterotonics.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI		IV, Random, 95% CI
Abdel-Aleem 2010	41/1260	55/641		-	19.26%	0.38[0.26,0.56]
Al-Sawaf 2013	2/37	8/39			3.59%	0.26[0.06,1.16]
Butwick 2010	8/59	7/15			8.83%	0.29[0.13,0.67]
de Groot 1996	14/78	26/143			13.67%	0.99[0.55,1.78]
Jans 2017	79/842	195/830		+	24.2%	0.4[0.31,0.51]
Nordstrom 1997	40/513	67/487			19.97%	0.57[0.39,0.82]
Poeschmann 1991	0/28	2/24	\leftarrow		0.97%	0.17[0.01,3.42]
Rosseland 2013	5/26	23/25			9.51%	0.21[0.09,0.46]
Total (95% CI)	2843	2204		•	100%	0.43[0.32,0.58]
Total events: 189 (Oxytocin), 383	(Placebo or no treatmer	nt)				
Heterogeneity: Tau ² =0.08; Chi ² =1	15.29, df=7(P=0.03); I ² =54	.22%				
Test for overall effect: Z=5.53(P<0	0.0001)					
		Favours Oxytocin	0.01	0.1 1 10	100 Favours Placebo or I	no treatment

Analysis 1.8. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 8 Blood loss.

Study or subgroup	0	xytocin		Placebo or Mean Differen		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Al-Sawaf 2013	37	314.7 (94.6)	39	438.6 (130.2)		20.14%	-123.9[-174.88,-72.92]
Benchimol 2001	196	278 (254)	220	382 (269.5)		20.66%	-104[-154.32,-53.68]
Butwick 2010	59	750.8 (160.9)	15	800 (256)		2.83%	-49.22[-185.12,86.68]
de Groot 1996	78	499 (454)	143	520 (419)	+	3.52%	-21[-142.93,100.93]
Jangsten 2011	810	535 (414.5)	821	680 (486.7)		27.21%	-145[-188.85,-101.15]
Nordstrom 1997	513	409 (345)	487	527 (412)		23.46%	-118[-165.23,-70.77]
			Fav	ours Oxytocin	-200 -100 0 100 20	DO Favours Pla	cebo or no treatment





Analysis 1.9. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 9 Change in haemoglobin.

Study or subgroup	O	cytocin		cebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Al-Sawaf 2013	37	12 (9)	39	13 (6)	-+-	17.27%	-1[-4.46,2.46]
Benchimol 2001	196	6.7 (20.1)	220	10.2 (23.9)	+	13.08%	-3.5[-7.73,0.73]
Jangsten 2011	810	10.6 (13.4)	821	13.5 (13.2)	-	38.86%	-2.95[-4.24,-1.66]
Jerbi 2007	65	5.1 (12.3)	65	12 (13)		12.54%	-6.9[-11.25,-2.55]
Rosseland 2013	26	8.2 (6.7)	25	8.4 (5.3)		18.26%	-0.2[-3.51,3.11]
Total ***	1134		1170		•	100%	-2.68[-4.47,-0.89]
Heterogeneity: Tau ² =1.71; Cl	hi²=6.97, df=4(P=	0.14); I ² =42.64%	ı				
Test for overall effect: Z=2.94	4(P=0)						
			Fave	ours Oxytocin	-10 -5 0 5 10	Favours Pla	cebo or no treatment

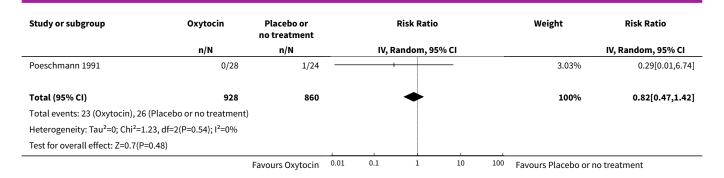
Analysis 1.10. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 10 Breastfeeding.

Study or subgroup	subgroup Oxytocin Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Jans 2017	618/779	606/761	-	100%	1[0.95,1.05]
Total (95% CI)	779	761	•	100%	1[0.95,1.05]
Total events: 618 (Oxytocin), 6	06 (Placebo or no treatmen	t)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.15(F	P=0.88)				
		Favours Oxytocin	1	Favours Placebo or i	no treatment

Analysis 1.11. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 11 Nausea.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, I	Random, 95	% CI			IV, Random, 95% CI
Butwick 2010	1/59	1/15				_		4.1%	0.25[0.02,3.83]
Jans 2017	22/841	24/821			-			92.86%	0.89[0.51,1.58]
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or r	no treatment





Analysis 1.12. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 12 Vomiting.

Study or subgroup	Oxytocin	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, F	andom, 95%	6 CI			IV, Random, 95% CI
Benchimol 2001	1/196	1/220		-				17.23%	1.12[0.07,17.83]
Butwick 2010	0/59	0/15							Not estimable
Jans 2017	6/840	4/820			-	_		82.77%	1.46[0.41,5.17]
Total (95% CI)	1095	1055				-		100%	1.4[0.44,4.41]
Total events: 7 (Oxytocin), 5 (Pl	acebo or no treatment)								
Heterogeneity: Tau ² =0; Chi ² =0.	03, df=1(P=0.86); I ² =0%								
Test for overall effect: Z=0.57(P	=0.57)								
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or r	no treatment

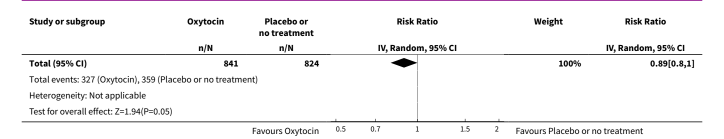
Analysis 1.13. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 13 Headache.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95% CI				IV, Random, 95% CI	
Jans 2017	18/836	14/817			-			86.96%	1.26[0.63,2.51]	
Rosseland 2013	3/26	0/25			+		•	13.04%	6.74[0.37,124.21]	
Total (95% CI)	862	842						100%	1.56[0.52,4.74]	
Total events: 21 (Oxytocin), 14	(Placebo or no treatment)									
Heterogeneity: Tau ² =0.24; Chi	² =1.21, df=1(P=0.27); l ² =17.2	26%								
Test for overall effect: Z=0.79(I	P=0.43)					1				
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or r	o treatment	

Analysis 1.14. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 14 Abdominal pain.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, F	Random, 95	% CI			IV, Random, 95% CI
Jans 2017	327/841	359/824		-				100%	0.89[0.8,1]
		Favours Oxytocin	0.5	0.7	1	1.5	2	Favours Placebo or i	no treatment





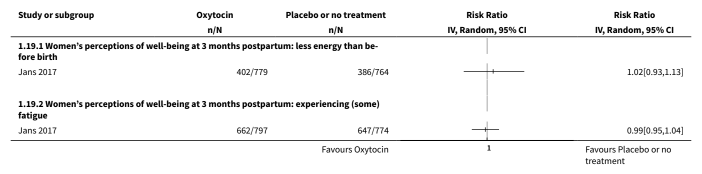
Analysis 1.16. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 16 Shivering.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Benchimol 2001	0/196	0/220							Not estimable
Total (95% CI)	196	220							Not estimable
Total events: 0 (Oxytocin), 0 (Plac	cebo or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applica	able					1			
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or	no treatment

Analysis 1.17. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 17 Fever.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio Weight			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Benchimol 2001	0/196	0/220							Not estimable
Total (95% CI)	196	220							Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	o or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	e					1			
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo o	r no treatment

Analysis 1.19. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 19 Maternal sense of well-being.





Analysis 1.20. Comparison 1 Oxytocin vs Placebo or no treatment, Outcome 20 Maternal satisfaction.

Study or subgroup	Oxytocin	Placebo or no treatment			Risk Ratio			Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI		IV, Random, 95% CI
1.20.1 Did management influer	nce positively the childbirth ex	xperiences of the mothers?						
Jangsten 2011	308/810	308/821			+			1.01[0.89,1.15]
1.20.2 Did management influer	nce negatively the childbirth e	experiences of the mothers?						
Jangsten 2011	33/810	46/821			+			0.73[0.47,1.13]
1.20.3 Did management make	no difference in the childbirth	experiences of the mothers?						
Jangsten 2011	318/810	316/821			+			1.02[0.9,1.15]
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or no treatment

Comparison 2. Carbetocin vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.75 [0.30, 1.85]
7 Additional uterotonics	2	169	Risk Ratio (IV, Random, 95% CI)	0.19 [0.12, 0.32]
8 Blood loss	1	50	Mean Difference (IV, Random, 95% CI)	-274.0 [-591.60, 43.60]
9 Change in haemoglobin	1	50	Mean Difference (IV, Random, 95% CI)	-3.40 [-7.23, 0.43]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 2.2. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Placebo or no treatment		Risk Ratio		Risk Ratio			Weight Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI		IV, Random, 95% CI	
Rosseland 2013	0/25	0/25						Not estimable	
Total (95% CI)	25	25						Not estimabl	
Total events: 0 (Carbetocin), 0 (P	Placebo or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applic	cable					1			
	Fa	avours Carbetocin	0.01	0.1	1	10	100	Favours Placebo or no treatment	

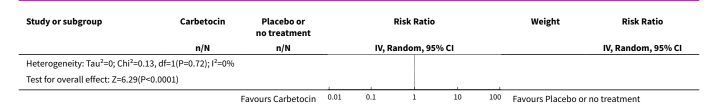
Analysis 2.6. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV,	Random, 95% CI				IV, Random, 95% CI
Rosseland 2013	6/25	8/25						100%	0.75[0.3,1.85]
Total (95% CI)	25	25						100%	0.75[0.3,1.85]
Total events: 6 (Carbetocin), 8 (Place	ebo or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.63(P=0.53	3)					1	1		
	Fa	avours Carbetocin	0.01	0.1	1	10	100	Favours Placebo or n	o treatment

Analysis 2.7. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Rand	om, 95% CI			IV, Random, 95% CI
Barton 1996	8/62	41/57		-			58.55%	0.18[0.09,0.35]
Rosseland 2013	5/25	23/25		-			41.45%	0.22[0.1,0.48]
Total (95% CI)	87	82		•			100%	0.19[0.12,0.32]
Total events: 13 (Carbetocin),	64 (Placebo or no treatment)							
	Favo	urs Carbetocin	0.01	0.1	1 10	100	Favours Placebo or i	no treatment





Analysis 2.8. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 8 Blood loss.

Study or subgroup	Carbetocin		Placebo or no treatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Rosseland 2013	25	579 (623)	25	853 (518)		100%	-274[-591.6,43.6]
Total ***	25		25			100%	-274[-591.6,43.6]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.69(P=0.09)							
			Favou	rs Carbetocin	-500 -250 0 250 500	Favours Pla	cebo or no treatment

Analysis 2.9. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 9 Change in haemoglobin.

Study or subgroup	Cai	betocin Placebo or no treatment				Mean Difference				Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ındom, 95% (CI			Random, 95% CI
Rosseland 2013	25	5 (8.2)	25	8.4 (5.3)			+			100%	-3.4[-7.23,0.43]
Total ***	25		25				•			100%	-3.4[-7.23,0.43]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.74(P=0.08)											
			Favou	rs Carbetocin	-100	-50	0	50	100	Favours Pla	cebo or no treatment

Analysis 2.13. Comparison 2 Carbetocin vs Placebo or no treatment, Outcome 13 Headache.

Study or subgroup	Carbetocin	Placebo or no treatment		F	lisk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ndom, 95%	6 CI			IV, Random, 95% CI	
Rosseland 2013	2/25	0/25		_		+		100%	5[0.25,99.16]	
Total (95% CI)	25	25		_				100%	5[0.25,99.16]	
Total events: 2 (Carbetocin), 0 (Pla	cebo or no treatment)									
Heterogeneity: Not applicable										
Test for overall effect: Z=1.06(P=0.	29)									
	Fa	avours Carbetocin	0.01	0.1	1	10	100	Favours Placebo or no	o treatment	



Comparison 3. Misoprostol vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3497	Risk Ratio (IV, Random, 95% CI)	1.00 [0.10, 9.59]
2 PPH >= 1000 mL	8	5467	Risk Ratio (IV, Random, 95% CI)	0.73 [0.56, 0.95]
3 Blood transfusion	6	3134	Risk Ratio (IV, Random, 95% CI)	0.46 [0.15, 1.47]
4 Severe maternal morbidity: intensive care admissions	1	1620	Risk Ratio (IV, Random, 95% CI)	1.00 [0.14, 7.05]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	7	4047	Risk Ratio (IV, Random, 95% CI)	0.75 [0.59, 0.94]
7 Additional uterotonics	8	3746	Risk Ratio (IV, Random, 95% CI)	0.67 [0.52, 0.87]
8 Blood loss	8	4146	Mean Difference (IV, Random, 95% CI)	-42.07 [-52.47, -31.68]
9 Change in haemoglo- bin	5	2334	Mean Difference (IV, Random, 95% CI)	-2.53 [-3.53, -1.52]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3997	Risk Ratio (IV, Random, 95% CI)	1.18 [0.78, 1.78]
12 Vomiting	6	4949	Risk Ratio (IV, Random, 95% CI)	1.41 [0.92, 2.16]
13 Headache	1	1116	Risk Ratio (IV, Random, 95% CI)	0.94 [0.32, 2.77]
14 Abdominal pain	4	1894	Risk Ratio (IV, Random, 95% CI)	0.98 [0.29, 3.37]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	5286	Risk Ratio (IV, Random, 95% CI)	2.94 [2.38, 3.64]
17 Fever	5	4396	Risk Ratio (IV, Random, 95% CI)	4.09 [2.01, 8.32]
18 Diarrhoea	6	4791	Risk Ratio (IV, Random, 95% CI)	2.80 [1.21, 6.49]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Analysis 3.1. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 1 Death.

Study or subgroup	Misoprostol	no treatment		Weight	Risk Ratio				
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Derman 2006	0/812	1/808		-	-			49.97%	0.33[0.01,8.13]
Hoj 2005	1/330	0/331			-	•		50.03%	3.01[0.12,73.6]
Mobeen 2011	0/533	0/583							Not estimable
Supe 2016	0/50	0/50							Not estimable
Total (95% CI)	1725	1772						100%	1[0.1,9.59]
Total events: 1 (Misoprostol),	1 (Placebo or no treatment)								
Heterogeneity: Tau ² =0; Chi ² =0	0.91, df=1(P=0.34); I ² =0%								
Test for overall effect: Z=0(P=	1)								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or i	no treatment

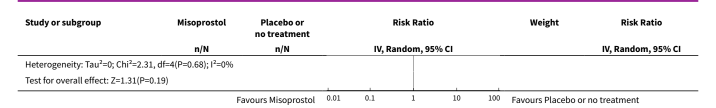
Analysis 3.2. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
Al-Sawaf 2013	2/28	6/39		2.78%	0.46[0.1,2.13]	
Bamigboye 1998a	13/270	19/272		11.94%	0.69[0.35,1.37]	
Benchimol 2001	16/186	13/220	+-	11.37%	1.46[0.72,2.95]	
Derman 2006	2/812	10/808		2.82%	0.2[0.04,0.91]	
Hofmeyr 1998	15/250	23/250	-+ 	13.82%	0.65[0.35,1.22]	
Hofmeyr 2001	27/300	29/299	-	19.58%	0.93[0.56,1.53]	
Hoj 2005	37/330	56/331		27.59%	0.66[0.45,0.98]	
Mobeen 2011	10/514	19/558		10.1%	0.57[0.27,1.22]	
Total (95% CI)	2690	2777	•	100%	0.73[0.56,0.95]	
Total events: 122 (Misoprostol),	175 (Placebo or no treatr	ment)				
Heterogeneity: Tau ² =0.02; Chi ² =	8.52, df=7(P=0.29); l ² =17.8	89%				
Test for overall effect: Z=2.36(P=	0.02)					

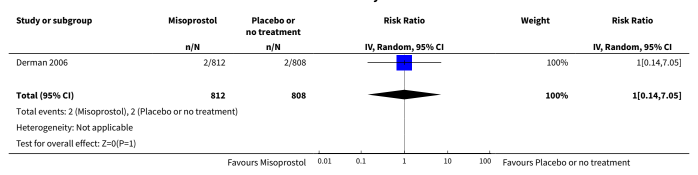
Analysis 3.3. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
Al-Sawaf 2013	0/28	1/39		•			13.34%	0.46[0.02,10.89]
Derman 2006	1/812	7/808		-	<u> </u> 		30.51%	0.14[0.02,1.15]
Hofmeyr 1998	1/250	1/250	-				17.46%	1[0.06,15.9]
Hofmeyr 2001	1/299	2/300	_	•			23.3%	0.5[0.05,5.5]
Supe 2016	0/50	0/50						Not estimable
Uncu 2015	3/199	0/49			+	_	15.39%	1.75[0.09,33.33]
Total (95% CI)	1638	1496		•			100%	0.46[0.15,1.47]
Total events: 6 (Misoprostol),	11 (Placebo or no treatment	t)						
	Fav	vours Misoprostol	0.01	0.1	1 10	100	Favours Placebo or r	no treatment





Analysis 3.4. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 4 Severe maternal morbidity: intensive care admissions.



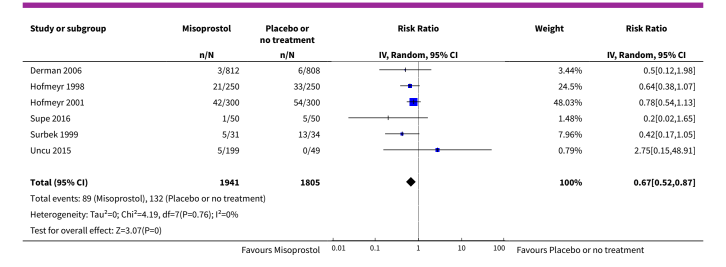
Analysis 3.6. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Al-Sawaf 2013	3/28	8/39				3.16%	0.52[0.15,1.8]
Benchimol 2001	52/186	60/220		+		20.35%	1.03[0.75,1.41]
Derman 2006	52/812	97/808				20.03%	0.53[0.39,0.74]
Fu 2003	2/80	9/76				2.21%	0.21[0.05,0.95]
Hoj 2005	150/330	170/331		=		28.47%	0.89[0.76,1.04]
Mobeen 2011	85/514	122/558		-		23.74%	0.76[0.59,0.97]
Surbek 1999	2/31	5/34	-			2.04%	0.44[0.09,2.1]
Total (95% CI)	1981	2066		•		100%	0.75[0.59,0.94]
Total events: 346 (Misoprosto	ol), 471 (Placebo or no treatr	ment)					
Heterogeneity: Tau ² =0.04; Ch	i ² =14.23, df=6(P=0.03); l ² =57	7.85%					
Test for overall effect: Z=2.49((P=0.01)				1		
	Fa	vours Misoprostol	0.01 0.1	. 1 10	100	Favours Placebo or	no treatment

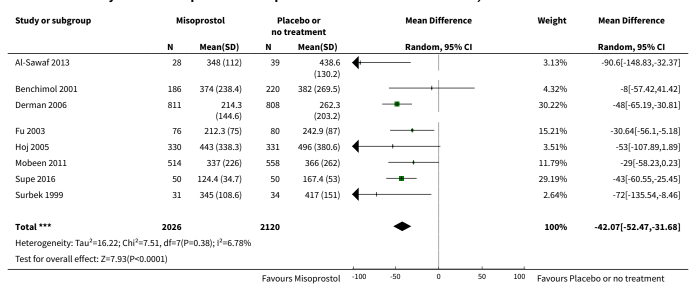
Analysis 3.7. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Al-Sawaf 2013	3/28	8/39			+-			4.32%	0.52[0.15,1.8]
Bamigboye 1998a	9/271	13/275			-			9.48%	0.7[0.31,1.62]
	Fa	Favours Misoprostol			1	10	100	Favours Placebo or r	no treatment





Analysis 3.8. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 8 Blood loss.



Analysis 3.9. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol		reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Al-Sawaf 2013	28	13 (9)	39	13 (6)	-	6.13%	0[-3.83,3.83]
Benchimol 2001	186	9.6 (23.5)	220	10.2 (23.9)		4.35%	-0.6[-5.22,4.02]
Hoj 2005	330	4.7 (18.2)	331	7.2 (18.2)	-+-	10.63%	-2.5[-5.27,0.27]
Mobeen 2011	528	11 (12)	572	13 (14)		24.13%	-2[-3.54,-0.46]
Supe 2016	50	4.9 (0.5)	50	8.1 (0.2)	•	54.76%	-3.2[-3.35,-3.05]
Total ***	1122		1212		•	100%	-2.53[-3.53,-1.52]
Heterogeneity: Tau ² =0.47; Cl	ni²=6.41, df=4(P=	0.17); I ² =37.59%					
			Favour	s Misoprostol	-10 -5 0 5 10	Favours Pla	cebo or no treatment



Study or subgroup	Mi	soprostol	Placebo or no treatment			Mear	n Diffei	ence		Weight Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rand	lom, 9!	5% CI		Random, 95% Cl
Test for overall effect: Z=4.93(P<0	.0001)									
			Favou	ırs Misoprostol	-10	-5	0	5	10	Favours Placebo or no treatment

Analysis 3.11. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 11 Nausea.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Derman 2006	35/812	29/808		+		72.56%	1.2[0.74,1.95]
Hofmeyr 2001	5/300	1/300		+		3.68%	5[0.59,42.54]
Hoj 2005	2/330	4/331	-	+		5.91%	0.5[0.09,2.72]
Mobeen 2011	8/533	8/583		-		17.84%	1.09[0.41,2.89]
Total (95% CI)	1975	2022		•		100%	1.18[0.78,1.78]
Total events: 50 (Misoprostol), 42 (Placebo or no treatme	nt)					
Heterogeneity: Tau ² =0; Chi ² =	2.76, df=3(P=0.43); I ² =0%						
Test for overall effect: Z=0.8(F	P=0.42)						
	Fa	vours Misoprostol	0.01 0.	1 1 10	100	Favours Placebo or i	no treatment

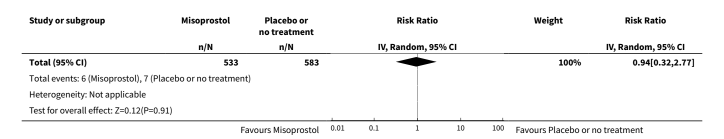
Analysis 3.12. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV	, Random, 95% CI		IV, Random, 95% CI
Bamigboye 1998a	1/271	1/275			2.41%	1.01[0.06,16.14]
Benchimol 2001	7/186	1/220		 	4.24%	8.28[1.03,66.68]
Derman 2006	28/812	25/808		-	65.64%	1.11[0.66,1.89]
Hofmeyr 2001	4/300	2/300		+	6.47%	2[0.37,10.84]
Hoj 2005	10/330	4/331		+	13.98%	2.51[0.79,7.92]
Mobeen 2011	3/533	3/583			7.25%	1.09[0.22,5.4]
Total (95% CI)	2432	2517		•	100%	1.41[0.92,2.16]
Total events: 53 (Misoprostol)), 36 (Placebo or no treatme	nt)				
Heterogeneity: Tau ² =0; Chi ² =	4.8, df=5(P=0.44); I ² =0%					
Test for overall effect: Z=1.56	(P=0.12)			į ,		
	Fa	vours Misoprostol	0.01 0.1	1 10	100 Favours Placebo or r	no treatment

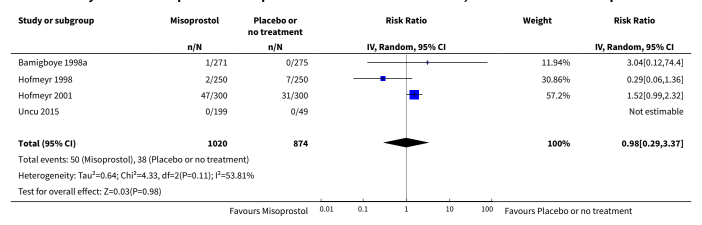
Analysis 3.13. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 13 Headache.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, F	andom, 95%	% CI			IV, Random, 95% CI
Mobeen 2011	6/533	7/583					-	100%	0.94[0.32,2.77]
	Fa	Favours Misoprostol		0.1	1	10	100	Favours Placebo or r	no treatment





Analysis 3.14. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 14 Abdominal pain.



Analysis 3.16. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 16 Shivering.

Study or subgroup	Misoprostol Placebo or Risk Ratio no treatment		Weight	Risk Ratio		
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
Bamigboye 1998a	1/34	4/36		0.96%	0.26[0.03,2.25]	
Benchimol 2001	5/186	0/220	+	0.53%	13[0.72,233.56]	
Derman 2006	419/812	140/808	•	29.5%	2.98[2.53,3.51]	
Hofmeyr 1998	48/250	13/250		9.6%	3.69[2.05,6.64]	
Hofmeyr 2001	133/300	33/300	-	18.39%	4.03[2.85,5.7]	
Hoj 2005	189/330	78/331		26.21%	2.43[1.96,3.01]	
Mobeen 2011	50/533	23/583		12.69%	2.38[1.47,3.84]	
Surbek 1999	7/31	1/34	+	1.06%	7.68[1,58.92]	
Uncu 2015	10/199	1/49		1.06%	2.46[0.32,18.78]	
Total (95% CI)	2675	2611	•	100%	2.94[2.38,3.64]	
Total events: 862 (Misoprosto	ol), 293 (Placebo or no treati	ment)				
Heterogeneity: Tau ² =0.03; Ch	i ² =14.21, df=8(P=0.08); l ² =43	3.69%				
Test for overall effect: Z=9.94((P<0.0001)					
	Fa	avours Misoprostol 0.0	1 0.1 1 10 1	.00 Favours Placebo or	no treatment	



Analysis 3.17. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 17 Fever.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IN	, Random, 95% CI		IV, Random, 95% CI	
Benchimol 2001	6/186	0/220		+	5.14%	15.36[0.87,270.93]	
Derman 2006	34/812	9/808		-	24.09%	3.76[1.81,7.79]	
Hofmeyr 2001	114/299	18/294		-	28.27%	6.23[3.89,9.97]	
Hoj 2005	78/330	11/331		-	26.02%	7.11[3.85,13.12]	
Mobeen 2011	4/533	7/583		-+-	16.48%	0.63[0.18,2.12]	
Total (95% CI)	2160	2236		•	100%	4.09[2.01,8.32]	
Total events: 236 (Misoprosto	ol), 45 (Placebo or no treatm	ent)					
Heterogeneity: Tau ² =0.41; Ch	i ² =14.39, df=4(P=0.01); l ² =72	.19%					
Test for overall effect: Z=3.89((P<0.0001)						
	Fa	vours Misoprostol	0.01 0.1	1 10	100 Favours Placebo or i	no treatment	

Analysis 3.18. Comparison 3 Misoprostol vs Placebo or no treatment, Outcome 18 Diarrhoea.

		Weight	Risk Ratio
n/N	IV, Random, 95% CI		IV, Random, 95% CI
0/275			Not estimable
2/808		30.27%	4.48[0.97,20.66]
1/300		9.24%	1[0.06,15.91]
4/331		53.57%	2.51[0.79,7.92]
0/583		- 6.92%	3.28[0.13,80.36]
0/49			Not estimable
2346	•	100%	2.8[1.21,6.49]
t)			
	avours Misoprostol 0.01	avours Misoprostol 0.01 0.1 1 10 1	avours Misoprostol 0.01 0.1 1 10 100 Favours Placebo or r

Comparison 4. Injectable prostaglandins vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.04, 3.24]
3 Blood transfusion	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6 PPH >= 500 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.55 [0.22, 1.35]
7 Additional uterotonics	2	146	Risk Ratio (IV, Random, 95% CI)	0.66 [0.21, 2.09]
8 Blood loss	2	146	Mean Difference (IV, Random, 95% CI)	-95.17 [-296.09, 105.75]
9 Change in haemoglobin	1	100	Mean Difference (IV, Random, 95% CI)	0.90 [0.56, 1.24]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.02, 8.46]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 4.1. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	•		Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% (CI	IV, Random, 95% CI
Supe 2016	0/50	0/50			Not estimable
Total (95% CI)	50	50			Not estimable
Total events: 0 (Injectable pros	staglandins), 0 (Placebo or	no treatment)			
Heterogeneity: Not applicable			ĺ		
Test for overall effect: Not appl	licable				
	Favours Injects	blo prostaglanding 0	.01 0.1 1	10 100 Favours Place	ho or no troatment



Analysis 4.2. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		i	Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95% C	:1			IV, Random, 95% CI
Poeschmann 1991	1/22	3/24						100%	0.36[0.04,3.24]
Total (95% CI)	22	24						100%	0.36[0.04,3.24]
Total events: 1 (Injectable pro	ostaglandins), 3 (Placebo or	no treatment)							
Heterogeneity: Not applicable	e								
Test for overall effect: Z=0.91((P=0.36)								
	Favours Injecta	ble prostaglandins	0.01	0.1	1	10	100	Favours Placebo or i	no treatment

Analysis 4.3. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	• • • • • • • • • • • • • • • • • • • •		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Supe 2016	0/50	0/50							Not estimable
Total (95% CI)	50	50							Not estimable
Total events: 0 (Injectable pr	ostaglandins), 0 (Placebo or	no treatment)							
Heterogeneity: Not applicab	le								
Test for overall effect: Not ap	plicable								
	Favours Injectal	ble prostaglandins	0.01	0.1	1	10	100	Favours Placebo or	no treatment

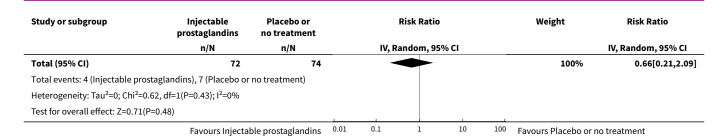
Analysis 4.6. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI	l			IV, Random, 95% CI
Poeschmann 1991	5/22	10/24		_	-			100%	0.55[0.22,1.35]
Total (95% CI)	22	24		-				100%	0.55[0.22,1.35]
Total events: 5 (Injectable pros	staglandins), 10 (Placebo o	r no treatment)							
Heterogeneity: Not applicable									
Test for overall effect: Z=1.31(P	P=0.19)					1			
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Placebo or r	no treatment

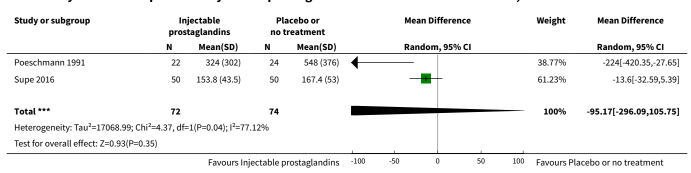
Analysis 4.7. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Poeschmann 1991	0/22	2/24		•		-		15.04%	0.22[0.01,4.29]
Supe 2016	4/50	5/50		_				84.96%	0.8[0.23,2.81]
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Placebo or n	o treatment

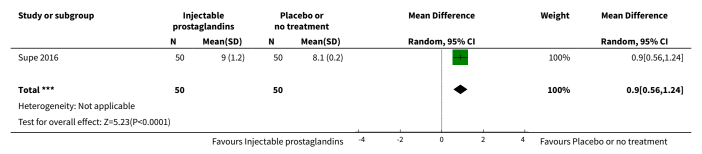




Analysis 4.8. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 8 Blood loss.



Analysis 4.9. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 9 Change in haemoglobin.



Analysis 4.11. Comparison 4 Injectable prostaglandins vs Placebo or no treatment, Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	CI			IV, Random, 95% CI
Poeschmann 1991	0/22	1/24		-	_		100%	0.36[0.02,8.46]
Total (95% CI)	22	24	_		_		100%	0.36[0.02,8.46]
Total events: 0 (Injectable prostaglar	ndins), 1 (Placebo or	no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Z=0.63(P=0.53))							
	Favours Injectal	ole prostaglandins	0.01	0.1 1	10	100	Favours Placebo or n	o treatment



Comparison 5. Ergometrine vs Placebo or no treatment

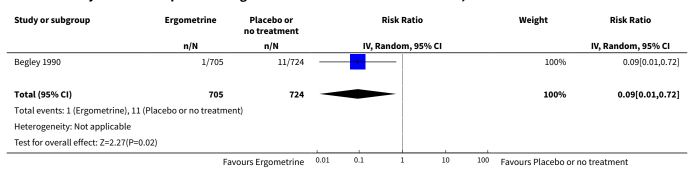
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.09 [0.01, 0.72]
3 Blood transfusion	2	1529	Risk Ratio (IV, Random, 95% CI)	0.34 [0.04, 3.28]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.42]
7 Additional uterotonics	2	1529	Risk Ratio (IV, Random, 95% CI)	0.18 [0.09, 0.37]
8 Blood loss	2	1529	Mean Difference (IV, Random, 95% CI)	-50.64 [-119.92, 18.65]
9 Change in haemoglo- bin	1	100	Mean Difference (IV, Random, 95% CI)	-0.5 [-0.58, -0.42]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	1429	Risk Ratio (IV, Random, 95% CI)	42.10 [2.55, 694.80]
12 Vomiting	1	1429	Risk Ratio (IV, Random, 95% CI)	25.67 [1.52, 432.78]
13 Headache	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [0.37, 138.91]
14 Abdominal pain	1	1429	Risk Ratio (IV, Random, 95% CI)	2.05 [1.04, 4.08]
15 Hypertension	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [2.83, 18.24]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Analysis 5.1. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 1 Death.

Study or subgroup	Ergometrine	Placebo or no treatment			Risk Ratio	Ratio Weight		Risk Ratio	
	n/N	n/N		IV, R	andom, 959	% CI			IV, Random, 95% CI
Supe 2016	0/50	0/50							Not estimable
Total (95% CI)	50	50							Not estimable
Total events: 0 (Ergometrine), 0 (Pl	acebo or no treatment	:)							
Heterogeneity: Not applicable									
Test for overall effect: Not applicab	le								
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or	no treatment

Analysis 5.2. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.



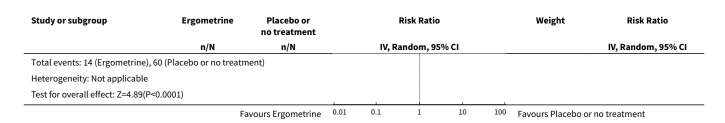
Analysis 5.3. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Begley 1990	1/705	3/724						100%	0.34[0.04,3.28]
Supe 2016	0/50	0/50			_				Not estimable
Total (95% CI)	755	774						100%	0.34[0.04,3.28]
Total events: 1 (Ergometrine), 3 (Place	ebo or no treatment)							
Heterogeneity: Not applicable									
Test for overall effect: Z=0.93(P=0.35)									
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or r	no treatment

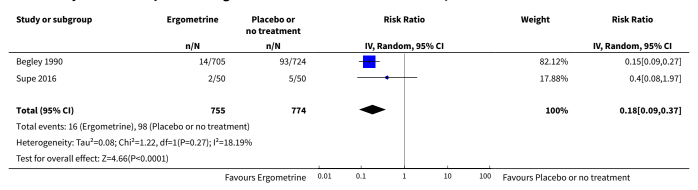
Analysis 5.6. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine	Placebo or no treatment	ı	Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Ra	ındom, 95% CI			IV, Random, 95% CI
Begley 1990	14/705	60/724	-	-		100%	0.24[0.14,0.42]
Total (95% CI)	705	724			1	100%	0.24[0.14,0.42]
	Favo	ours Ergometrine 0	.01 0.1	1 10	100	Favours Placebo or r	no treatment





Analysis 5.7. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 7 Additional uterotonics.



Analysis 5.8. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine		cebo or reatment		Me	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
Begley 1990	705	148.9 (127.1)	724	234.8 (223.9)		-			50.13%	-85.9[-104.72,-67.08]
Supe 2016	50	152.2 (49.3)	50	167.4 (53)		_	-		49.87%	-15.2[-35.25,4.85]
Total ***	755		774						100%	-50.64[-119.92,18.65]
Heterogeneity: Tau ² =2400.83	; Chi²=25.4, df=1	.(P<0.0001); I ² =9	6.06%							
Test for overall effect: Z=1.43	(P=0.15)									
			Favours	Ergometrine	-100	-50	0 50	100	Favours Pla	acebo or no treatment

Analysis 5.9. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 9 Change in haemoglobin.

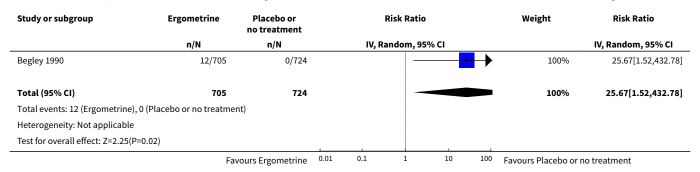
Study or subgroup	Erg	ometrine	Placebo or no treatment			Mean Difference				Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ran	ndom, 95% C	I			Random, 95% CI
Supe 2016	50	7.6 (0.2)	50	8.1 (0.2)						100%	-0.5[-0.58,-0.42]
Total ***	50		50							100%	-0.5[-0.58,-0.42]
Heterogeneity: Not applicable											
Test for overall effect: Z=12.5(P<0.0	001)										
			Favours	Ergometrine	-100	-50	0	50	100	Favours Pla	cebo or no treatment



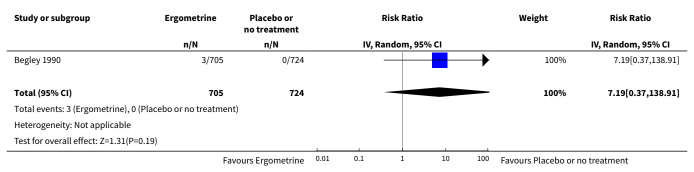
Analysis 5.11. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 11 Nausea.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ndom, 95	% CI			IV, Random, 95% CI
Begley 1990	20/705	0/724			-		—	100%	42.1[2.55,694.8]
Total (95% CI)	705	724						100%	42.1[2.55,694.8]
Total events: 20 (Ergometrine), 0 (Pl	acebo or no treatmen	t)							
Heterogeneity: Not applicable									
Test for overall effect: Z=2.61(P=0.01	.)					1			
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or r	no treatment

Analysis 5.12. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 12 Vomiting.



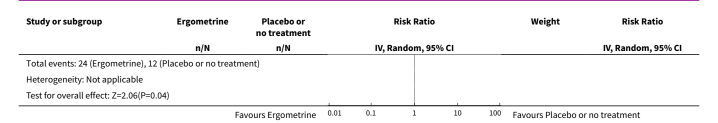
Analysis 5.13. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 13 Headache.



Analysis 5.14. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 14 Abdominal pain.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95°	% CI			IV, Random, 95% CI
Begley 1990	24/705	12/724			1	_		100%	2.05[1.04,4.08]
Total (95% CI)	705	724			•	-		100%	2.05[1.04,4.08]
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or r	no treatment





Analysis 5.15. Comparison 5 Ergometrine vs Placebo or no treatment, Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio		
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Begley 1990	35/705	5/724				-		100%	7.19[2.83,18.24]
Total (95% CI)	705	724				•		100%	7.19[2.83,18.24]
Total events: 35 (Ergometrine), 5 (Plac	ebo or no treatmen	t)							
Heterogeneity: Not applicable									
Test for overall effect: Z=4.15(P<0.000	1)					i			
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or r	no treatment

Comparison 6. Ergometrine plus oxytocin vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.44 [0.18, 1.05]
3 Blood transfusion	3	3400	Risk Ratio (IV, Random, 95% CI)	0.34 [0.18, 0.66]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.37 [0.30, 0.46]
7 Additional uterotonics	3	3400	Risk Ratio (IV, Random, 95% CI)	0.19 [0.15, 0.24]
8 Blood loss	2	1705	Mean Difference (IV, Random, 95% CI)	-35.02 [-101.63, 31.59]
9 Change in haemoglobin	2	1454	Mean Difference (IV, Random, 95% CI)	-3.57 [-6.50, -0.63]
10 Breastfeeding	2	3207	Risk Ratio (IV, Random, 95% CI)	1.03 [0.99, 1.07]
11 Nausea	1	1512	Risk Ratio (IV, Random, 95% CI)	1.95 [1.38, 2.76]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12 Vomiting	2	3207	Risk Ratio (IV, Random, 95% CI)	2.15 [1.46, 3.18]
13 Headache	2	3207	Risk Ratio (IV, Random, 95% CI)	1.65 [0.78, 3.48]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 General health at 6 weeks postpartum (Worse than prepregnancy)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 General health at 6 weeks postpartum (Exhausted since birth)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.3 General health at 6 weeks postpartum (Exhausted at 6 weeks)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.4 General health at 6 weeks postpartum (Blues)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.5 General health at 6 weeks postpartum (Depressed)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.6 General health at 6 weeks postpartum (Help for depression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.7 General health at 6 weeks postpartum (Admission to hospital for depression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.8 General health at 6 weeks postpartum (No health problems reported)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Satisfied with third-stage management	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Felt in control during third stage	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 6.2. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ra	andom, 95% CI				IV, Random, 95% CI
Prendiville 1988	7/846	26/849		-	_			46.62%	0.27[0.12,0.62]
Rogers 1998	13/748	20/764			-			53.38%	0.66[0.33,1.32]
Total (95% CI)	1594	1613		⋖	-			100%	0.44[0.18,1.05]
Total events: 20 (Ergometrine	e plus oxytocin), 46 (Placebo	o or no treatment)							
Heterogeneity: Tau ² =0.25; Ch	ni²=2.67, df=1(P=0.1); I²=62.4	8%							
Test for overall effect: Z=1.85	(P=0.06)					1			
	Favours Ergome	trine plus oxvtocin	0.01	0.1	1	10 1	100	Favours Placebo or n	o treatment

Analysis 6.3. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		R	isk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Raı	ndom, 95	% CI			IV, Random, 95% CI
Prendiville 1988	18/846	48/849		-	-			66.88%	0.38[0.22,0.64]
Rogers 1998	4/748	20/764			-			28.96%	0.2[0.07,0.59]
Thilaganathan 1993	1/103	0/90			+			4.16%	2.63[0.11,63.64]
Total (95% CI)	1697	1703		•	>			100%	0.34[0.18,0.66]
Total events: 23 (Ergometrine	e plus oxytocin), 68 (Placebo	o or no treatment)							
Heterogeneity: Tau ² =0.1; Chi	² =2.58, df=2(P=0.28); I ² =22.4	8%							
Test for overall effect: Z=3.18	s(P=0)			1			1		
	Favours Ergome	trine plus oxytocin	0.01	0.1	1	10	100	Favours Placebo or	no treatment

Analysis 6.6. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% C	ı			IV, Random, 95% CI
Prendiville 1988	50/846	152/849		+	-			50.67%	0.33[0.24,0.45]
Rogers 1998	51/748	126/764			-			49.33%	0.41[0.3,0.56]
Total (95% CI)	1594	1613			•			100%	0.37[0.3,0.46]
Total events: 101 (Ergometrin ment)	ne plus oxytocin), 278 (Place	ebo or no treat-							
Heterogeneity: Tau ² =0; Chi ² =	:1.03, df=1(P=0.31); I ² =3%								
Test for overall effect: Z=8.86	(P<0.0001)								
	Favours Ergome	trine plus oxytocin	0.01	0.1	1	10	100	Favours Placebo or i	no treatment



Analysis 6.7. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	CI			IV, Random, 95% CI
Prendiville 1988	54/846	252/849		-			68.36%	0.22[0.16,0.28]
Rogers 1998	24/748	161/764		-			30.42%	0.15[0.1,0.23]
Thilaganathan 1993	1/103	7/90					1.23%	0.12[0.02,1]
Total (95% CI)	1697	1703		•			100%	0.19[0.15,0.24]
Total events: 79 (Ergometrine ment)	e plus oxytocin), 420 (Placel	oo or no treat-						
Heterogeneity: Tau ² =0; Chi ² =	1.99, df=2(P=0.37); I ² =0%							
Test for overall effect: Z=14.00	6(P<0.0001)							
	Favours Froome	trine plus oxytocin	0.01	0.1 1	10	100	Favours Placebo or i	no treatment

Analysis 6.8. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 8 Blood loss.

Study or subgroup	Ū	ometrine oxytocin		acebo or reatment		Mea	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ran	ıdom, 95% CI			Random, 95% CI
Rogers 1998	748	268.5 (246.1)	764	336.5 (243.2)	-	_			51.5%	-68[-92.67,-43.33]
Thilaganathan 1993	103	200 (74.1)	90	200 (148.3)		_	-		48.5%	0[-33.81,33.81]
Total ***	851		854						100%	-35.02[-101.63,31.59]
Heterogeneity: Tau ² =2083.99;	Chi ² =10.14, df=	1(P=0); I ² =90.14 ⁰	%							
Test for overall effect: Z=1.03(F	P=0.3)									
		Favours Ergo	ometrine	plus oxytocin	-100	-50	0 50	100	Favours Pla	acebo or no treatment

Analysis 6.9. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 9 Change in haemoglobin.

Study or subgroup	-	ometrine oxytocin	Placebo or no treatment			Mean Difference			Weight		Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95% (:1			Random, 95% CI
Prendiville 1988	634	1 (21)	627	6 (13)			+			52.23%	-5[-6.93,-3.07]
Thilaganathan 1993	103	5 (8.1)	90	7 (8.1)						47.77%	-2[-4.29,0.29]
Total ***	737		717				•			100%	-3.57[-6.5,-0.63]
Heterogeneity: Tau ² =3.33; Ch	i ² =3.86, df=1(P=	0.05); I ² =74.1%									
Test for overall effect: Z=2.38((P=0.02)										
		Favours Erge	ometrine	plus oxytocin	-100	-50	0	50	100	Favours Pla	cebo or no treatment



Analysis 6.10. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 10 Breastfeeding.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Prendiville 1988	637/846	632/849		•		57.43%	1.01[0.96,1.07]
Rogers 1998	546/748	531/764		•		42.57%	1.05[0.99,1.12]
Total (95% CI)	1594	1613				100%	1.03[0.99,1.07]
Total events: 1183 (Ergometri ment)	ine plus oxytocin), 1163 (Pla	acebo or no treat-					
Heterogeneity: Tau ² =0; Chi ² =	0.76, df=1(P=0.38); I ² =0%						
Test for overall effect: Z=1.29	(P=0.2)						
	Favours Frgome	trine plus oxytocin	0.01	0.1 1 10	100	Favours Placebo or	no treatment

Analysis 6.11. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 11 Nausea.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, F	Random, 95%	6 CI			IV, Random, 95% CI
Rogers 1998	86/748	45/764			-			100%	1.95[1.38,2.76]
Total (95% CI)	748	764			•			100%	1.95[1.38,2.76]
Total events: 86 (Ergometrine plus	s oxytocin), 45 (Placebo	o or no treatment)							
Heterogeneity: Not applicable									
Test for overall effect: Z=3.79(P=0)									
	Favours Ergome	trine plus oxytocin	0.01	0.1	1	10	100	Favours Placebo or i	no treatment

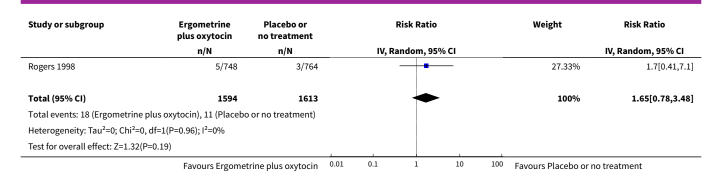
Analysis 6.12. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 12 Vomiting.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
Prendiville 1988	102/846	55/849			-		64.91%	1.86[1.36,2.55]
Rogers 1998	47/748	17/764			-		35.09%	2.82[1.64,4.87]
Total (95% CI)	1594	1613			•		100%	2.15[1.46,3.18]
Total events: 149 (Ergometrin ment)	ne plus oxytocin), 72 (Placeb	oo or no treat-						
Heterogeneity: Tau ² =0.04; Ch	ni ² =1.69, df=1(P=0.19); l ² =40.	.7%						
Test for overall effect: Z=3.86	(P=0)							
	Favours Ergome	trine plus oxytocin	0.01	0.1	1 10	100	Favours Placebo or	no treatment

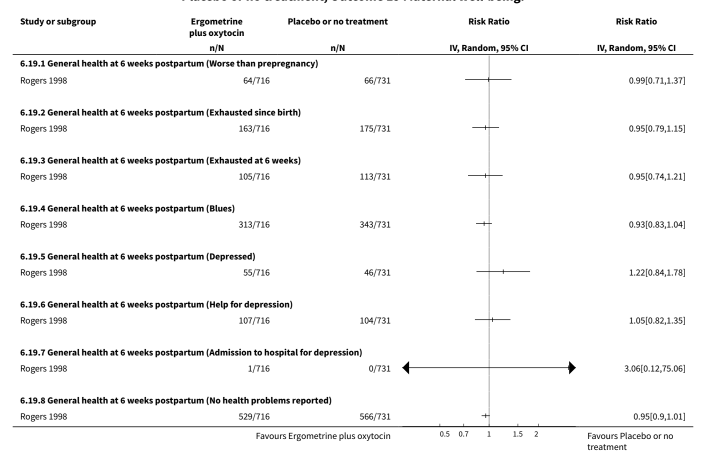
Analysis 6.13. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 13 Headache.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, R	andom, 95°	% CI			IV, Random, 95% CI
Prendiville 1988	13/846	8/849			+	-		72.67%	1.63[0.68,3.91]
	Favours Ergomet	rine plus oxytocin	0.01	0.1	1	10	100	Favours Placebo or r	no treatment





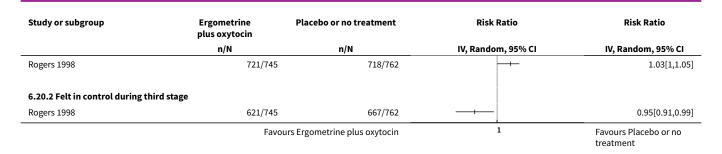
Analysis 6.19. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 19 Maternal well-being.



Analysis 6.20. Comparison 6 Ergometrine plus oxytocin vs Placebo or no treatment, Outcome 20 Maternal satisfaction.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio	Risk Ratio
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI
6.20.1 Satisfied with third-stag	e management			
	Fa	vours Ergometrine plus oxytocin	1	Favours Placebo or no treatment





Comparison 7. Misoprostol plus oxytocin vs Placebo or no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Comparison 8. Misoprostol vs Oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	24	28520	Risk Ratio (IV, Random, 95% CI)	0.62 [0.14, 2.74]
2 PPH >= 1000 mL	38	34261	Risk Ratio (IV, Random, 95% CI)	1.26 [1.11, 1.43]
3 Blood transfusion	42	35467	Risk Ratio (IV, Random, 95% CI)	0.81 [0.65, 1.00]
4 Severe maternal morbidity: intensive care admissions	10	21698	Risk Ratio (IV, Random, 95% CI)	1.16 [0.55, 2.43]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	44	36920	Risk Ratio (IV, Random, 95% CI)	1.08 [0.94, 1.24]
7 Additional uterotonics	47	35981	Risk Ratio (IV, Random, 95% CI)	1.01 [0.85, 1.20]
8 Blood loss	43	35239	Mean Difference (IV, Random, 95% CI)	-8.90 [-23.45, 5.65]
9 Change in haemoglobin	31	12028	Mean Difference (IV, Random, 95% CI)	-0.14 [-0.74, 0.47]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	33	29732	Risk Ratio (IV, Random, 95% CI)	1.22 [0.93, 1.60]
12 Vomiting	41	32687	Risk Ratio (IV, Random, 95% CI)	1.51 [1.19, 1.91]
13 Headache	10	4079	Risk Ratio (IV, Random, 95% CI)	0.88 [0.54, 1.42]
14 Abdominal pain	8	3382	Risk Ratio (IV, Random, 95% CI)	0.91 [0.79, 1.06]
15 Hypertension	3	1028	Risk Ratio (IV, Random, 95% CI)	3.64 [0.60, 22.27]
16 Shivering	49	34865	Risk Ratio (IV, Random, 95% CI)	4.02 [3.23, 4.99]
17 Fever	41	33008	Risk Ratio (IV, Random, 95% CI)	3.75 [2.73, 5.15]
18 Diarrhoea	26	30733	Risk Ratio (IV, Random, 95% CI)	2.13 [1.55, 2.93]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20.1 Satisfied or very satisfied with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Complaints about or problems with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Would take drug again after subsequent deliveries	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.4 Would recommend drug to a friend	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 8.1. Comparison 8 Misoprostol vs Oxytocin, Outcome 1 Death.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Afolabi 2010	0/100	0/100			Not estimable
Alwani 2014	0/100	1/100 —	•	21.54%	0.33[0.01,8.09]
Amin 2014	0/100	0/100			Not estimable
Atukunda 2014	0/570	0/570			Not estimable
Baskett 2007	0/311	0/311			Not estimable
Bellad 2012	0/321	0/331			Not estimable
Bhatti 2014	0/60	0/60			Not estimable
Chaudhuri 2010	0/96	0/94			Not estimable
Chaudhuri 2012	0/265	0/265			Not estimable
Diop 2016	0/252	0/141			Not estimable
Elbohoty 2016	0/89	0/86			Not estimable
Gulmezoglu 2001	2/9225	2/9230		57.02%	1[0.14,7.1]
Kundodyiwa 2001	0/243	0/256			Not estimable
Lumbiganon 1999	0/397	0/200			Not estimable
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	0/257	0/257			Not estimable
Oboro 2003	0/247	0/249			Not estimable
Parsons 2006	0/225	0/225			Not estimable
Parsons 2007	0/224	1/226 —	•	21.45%	0.34[0.01,8.21]
Perez-Rumbos 2017	0/195	0/197			Not estimable
Shrestha 2011	0/100	0/100			Not estimable
Tewatia 2014	0/50	0/50			Not estimable
Walley 2000	0/202	0/196			Not estimable
Zachariah 2006	0/730	0/617			Not estimable
Total (95% CI)	14459	14061		100%	0.62[0.14,2.74]
Total events: 2 (Misoprostol), 4 (Ox	ytocin)				
Heterogeneity: Tau²=0; Chi²=0.52,	df=2(P=0.77); I ² =0%				
Test for overall effect: Z=0.62(P=0.5	53)				



Analysis 8.2. Comparison 8 Misoprostol vs Oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Acharya 2001	1/30	1/30		0.21%	1[0.07,15.26
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	2/28	1/37		0.29%	2.64[0.25,27.7
Atukunda 2014	18/570	14/570	+-	3.32%	1.29[0.65,2.56
Baskett 2007	14/311	7/311	+	1.97%	2[0.82,4.89
Bellad 2012	0/321	0/331			Not estimable
Benchimol 2001	16/186	12/196	+	3.03%	1.41[0.68,2.89
Bhatti 2014	0/60	0/60			Not estimable
Bugalho 2001	0/323	1/339 —	+	0.15%	0.35[0.01,8.56
Caliskan 2002	17/396	14/407	+	3.27%	1.25[0.62,2.5
Caliskan 2003	14/388	15/384	-	3.08%	0.92[0.45,1.89
Chaudhuri 2010	1/96	6/94 -		0.36%	0.16[0.02,1.33
Chaudhuri 2012	1/265	2/265		0.27%	0.5[0.05,5.48
Cook 1999	13/424	0/129	+	0.2%	8.26[0.49,137.98
Diallo 2017	2/154	4/150		0.56%	0.49[0.09,2.62
Elbohoty 2016	7/89	5/86	 _	1.28%	1.35[0.45,4.1
Gavilanes 2016	12/50	13/50	-	3.41%	0.92[0.47,1.82
Gerstenfeld 2001	15/154	14/161		3.27%	1.12[0.56,2.24
Gulmezoglu 2001	366/9214	263/9228	•	64.84%	1.39[1.19,1.63
Gupta 2006	0/100	0/100			Not estimable
Kundodyiwa 2001	9/243	5/256		1.35%	1.9[0.64,5.58
Lokugamage 2001	3/20	3/20		0.72%	1[0.23,4.37
Lumbiganon 1999	22/397	13/200		3.57%	0.85[0.44,1.66
Modi 2014	0/25	0/25			Not estimable
Nasr 2009	0/257	0/257			Not estimable
Oboro 2003	0/247	0/249			Not estimable
Owonikoko 2011	4/50	5/50		1%	0.8[0.23,2.81
Parsons 2006	0/225	0/225			Not estimable
Parsons 2007	0/217	1/224 —		0.15%	0.34[0.01,8.4
Penaranda 2002	1/25	3/25		0.33%	0.33[0.04,2.99
Perez-Rumbos 2017	0/195	3/197		0.18%	0.14[0.01,2.78
Sadiq 2011	0/900	0/900			Not estimable
Sultana 2007	5/210	3/190		0.78%	1.51[0.37,6.23
Tewatia 2014	0/50	0/50		0.1070	Not estimable
Vagge 2014	1/100	2/100		0.28%	0.5[0.05,5.43
Vimala 2006	6/50	10/50		1.81%	0.6[0.24,1.53
Walley 2000	0/202	0/196	·	1.0170	Not estimable
Zachariah 2006	1/730	4/617		0.33%	0.21[0.02,1.89
Luchanan 2000	1/130	4/011	·	0.53%	0.21[0.02,1.89
Total (95% CI)	17402	16859	 	100%	1.26[1.11,1.43
Total events: 551 (Misoprosto	· · · · · · · · · · · · · · · · · · ·				
Heterogeneity: Tau ² =0; Chi ² =2					
Test for overall effect: Z=3.58((P=0)				



Analysis 8.3. Comparison 8 Misoprostol vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Acharya 2001	1/30	1/30		0.6%	1[0.07,15.26]
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	0/28	0/37			Not estimable
Alwani 2014	2/100	5/100		1.71%	0.4[0.08,2.01]
Atukunda 2014	7/570	16/570	-	5.75%	0.44[0.18,1.06]
Baskett 2007	0/311	0/311			Not estimable
Bellad 2012	1/321	1/331		0.58%	1.03[0.06,16.42]
Bhatti 2014	1/60	1/60		0.59%	1[0.06,15.62]
Bugalho 2001	2/323	1/339	-	0.78%	2.1[0.19,23.04]
Caliskan 2002	12/396	13/407	- 	7.48%	0.95[0.44,2.05]
Caliskan 2003	14/388	13/384	-	8.11%	1.07[0.51,2.24]
Chaudhuri 2010	0/96	3/94	+	0.51%	0.14[0.01,2.67]
Chaudhuri 2012	5/265	3/265	- +	2.21%	1.67[0.4,6.9]
Cook 1999	5/424	2/129		1.68%	0.76[0.15,3.87]
Diallo 2017	5/154	7/150		3.52%	0.7[0.23,2.14]
Dutta 2016	5/200	4/200		2.64%	1.25[0.34,4.59]
Elbohoty 2016	1/89	1/86		0.59%	0.97[0.06,15.21]
Fazel 2013	0/50	0/50			Not estimable
Gerstenfeld 2001	2/159	0/166	-	0.49%	5.22[0.25,107.86]
Gulmezoglu 2001	72/9221	97/9226	-	48.43%	0.74[0.55,1.01]
Gupta 2006	0/100	0/100			Not estimable
Karkanis 2002	0/110	0/113			Not estimable
Kundodyiwa 2001	2/243	1/256		0.78%	2.11[0.19,23.09]
Lokugamage 2001	1/20	0/20	-	0.45%	3[0.13,69.52]
Lumbiganon 1999	0/397	0/200		•	Not estimable
Modi 2014	0/25	0/25			Not estimable
Nankaly 2016	1/122	5/63		0.99%	0.1[0.01,0.87]
Nasr 2009	8/257	4/257		3.16%	2[0.61,6.56]
Oboro 2003	0/247	0/249			Not estimable
Owonikoko 2011	1/50	0/50		0.44%	3[0.13,71.92]
Parsons 2006	1/222	2/221		0.78%	0.5[0.05,5.45]
Parsons 2007	1/217	5/221		0.98%	0.2[0.02,1.73]
Perez-Rumbos 2017	2/195	3/197		1.41%	0.67[0.11,3.99]
Rajaei 2014	1/200	4/200 —		0.94%	0.25[0.03,2.22]
Sadiq 2011	0/900	0/884		0.0 170	Not estimable
Shady 2017	13/120	0/120		0.56%	27[1.62,449.1]
Singh 2009	0/150	0/75		0.3070	Not estimable
Sultana 2007	4/210	3/190		2.03%	1.21[0.27,5.32]
Tewatia 2014	0/50	0/50		2.03 /0	Not estimable
Vagge 2014	1/100	1/100		0.59%	1[0.06,15.77]
Walley 2000	0/136	1/138		0.44%	0.34[0.01,8.23]
Zachariah 2006	1/730	2/617	•	0.78%	0.42[0.04,4.65]
Total (95% CI)	18086	17381	•	100%	0.81[0.65,1]
Total events: 172 (Misoprostol			•		<u>.</u>
Heterogeneity: Tau ² =0; Chi ² =2					
Test for overall effect: Z=2(P=0					



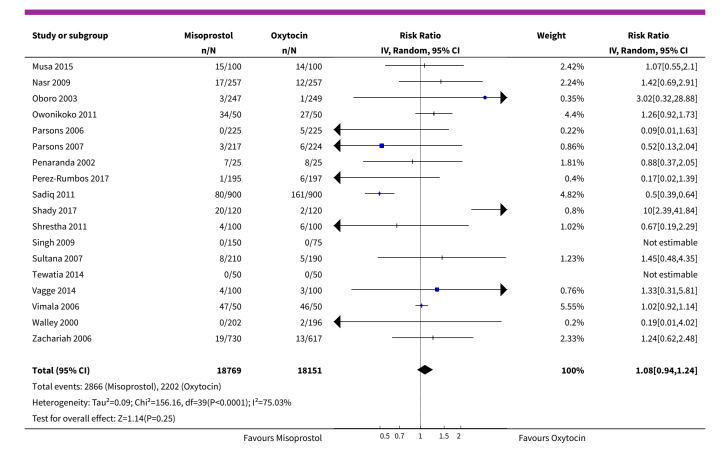
Analysis 8.4. Comparison 8 Misoprostol vs Oxytocin, Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Afolabi 2010	0/100	0/100			Not estimable
Amin 2014	0/100	0/100			Not estimable
Atukunda 2014	11/570	8/570	- -	67.92%	1.38[0.56,3.39]
Chaudhuri 2010	0/96	0/94			Not estimable
Gulmezoglu 2001	4/9224	5/9231		32.08%	0.8[0.22,2.98]
Kundodyiwa 2001	0/243	0/256			Not estimable
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	0/257	0/257			Not estimable
Shrestha 2011	0/100	0/100			Not estimable
Tewatia 2014	0/50	0/50			Not estimable
Total (95% CI)	10840	10858	•	100%	1.16[0.55,2.43]
Total events: 15 (Misoprostol), 13	(Oxytocin)				
Heterogeneity: Tau²=0; Chi²=0.44,	df=1(P=0.51); I ² =0%				
Test for overall effect: Z=0.38(P=0.	7)				
	Fav	ours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

Analysis 8.6. Comparison 8 Misoprostol vs Oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	3/28	2/37	+	0.58%	1.98[0.35,11.08]
Amin 2014	4/100	3/100	•	0.76%	1.33[0.31,5.81]
Asmat 2017	123/839	120/839		4.94%	1.02[0.81,1.29]
Atukunda 2014	89/570	57/570		4.44%	1.56[1.14,2.13]
Bellad 2012	10/321	30/331	— —	2.32%	0.34[0.17,0.69]
Benchimol 2001	52/186	29/196		3.83%	1.89[1.26,2.84]
Bhatti 2014	2/60	5/60	-	0.66%	0.4[0.08,1.98]
Bugalho 2001	10/323	15/339		2.01%	0.7[0.32,1.53]
Caliskan 2002	39/396	33/407		3.61%	1.21[0.78,1.89]
Caliskan 2003	35/388	28/384		3.41%	1.24[0.77,1.99]
Chaudhuri 2010	38/96	51/94		4.46%	0.73[0.54,0.99]
Chaudhuri 2012	16/265	15/265		2.38%	1.07[0.54,2.11]
Cook 1999	63/424	1/129	_	0.45%	19.17[2.68,136.84]
Diallo 2017	10/154	14/150		2.02%	0.7[0.32,1.52]
Dutta 2016	6/200	4/200		1%	1.5[0.43,5.23]
Elbohoty 2016	49/89	34/86		4.38%	1.39[1.01,1.92]
Gavilanes 2016	33/50	26/50	++-	4.31%	1.27[0.91,1.77]
Gerstenfeld 2001	70/154	61/161	+-	4.76%	1.2[0.92,1.56]
Gulmezoglu 2001	1793/9213	1248/9227	+	5.67%	1.44[1.35,1.54]
Gupta 2006	1/100	0/100	+ +	0.18%	3[0.12,72.77]
Kundodyiwa 2001	37/243	34/256		3.68%	1.15[0.74,1.76]
Lokugamage 2001	17/20	17/20		4.77%	1[0.77,1.3]
Lumbiganon 1999	96/397	52/200		4.57%	0.93[0.69,1.24]
Modi 2014	0/25	0/25			Not estimable
Mukta 2013	8/100	6/100		1.38%	1.33[0.48,3.7]
	Fav	ours Misoprostol	0.5 0.7 1 1.5 2	Favours Oxytocin	

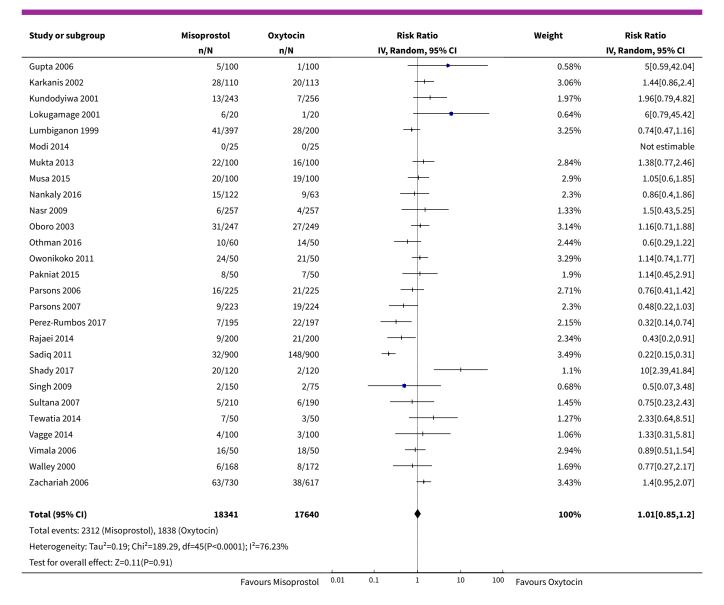




Analysis 8.7. Comparison 8 Misoprostol vs Oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Acharya 2001	2/30	3/30		0.83%	0.67[0.12,3.71]
Afolabi 2010	3/100	4/100		1.06%	0.75[0.17,3.27]
Al-Sawaf 2013	3/28	2/37		0.83%	1.98[0.35,11.08]
Alwani 2014	4/100	9/100		1.5%	0.44[0.14,1.4]
Atukunda 2014	47/570	31/570	 + -	3.28%	1.52[0.98,2.35]
Baskett 2007	159/311	126/311	+	3.97%	1.26[1.06,1.5]
Bellad 2012	1/321	8/331		0.61%	0.13[0.02,1.02]
Bhatti 2014	1/60	3/60		0.54%	0.33[0.04,3.11]
Bugalho 2001	7/323	7/339		1.69%	1.05[0.37,2.96]
Caliskan 2002	33/396	26/407	+-	3.11%	1.3[0.8,2.14]
Caliskan 2003	23/388	26/384	 -	2.96%	0.88[0.51,1.51]
Chaudhuri 2010	11/96	14/94		2.39%	0.77[0.37,1.61]
Chaudhuri 2012	20/265	23/265	 -	2.86%	0.87[0.49,1.54]
Cook 1999	95/424	6/129		2.22%	4.82[2.16,10.73]
Diallo 2017	7/154	6/150	 +	1.63%	1.14[0.39,3.3]
Eftekhari 2009	7/50	16/50		2.23%	0.44[0.2,0.97]
Elbohoty 2016	20/89	11/86	 	2.57%	1.76[0.9,3.45]
Gavilanes 2016	10/50	12/50		2.37%	0.83[0.4,1.75]
Gerstenfeld 2001	36/159	18/166		3.02%	2.09[1.24,3.52]
Gulmezoglu 2001	1398/9225	1002/9228	+	4.1%	1.4[1.29,1.51]
	Fav	ours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

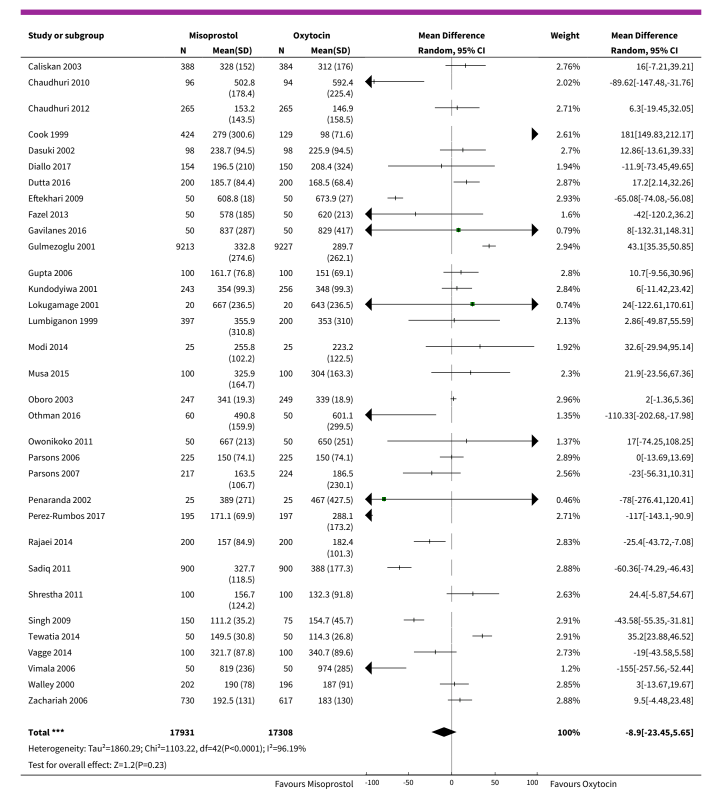




Analysis 8.8. Comparison 8 Misoprostol vs Oxytocin, Outcome 8 Blood loss.

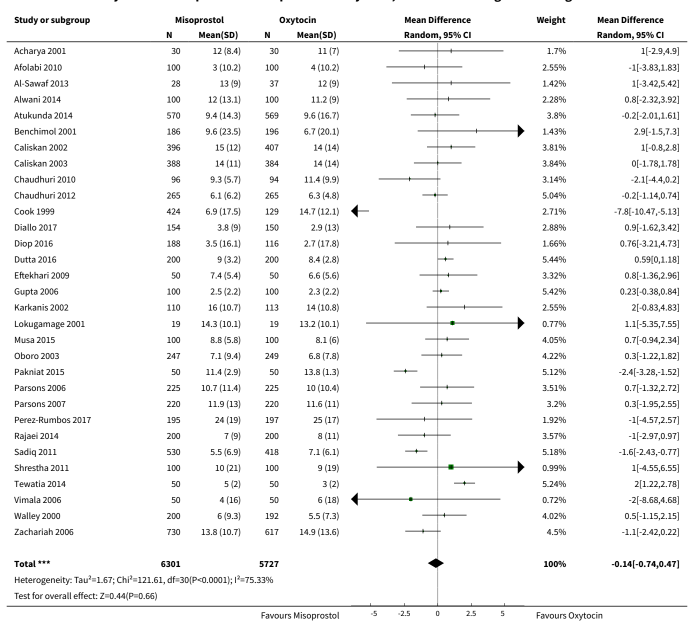
Study or subgroup	Mis	oprostol	0	xytocin		Mean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI			Random, 95% CI
Acharya 2001	30	545 (192.8)	30	533 (296.2)	\leftarrow	-	—	0.91%	12[-114.47,138.47]
Afolabi 2010	100	153.2 (58)	100	155.6 (58)				2.86%	-2.4[-18.47,13.67]
Al-Sawaf 2013	28	348 (112)	37	314.7 (94.6)		+		2.16%	33.3[-18.18,84.78]
Amin 2014	100	300 (262.8)	100	250 (262.8)		+		1.7%	50[-22.83,122.83]
Asmat 2017	839	322 (199.9)	839	337 (211.4)				2.81%	-15[-34.69,4.69]
Atukunda 2014	570	341.5 (206.2)	570	304.2 (190.8)			-	2.76%	37.3[14.24,60.36]
Bellad 2012	321	192 (124)	331	366 (135.9)	◀			2.81%	-174[-193.96,-154.04]
Benchimol 2001	186	374 (238.4)	196	278 (254)		_		2.21%	96[46.63,145.37]
Bhatti 2014	60	200 (125)	60	360 (136)	◀			2.27%	-160[-206.74,-113.26]
Bugalho 2001	323	155 (122)	339	157.3 (138.7)		-		2.81%	-2.3[-22.17,17.57]
			Favou	rs Misoprostol	-100	-50 0 50	100	Favours Ox	ytocin







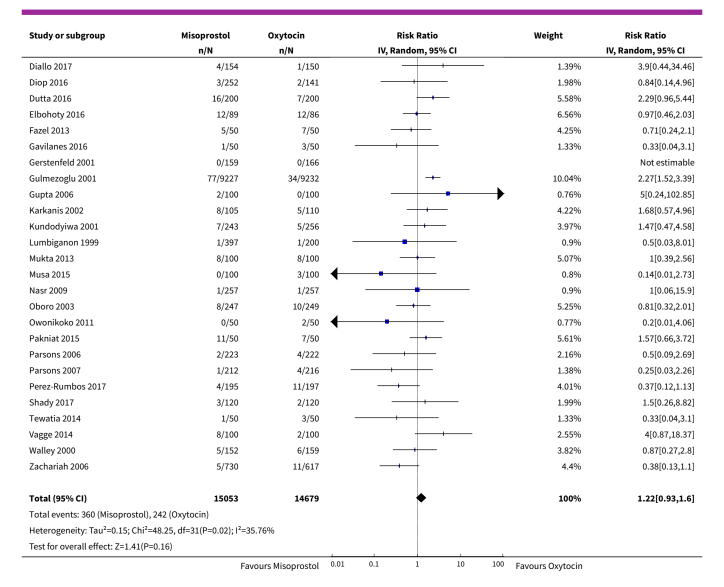
Analysis 8.9. Comparison 8 Misoprostol vs Oxytocin, Outcome 9 Change in haemoglobin.



Analysis 8.11. Comparison 8 Misoprostol vs Oxytocin, Outcome 11 Nausea.

Study or subgroup	Misoprostol	Oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
Adanikin 2013	2/25	1/25		_		_	1.23%	2[0.19,20.67]
Afolabi 2010	4/100	0/100			-		0.82%	9[0.49,165]
Alwani 2014	1/100	2/100			+		1.18%	0.5[0.05,5.43]
Atukunda 2014	138/570	86/570			+		11.67%	1.6[1.26,2.05]
Bellad 2012	12/321	0/331				-	0.87%	25.78[1.53,433.55]
Bhatti 2014	4/60	0/60			-		0.82%	9[0.5,163.58]
Chaudhuri 2012	6/265	2/265					2.38%	3[0.61,14.73]
	Fav	ours Misoprostol	0.01	0.1	1 10	100	Favours Oxytocin	

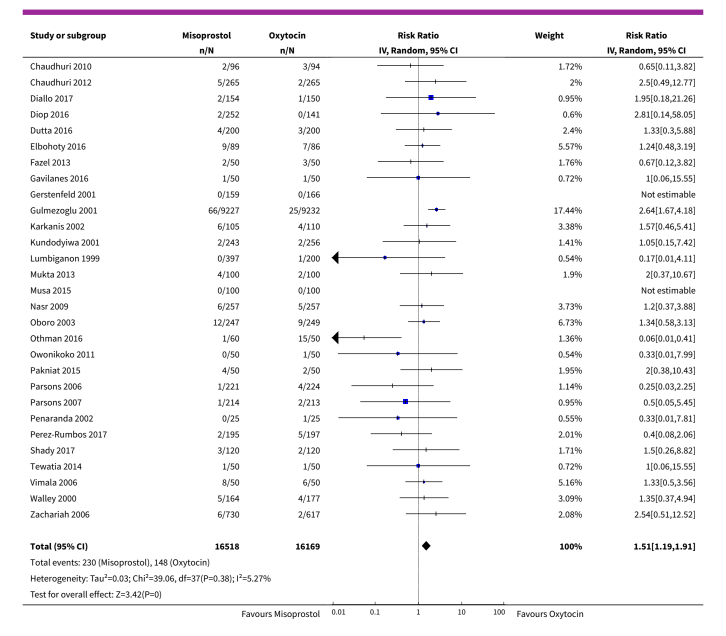




Analysis 8.12. Comparison 8 Misoprostol vs Oxytocin, Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Acharya 2001	2/30	3/30		1.81%	0.67[0.12,3.71]
Adanikin 2013	2/25	2/25		1.52%	1[0.15,6.55]
Afolabi 2010	0/100	0/100			Not estimable
Alwani 2014	1/100	1/100		0.72%	1[0.06,15.77]
Amin 2014	12/100	2/100		2.44%	6[1.38,26.12]
Atukunda 2014	35/569	19/570		13.68%	1.85[1.07,3.19]
Bellad 2012	5/321	0/331	+ +	0.65%	11.34[0.63,204.28]
Benchimol 2001	7/186	1/196	 	1.24%	7.38[0.92,59.38]
Bhatti 2014	3/60	1/60	- +	1.09%	3[0.32,28.03]
Bugalho 2001	2/323	1/337		0.95%	2.09[0.19,22.9]
Caliskan 2002	2/396	2/407		1.41%	1.03[0.15,7.26]
Caliskan 2003	4/388	3/384		2.38%	1.32[0.3,5.86]
	Fav	ours Misoprostol	0.01 0.1 1 10 10	⁰ Favours Oxytocin	

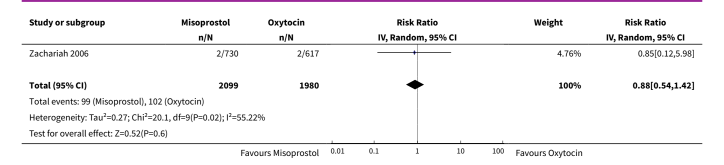




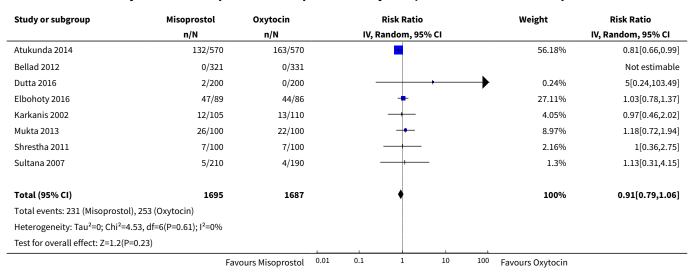
Analysis 8.13. Comparison 8 Misoprostol vs Oxytocin, Outcome 13 Headache.

Study or subgroup	Misoprostol	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Atukunda 2014	10/570	11/570				13.25%	0.91[0.39,2.12]
Dutta 2016	5/200	3/200				7.61%	1.67[0.4,6.88]
Elbohoty 2016	24/89	24/86		-		18.39%	0.97[0.6,1.56]
Gavilanes 2016	0/50	1/50		+		2.08%	0.33[0.01,7.99]
Karkanis 2002	9/105	4/110		+-		9.87%	2.36[0.75,7.42]
Othman 2016	3/60	20/50				9.81%	0.13[0.04,0.4]
Owonikoko 2011	1/50	3/50	-			3.85%	0.33[0.04,3.1]
Perez-Rumbos 2017	39/195	26/197		-		18.77%	1.52[0.96,2.39]
Vimala 2006	6/50	8/50				11.6%	0.75[0.28,2]
	Fav	ours Misoprostol	0.01	0.1 1 10	100	Favours Oxytocin	

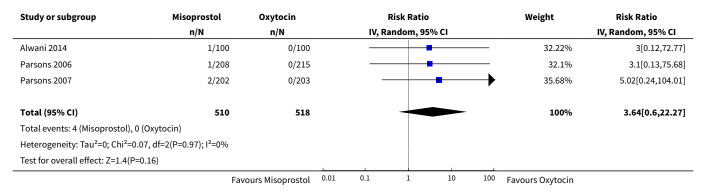




Analysis 8.14. Comparison 8 Misoprostol vs Oxytocin, Outcome 14 Abdominal pain.



Analysis 8.15. Comparison 8 Misoprostol vs Oxytocin, Outcome 15 Hypertension.

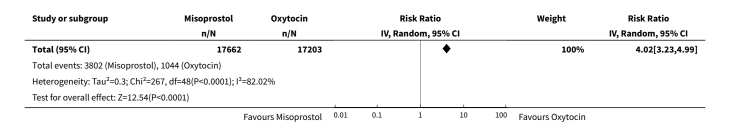




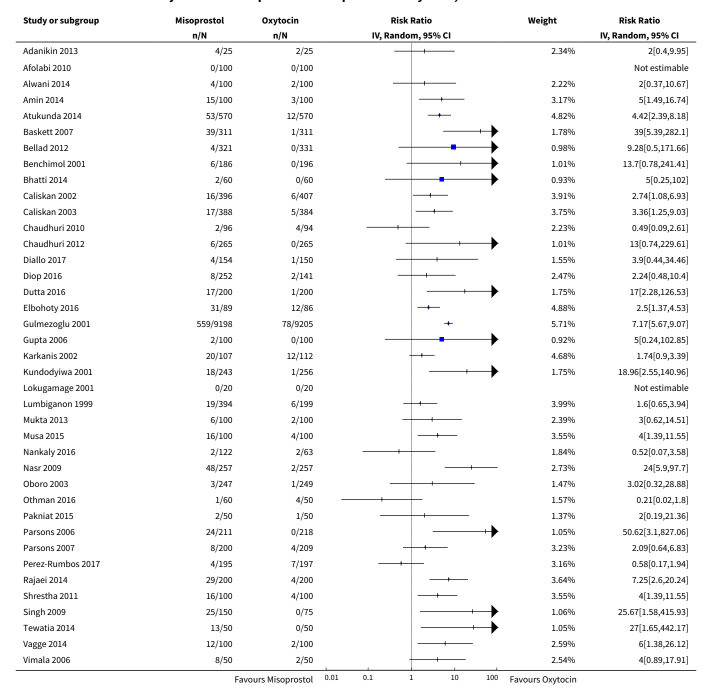
Analysis 8.16. Comparison 8 Misoprostol vs Oxytocin, Outcome 16 Shivering.

	Misoprostol n/N	Oxytocin n/N	Risk Ratio IV, Random, 95% CI	Weight	Risk Ratio IV, Random, 95% CI	
Acharya 2001	2/30	2/30		1%	1[0.15,6.64]	
Adanikin 2013	1/25	2/25		0.72%	0.5[0.05,5.17]	
Afolabi 2010	4/100	2/100		1.2%	2[0.37,10.67]	
Alwani 2014	7/100	1/100	-	0.86%	7[0.88,55.86]	
Amin 2014	25/100	4/100		2.16%	6.25[2.26,17.31]	
Atukunda 2014	321/570	168/570	+	4.04%	1.91[1.65,2.21]	
Baskett 2007	21/311	0/311		0.53%	43[2.62,706.74]	
Bellad 2012	173/321	14/331	ļ 	3.33%	12.74[7.56,21.49]	
Benchimol 2001	5/186	0/196	-	0.5%	11.59[0.65,208.12]	
Bhatti 2014	32/60	3/60	<u> </u>	1.95%	10.67[3.45,32.96]	
Bugalho 2001	123/323	51/337	+	3.84%	2.52[1.89,3.36]	
Caliskan 2002	47/396	16/407		3.26%	3.02[1.74,5.23]	
Caliskan 2003	44/388	19/384		3.34%	2.29[1.36,3.85]	
Chaudhuri 2010	8/96	1/94		0.88%	7.83[1,61.42]	
Chaudhuri 2012	51/265	2/265		1.52%	25.5[6.27,103.67]	
Dasuki 2002	13/98	2/98		1.44%	6.5[1.51,28.05]	
Diallo 2017	11/154	3/150		1.73%	3.57[1.02,12.55]	
Diop 2016	154/191	7/140		2.82%	16.13[7.81,33.31]	
Diop 2016 Dutta 2016	41/200			2.18%		
		4/200	<u> </u>		10.25[3.74,28.08]	
Elbohoty 2016	28/89	16/86	_	3.29%	1.69[0.99,2.9]	
Fazel 2013	8/50	1/50		0.89%	8[1.04,61.62]	
Gavilanes 2016	33/50	1/50		0.95%	33[4.69,232.05]	
Gerstenfeld 2001	7/159	7/166	 .	2.15%	1.04[0.37,2.91]	
Gulmezoglu 2001	1620/9227	466/9232		4.09%	3.48[3.15,3.84]	
Gupta 2006	16/100	13/100	T.	2.94%	1.23[0.63,2.42]	
Karkanis 2002	26/105	15/110		3.19%	1.82[1.02,3.23]	
Kundodyiwa 2001	106/243	78/256	+	3.93%	1.43[1.13,1.81]	
Lokugamage 2001	13/20	8/20	 	3.07%	1.63[0.87,3.04]	
Lumbiganon 1999	94/397	25/200	—	3.6%	1.89[1.26,2.85]	
Mukta 2013	50/100	8/100		2.9%	6.25[3.13,12.5]	
Musa 2015	42/100	15/100		3.33%	2.8[1.66,4.71]	
Nankaly 2016	2/122	0/63	-	0.46%	2.6[0.13,53.38]	
Nasr 2009	80/257	0/257		0.53%	161[10.04,2582.33]	
Oboro 2003	141/247	35/249	+	3.77%	4.06[2.93,5.62]	
Othman 2016	36/60	9/50		3.07%	3.33[1.78,6.24]	
Owonikoko 2011	27/50	1/50		0.95%	27[3.81,191.12]	
Pakniat 2015	3/50	0/50	-	0.48%	7[0.37,132.1]	
Parsons 2006	180/223	8/223		2.93%	22.5[11.36,44.56]	
Parsons 2007	16/213	2/213		1.44%	8[1.86,34.37]	
Penaranda 2002	1/25	0/25	+	0.43%	3[0.13,70.3]	
Perez-Rumbos 2017	11/195	2/197		1.4%	5.56[1.25,24.74]	
Rajaei 2014	2/200	2/200		0.95%	1[0.14,7.03]	
Singh 2009	19/150	0/75	+	0.53%	19.63[1.2,320.72]	
Sultana 2007	13/210	2/190		1.42%	5.88[1.34,25.72]	
Tewatia 2014	10/50	0/50		0.52%	21[1.26,348.93]	
Vagge 2014	15/100	3/100		1.81%	5[1.49,16.74]	
Vimala 2006	13/50	2/50		1.47%	6.5[1.55,27.33]	
	39/176	10/176		2.98%	3.9[2.01,7.57]	
Walley 2000						

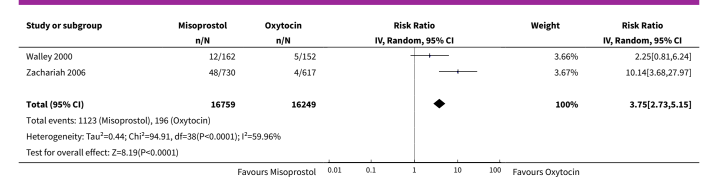




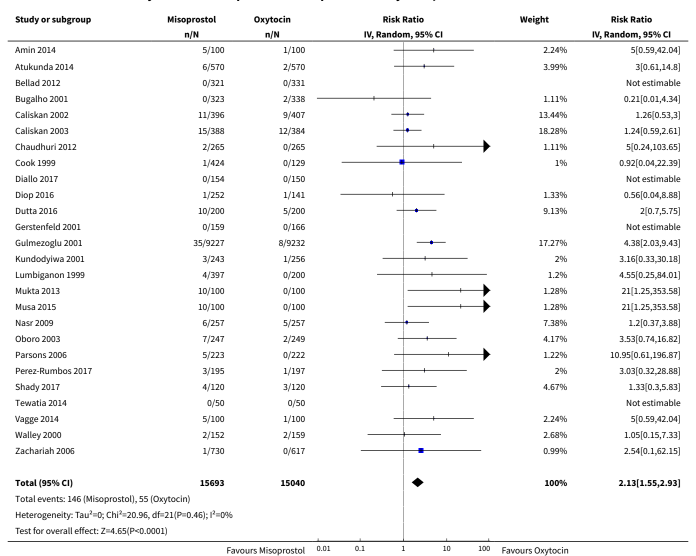
Analysis 8.17. Comparison 8 Misoprostol vs Oxytocin, Outcome 17 Fever.







Analysis 8.18. Comparison 8 Misoprostol vs Oxytocin, Outcome 18 Diarrhoea.





Analysis 8.20. Comparison 8 Misoprostol vs Oxytocin, Outcome 20 Maternal satisfaction.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Risk Ratio
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI
8.20.1 Satisfied or very satisfi	ed with drug			
Diop 2016	854/854	480/484	+	1.01[1,1.02]
8.20.2 Complaints about or pr	oblems with drug			
Diop 2016	18/858	28/481		0.36[0.2,0.64]
8.20.3 Would take drug again	after subsequent deliveries			
Diop 2016	833/834	457/461	+	1.01[1,1.02]
8.20.4 Would recommend dru	g to a friend			
Diop 2016	812/812	451/454	+	1.01[1,1.02]
		Favours Misoprostol	1	Favours Oxytocin

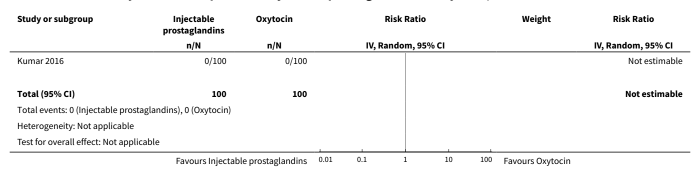
Comparison 9. Injectable prostaglandins vs Oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	100	Risk Ratio (IV, Random, 95% CI)	1.43 [0.20, 10.31]
3 Blood transfusion	2	250	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.65]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	4	500	Risk Ratio (IV, Random, 95% CI)	0.84 [0.26, 2.71]
7 Additional uterotonics	4	500	Risk Ratio (IV, Random, 95% CI)	0.29 [0.09, 0.94]
8 Blood loss	4	500	Mean Difference (IV, Random, 95% CI)	-15.83 [-152.28, 120.62]
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	3	450	Risk Ratio (IV, Random, 95% CI)	1.17 [0.40, 3.41]
12 Vomiting	2	400	Risk Ratio (IV, Random, 95% CI)	2.48 [0.57, 10.73]
13 Headache	1	200	Risk Ratio (IV, Random, 95% CI)	0.20 [0.01, 4.11]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	400	Risk Ratio (IV, Random, 95% CI)	0.91 [0.11, 7.73]
17 Fever	1	200	Risk Ratio (IV, Random, 95% CI)	2.0 [0.18, 21.71]
18 Diarrhoea	2	400	Risk Ratio (IV, Random, 95% CI)	10.38 [1.96, 54.98]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 9.1. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 1 Death.



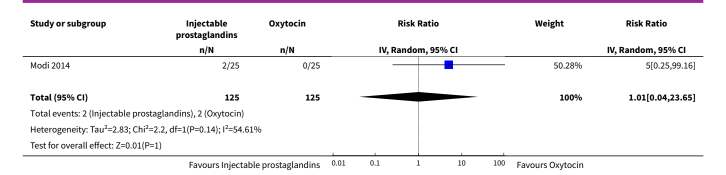
Analysis 9.2. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Injectable prostaglandins	Oxytocin	Risk	Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI			IV, Random, 95% CI
Modi 2014	2/25	0/25		•		39.36%	5[0.25,99.16]
Poeschmann 1991	1/22	2/28	-			60.64%	0.64[0.06,6.57]
Total (95% CI)	47	53				100%	1.43[0.2,10.31]
Total events: 3 (Injectable pr	rostaglandins), 2 (Oxytocin)						
Heterogeneity: Tau ² =0.25; Ch	ni²=1.14, df=1(P=0.29); I²=11.9	4%					
Test for overall effect: Z=0.36	6(P=0.72)	1					
	Favours Injectab	e prostaglandins 0	0.01 0.1	1 10	100	Favours Oxytocin	

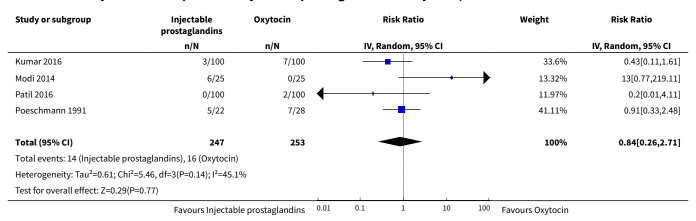
Analysis 9.3. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Oxytocin	rtocin Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ındom, 95	5% CI			IV, Random, 95% CI
Kumar 2016	0/100	2/100	•	-		_		49.72%	0.2[0.01,4.11]
	Favours Injectable	prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	





Analysis 9.6. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 6 PPH >= 500 mL.



Analysis 9.7. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 7 Additional uterotonics.

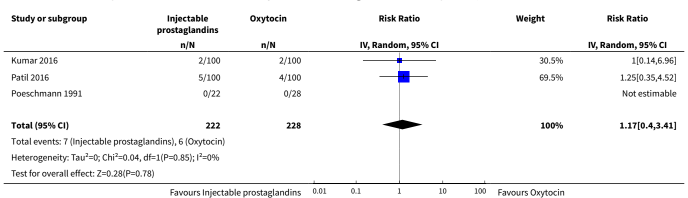
Study or subgroup	Injectable prostaglandins	Oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI
	n/N	n/N	IV, Random, 95% CI		
Kumar 2016	4/100	21/100		43.68%	0.19[0.07,0.53]
Modi 2014	2/25	0/25	+	12.64%	5[0.25,99.16]
Patil 2016	4/100	21/100		43.68%	0.19[0.07,0.53]
Poeschmann 1991	0/22	0/28			Not estimable
Total (95% CI)	247	253	•	100%	0.29[0.09,0.94]
Total events: 10 (Injectable p	orostaglandins), 42 (Oxytocin)				
Heterogeneity: Tau ² =0.55; Ch	hi²=4.34, df=2(P=0.11); I²=53.8	9%			
Test for overall effect: Z=2.06	6(P=0.04)				
	Favours Injectab	le prostaglandins 0.0	1 0.1 1 10 1	00 Favours Oxytocin	



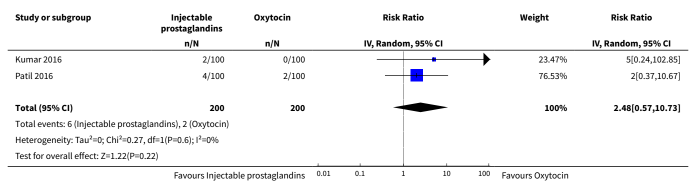
Analysis 9.8. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 8 Blood loss.

Study or subgroup		Injectable prostaglandins		Oxytocin		Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
Kumar 2016	100	170.2 (197.4)	100	281.1 (197.4)	←				26.67%	-110.85[-165.57,-56.13]
Modi 2014	25	435 (147.6)	25	223.2 (122.5)				•	25.69%	211.8[136.61,286.99]
Patil 2016	100	170.2 (50.2)	100	281.1 (84.8)	◀				27.71%	-110.85[-130.17,-91.53]
Poeschmann 1991	22	324 (302)	28	374 (279)	←	-			19.93%	-50[-213.11,113.11]
Total ***	247		253						100%	-15.83[-152.28,120.62]
Heterogeneity: Tau ² =17392.8;	Chi ² =67.08, df=	3(P<0.0001); I ² =9	95.53%							
Test for overall effect: Z=0.23(P=0.82)									
		Favours Inj	ectable p	rostaglandins	-100	-50	0 50	100	Favours Ox	kytocin

Analysis 9.11. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 11 Nausea.



Analysis 9.12. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 12 Vomiting.

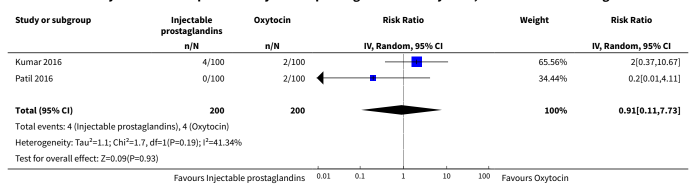




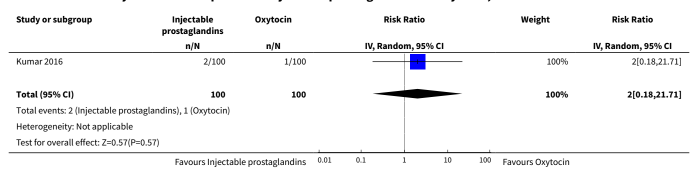
Analysis 9.13. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 13 Headache.

Study or subgroup	Injectable prostaglandins	• • • • • • • • • • • • • • • • • • • •			isk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ndom, 95%	6 CI			IV, Random, 95% CI	
Patil 2016	0/100	2/100	←	1		-		100%	0.2[0.01,4.11]	
Total (95% CI)	100	100	_			-		100%	0.2[0.01,4.11]	
Total events: 0 (Injectable pr	ostaglandins), 2 (Oxytocin)									
Heterogeneity: Tau ² =0; Chi ² =	:0, df=0(P<0.0001); I ² =100%									
Test for overall effect: Z=1.04	(P=0.3)					1				
	Favours Injectable	prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin		

Analysis 9.16. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 16 Shivering.



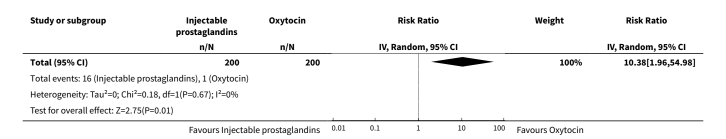
Analysis 9.17. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 17 Fever.



Analysis 9.18. Comparison 9 Injectable prostaglandins vs Oxytocin, Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	•			Risk Ratio	•		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Kumar 2016	8/100	1/100				-		65.5%	8[1.02,62.79]
Patil 2016	8/100	0/100				-		34.5%	17[0.99,290.62]
						1			
	Favours Injectabl	e prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	





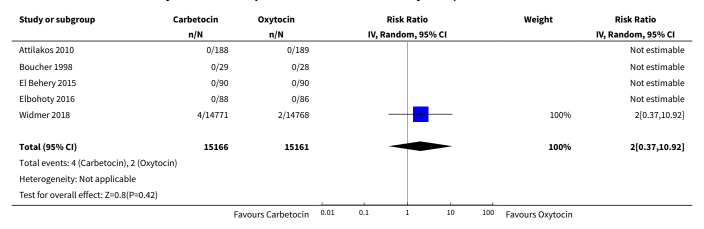
Comparison 10. Carbetocin vs Oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	5	30327	Risk Ratio (IV, Random, 95% CI)	2.00 [0.37, 10.92]
2 PPH >= 1000 mL	10	30633	Risk Ratio (IV, Random, 95% CI)	0.73 [0.45, 1.19]
3 Blood transfusion	16	32115	Risk Ratio (IV, Random, 95% CI)	0.68 [0.38, 1.22]
4 Severe maternal morbidity: intensive care admissions	2	29847	Risk Ratio (IV, Random, 95% CI)	1.16 [0.67, 2.02]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	30633	Risk Ratio (IV, Random, 95% CI)	0.75 [0.58, 0.98]
7 Additional uterotonics	22	32699	Risk Ratio (IV, Random, 95% CI)	0.48 [0.34, 0.68]
8 Blood loss	16	2115	Mean Difference (IV, Random, 95% CI)	-92.73 [-154.97, -30.49]
9 Change in haemoglo- bin	12	2096	Mean Difference (IV, Random, 95% CI)	-1.66 [-3.81, 0.50]
10 Breastfeeding	2	190	Risk Ratio (IV, Random, 95% CI)	0.94 [0.86, 1.03]
11 Nausea	12	2288	Risk Ratio (IV, Random, 95% CI)	1.11 [0.78, 1.56]
12 Vomiting	13	31833	Risk Ratio (IV, Random, 95% CI)	0.90 [0.53, 1.50]
13 Headache	15	2620	Risk Ratio (IV, Random, 95% CI)	0.84 [0.63, 1.12]
14 Abdominal pain	10	31293	Risk Ratio (IV, Random, 95% CI)	1.18 [0.97, 1.44]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	1998	Risk Ratio (IV, Random, 95% CI)	0.78 [0.49, 1.23]
17 Fever	4	466	Risk Ratio (IV, Random, 95% CI)	1.58 [0.27, 9.35]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 10.1. Comparison 10 Carbetocin vs Oxytocin, Outcome 1 Death.



Analysis 10.2. Comparison 10 Carbetocin vs Oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Attilakos 2010	9/188	9/189		17.68%	1.01[0.41,2.48]
Boucher 1998	0/29	0/28			Not estimable
El Behery 2015	2/90	12/90		8.81%	0.17[0.04,0.72]
Elbohoty 2016	3/88	5/86		9.51%	0.59[0.14,2.38]
Fenix 2012	0/30	0/30			Not estimable
Kabir 2015	0/47	4/47	←	2.65%	0.11[0.01,2.01]
Maged 2016	0/100	1/100	+	2.21%	0.33[0.01,8.09]
Rosseland 2013	0/25	0/26			Not estimable
Whigham 2016	7/59	8/53		16.7%	0.79[0.31,2.02]
Widmer 2018	222/14651	212/14677	•	42.44%	1.05[0.87,1.26]
Total (95% CI)	15307	15326	•	100%	0.73[0.45,1.19]
Total events: 243 (Carbetocin),	251 (Oxytocin)				
Heterogeneity: Tau ² =0.14; Chi ²	=9.44, df=6(P=0.15); I ² =36.4	1%			
Test for overall effect: Z=1.25(P	=0.21)				
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	



Analysis 10.3. Comparison 10 Carbetocin vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Attilakos 2010	4/188	5/189		13.69%	0.8[0.22,2.95]
Boucher 1998	0/29	0/28			Not estimable
Carrillo-Gaucin 2016	1/60	2/57		5.31%	0.48[0.04,5.1]
Dansereau 1999	2/317	2/318		7.4%	1[0.14,7.08]
El Behery 2015	0/90	14/90	—	3.95%	0.03[0,0.57]
Elbohoty 2016	0/88	1/86		3.12%	0.33[0.01,7.89]
Fahmy 2015	0/50	0/50			Not estimable
Fahmy 2016	1/30	4/30		6.39%	0.25[0.03,2.11]
Fenix 2012	0/30	0/30			Not estimable
Kabir 2015	0/47	3/47	—	3.63%	0.14[0.01,2.69]
Maged 2016	1/100	2/100		5.27%	0.5[0.05,5.43]
Reyes 2011	1/45	0/90		3.13%	5.93[0.25,142.84]
Reyes, Gonzalez 2011	0/26	3/29	—	3.67%	0.16[0.01,2.93]
Taheripanah 2018	0/110	0/110			Not estimable
Whigham 2016	1/59	1/53		4.1%	0.9[0.06,14.01]
Widmer 2018	229/14771	198/14768	•	40.33%	1.16[0.96,1.4]
Total (95% CI)	16040	16075	•	100%	0.68[0.38,1.22]
Total events: 240 (Carbetocin), 23	5 (Oxytocin)				
Heterogeneity: Tau ² =0.21; Chi ² =14	I.31, df=11(P=0.22); I ² =23	3.13%			
Test for overall effect: Z=1.31(P=0.	19)				
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	

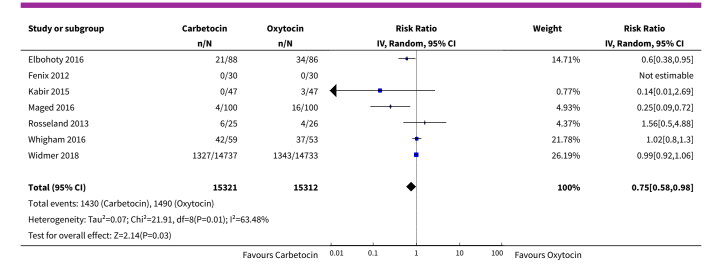
Analysis 10.4. Comparison 10 Carbetocin vs Oxytocin, Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Carbetocin	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Attilakos 2010	1/188	0/189						2.99%	3.02[0.12,73.56]
Widmer 2018	26/14737	23/14733			-			97.01%	1.13[0.65,1.98]
Total (95% CI)	14925	14922			•			100%	1.16[0.67,2.02]
Total events: 27 (Carbetocin),	23 (Oxytocin)								
Heterogeneity: Tau ² =0; Chi ² =0	0.35, df=1(P=0.55); I ² =0%								
Test for overall effect: Z=0.54((P=0.59)						1		
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

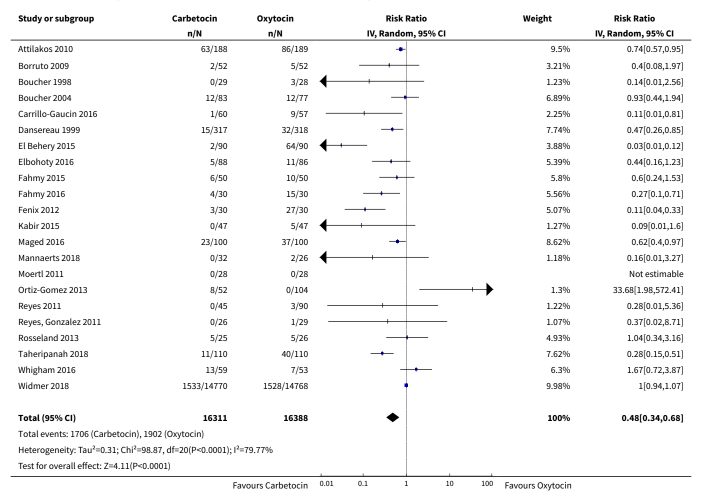
Analysis 10.6. Comparison 10 Carbetocin vs Oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI			IV, Random, 95% CI
Borruto 2009	12/52	23/52	-+-		11.45%	0.52[0.29,0.93]
Boucher 1998	0/29	0/28				Not estimable
Boucher 2004	10/64	11/67	-		7.81%	0.95[0.43,2.09]
El Behery 2015	8/90	19/90			7.98%	0.42[0.19,0.91]
	Fa	vours Carbetocin 0.0	01 0.1 1 1	.0 100	Favours Oxytocin	





Analysis 10.7. Comparison 10 Carbetocin vs Oxytocin, Outcome 7 Additional uterotonics.





Analysis 10.8. Comparison 10 Carbetocin vs Oxytocin, Outcome 8 Blood loss.

Study or subgroup	Ca	rbetocin	0:	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Ahmed 2014	40	323 (542.2)	40	673 (542.2)		3.57%	-350[-587.61,-112.39]
Attilakos 2010	188	500 (222.4)	189	500 (148.3)		7.25%	0[-38.18,38.18]
Borruto 2009	52	370.1 (226)	52	400.5 (226)	•	6.51%	-30.4[-117.27,56.47]
Boucher 1998	29	159 (92)	28	188 (115)		7.06%	-29[-83.18,25.18]
Boucher 2004	64	413.3 (197.5)	67	410.4 (194.1)	+	6.86%	2.9[-64.19,69.99]
Carrillo-Gaucin 2016	60	482.5 (126.5)	57	464 (180.7)		7.02%	18.46[-38.33,75.25]
El Behery 2015	90	689 (580)	90	1027 (659)		4.57%	-338[-519.37,-156.63]
Fahmy 2015	50	398.7 (60.4)	50	449 (69)		7.36%	-50.3[-75.7,-24.9]
Fahmy 2016	30	437 (45)	30	721 (50)		7.37%	-284[-308.07,-259.93]
Fenix 2012	30	296 (183.3)	30	493.3 (183.3)		6.4%	-197.3[-290.04,-104.56]
Kabir 2015	47	325 (306)	47	389 (366)	+	5.49%	-64[-200.39,72.39]
Maged 2016	100	337.7 (118.8)	100	378 (143.2)		7.27%	-40.27[-76.73,-3.81]
Mohamed 2015	86	366.4 (165)	86	434.7 (171.7)		7.11%	-68.3[-118.63,-17.97]
Rosseland 2013	25	579 (623)	26	841 (556)		2.46%	-262[-586.52,62.52]
Taheripanah 2018	110	430.7 (118)	110	552.6 (156)	-	7.27%	-121.92[-158.47,-85.37]
Whigham 2016	59	586 (245.1)	53	561 (245.1)	+	6.43%	25[-65.92,115.92]
Total ***	1060		1055			100%	-92.73[-154.97,-30.49]
Heterogeneity: Tau ² =13525.63	; Chi²=336.03, d	df=15(P<0.0001)	l ² =95.54 ⁹	%			
Test for overall effect: Z=2.92(F	P=0)						

Analysis 10.9. Comparison 10 Carbetocin vs Oxytocin, Outcome 9 Change in haemoglobin.

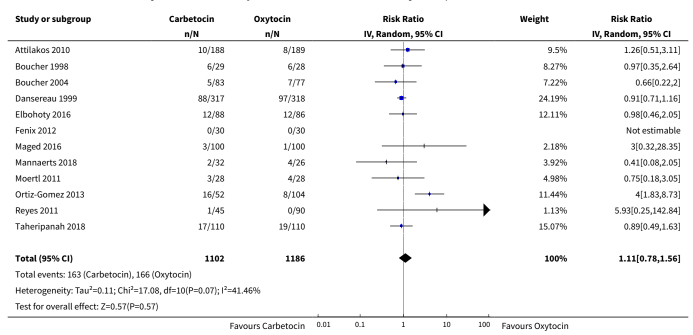
Study or subgroup	Car	rbetocin	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Attilakos 2010	183	16 (2.2)	185	16 (2.2)	•	10.49%	0[-0.45,0.45]
Boucher 2004	82	12.8 (10.8)	73	15.9 (11.6)	+	8.2%	-3.1[-6.64,0.44]
Dansereau 1999	317	7.5 (10)	318	8.3 (10)	•	9.99%	-0.8[-2.36,0.76]
El Behery 2015	90	17.4 (8.7)	90	9.4 (6.7)	•	9.44%	8[5.73,10.27]
Fenix 2012	30	6 (4.6)	30	11 (4.6)	+	9.37%	-5[-7.35,-2.65]
Maged 2016	100	5.5 (3.5)	100	9.6 (6.2)	+	10.09%	-4.1[-5.5,-2.7]
Mannaerts 2018	32	14.5 (11)	26	15 (9)	+	6.58%	-0.5[-5.65,4.65]
Moertl 2011	28	11 (9.9)	28	11.4 (7.6)	+	7.1%	-0.4[-5.02,4.22]
Reyes, Gonzalez 2011	26	12.4 (8.7)	29	14.1 (11.2)	+	6.46%	-1.7[-6.97,3.57]
Rosseland 2013	25	5 (8.2)	26	8.2 (6.7)	+	7.61%	-3.2[-7.32,0.92]
Taheripanah 2018	110	11 (10.1)	110	20.5 (12)	*	8.82%	-9.5[-12.43,-6.57]
Whigham 2016	30	22 (11.5)	28	21 (11.5)	+	5.86%	1[-4.94,6.94]
Total ***	1053		1043		•	100%	-1.66[-3.81,0.5]
Heterogeneity: Tau ² =11.48; Chi	² =138.97, df=1	1(P<0.0001); I ² =9	92.08%				
Test for overall effect: Z=1.51(P:	=0.13)						



Analysis 10.10. Comparison 10 Carbetocin vs Oxytocin, Outcome 10 Breastfeeding.

Study or subgroup	Carbetocin	Oxytocin			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI			% CI			IV, Random, 95% CI	
Reyes 2011	41/45	88/90			+			83.79%	0.93[0.85,1.03]	
Reyes, Gonzalez 2011	22/26	25/29			+			16.21%	0.98[0.79,1.22]	
Total (95% CI)	71	119			•			100%	0.94[0.86,1.03]	
Total events: 63 (Carbetocin), 1	13 (Oxytocin)									
Heterogeneity: Tau ² =0; Chi ² =0.1	L8, df=1(P=0.67); I ² =0%									
Test for overall effect: Z=1.38(P=	=0.17)									
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin		

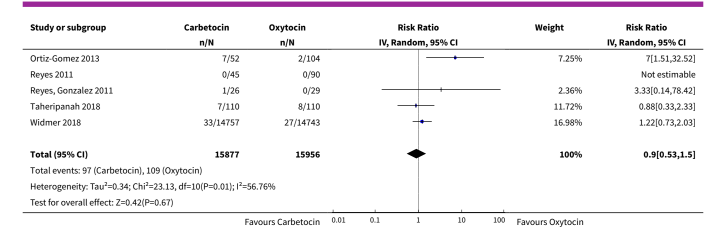
Analysis 10.11. Comparison 10 Carbetocin vs Oxytocin, Outcome 11 Nausea.



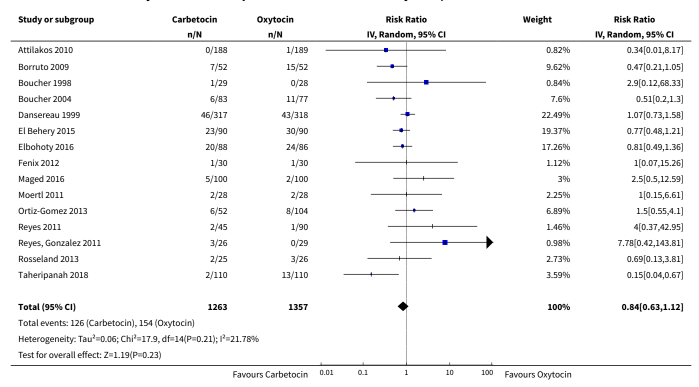
Analysis 10.12. Comparison 10 Carbetocin vs Oxytocin, Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Oxytocin		Ris	sk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 959	% CI			IV, Random, 95% CI
Attilakos 2010	5/188	5/189			+			9.48%	1.01[0.3,3.42]
Borruto 2009	4/52	20/52						11.49%	0.2[0.07,0.54]
Boucher 1998	2/29	5/28		-				7.13%	0.39[0.08,1.83]
Boucher 2004	0/83	6/77	\leftarrow	-	+			2.8%	0.07[0,1.25]
Dansereau 1999	30/317	29/318			+			17.22%	1.04[0.64,1.69]
Elbohoty 2016	6/88	7/86		_	+			11.03%	0.84[0.29,2.39]
Fenix 2012	0/30	0/30							Not estimable
Maged 2016	2/100	0/100				+ -	—	2.54%	5[0.24,102.85]
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	





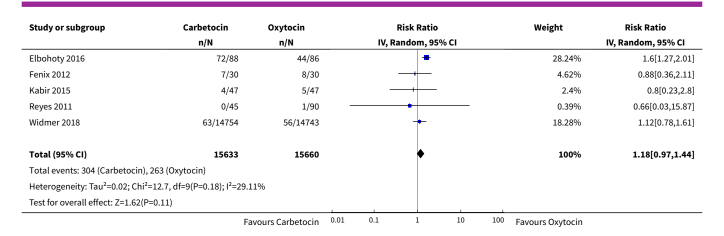
Analysis 10.13. Comparison 10 Carbetocin vs Oxytocin, Outcome 13 Headache.



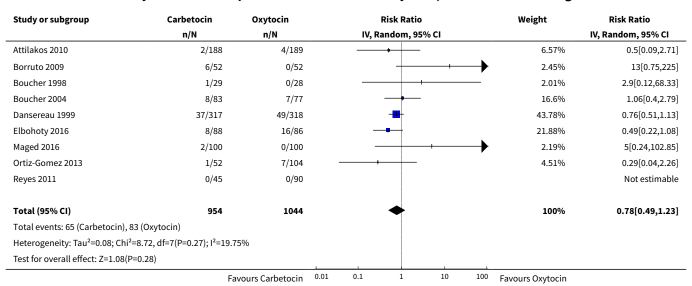
Analysis 10.14. Comparison 10 Carbetocin vs Oxytocin, Outcome 14 Abdominal pain.

Study or subgroup	Carbetocin	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	% CI			IV, Random, 95% CI
Attilakos 2010	1/188	1/189			+			0.51%	1.01[0.06,15.95]
Borruto 2009	21/52	20/52			+			12.59%	1.05[0.65,1.69]
Boucher 1998	0/29	1/28			•			0.39%	0.32[0.01,7.59]
Boucher 2004	5/83	0/77			-	•	\rightarrow	0.47%	10.21[0.57,181.71]
Dansereau 1999	131/317	127/318			+	1	1	32.1%	1.03[0.86,1.25]
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

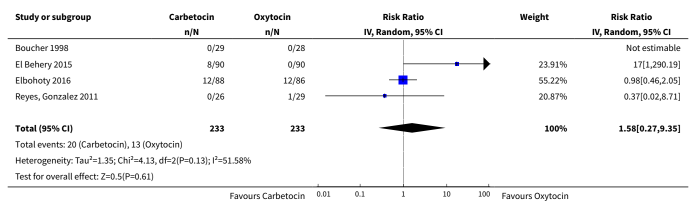




Analysis 10.16. Comparison 10 Carbetocin vs Oxytocin, Outcome 16 Shivering.



Analysis 10.17. Comparison 10 Carbetocin vs Oxytocin, Outcome 17 Fever.





Comparison 11. Ergometrine vs Oxytocin

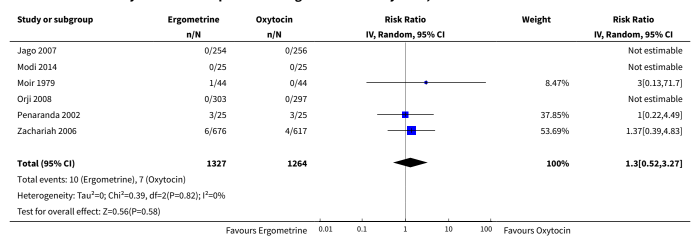
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	1293	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	6	2591	Risk Ratio (IV, Random, 95% CI)	1.30 [0.52, 3.27]
3 Blood transfusion	5	1893	Risk Ratio (IV, Random, 95% CI)	1.44 [0.20, 10.23]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	10	3221	Risk Ratio (IV, Random, 95% CI)	1.31 [0.86, 1.99]
7 Additional uterotonics	6	2493	Risk Ratio (IV, Random, 95% CI)	1.46 [0.61, 3.48]
8 Blood loss	10	3221	Mean Difference (IV, Random, 95% CI)	8.09 [-17.83, 34.00]
9 Change in haemoglo- bin	2	1893	Mean Difference (IV, Random, 95% CI)	0.42 [-0.30, 1.13]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	6	2529	Risk Ratio (IV, Random, 95% CI)	4.56 [1.13, 18.44]
12 Vomiting	5	2343	Risk Ratio (IV, Random, 95% CI)	3.83 [1.10, 13.28]
13 Headache	4	2293	Risk Ratio (IV, Random, 95% CI)	5.63 [0.93, 33.96]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	1410	Risk Ratio (IV, Random, 95% CI)	13.39 [2.01, 89.44]
16 Shivering	3	1493	Risk Ratio (IV, Random, 95% CI)	1.73 [0.93, 3.25]
17 Fever	2	1443	Risk Ratio (IV, Random, 95% CI)	2.97 [0.97, 9.05]
18 Diarrhoea	2	1393	Risk Ratio (IV, Random, 95% CI)	3.74 [0.42, 33.53]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Analysis 11.1. Comparison 11 Ergometrine vs Oxytocin, Outcome 1 Death.

Study or subgroup	Ergometrine	Ergometrine Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI					IV, Random, 95% CI	
Zachariah 2006	0/676	0/617							Not estimable
Total (95% CI)	676	617							Not estimable
Total events: 0 (Ergometrine),	, 0 (Oxytocin)								
Heterogeneity: Not applicable	2								
Test for overall effect: Not app	olicable			1		1			
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 11.2. Comparison 11 Ergometrine vs Oxytocin, Outcome 2 PPH >= 1000 mL.



Analysis 11.3. Comparison 11 Ergometrine vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Dhananjaya 2014	4/50	0/50	*	21.32%	9[0.5,162.89]
Ezeama 2014	1/149	9/151		27.74%	0.11[0.01,0.88]
Modi 2014	0/25	0/25			Not estimable
Singh 2009	3/75	0/75	-	20.98%	7[0.37,133.22]
Zachariah 2006	3/676	2/617		29.96%	1.37[0.23,8.17]
Total (95% CI)	975	918		100%	1.44[0.2,10.23]
Total events: 11 (Ergometrine	e), 11 (Oxytocin)				
Heterogeneity: Tau ² =2.51; Chi	i ² =8.33, df=3(P=0.04); l ² =63.9	98%			
Test for overall effect: Z=0.37((P=0.72)				
	Fav	ours Ergometrine 0	0.01 0.1 1 10 10	D Favours Oxytocin	



Analysis 11.6. Comparison 11 Ergometrine vs Oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine	Oxytocin	1	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Ra	ındom, 95% CI		IV, Random, 95% CI
Dhananjaya 2014	6/50	1/50		+	3.72%	6[0.75,48.05]
Ezeama 2014	3/149	12/151			9.07%	0.25[0.07,0.88]
Jago 2007	0/254	0/256				Not estimable
Modi 2014	0/25	0/25				Not estimable
Moir 1979	2/44	2/44			4.32%	1[0.15,6.79]
Moodie 1976	14/40	10/40		+	20.72%	1.4[0.71,2.77]
Orji 2008	18/303	12/297		+-	19.75%	1.47[0.72,3]
Penaranda 2002	12/25	8/25		+-	20.09%	1.5[0.74,3.03]
Singh 2009	2/75	0/75	_	- - 	1.84%	5[0.24,102.42]
Zachariah 2006	20/676	13/617		+	20.49%	1.4[0.7,2.8]
Total (95% CI)	1641	1580		•	100%	1.31[0.86,1.99]
Total events: 77 (Ergometrine	e), 58 (Oxytocin)					
Heterogeneity: Tau ² =0.1; Chi ²	=9.89, df=7(P=0.19); I ² =29.22	2%				
Test for overall effect: Z=1.26((P=0.21)					
<u> </u>		ours Ergometrine	0.01 0.1	1 10	100 Favours Oxytocin	

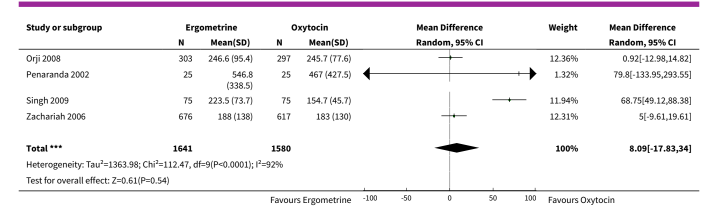
Analysis 11.7. Comparison 11 Ergometrine vs Oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine	Oxytocin			Risk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95% C	1		IV, Random, 95% CI	
Dhananjaya 2014	9/50	0/50				\longrightarrow	7.16%	19[1.14,317.87]	
Ezeama 2014	11/149	35/151		_	-		24.74%	0.32[0.17,0.6]	
Modi 2014	0/25	0/25						Not estimable	
Orji 2008	30/303	18/297			-		25.5%	1.63[0.93,2.87]	
Singh 2009	11/75	2/75					15.72%	5.5[1.26,23.97]	
Zachariah 2006	51/676	38/617			-		26.87%	1.22[0.82,1.84]	
Total (95% CI)	1278	1215			•		100%	1.46[0.61,3.48]	
Total events: 112 (Ergometrin	ne), 93 (Oxytocin)								
Heterogeneity: Tau ² =0.69; Ch	ni ² =25.16, df=4(P<0.0001); l ² =	84.1%							
Test for overall effect: Z=0.84	(P=0.4)								
	Fav	ours Ergometrine	0.01	0.1	1	10 100	Favours Oxytocin		

Analysis 11.8. Comparison 11 Ergometrine vs Oxytocin, Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Dhananjaya 2014	50	345 (109.5)	50	219 (86.3)		9.97%	126[87.35,164.65]
Ezeama 2014	149	287.1 (84.4)	151	301.8 (109.2)	-+-	11.73%	-14.7[-36.77,7.37]
Jago 2007	254	150.2 (63.6)	256	171.9 (81.6)		12.43%	-21.7[-34.39,-9.01]
Modi 2014	25	131 (72)	25	223.2 (122.5)	←	8.05%	-92.2[-147.92,-36.48]
Moir 1979	44	201 (50)	44	208 (58)		11.68%	-7[-29.63,15.63]
Moodie 1976	40	369 (118)	40	391 (129)		8.21%	-22[-76.18,32.18]
			Favour	s Ergometrine	-100 -50 0 50	100 Favours Oxy	/tocin





Analysis 11.9. Comparison 11 Ergometrine vs Oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup	Erge	ometrine	0:	kytocin		Mea	n Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ran	dom, 95% CI			Random, 95% CI
Orji 2008	303	15.1 (5.2)	297	14.5 (5.2)			+		73.89%	0.6[-0.23,1.43]
Zachariah 2006	676	14.8 (11.9)	617	14.9 (13.6)		_	-		26.11%	-0.1[-1.5,1.3]
Total ***	979		914				•		100%	0.42[-0.3,1.13]
Heterogeneity: Tau ² =0; Chi ² =	0.71, df=1(P=0.4)); I ² =0%								
Test for overall effect: Z=1.14	(P=0.25)									
			Favours	Ergometrine	-5	-2.5	0 2.	5 5	Favours Oxytoc	in

Analysis 11.11. Comparison 11 Ergometrine vs Oxytocin, Outcome 11 Nausea.

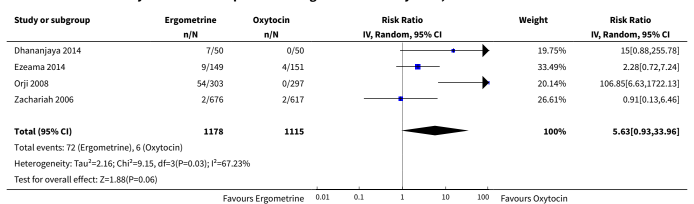
Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% C	1	IV, Random, 95% CI
Dhananjaya 2014	1/50	0/50		10.73%	3[0.13,71.92]
Ezeama 2014	9/149	3/151	+		3.04[0.84,11.01]
Moir 1979	6/44	0/44		12.05%	13[0.75,223.98]
Moodie 1976	35/78	0/70	<u> </u>	12.37%	63.81[3.99,1021.17]
Orji 2008	132/303	15/297	-	23.34%	8.63[5.18,14.36]
Zachariah 2006	6/676	11/617	-+-	21.5%	0.5[0.19,1.34]
Total (95% CI)	1300	1229		100%	4.56[1.13,18.44]
Total events: 189 (Ergometri	ne), 29 (Oxytocin)				
Heterogeneity: Tau ² =2.11; Ch	ni ² =29.64, df=5(P<0.0001); I ² =8	83.13%			
Test for overall effect: Z=2.13	(P=0.03)				
	Fav	ours Ergometrine	0.01 0.1 1	10 100 Favours Oxytocin	



Analysis 11.12. Comparison 11 Ergometrine vs Oxytocin, Outcome 12 Vomiting.

Study or subgroup	Ergometrine	Oxytocin		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
Dhananjaya 2014	4/50	0/50		_	+	\rightarrow	12.59%	9[0.5,162.89]
Ezeama 2014	4/149	1/151			+	_	17.87%	4.05[0.46,35.85]
Orji 2008	132/303	12/297			-		36.56%	10.78[6.1,19.04]
Penaranda 2002	0/25	1/25		+			11.17%	0.33[0.01,7.81]
Zachariah 2006	3/676	2/617			<u> </u>		21.8%	1.37[0.23,8.17]
Total (95% CI)	1203	1140			-		100%	3.83[1.1,13.28]
Total events: 143 (Ergometrin	ne), 16 (Oxytocin)							
Heterogeneity: Tau ² =1.02; Ch	i ² =9.08, df=4(P=0.06); l ² =55.9	7%						
Test for overall effect: Z=2.11((P=0.03)			1				
	Fav	ours Ergometrine	0.01	0.1	1 10	100	Favours Oxytocin	

Analysis 11.13. Comparison 11 Ergometrine vs Oxytocin, Outcome 13 Headache.



Analysis 11.15. Comparison 11 Ergometrine vs Oxytocin, Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N n/N IV, Rand		andom, 95	% CI			IV, Random, 95% CI		
Ezeama 2014	7/149	0/151			-		\rightarrow	23.98%	15.2[0.88,263.78]
Jago 2007	134/254	29/256				-		51.35%	4.66[3.24,6.69]
Orji 2008	54/303	0/297					→	24.66%	106.85[6.63,1722.13]
Total (95% CI)	706	704			-		_	100%	13.39[2.01,89.44]
Total events: 195 (Ergometrin	ne), 29 (Oxytocin)								
Heterogeneity: Tau ² =1.79; Ch	i ² =5.39, df=2(P=0.07); I ² =62.9	1%							
Test for overall effect: Z=2.68((P=0.01)								
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	



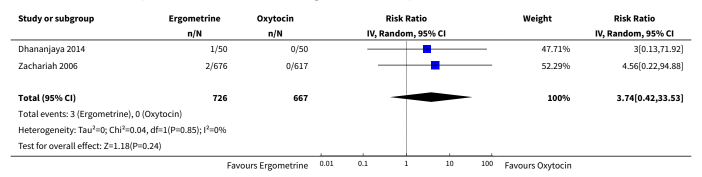
Analysis 11.16. Comparison 11 Ergometrine vs Oxytocin, Outcome 16 Shivering.

Study or subgroup	Ergometrine	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, I	Random, 95%	% CI			IV, Random, 95% CI
Penaranda 2002	1/25	0/25			+			3.96%	3[0.13,70.3]
Singh 2009	0/75	0/75			İ				Not estimable
Zachariah 2006	26/676	14/617						96.04%	1.7[0.89,3.22]
Total (95% CI)	776	717			•			100%	1.73[0.93,3.25]
Total events: 27 (Ergometrine	e), 14 (Oxytocin)				İ				
Heterogeneity: Tau ² =0; Chi ² =	0.12, df=1(P=0.73); I ² =0%				İ				
Test for overall effect: Z=1.72	(P=0.09)					1			
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 11.17. Comparison 11 Ergometrine vs Oxytocin, Outcome 17 Fever.

Study or subgroup	Ergometrine	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Singh 2009	0/75	0/75							Not estimable
Zachariah 2006	13/676	4/617						100%	2.97[0.97,9.05]
Total (95% CI)	751	692				>		100%	2.97[0.97,9.05]
Total events: 13 (Ergometrine), 4 (Oxytocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.91(P=0.0	06)					1			
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 11.18. Comparison 11 Ergometrine vs Oxytocin, Outcome 18 Diarrhoea.



Comparison 12. Ergometine plus oxytocin vs Oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	2	1314	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	10	10990	Risk Ratio (IV, Random, 95% CI)	0.73 [0.57, 0.93]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3 Blood transfusion	11	10985	Risk Ratio (IV, Random, 95% CI)	0.88 [0.53, 1.44]
4 Severe maternal morbidity: intensive care admissions	1	991	Risk Ratio (IV, Random, 95% CI)	2.99 [0.12, 73.32]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	11090	Risk Ratio (IV, Random, 95% CI)	0.72 [0.57, 0.91]
7 Additional uterotonics	10	8968	Risk Ratio (IV, Random, 95% CI)	0.79 [0.59, 1.07]
8 Blood loss	10	4248	Mean Difference (IV, Random, 95% CI)	-10.31 [-40.32, 19.70]
9 Change in haemoglobin	6	4324	Mean Difference (IV, Random, 95% CI)	-2.23 [-5.24, 0.77]
10 Breastfeeding	1	3483	Risk Ratio (IV, Random, 95% CI)	0.99 [0.96, 1.01]
11 Nausea	7	6931	Risk Ratio (IV, Random, 95% CI)	1.72 [0.84, 3.53]
12 Vomiting	9	10227	Risk Ratio (IV, Random, 95% CI)	3.05 [1.76, 5.29]
13 Headache	5	5105	Risk Ratio (IV, Random, 95% CI)	1.26 [0.79, 1.99]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	1362	Risk Ratio (IV, Random, 95% CI)	2.00 [0.29, 13.97]
16 Shivering	2	1591	Risk Ratio (IV, Random, 95% CI)	0.96 [0.60, 1.53]
17 Fever	2	1591	Risk Ratio (IV, Random, 95% CI)	1.08 [0.48, 2.43]
18 Diarrhoea	3	2030	Risk Ratio (IV, Random, 95% CI)	1.26 [0.72, 2.22]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

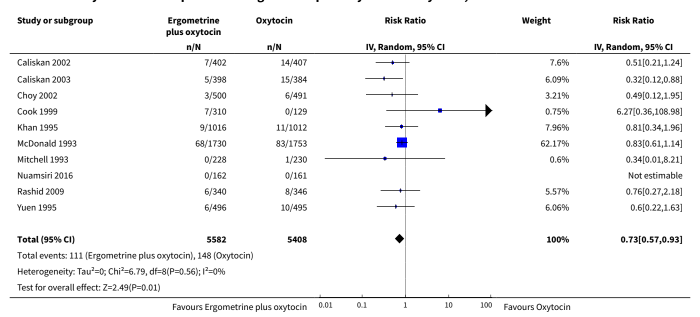
Analysis 12.1. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 1 Death.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Nuamsiri 2016	0/162	0/161							Not estimable
Yuen 1995	0/496	0/495							Not estimable
Total (95% CI)	658	656							Not estimable
Total events: 0 (Ergometrine plu	s oxytocin), 0 (Oxytocin)								
Heterogeneity: Not applicable						1			
	Favours Ergometr	ine plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	



Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Test for overall effect: Not applicable									
	Favours Ergomet	rine plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 12.2. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 2 PPH >= 1000 mL.

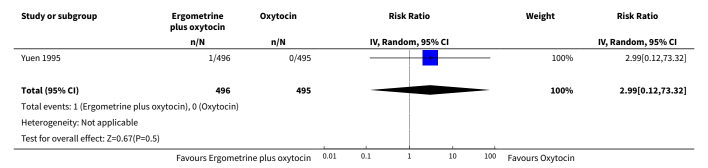


Analysis 12.3. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Balki 2008	0/24	0/24			Not estimable
Caliskan 2002	4/402	13/407		10.62%	0.31[0.1,0.95]
Caliskan 2003	6/398	13/384		12.33%	0.45[0.17,1.16]
Choy 2002	13/493	7/487	+-	12.9%	1.83[0.74,4.56]
Cook 1999	3/310	2/129		5.79%	0.62[0.11,3.69]
Khan 1995	2/1016	1/1012		3.59%	1.99[0.18,21.93]
Koen 2016	7/202	19/214		13.73%	0.39[0.17,0.91]
McDonald 1993	24/1730	16/1753	+-	16.71%	1.52[0.81,2.85]
Nuamsiri 2016	2/162	1/161		3.61%	1.99[0.18,21.7]
Rashid 2009	6/340	2/346	++	6.79%	3.05[0.62,15.02]
Yuen 1995	10/496	12/495		13.93%	0.83[0.36,1.91]
Total (95% CI)	5573	5412	•	100%	0.88[0.53,1.44]
Total events: 77 (Ergometrine	plus oxytocin), 86 (Oxytocir	n)			
Heterogeneity: Tau ² =0.28; Chi	² =17.64, df=9(P=0.04); l ² =48	.98%			
Test for overall effect: Z=0.52(P=0.6)				
	Favours Ergomet	rine plus oxytocin 0.01	0.1 1 10	100 Favours Oxytocin	



Analysis 12.4. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 4 Severe maternal morbidity: intensive care admissions.



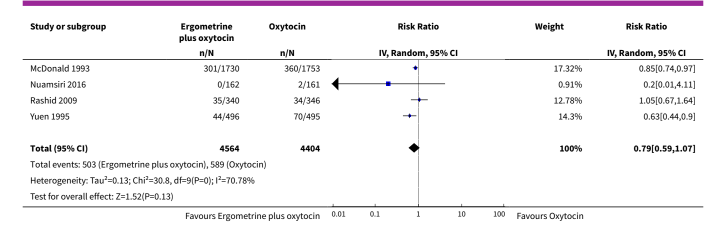
Analysis 12.6. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Caliskan 2002	14/402	33/407		9.06%	0.43[0.23,0.79]
Caliskan 2003	14/398	28/384		8.78%	0.48[0.26,0.9]
Choy 2002	23/500	26/491	+	10.24%	0.87[0.5,1.5]
Cook 1999	23/310	1/129		1.35%	9.57[1.31,70.13]
Khan 1995	36/1016	41/1012	+	12.66%	0.87[0.56,1.36]
McDonald 1993	286/1730	316/1753	+	20.46%	0.92[0.79,1.06]
Mitchell 1993	6/228	17/230		5.23%	0.36[0.14,0.89]
Nuamsiri 2016	0/162	1/161	+	0.55%	0.33[0.01,8.07]
Rashid 2009	8/340	9/346		4.99%	0.9[0.35,2.32]
Un Nisa 2012	20/50	26/50	-+ +	12.85%	0.77[0.5,1.18]
Yuen 1995	36/496	60/495	+	13.82%	0.6[0.4,0.89]
Total (95% CI)	5632	5458	•	100%	0.72[0.57,0.91]
Total events: 466 (Ergometrin	e plus oxytocin), 558 (Oxyto	cin)			
Heterogeneity: Tau²=0.07; Chi	² =21.42, df=10(P=0.02); I ² =5	3.31%			
Test for overall effect: Z=2.7(P	=0.01)				
	Favours Ergometi	rine nlus oxytocin	0.01 0.1 1 10	100 Favours Oxytocin	

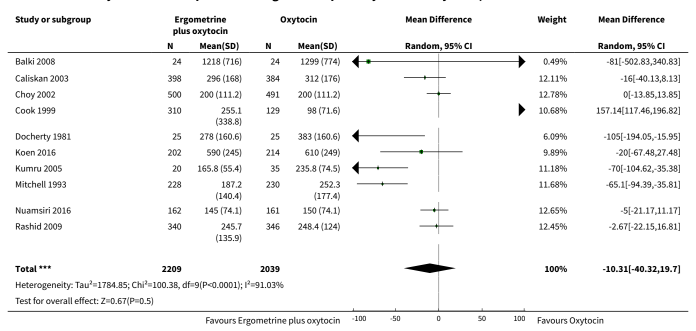
Analysis 12.7. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Balki 2008	5/24	13/24	-+-	7.16%	0.38[0.16,0.91]
Caliskan 2002	9/402	26/407	-+-	8.44%	0.35[0.17,0.74]
Caliskan 2003	9/398	26/384		8.45%	0.33[0.16,0.7]
Choy 2002	52/500	36/491	+	13.47%	1.42[0.94,2.13]
Cook 1999	28/310	6/129	+-	7.21%	1.94[0.82,4.58]
Koen 2016	20/202	16/214	-	9.97%	1.32[0.71,2.48]
	Favours Ergometi	ine plus oxytocin 0.0	01 0.1 1 10	100 Favours Oxytocin	





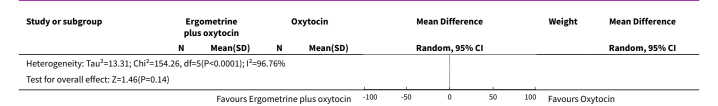
Analysis 12.8. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 8 Blood loss.



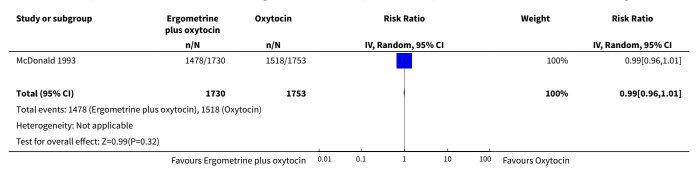
Analysis 12.9. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup		ometrine oxytocin	0	xytocin		M	ean Differen	:e		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	andom, 95%	CI			Random, 95% CI
Caliskan 2002	402	15 (12)	407	14 (14)			•			16.59%	1[-0.8,2.8]
Caliskan 2003	398	15 (12)	384	14 (14)			•			16.55%	1[-0.83,2.83]
Choy 2002	493	10 (11.1)	487	8 (11.9)			•			16.95%	2[0.56,3.44]
Cook 1999	310	0.5 (14.1)	129	14.7 (12.1)			+			15.56%	-14.2[-16.81,-11.59]
Nuamsiri 2016	162	8 (7.4)	161	9 (7.4)			+			16.77%	-1[-2.62,0.62]
Yuen 1995	496	12 (2.8)	495	15 (2.8)			•			17.59%	-3[-3.35,-2.65]
Total ***	2261		2063		1		•			100%	-2.23[-5.24,0.77]
		Favours Erg	ometrine	plus oxytocin	-100	-50	0	50	100	Favours Oxy	rtocin





Analysis 12.10. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 10 Breastfeeding.



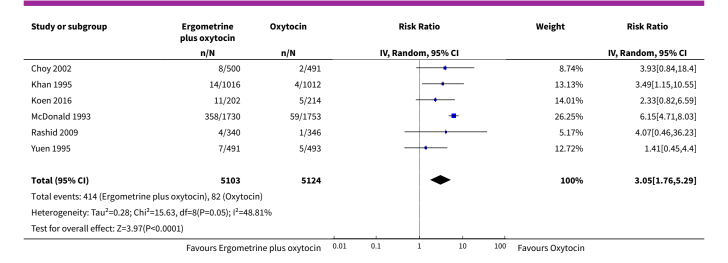
Analysis 12.11. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 11 Nausea.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Ri	Risk Ratio		Risk Ratio
	n/N	n/N	IV, Ran	dom, 95% CI		IV, Random, 95% CI
Balki 2008	10/24	2/24			10.95%	5[1.22,20.46]
Choy 2002	198/500	186/491		+	18.75%	1.05[0.89,1.22]
Koen 2016	26/202	23/214			17.17%	1.2[0.71,2.03]
McDonald 1993	467/1730	117/1753		+	18.67%	4.04[3.34,4.9]
Nuamsiri 2016	1/162	2/161			6.11%	0.5[0.05,5.43]
Rashid 2009	12/340	10/346		+	15.14%	1.22[0.53,2.79]
Yuen 1995	9/491	5/493		+-	13.21%	1.81[0.61,5.35]
Total (95% CI)	3449	3482		•	100%	1.72[0.84,3.53]
Total events: 723 (Ergometrin	ne plus oxytocin), 345 (Oxyto	cin)				
Heterogeneity: Tau ² =0.71; Ch	ni ² =121.03, df=6(P<0.0001); I ² =	=95.04%				
Test for overall effect: Z=1.48	(P=0.14)					
	Favours Ergometr	ine plus oxytocin	0.01 0.1	1 10	¹⁰⁰ Favours Oxytocin	

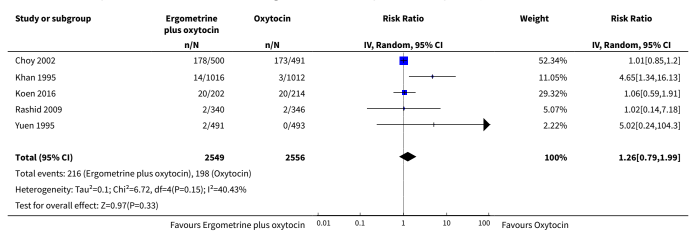
Analysis 12.12. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 12 Vomiting.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio			Weight	Risk Ratio
	n/N	n/N	IV, Rand	lom, 95% CI			IV, Random, 95% CI
Balki 2008	6/24	1/24		+		5.78%	6[0.78,46.14]
Caliskan 2002	1/402	2/407		 		4.44%	0.51[0.05,5.56]
Caliskan 2003	5/398	3/384	. –	 • 		9.75%	1.61[0.39,6.68]
	Favours Ergometr	ine plus oxytocin	0.01 0.1	1 10	100	Favours Oxytocin	





Analysis 12.13. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 13 Headache.



Analysis 12.15. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 15 Hypertension.

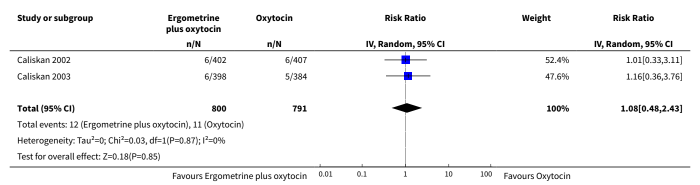
Study or subgroup	Ergometrine plus oxytocin	Oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 959	% CI			IV, Random, 95% CI
Balki 2008	1/24	4/24	_	-				31.09%	0.25[0.03,2.08]
Choy 2002	17/500	7/491			-	_		44.78%	2.38[1,5.7]
Nuamsiri 2016	10/162	0/161				•	—	24.12%	20.87[1.23,353.21]
Total (95% CI)	686	676						100%	2[0.29,13.97]
Total events: 28 (Ergometrine	e plus oxytocin), 11 (Oxytocin)							
Heterogeneity: Tau ² =2; Chi ² =	6.5, df=2(P=0.04); I ² =69.21%								
Test for overall effect: Z=0.7(I	P=0.49)								
	Favours Ergometr	ine plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	



Analysis 12.16. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 16 Shivering.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV	, Random, 95% CI			IV, Random, 95% CI
Caliskan 2002	19/402	16/407		-		50.89%	1.2[0.63,2.3]
Caliskan 2003	15/398	19/384		-		49.11%	0.76[0.39,1.48]
Total (95% CI)	800	791		•		100%	0.96[0.6,1.53]
Total events: 34 (Ergometrine	plus oxytocin), 35 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0	0.93, df=1(P=0.34); I ² =0%						
Test for overall effect: Z=0.17((P=0.87)				1		
	Favours Ergometr	ine plus oxytocin	0.01 0.1	1 10	100	Favours Oxytocin	

Analysis 12.17. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 17 Fever.



Analysis 12.18. Comparison 12 Ergometine plus oxytocin vs Oxytocin, Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine Oxytocin plus oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N	IV,	Random, 95% CI			IV, Random, 95% CI
Caliskan 2002	10/402	9/407				39.93%	1.12[0.46,2.74]
Caliskan 2003	17/398	12/384		-		60.07%	1.37[0.66,2.82]
Cook 1999	0/310	0/129					Not estimable
Total (95% CI)	1110	920		•		100%	1.26[0.72,2.22]
Total events: 27 (Ergometrine	e plus oxytocin), 21 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =	0.11, df=1(P=0.74); I ² =0%						
Test for overall effect: Z=0.82	(P=0.41)						
	Favours Ergometr	ine plus oxytocin	0.01 0.1	1 10	100	Favours Oxytocin	

Comparison 13. Misoprostol plus oxytocin vs Oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
1 Death	9	4737	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2 PPH >= 1000 mL	17	8514	Risk Ratio (IV, Random, 95% CI)	0.87 [0.69, 1.09]
3 Blood transfusion	19	8742	Risk Ratio (IV, Random, 95% CI)	0.50 [0.37, 0.67]
4 Severe maternal morbidity: intensive care admissions	3	1886	Risk Ratio (IV, Random, 95% CI)	0.50 [0.05, 5.47]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	14	8148	Risk Ratio (IV, Random, 95% CI)	0.71 [0.59, 0.85]
7 Additional uterotonics	18	8391	Risk Ratio (IV, Random, 95% CI)	0.54 [0.44, 0.67]
8 Blood loss	17	8690	Mean Difference (IV, Random, 95% CI)	-87.26 [-157.83, -16.69]
9 Change in haemoglo- bin	15	7929	Mean Difference (IV, Random, 95% CI)	-2.59 [-3.70, -1.48]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	7	3798	Risk Ratio (IV, Random, 95% CI)	2.21 [1.19, 4.10]
12 Vomiting	11	6718	Risk Ratio (IV, Random, 95% CI)	2.24 [1.52, 3.31]
13 Headache	2	303	Risk Ratio (IV, Random, 95% CI)	1.26 [0.26, 6.23]
14 Abdominal pain	1	366	Risk Ratio (IV, Random, 95% CI)	1.93 [1.01, 3.67]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	19	9458	Risk Ratio (IV, Random, 95% CI)	3.38 [2.50, 4.57]
17 Fever	17	8607	Risk Ratio (IV, Random, 95% CI)	2.99 [2.00, 4.45]
18 Diarrhoea	7	5649	Risk Ratio (IV, Random, 95% CI)	2.08 [0.99, 4.38]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 13.1. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 1 Death.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Badejoko 2012	0/126	0/129							Not estimable
Bhullar 2004	0/377	0/379							Not estimable
Carbonell 2009	0/702	0/698							Not estimable
	Favours Misopros	tol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	



Study or subgroup	Misoprostol plus oxytocin				Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Chaudhuri 2015	0/198	0/198							Not estimable
Chaudhuri 2016	0/144	0/144							Not estimable
El Tahan 2012	0/179	0/187							Not estimable
Hofmeyr 2011	0/546	0/557							Not estimable
Lapaire 2006	0/28	0/25							Not estimable
Ugwu 2014	0/60	0/60							Not estimable
Total (95% CI)	2360	2377							Not estimable
Total events: 0 (Misoprostol plus	s oxytocin), 0 (Oxytocin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applic	cable								
	Favours Misopros	stol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 13.2. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Adanikin 2012	0/109	0/109			Not estimable
Badejoko 2012	3/126	5/129		2.55%	0.61[0.15,2.52]
Caliskan 2002	11/401	14/407	+ -	8.39%	0.8[0.37,1.74]
Caliskan 2003	6/404	15/384		5.78%	0.38[0.15,0.97]
Carbonell 2009	13/702	11/698	-	8%	1.18[0.53,2.6]
Chaudhuri 2015	5/198	3/198		2.52%	1.67[0.4,6.88]
Chaudhuri 2016	2/144	4/144		1.79%	0.5[0.09,2.69]
Elsedeek 2012	0/200	0/200			Not estimable
Fekih 2009	19/125	24/125	-+	16.86%	0.79[0.46,1.37]
Hamm 2005	24/173	22/179	-	17.42%	1.13[0.66,1.94]
Hernandez-Castro 2016	3/60	7/60		2.98%	0.43[0.12,1.58]
Hofmeyr 2011	5/546	1/553	+	1.1%	5.06[0.59,43.2]
Lapaire 2006	13/24	11/19		17.95%	0.94[0.55,1.59]
Quibel 2016	13/806	17/797		9.9%	0.76[0.37,1.55]
Sitaula 2017	0/100	1/100 —		0.5%	0.33[0.01,8.09]
Sood 2012	6/90	4/84	- +	3.35%	1.4[0.41,4.79]
Ugwu 2014	1/60	2/60		0.9%	0.5[0.05,5.37]
Total (95% CI)	4268	4246	•	100%	0.87[0.69,1.09]
Total events: 124 (Misoprostol	plus oxytocin), 141 (Oxytoci	n)			
Heterogeneity: Tau ² =0; Chi ² =11	14, df=14(P=0.67); I ² =0%				
Test for overall effect: Z=1.22(P	=0.22)				
Test for overall effect: Z=1.22(P	=0.22) Favours Misopros	tol plus oxytocin 0.01	0.1 1 10	100 Favours Oxytocin	



Analysis 13.3. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol Oxytocin plus oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Badejoko 2012	1/126	6/129		1.89%	0.17[0.02,1.4]
Bhullar 2004	3/377	6/379		4.41%	0.5[0.13,2]
Caliskan 2002	4/401	13/407		6.77%	0.31[0.1,0.95]
Caliskan 2003	5/404	13/384		8.02%	0.37[0.13,1.02]
Carbonell 2009	5/702	13/698		7.96%	0.38[0.14,1.07]
Chaudhuri 2015	10/198	15/198		13.93%	0.67[0.31,1.45]
Chaudhuri 2016	5/144	12/144		8.09%	0.42[0.15,1.15]
El Tahan 2012	0/179	11/187		1.05%	0.05[0,0.76]
Elsedeek 2012	0/200	0/200			Not estimable
Fekih 2009	0/125	4/125		0.99%	0.11[0.01,2.04]
Hamm 2005	3/173	3/179		3.33%	1.03[0.21,5.06]
Hernandez-Castro 2016	0/60	5/60		1.01%	0.09[0.01,1.61]
Hong 2007	11/96	13/118		14.64%	1.04[0.49,2.22]
Lapaire 2006	0/28	0/25			Not estimable
Nayak 2017	9/100	23/100		16.2%	0.39[0.19,0.8]
Quibel 2016	5/806	9/797		7.07%	0.55[0.18,1.63]
Sitaula 2017	0/100	1/100 -		0.82%	0.33[0.01,8.09]
Sood 2012	3/90	2/84		2.69%	1.4[0.24,8.17]
Ugwu 2014	1/60	1/59		1.11%	0.98[0.06,15.36]
Total (95% CI)	4369	4373	•	100%	0.5[0.37,0.67]
Total events: 65 (Misoprostol plu	s oxytocin), 150 (Oxytocin)				
Heterogeneity: Tau²=0; Chi²=14.6	5, df=16(P=0.55); I ² =0%				
Test for overall effect: Z=4.7(P<0.	0001)				

Analysis 13.4. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Misoprostol Oxytocin plus oxytocin			Ris	sk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Rand	dom, 95% C	ı			IV, Random, 95% CI
Carbonell 2009	1/702	2/698		-	-			100%	0.5[0.05,5.47]
El Tahan 2012	0/179	0/187			_				Not estimable
Ugwu 2014	0/60	0/60							Not estimable
Total (95% CI)	941	945						100%	0.5[0.05,5.47]
Total events: 1 (Misoprostol plus oxy	tocin), 2 (Oxytocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.57(P=0.57)									
	Favours Misopros	stol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	



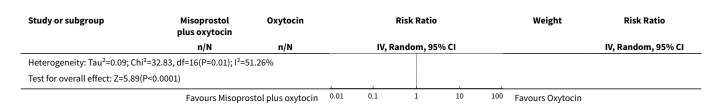
Analysis 13.6. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Adanikin 2012	0/109	0/109			Not estimable
Badejoko 2012	28/126	27/129	+	7.6%	1.06[0.66,1.7]
Bhullar 2004	13/377	20/379		4.82%	0.65[0.33,1.29]
Caliskan 2002	28/401	33/407	-+	7.34%	0.86[0.53,1.4]
Caliskan 2003	13/404	28/384		5.24%	0.44[0.23,0.84]
Carbonell 2009	28/702	50/698	-+-	7.9%	0.56[0.35,0.87]
Chaudhuri 2015	79/198	132/198	+	13.09%	0.6[0.49,0.73]
Chaudhuri 2016	7/144	19/144		3.6%	0.37[0.16,0.85]
Hofmeyr 2011	22/546	35/553	-+ 	6.79%	0.64[0.38,1.07]
Lapaire 2006	18/24	15/19	+	10.27%	0.95[0.68,1.32]
Nayak 2017	2/100	9/100		1.32%	0.22[0.05,1]
Quibel 2016	68/806	66/797	+	10.34%	1.02[0.74,1.41]
Sood 2012	73/90	77/84	+	14.53%	0.88[0.79,1]
Ugwu 2014	15/60	33/60		7.18%	0.45[0.28,0.75]
Total (95% CI)	4087	4061	•	100%	0.71[0.59,0.85]
Total events: 394 (Misoprosto	ol plus oxytocin), 544 (Oxytoc	in)			
Heterogeneity: Tau ² =0.05; Ch	ni ² =33.44, df=12(P=0); l ² =64.1	1%			
Test for overall effect: Z=3.77	(P=0)				
	Favours Misopro	stol plus oxytocin 0.	.01 0.1 1 10	100 Favours Oxytocin	

Analysis 13.7. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Badejoko 2012	6/126	5/129		2.48%	1.23[0.38,3.92]
Bhullar 2004	10/377	13/379		4.2%	0.77[0.34,1.74]
Caliskan 2002	17/401	26/407	+	6.08%	0.66[0.37,1.2]
Caliskan 2003	10/404	26/384		4.94%	0.37[0.18,0.75]
Carbonell 2009	33/702	54/698	-+-	8.23%	0.61[0.4,0.92]
Chaudhuri 2015	18/198	45/198	→	7.06%	0.4[0.24,0.67]
Chaudhuri 2016	12/144	22/144		5.39%	0.55[0.28,1.06]
El Tahan 2012	12/179	52/187		6.1%	0.24[0.13,0.44]
Elsedeek 2012	14/200	36/200		6.19%	0.39[0.22,0.7]
Hamm 2005	45/173	76/179		9.88%	0.61[0.45,0.83]
Hernandez-Castro 2016	6/60	24/60		4.15%	0.25[0.11,0.57]
Hong 2007	28/96	31/118	- +	8.04%	1.11[0.72,1.71]
Lapaire 2006	0/28	0/25			Not estimable
Nayak 2017	4/100	7/100		2.36%	0.57[0.17,1.89]
Pakniat 2015	7/50	7/50		3.26%	1[0.38,2.64]
Quibel 2016	19/806	25/797	-+ 	6.16%	0.75[0.42,1.35]
Sood 2012	20/90	36/84		7.71%	0.52[0.33,0.82]
Ugwu 2014	16/58	40/60		7.78%	0.41[0.26,0.65]
Total (95% CI)	4192	4199	•	100%	0.54[0.44,0.67]
Total events: 277 (Misoprostol	plus oxytocin), 525 (Oxytoc	in)			
	Favours Misopro	stol plus oxytocin 0.0	01 0.1 1 10	100 Favours Oxytocin	





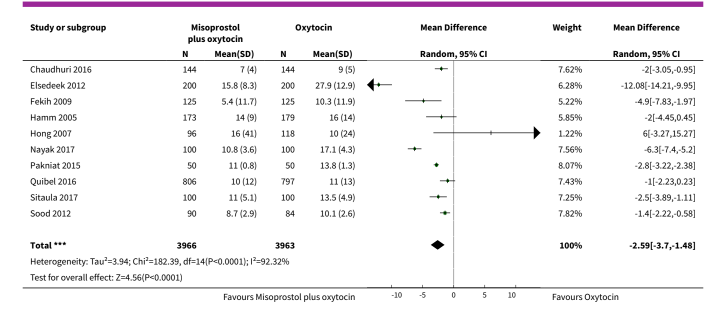
Analysis 13.8. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 8 Blood loss.

Study or subgroup		oprostol oxytocin	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Badejoko 2012	126	387.3 (203.1)	129	386.7 (298.5)	+	5.97%	0.55[-61.99,63.09]
Bhullar 2004	377	322 (114)	379	329 (123)	+	6.24%	-7[-23.9,9.9]
Caliskan 2003	404	280 (182)	384	312 (176)	+	6.22%	-32[-57,-7]
Carbonell 2009	702	243.6 (181.2)	698	240.9 (145.8)	+	6.24%	2.7[-14.53,19.93]
Chaudhuri 2015	198	505.4 (215.5)	198	587.3 (201.5)	+	6.13%	-81.9[-122.99,-40.81]
Chaudhuri 2016	144	225.8 (156.7)	144	302.4 (230.3)	+	6.11%	-76.6[-122.1,-31.1]
El Tahan 2012	179	324 (97.4)	187	894 (160.9)	+	6.21%	-570[-597.12,-542.88]
Elsedeek 2012	200	429 (234)	200	620 (375)		5.98%	-191[-252.26,-129.74]
Fekih 2009	125	669.7 (333)	125	852.5 (295.1)		5.82%	-182.84[-260.84,-104.84]
Hamm 2005	173	749 (173)	179	725 (212)	+	6.14%	24[-16.36,64.36]
Hofmeyr 2011	540	189 (288.1)	549	199 (290.5)	+	6.17%	-10[-44.37,24.37]
Lapaire 2006	28	1083 (920)	25	970 (560)		2.04%	113[-292.35,518.35]
Nayak 2017	100	363.4 (77.7)	100	481.3 (116.6)	+	6.21%	-117.9[-145.36,-90.44]
Quibel 2016	806	150 (122.3)	797	150 (111.2)	 	6.26%	0[-11.44,11.44]
Sitaula 2017	100	326.9 (116.2)	100	397.7 (110.1)	+	6.19%	-70.8[-102.17,-39.43]
Sood 2012	90	595 (108)	84	651 (118)	+	6.18%	-56[-89.68,-22.32]
Ugwu 2014	60	451.3 (204)	60	551.2 (192)	+	5.89%	-99.9[-170.78,-29.02]
Total ***	4352		4338		•	100%	-87.26[-157.83,-16.69]
Heterogeneity: Tau ² =20689.5	51; Chi²=1630.37,	df=16(P<0.0001); I ² =99.02	2%			
Test for overall effect: Z=2.42	2(P=0.02)						

Analysis 13.9. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup		oprostol oxytocin	0:	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Bhullar 2004	377	10 (11)	379	11 (12)	-+ 	6.93%	-1[-2.64,0.64]
Caliskan 2002	401	15 (13)	407	14 (14)	 	6.65%	1[-0.86,2.86]
Caliskan 2003	404	14 (13)	384	14 (14)	-	6.61%	0[-1.89,1.89]
Carbonell 2009	702	10.5 (7.5)	698	11.5 (8.2)	-+-	7.82%	-1[-1.82,-0.18]
Chaudhuri 2015	198	10 (4)	198	13 (6)		7.66%	-3[-4,-2]
		Favours Mis	oprostol	plus oxytocin	-10 -5 0 5 10	Favours Ox	ytocin





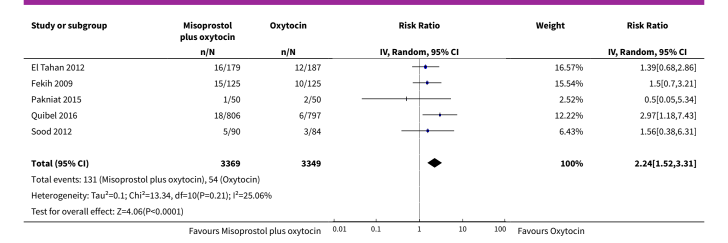
Analysis 13.11. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 11 Nausea.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	•		Weight	Risk Ratio	
	n/N	n/N	IV, Rando	om, 95% CI		IV, Random, 95% CI	
Adanikin 2012	7/109	2/109		+	10.46%	3.5[0.74,16.47]	
Carbonell 2009	14/702	2/698			11.11%	6.96[1.59,30.51]	
Fekih 2009	26/125	6/125			19.32%	4.33[1.85,10.16]	
Lapaire 2006	0/28	1/25	+		3.4%	0.3[0.01,7.02]	
Pakniat 2015	6/50	7/50		•	16.71%	0.86[0.31,2.37]	
Quibel 2016	22/806	8/797		-	20.15%	2.72[1.22,6.07]	
Sood 2012	10/90	8/84	_	-	18.85%	1.17[0.48,2.82]	
Total (95% CI)	1910	1888		•	100%	2.21[1.19,4.1]	
Total events: 85 (Misoprostol	l plus oxytocin), 34 (Oxytocin)	ı					
Heterogeneity: Tau ² =0.32; Ch	ni²=12.2, df=6(P=0.06); I²=50.8	1%					
Test for overall effect: Z=2.52	(P=0.01)						
	Favours Misopros	stol plus oxytocin	0.01 0.1	1 10	100 Favours Oxytocin		

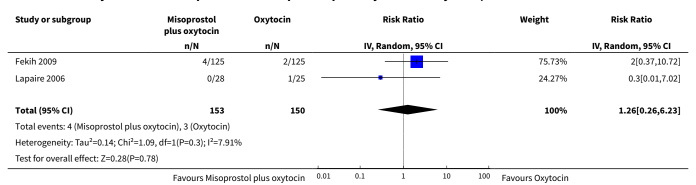
Analysis 13.12. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	ľ	V, Random, 95% CI			IV, Random, 95% CI
Adanikin 2012	8/109	3/109		++-		7.26%	2.67[0.73,9.79]
Badejoko 2012	29/126	7/129		_ 		14.9%	4.24[1.93,9.33]
Bhullar 2004	5/377	2/379		+		4.94%	2.51[0.49,12.87]
Caliskan 2002	3/401	2/407		+		4.23%	1.52[0.26,9.06]
Caliskan 2003	3/404	3/384				5.16%	0.95[0.19,4.68]
Carbonell 2009	28/702	4/698				10.22%	6.96[2.45,19.74]
	Favours Misopros	tol plus oxytocin	0.01 0.1	1 10	100	Favours Oxytocin	





Analysis 13.13. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 13 Headache.



Analysis 13.14. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol plus oxytocin	•			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI	
El Tahan 2012	24/179	13/187						100%	1.93[1.01,3.67]	
Total (95% CI)	179	187			•			100%	1.93[1.01,3.67]	
Total events: 24 (Misoprostol plus o	oxytocin), 13 (Oxytocin)									
Heterogeneity: Not applicable										
Test for overall effect: Z=2(P=0.05)										
	Favours Misopros	stol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin		



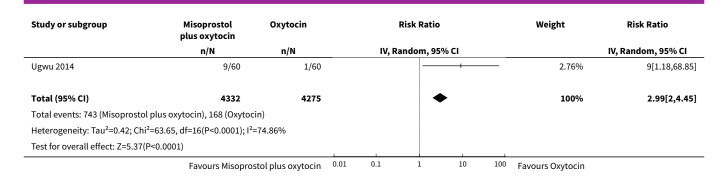
Analysis 13.16. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 16 Shivering.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
Adanikin 2012	11/109	8/109		5.84%	1.38[0.58,3.29]	
Badejoko 2012	34/126	17/129		8.46%	2.05[1.21,3.47]	
Bhullar 2004	2/377	1/379		1.39%	2.01[0.18,22.08]	
Caliskan 2002	52/401	16/407		8.33%	3.3[1.92,5.68]	
Caliskan 2003	49/404	19/384		8.6%	2.45[1.47,4.09]	
Carbonell 2009	390/702	81/698	+	10.8%	4.79[3.86,5.94]	
Cayan 2010	6/120	2/40		2.81%	1[0.21,4.76]	
Chaudhuri 2015	42/198	11/198		7.59%	3.82[2.03,7.2]	
Chaudhuri 2016	39/144	6/144	_ 	6.13%	6.5[2.84,14.88]	
El Tahan 2012	17/179	6/187		5.61%	2.96[1.19,7.34]	
Elsedeek 2012	0/200	0/200			Not estimable	
Fekih 2009	7/125	1/125	 	1.77%	7[0.87,56.06]	
Hofmeyr 2011	142/544	67/556	+	10.5%	2.17[1.66,2.82]	
Hong 2007	1/96	0/118		- 0.83%	3.68[0.15,89.33]	
Lapaire 2006	10/28	2/25		3.23%	4.46[1.08,18.45]	
Pakniat 2015	0/50	0/50			Not estimable	
Quibel 2016	87/806	5/806		5.68%	17.4[7.1,42.63]	
Sood 2012	19/90	8/84	├	6.54%	2.22[1.03,4.79]	
Ugwu 2014	37/60	5/60		5.9%	7.4[3.12,17.53]	
Total (95% CI)	4759	4699	•	100%	3.38[2.5,4.57]	
Total events: 945 (Misoprosto	ol plus oxytocin), 255 (Oxytoc	in)				
Heterogeneity: Tau ² =0.21; Ch	ni ² =52.87, df=16(P<0.0001); l ²	=69.74%				
Test for overall effect: Z=7.93	(P<0.0001)					
	Favours Misopro	stol plus oxytocin 0.01	0.1 1 10 1	00 Favours Oxytocin		

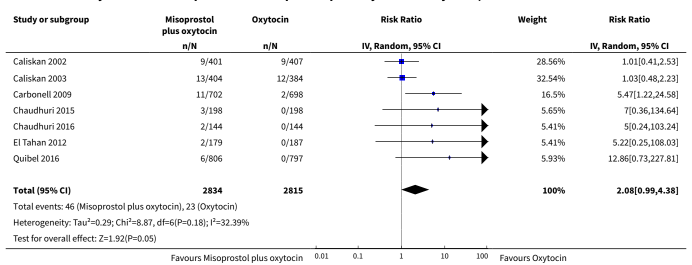
Analysis 13.17. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Adanikin 2012	8/109	6/109		5.95%	1.33[0.48,3.71]
Badejoko 2012	28/126	7/129	_ 	7.07%	4.1[1.86,9.03]
Caliskan 2002	19/401	6/407		6.5%	3.21[1.3,7.96]
Caliskan 2003	16/404	5/384		6.09%	3.04[1.13,8.22]
Carbonell 2009	272/702	23/698		8.86%	11.76[7.78,17.76]
Cayan 2010	6/120	1/40		2.66%	2[0.25,16.11]
Chaudhuri 2015	13/198	4/198		5.6%	3.25[1.08,9.79]
Chaudhuri 2016	8/144	2/144	 	4.01%	4[0.86,18.51]
El Tahan 2012	16/179	8/187	 • -	6.91%	2.09[0.92,4.76]
Elsedeek 2012	11/200	13/200		7.13%	0.85[0.39,1.84]
Fekih 2009	9/125	2/125		4.07%	4.5[0.99,20.41]
Hofmeyr 2011	61/522	28/536	-	8.78%	2.24[1.45,3.44]
Hong 2007	10/96	5/118	 	5.88%	2.46[0.87,6.95]
Pakniat 2015	2/50	1/50		2.2%	2[0.19,21.36]
Quibel 2016	245/806	50/806	+	9.31%	4.9[3.67,6.54]
Sood 2012	10/90	6/84	· · · · · · · · · · · · · · · · · · ·	6.21%	1.56[0.59,4.09]
	Favours Misopro	stol plus oxvtocin	0.01 0.1 1 10	100 Favours Oxvtocin	





Analysis 13.18. Comparison 13 Misoprostol plus oxytocin vs Oxytocin, Outcome 18 Diarrhoea.



Comparison 14. Injectable prostaglandins vs Misoprostol

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	170	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
3 Blood transfusion	4	403	Risk Ratio (IV, Random, 95% CI)	0.88 [0.14, 5.39]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	303	Risk Ratio (IV, Random, 95% CI)	1.60 [0.57, 4.52]
7 Additional uterotonics	4	403	Risk Ratio (IV, Random, 95% CI)	1.02 [0.28, 3.69]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
8 Blood loss	3	270	Mean Difference (IV, Random, 95% CI)	52.99 [-39.74, 145.71]
9 Change in haemoglo- bin	2	220	Mean Difference (IV, Random, 95% CI)	2.28 [-1.35, 5.90]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	2	233	Risk Ratio (IV, Random, 95% CI)	0.23 [0.06, 0.92]
12 Vomiting	2	233	Risk Ratio (IV, Random, 95% CI)	1.31 [0.01, 163.15]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	0.99 [0.20, 4.80]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	353	Risk Ratio (IV, Random, 95% CI)	0.04 [0.01, 0.21]
17 Fever	2	233	Risk Ratio (IV, Random, 95% CI)	0.08 [0.01, 0.39]
18 Diarrhoea	2	233	Risk Ratio (IV, Random, 95% CI)	9.03 [1.70, 48.00]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

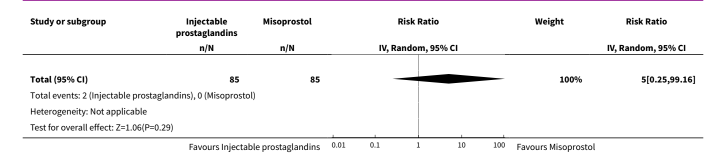
Analysis 14.1. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Misoprostol			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Supe 2016	0/50	0/50							Not estimable
Total (95% CI)	50	50							Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Misoprosto	ol)							
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	le					1			
	Favours Injectal	ble prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	

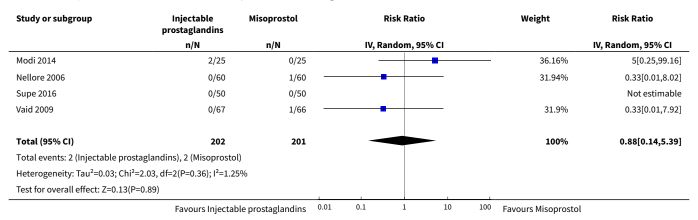
Analysis 14.2. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio					Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95	% CI			IV, Random, 95% CI
Modi 2014	2/25	0/25		_		-		100%	5[0.25,99.16]
Nellore 2006	0/60	0/60							Not estimable
	Favours Injectabl	le prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	





Analysis 14.3. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 3 Blood transfusion.



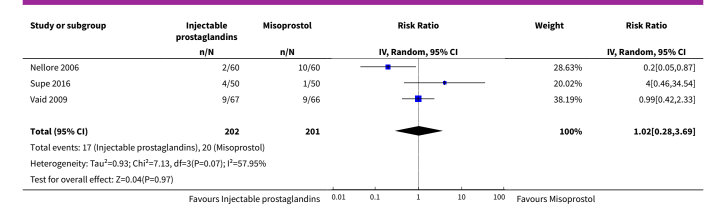
Analysis 14.6. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	•			Risk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, F	andom, 95%	6 CI			IV, Random, 95% CI
Modi 2014	6/25	0/25			+	+	\rightarrow	11.65%	13[0.77,219.11]
Nellore 2006	3/60	4/60		_				31.99%	0.75[0.18,3.21]
Vaid 2009	13/67	8/66			+			56.36%	1.6[0.71,3.61]
Total (95% CI)	152	151				-		100%	1.6[0.57,4.52]
Total events: 22 (Injectable p	prostaglandins), 12 (Misopros	tol)							
Heterogeneity: Tau ² =0.32; Cl	hi²=3.14, df=2(P=0.21); l²=36.	31%			İ				
Test for overall effect: Z=0.89	9(P=0.37)								
	Favours Injectal	le prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	

Analysis 14.7. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio				Weight	Risk Ratio
	n/N	n/N	IN	/, Random, 9	95% CI			IV, Random, 95% CI
Modi 2014	2/25	0/25		-	+ ,		13.17%	5[0.25,99.16]
	Favours Injectab	le prostaglandins	0.01 0.1	1	10	100	Favours Misoprostol	





Analysis 14.8. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 8 Blood loss.

Study or subgroup		Injectable prostaglandins		oprostol	Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Rando	m, 95% CI			Random, 95% CI
Modi 2014	25	435 (147.6)	25	255.8 (102.2)			•	30.72%	179.2[108.84,249.56]
Nellore 2006	60	205 (175)	60	245 (158)				32.33%	-40[-99.66,19.66]
Supe 2016	50	153.8 (43.5)	50	124.4 (34.7)		-		36.95%	29.4[13.98,44.82]
Total ***	135		135					100%	52.99[-39.74,145.71]
Heterogeneity: Tau ² =5996.41	; Chi ² =22.52, df=	2(P<0.0001); I ² =9	91.12%						
Test for overall effect: Z=1.12	(P=0.26)								
		Favours Ini	ectable p	rostaglandins	-100 -50	0 50	100	Favours Mis	oprostol

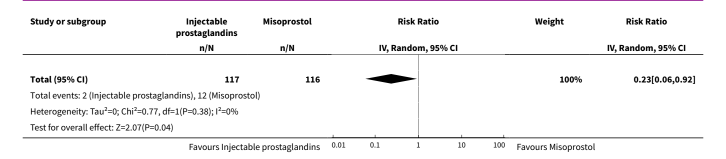
Analysis 14.9. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 9 Change in haemoglobin.

Study or subgroup		ectable aglandins	Mis	oprostol		Ме	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Raı	ndom, 95% CI			Random, 95% CI
Nellore 2006	60	6.2 (2.8)	60	5.8 (2.3)					49.32%	0.4[-0.52,1.32]
Supe 2016	50	9 (1.2)	50	4.9 (0.5)			•		50.68%	4.1[3.74,4.46]
Total ***	110		110				*		100%	2.28[-1.35,5.9]
Heterogeneity: Tau ² =6.72; Ch	ii ² =54.19, df=1(P	<0.0001); I ² =98.1	5%							
Test for overall effect: Z=1.23	(P=0.22)									
		Favours Inj	ectable p	rostaglandins	-100	-50	0 50) 100	Favours Mis	oprostol

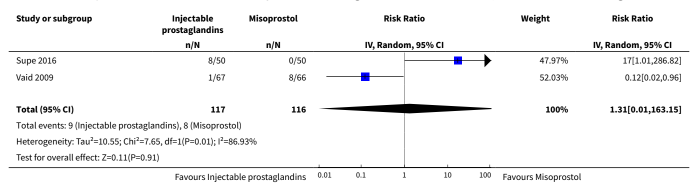
Analysis 14.11. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 11 Nausea.

Study or subgroup	Injectable Misoprostol prostaglandins				Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Supe 2016	0/50	6/50	-	-				23.14%	0.08[0,1.33]
Vaid 2009	2/67	6/66			-	1		76.86%	0.33[0.07,1.57]
	Favours Injectab	le prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	

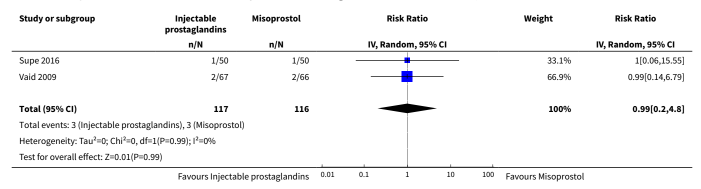




Analysis 14.12. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 12 Vomiting.



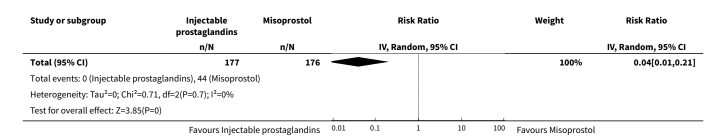
Analysis 14.14. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 14 Abdominal pain.



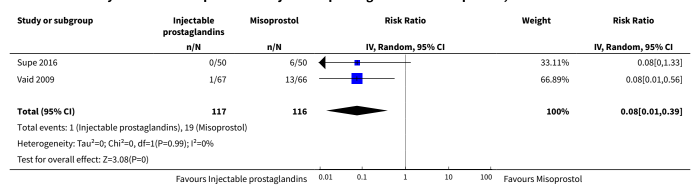
Analysis 14.16. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 16 Shivering.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio					Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Nellore 2006	0/60	5/60	+	-				32.08%	0.09[0.01,1.61]
Supe 2016	0/50	10/50	\leftarrow	•				33.53%	0.05[0,0.79]
Vaid 2009	0/67	29/66	+					34.39%	0.02[0,0.27]
						1			
	Favours Injectab	le prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	

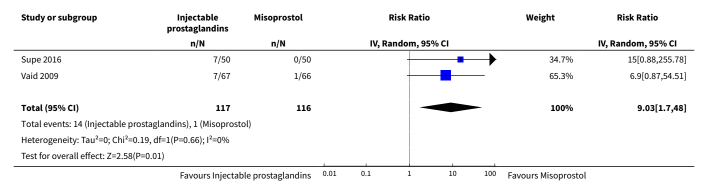




Analysis 14.17. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 17 Fever.



Analysis 14.18. Comparison 14 Injectable prostaglandins vs Misoprostol, Outcome 18 Diarrhoea.



Comparison 15. Misoprostol vs Carbetocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	177	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [0.62, 8.64]
3 Blood transfusion	1	177	Risk Ratio (IV, Random, 95% CI)	2.97 [0.12, 71.85]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [1.52, 3.50]
7 Additional uterotonics	1	177	Risk Ratio (IV, Random, 95% CI)	3.96 [1.55, 10.07]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	177	Risk Ratio (IV, Random, 95% CI)	0.99 [0.47, 2.08]
12 Vomiting	1	177	Risk Ratio (IV, Random, 95% CI)	1.48 [0.55, 3.99]
13 Headache	1	177	Risk Ratio (IV, Random, 95% CI)	1.19 [0.71, 1.99]
14 Abdominal pain	1	177	Risk Ratio (IV, Random, 95% CI)	0.65 [0.52, 0.80]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	177	Risk Ratio (IV, Random, 95% CI)	3.46 [1.67, 7.17]
17 Fever	1	177	Risk Ratio (IV, Random, 95% CI)	2.55 [1.41, 4.64]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 15.1. Comparison 15 Misoprostol vs Carbetocin, Outcome 1 Death.

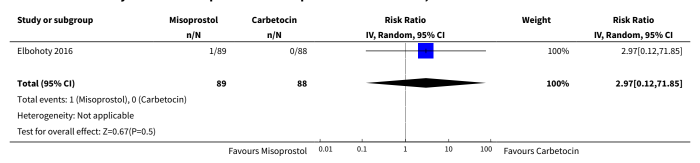
Study or subgroup	Misoprostol	Carbetocin			Risk Ratio			Weight Risk Ra	
	n/N	n/N	IV, Random, 95% CI			% CI			IV, Random, 95% CI
Elbohoty 2016	0/89	0/88							Not estimable
Total (95% CI)	89	88							Not estimable
Total events: 0 (Misoprostol),	0 (Carbetocin)								
Heterogeneity: Not applicable	e								
Test for overall effect: Not app	olicable								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	



Analysis 15.2. Comparison 15 Misoprostol vs Carbetocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Carbetocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Ra	andom, 95	% CI			IV, Random, 95% CI
Elbohoty 2016	7/89	3/88			+	_		100%	2.31[0.62,8.64]
Total (95% CI)	89	88				-		100%	2.31[0.62,8.64]
Total events: 7 (Misoprostol), 3 (Carb	oetocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.24(P=0.21)								
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 15.3. Comparison 15 Misoprostol vs Carbetocin, Outcome 3 Blood transfusion.



Analysis 15.6. Comparison 15 Misoprostol vs Carbetocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Carbetocin			Risk R	atio		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI						IV, Random, 95% CI	
Elbohoty 2016	49/89	21/88		-				100%	2.31[1.52,3.5]	
Total (95% CI)	89	88				•		100%	2.31[1.52,3.5]	
Total events: 49 (Misoprostol)), 21 (Carbetocin)									
Heterogeneity: Not applicable	e									
Test for overall effect: Z=3.92	(P<0.0001)					1	1			
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin		

Analysis 15.7. Comparison 15 Misoprostol vs Carbetocin, Outcome 7 Additional uterotonics.

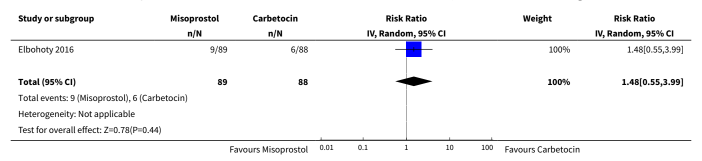
Study or subgroup	Misoprostol	Carbetocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
Elbohoty 2016	20/89	5/88					100%	3.96[1.55,10.07]
Total (95% CI)	89	88			•		100%	3.96[1.55,10.07]
Total events: 20 (Misoprostol), 5 (Ca	rbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Z=2.88(P=0)			1		1			
	Fav	ours Misoprostol	0.01).1	. 10	100	Favours Carbetocin	



Analysis 15.11. Comparison 15 Misoprostol vs Carbetocin, Outcome 11 Nausea.

Study or subgroup	Misoprostol	Carbetocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
Elbohoty 2016	12/89	12/88			-			100%	0.99[0.47,2.08]
Total (95% CI)	89	88			•			100%	0.99[0.47,2.08]
Total events: 12 (Misoprostol)), 12 (Carbetocin)								
Heterogeneity: Not applicable	e								
Test for overall effect: Z=0.03((P=0.98)								
	Far	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 15.12. Comparison 15 Misoprostol vs Carbetocin, Outcome 12 Vomiting.



Analysis 15.13. Comparison 15 Misoprostol vs Carbetocin, Outcome 13 Headache.

Study or subgroup	Misoprostol	Carbetocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
Elbohoty 2016	24/89	20/88			-			100%	1.19[0.71,1.99]
Total (95% CI)	89	88						100%	1.19[0.71,1.99]
Total events: 24 (Misoprostol), 20 (Carbetocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.65(P=0.5	2)					1			
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	

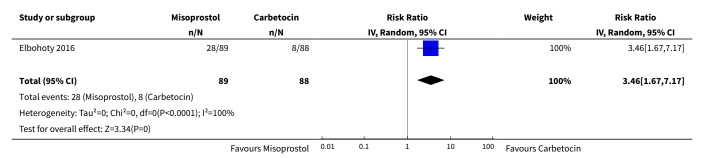
Analysis 15.14. Comparison 15 Misoprostol vs Carbetocin, Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Carbetocin		Risk Ratio IV, Random, 95% CI				Weight	Risk Ratio
	n/N	n/N							IV, Random, 95% CI
Elbohoty 2016	47/89	72/88			+			100%	0.65[0.52,0.8]
Total (95% CI)	89	88			•			100%	0.65[0.52,0.8]
Total events: 47 (Misoprostol),	72 (Carbetocin)								
Heterogeneity: Not applicable									
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	



Study or subgroup	Misoprostol n/N	Carbetocin n/N		Risk Ratio IV, Random, 95% CI				Weight	Risk Ratio IV, Random, 95% CI
Test for overall effect: Z=3.91(P<0.00	01)		_			1			
		Favours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 15.16. Comparison 15 Misoprostol vs Carbetocin, Outcome 16 Shivering.



Analysis 15.17. Comparison 15 Misoprostol vs Carbetocin, Outcome 17 Fever.

Study or subgroup	Misoprostol	Carbetocin			Risk Ratio			Weight Risk Rati		
	n/N	n/N		IV, R	andom, 95º	6 CI			IV, Random, 95% CI	
Elbohoty 2016	31/89	12/88			-	_		100%	2.55[1.41,4.64]	
Total (95% CI)	89	88			•	•		100%	2.55[1.41,4.64]	
Total events: 31 (Misoprostol), 3	12 (Carbetocin)									
Heterogeneity: Not applicable										
Test for overall effect: Z=3.08(P	=0)									
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin		

Comparison 16. Ergometrine vs Misoprostol

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	7	5254	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	11	5562	Risk Ratio (IV, Random, 95% CI)	1.25 [0.45, 3.48]
3 Blood transfusion	13	5308	Risk Ratio (IV, Random, 95% CI)	1.71 [0.65, 4.50]
4 Severe maternal morbidity: intensive care admissions	1	864	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.16]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	16	6459	Risk Ratio (IV, Random, 95% CI)	1.18 [0.79, 1.75]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7 Additional uterotonics	16	6565	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.60]
8 Blood loss	15	6337	Mean Difference (IV, Random, 95% CI)	11.66 [-9.85, 33.17]
9 Change in haemoglo- bin	8	4438	Mean Difference (IV, Random, 95% CI)	0.91 [-0.43, 2.26]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	13	5202	Risk Ratio (IV, Random, 95% CI)	1.53 [1.15, 2.04]
12 Vomiting	14	6136	Risk Ratio (IV, Random, 95% CI)	1.22 [0.81, 1.83]
13 Headache	8	3369	Risk Ratio (IV, Random, 95% CI)	1.70 [0.70, 4.12]
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	1.58 [0.04, 65.80]
15 Hypertension	2	300	Risk Ratio (IV, Random, 95% CI)	7.55 [0.94, 60.53]
16 Shivering	17	6860	Risk Ratio (IV, Random, 95% CI)	0.34 [0.26, 0.44]
17 Fever	14	6330	Risk Ratio (IV, Random, 95% CI)	0.24 [0.17, 0.33]
18 Diarrhoea	8	4165	Risk Ratio (IV, Random, 95% CI)	1.04 [0.46, 2.34]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 16.1. Comparison 16 Ergometrine vs Misoprostol, Outcome 1 Death.

Study or subgroup	Ergometrine	Misoprostol	Ris	sk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Ran	dom, 95% CI		IV, Random, 95% CI
Chandhiok 2006	0/600	0/600				Not estimable
Chhabra 2008	0/100	0/200				Not estimable
Enakpene 2007	0/432	0/432				Not estimable
Fawzy 2012	0/100	0/200				Not estimable
Soltan 2007	0/266	0/818				Not estimable
Supe 2016	0/50	0/50				Not estimable
Zachariah 2006	0/676	0/730				Not estimable
Total (95% CI)	2224	3030				Not estimable
Total events: 0 (Ergometrine), 0 (Misc	prostol)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable					1	
	Fav	ours Ergometrine	0.01 0.1	1 10	100 Favours Misoprostol	



Analysis 16.2. Comparison 16 Ergometrine vs Misoprostol, Outcome 2 PPH >= 1000 mL.

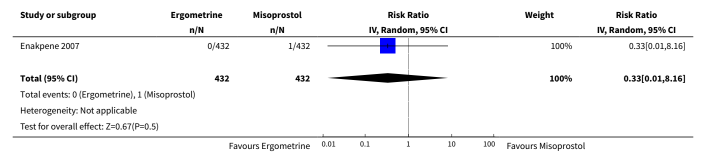
Study or subgroup	Ergometrine	Misoprostol	Ris	sk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Ran	dom, 95% CI		IV, Random, 95% CI
Amant 1999	0/93	1/96			9.75%	0.34[0.01,8.34]
Chandhiok 2006	0/600	1/600			9.69%	0.33[0.01,8.17]
Chhabra 2008	0/100	0/200				Not estimable
Enakpene 2007	1/432	3/432			18.3%	0.33[0.03,3.19]
Humera 2016	0/50	0/50				Not estimable
Modi 2014	0/25	0/25				Not estimable
Patil 2013	0/99	1/100			9.75%	0.34[0.01,8.17]
Penaranda 2002	3/25	1/25	_	+	19.27%	3[0.33,26.92]
Soltan 2007	1/266	1/818		+	— 12.68%	3.08[0.19,49]
Vimala 2004	0/60	0/60				Not estimable
Zachariah 2006	6/676	1/730		+		6.48[0.78,53.68]
Total (95% CI)	2426	3136	-	•	100%	1.25[0.45,3.48]
Total events: 11 (Ergometrine), 9 (Misoprostol)					
Heterogeneity: Tau ² =0.17; Chi	² =6.59, df=6(P=0.36); I ² =8.9	9%				
Test for overall effect: Z=0.42(P=0.67)					
	Fav	ours Ergometrine	0.01 0.1	1 10	100 Favours Misoprostol	

Analysis 16.3. Comparison 16 Ergometrine vs Misoprostol, Outcome 3 Blood transfusion.

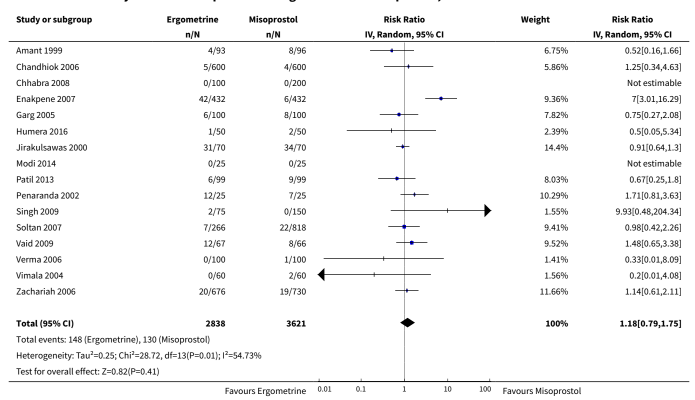
Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Amant 1999	1/100	1/100		12.34%	1[0.06,15.77]
Chandhiok 2006	0/600	1/600		9.17%	0.33[0.01,8.17]
Chhabra 2008	0/100	0/200			Not estimable
Humera 2016	0/50	0/50			Not estimable
Modi 2014	0/25	0/25			Not estimable
Patil 2013	0/99	1/100	•	9.23%	0.34[0.01,8.17]
Ray 2001	3/100	1/100	-	18.6%	3[0.32,28.35]
Singh 2009	3/75	0/150	+	10.78%	13.91[0.73,265.81]
Soltan 2007	1/266	1/809	+		3.04[0.19,48.46]
Supe 2016	0/50	0/50			Not estimable
Vaid 2009	0/67	1/66	+	9.26%	0.33[0.01,7.92]
Vimala 2004	0/60	0/60			Not estimable
Zachariah 2006	3/676	1/730	-	18.36%	3.24[0.34,31.07]
Total (95% CI)	2268	3040	•	100%	1.71[0.65,4.5]
Total events: 11 (Ergometrine), 7	(Misoprostol)				
Heterogeneity: Tau ² =0; Chi ² =5.83	, df=7(P=0.56); I ² =0%				
Test for overall effect: Z=1.08(P=0	.28)				
	Far	vours Frgometrine	0.01 0.1 1 10	100 Favours Misoprostol	



Analysis 16.4. Comparison 16 Ergometrine vs Misoprostol, Outcome 4 Severe maternal morbidity: intensive care admissions.



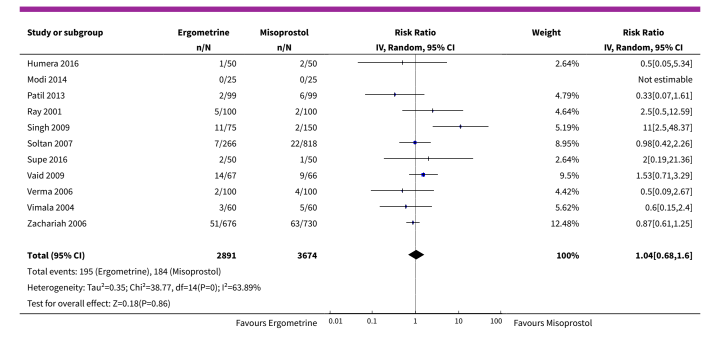
Analysis 16.6. Comparison 16 Ergometrine vs Misoprostol, Outcome 6 PPH >= 500 mL.



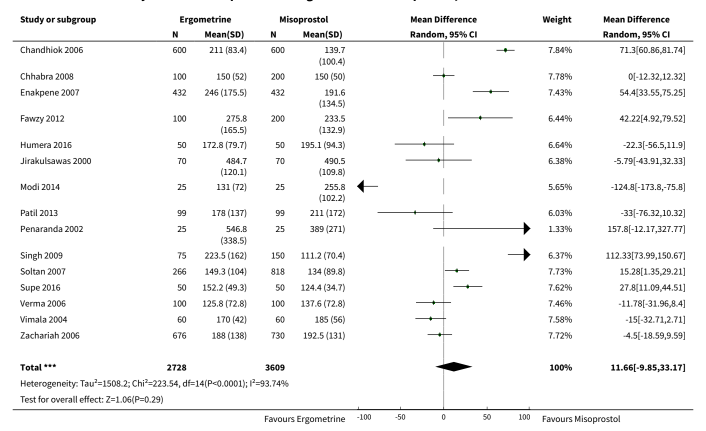
Analysis 16.7. Comparison 16 Ergometrine vs Misoprostol, Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine	Misoprostol		Risk Ratio			Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI				IV, Random, 95% CI	
Amant 1999	4/91	12/94	-	+			7.22%	0.34[0.12,1.03]
Chandhiok 2006	3/600	4/600					5.14%	0.75[0.17,3.34]
Chhabra 2008	3/100	9/200					6.13%	0.67[0.18,2.41]
Enakpene 2007	80/432	33/432		-			12.3%	2.42[1.65,3.56]
Garg 2005	7/100	10/100		-+-	1	1	8.34%	0.7[0.28,1.77]
	Fav	ours Ergometrine	0.01 0.1	1	10	100	Favours Misoprostol	





Analysis 16.8. Comparison 16 Ergometrine vs Misoprostol, Outcome 8 Blood loss.





Analysis 16.9. Comparison 16 Ergometrine vs Misoprostol, Outcome 9 Change in haemoglobin.

Study or subgroup	Erge	ometrine	Mis	oprostol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Chhabra 2008	100	8 (6.3)	200	7.8 (2)	•	11.87%	0.2[-1.07,1.47]
Enakpene 2007	432	4.1 (5.1)	432	1.4 (2)	•	13%	2.72[2.2,3.24]
Gore 2017	182	9.4 (0.7)	182	9.7 (0.8)	•	13.24%	-0.3[-0.45,-0.15]
Soltan 2007	266	4.6 (4)	818	3.6 (2.9)	•	13%	1[0.48,1.52]
Supe 2016	50	7.6 (0.2)	50	4.9 (0.5)	•	13.24%	2.7[2.55,2.85]
Verma 2006	100	2.5 (3)	100	3.1 (3)	•	12.62%	-0.6[-1.43,0.23]
Vimala 2004	60	8 (6.3)	60	7.6 (2)		11.01%	0.4[-1.27,2.07]
Zachariah 2006	676	14.8 (11.9)	730	13.8 (10.7)	į	12.02%	1[-0.19,2.19]
Total ***	1866		2572			100%	0.91[-0.43,2.26]
Heterogeneity: Tau ² =3.57; Cl	hi²=816.27, df=7(I	P<0.0001); I ² =99	.14%				
Test for overall effect: Z=1.33	3(P=0.18)						
			Favours	Ergometrine -100) -50 0 50	100 Favours Mis	oprostol

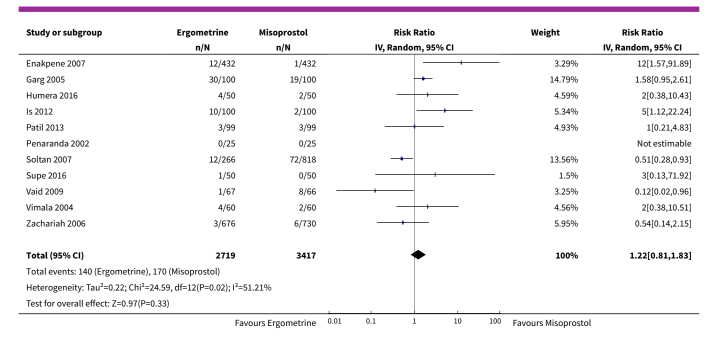
Analysis 16.11. Comparison 16 Ergometrine vs Misoprostol, Outcome 11 Nausea.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Amant 1999	30/94	20/87	+-	15.1%	1.39[0.85,2.25]
Chandhiok 2006	121/600	61/600		20.94%	1.98[1.49,2.64]
Chhabra 2008	16/100	24/200	+	12.63%	1.33[0.74,2.39]
Enakpene 2007	16/432	10/432		9.01%	1.6[0.73,3.49]
Garg 2005	30/100	20/100	 • -	14.88%	1.5[0.92,2.46]
Humera 2016	9/50	3/50	+	4.44%	3[0.86,10.43]
Is 2012	12/100	1/100		1.89%	12[1.59,90.56]
Patil 2013	8/99	4/99		4.95%	2[0.62,6.43]
Supe 2016	1/50	6/50	+	1.79%	0.17[0.02,1.33]
Vaid 2009	1/67	6/66		1.77%	0.16[0.02,1.33]
Verma 2006	2/100	4/100		2.66%	0.5[0.09,2.67]
Vimala 2004	8/60	4/60		5.1%	2[0.64,6.29]
Zachariah 2006	6/676	5/730		4.85%	1.3[0.4,4.23]
Total (95% CI)	2528	2674	•	100%	1.53[1.15,2.04]
Total events: 260 (Ergometrine), 1	68 (Misoprostol)				
Heterogeneity: Tau ² =0.08; Chi ² =19	, df=12(P=0.09); I ² =36.8	33%			
Test for overall effect: Z=2.91(P=0)					

Analysis 16.12. Comparison 16 Ergometrine vs Misoprostol, Outcome 12 Vomiting.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Amant 1999	18/94	13/87			+			12.83%	1.28[0.67,2.46]
Chandhiok 2006	34/600	30/600			+			15.13%	1.13[0.7,1.83]
Chhabra 2008	8/100	12/200			+	1	1	10.28%	1.33[0.56,3.16]
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	





Analysis 16.13. Comparison 16 Ergometrine vs Misoprostol, Outcome 13 Headache.

Study or subgroup	Ergometrine	Misoprostol	Ris	sk Ratio		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI				IV, Random, 95% CI	
Amant 1999	12/94	10/87	-	-	,	22.73%	1.11[0.51,2.44]	
Chhabra 2008	8/100	14/200	-	-		22.23%	1.14[0.5,2.63]	
Enakpene 2007	54/432	1/432				11.67%	54[7.5,388.59]	
Garg 2005	3/100	4/100		•		15.71%	0.75[0.17,3.27]	
Humera 2016	0/50	0/50					Not estimable	
Patil 2013	0/99	0/99					Not estimable	
Vimala 2004	4/60	3/60				15.87%	1.33[0.31,5.7]	
Zachariah 2006	2/676	2/730		 		11.78%	1.08[0.15,7.64]	
Total (95% CI)	1611	1758		•		100%	1.7[0.7,4.12]	
Total events: 83 (Ergometrine), 34	4 (Misoprostol)							
Heterogeneity: Tau ² =0.74; Chi ² =1	.4.47, df=5(P=0.01); I ² =65	.45%						
Test for overall effect: Z=1.17(P=0	0.24)							
	Fav	ours Ergometrine	0.01 0.1	1 10	100	Favours Misoprostol		

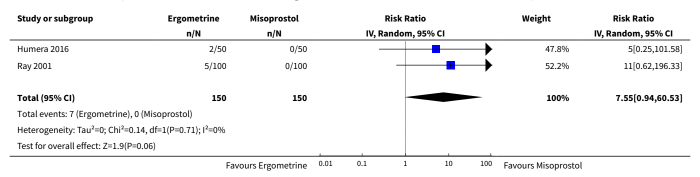
Analysis 16.14. Comparison 16 Ergometrine vs Misoprostol, Outcome 14 Abdominal pain.

Study or subgroup	Ergometrine	Misoprostol		Risk Ratio			Weight	Risk Ratio	
	n/N n/N		IV, Ra	ndom, 9	5% CI			IV, Random, 95% CI	
Supe 2016	9/50	1/50			-	1		54.45%	9[1.18,68.42]
Vaid 2009	0/67	2/66	←	-		_		45.55%	0.2[0.01,4.03]
Total (95% CI)	117	116						100%	1.58[0.04,65.8]
Total events: 9 (Ergometrine)	, 3 (Misoprostol)								
Heterogeneity: Tau ² =5.58; Ch	i ² =4.24, df=1(P=0.04); l ² =76.4	13%							
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	

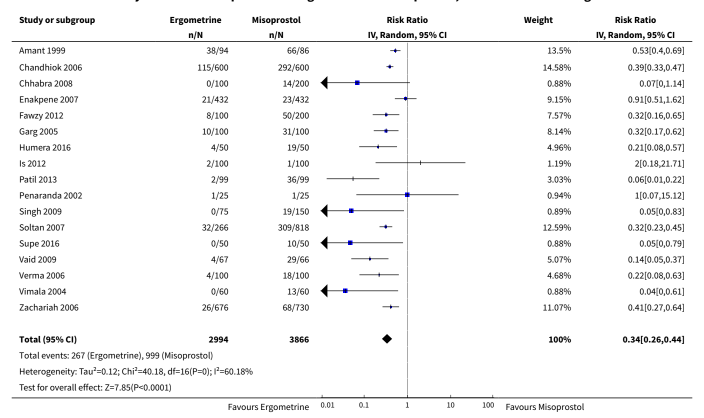


Study or subgroup	Ergometrine n/N	Misoprostol n/N			Risk Ratio			Weight	Risk Ratio IV, Random, 95% CI
Test for overall effect: Z=0.24(P=0.81)						1			
		Favours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	

Analysis 16.15. Comparison 16 Ergometrine vs Misoprostol, Outcome 15 Hypertension.



Analysis 16.16. Comparison 16 Ergometrine vs Misoprostol, Outcome 16 Shivering.





Analysis 16.17. Comparison 16 Ergometrine vs Misoprostol, Outcome 17 Fever.

n/N 3/100 26/600 0/100 7/432 7/100 0/50 0/99 0/75	n/N 34/100 58/600 10/200 31/432 29/100 2/50 2/99	IV, Random, 95% CI	6.89% 26.22% 1.27% 12.25% 13.02% 1.13%	1V, Random, 95% CI 0.09[0.03,0.28] 0.45[0.29,0.7] 0.09[0.01,1.6] 0.23[0.1,0.51] 0.24[0.11,0.53] 0.2[0.01,4.06]
26/600 0/100 7/432 7/100 0/50 0/99	58/600 10/200 31/432 29/100 2/50 2/99		26.22% 1.27% 12.25% 13.02% 1.13%	0.45[0.29,0.7] 0.09[0.01,1.6] 0.23[0.1,0.51] 0.24[0.11,0.53]
0/100 7/432 7/100 0/50 0/99	10/200 31/432 29/100 2/50 2/99		1.27% 12.25% 13.02% 1.13%	0.09[0.01,1.6] 0.23[0.1,0.51] 0.24[0.11,0.53]
7/432 7/100 0/50 0/99	31/432 29/100 2/50 2/99		12.25% 13.02% 1.13%	0.23[0.1,0.51] 0.24[0.11,0.53]
7/100 0/50 0/99	29/100 2/50 2/99		13.02% 1.13%	0.24[0.11,0.53]
0/50 0/99	2/50 2/99		1.13%	
0/99	2/99			0.2[0.01,4.06]
,	•	4	1 100/	
0/75			1.12%	0.2[0.01,4.11]
0/13	25/150	←	1.31%	0.04[0,0.63]
6/266	95/818		12.16%	0.19[0.09,0.44]
0/50	6/50		1.25%	0.08[0,1.33]
0/67	13/66	←	1.3%	0.04[0,0.6]
1/100	6/100		2.27%	0.17[0.02,1.36]
0/60	4/60		1.21%	0.11[0.01,2.02]
13/676	48/730		18.59%	0.29[0.16,0.53]
2775	3555	•	100%	0.24[0.17,0.33]
prostol)				
:13(P=0.29); I ² =1	4.72%			
	0/50 0/67 1/100 0/60 13/676 2775 prostol) :13(P=0.29); l ² =1	6/266 95/818 0/50 6/50 0/67 13/66 1/100 6/100 0/60 4/60 13/676 48/730	6/266 95/818	6/266 95/818 12.16% 0/50 6/50 0/67 13/66 1.3% 1/100 6/100 0/60 4/60 1.3676 48/730 18.59% 2775 3555

Analysis 16.18. Comparison 16 Ergometrine vs Misoprostol, Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine	Misoprostol		Risk Rat	io		Weight	Risk Ratio
	n/N	n/N		IV, Random,	95% CI			IV, Random, 95% CI
Amant 1999	0/94	1/86		+ +			6.44%	0.31[0.01,7.39]
Enakpene 2007	0/432	0/432						Not estimable
Garg 2005	3/100	3/100					26.34%	1[0.21,4.84]
Patil 2013	0/99	0/99						Not estimable
Soltan 2007	4/266	10/818		-			49.38%	1.23[0.39,3.89]
Supe 2016	0/50	0/50						Not estimable
Vaid 2009	0/67	1/66		++			6.46%	0.33[0.01,7.92]
Zachariah 2006	2/676	1/730		-	+		11.38%	2.16[0.2,23.76]
Total (95% CI)	1784	2381		•	•		100%	1.04[0.46,2.34]
Total events: 9 (Ergometrine),	16 (Misoprostol)							
Heterogeneity: Tau ² =0; Chi ² =1	.51, df=4(P=0.82); I ² =0%							
Test for overall effect: Z=0.1(P=	=0.92)				1			
	Fav	ours Ergometrine	0.01	0.1 1	10	100	Favours Misoprostol	

Comparison 17. Misoprostol vs Ergometrine plus oxytocin

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3258	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

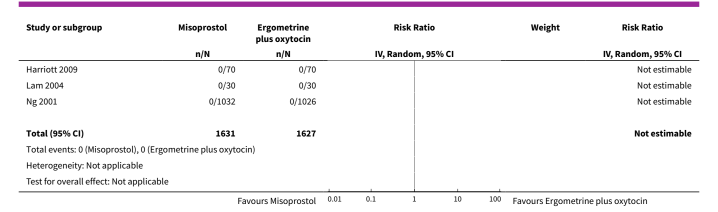


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
2 PPH >= 1000 mL	9	6028	Risk Ratio (IV, Random, 95% CI)	1.61 [1.08, 2.40]
3 Blood transfusion	8	6335	Risk Ratio (IV, Random, 95% CI)	1.41 [0.91, 2.21]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	
6 PPH >= 500 mL	10	6492	492 Risk Ratio (IV, Random, 95% CI)	
7 Additional uterotonics	9	6395	Risk Ratio (IV, Random, 95% CI)	
8 Blood loss	8	5634	4 Mean Difference (IV, Random, 95% CI)	
9 Change in haemoglobin	9	5588	Mean Difference (IV, Random, 95% CI)	1.09 [-0.49, 2.67]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3399	Risk Ratio (IV, Random, 95% CI)	0.71 [0.61, 0.84]
12 Vomiting	6	4983	Risk Ratio (IV, Random, 95% CI)	0.72 [0.50, 1.04]
13 Headache	3	3259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.47, 1.76]
14 Abdominal pain	1	846	Risk Ratio (IV, Random, 95% CI)	0.90 [0.79, 1.02]
15 Hypertension	3	2553	Risk Ratio (IV, Random, 95% CI)	0.50 [0.18, 1.41]
16 Shivering	6	4980	Risk Ratio (IV, Random, 95% CI)	2.71 [1.95, 3.76]
17 Fever	7	5045	Risk Ratio (IV, Random, 95% CI)	5.33 [2.87, 9.88]
18 Diarrhoea	6	3659	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.57]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	1		Mean Difference (IV, Random, 95% CI)	Totals not selected
20.1 Woman's satisfaction using an eight item Client Satisfaction Questionnaire	1		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

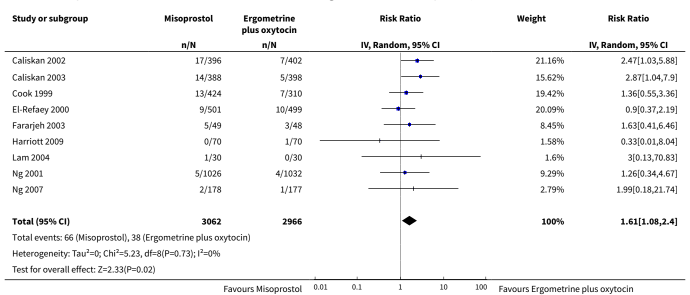
Analysis 17.1. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 1 Death.

Study or subgroup	Misoprostol	Ergometrine Risk Ratio plus oxytocin			Weight Risk Ratio			
	n/N	n/N		IV, R	andom, 95	% CI		IV, Random, 95% CI
El-Refaey 2000	0/499	0/501						Not estimable
	Fav	0.01	0.1	1	10	100	Favours Ergometrine plus oxytocin	





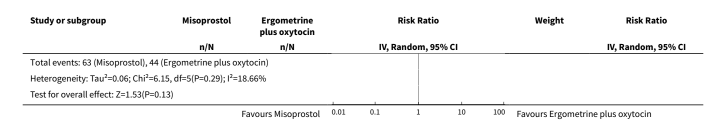
Analysis 17.2. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 2 PPH >= 1000 mL.



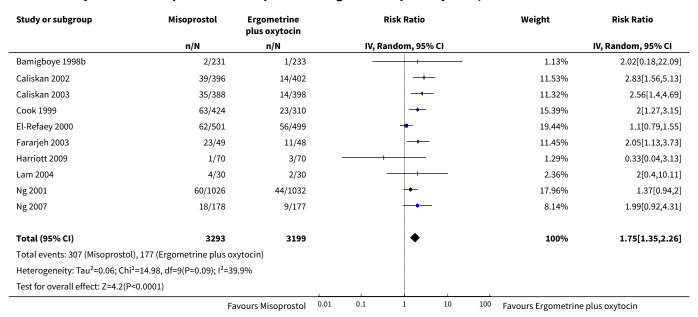
Analysis 17.3. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Ris	Risk Ratio		Weight	Risk Ratio	
	n/N	n/N	IV, Ran	dom, 95% CI			IV, Random, 95% CI	
Bamigboye 1998b	0/231	0/233					Not estimable	
Caliskan 2002	12/396	4/402		+		13.34%	3.05[0.99,9.36]	
Caliskan 2003	14/388	6/398		 • 		17.71%	2.39[0.93,6.16]	
Cook 1999	5/424	3/310	_			8.79%	1.22[0.29,5.06]	
El-Refaey 2000	9/501	11/499	_	-		20.13%	0.81[0.34,1.95]	
Harriott 2009	0/70	0/70					Not estimable	
Ng 2001	15/1026	16/1032	-	-		27.83%	0.94[0.47,1.9]	
Ng 2007	8/178	4/177		+-		12.21%	1.99[0.61,6.49]	
Total (95% CI)	3214	3121	1	•		100%	1.41[0.91,2.21]	
	Fa	avours Misoprostol	0.01 0.1	1 10	100	Favours Ergometrine	e plus oxytocin	





Analysis 17.6. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 6 PPH >= 500 mL.



Analysis 17.7. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Bamigboye 1998b	4/231	1/233		1.09%	4.03[0.45,35.83]
Caliskan 2002	33/396	9/402	_ -	7.89%	3.72[1.8,7.68]
Caliskan 2003	23/388	9/398	_ 	7.35%	2.62[1.23,5.59]
Cook 1999	95/424	28/310		17.13%	2.48[1.67,3.68]
El-Refaey 2000	68/501	50/499	+-	19.5%	1.35[0.96,1.91]
Harriott 2009	6/70	6/70		4.05%	1[0.34,2.95]
Lam 2004	3/30	0/30	+	0.62%	7[0.38,129.93]
Ng 2001	232/1026	144/1032	-	27.76%	1.62[1.34,1.96]
Ng 2007	41/178	24/177	-	14.6%	1.7[1.07,2.69]
Total (95% CI)	3244	3151	•	100%	1.87[1.49,2.36]
Total events: 505 (Misoprosto	l), 271 (Ergometrine plus o	kytocin)			
Heterogeneity: Tau ² =0.04; Chi	i ² =13.5, df=8(P=0.1); I ² =40.7	4%			
Test for overall effect: Z=5.31(P<0.0001)				
	Fa	avours Misoprostol C	0.01 0.1 1 10 10	Pavours Ergometrin	e plus oxytocin



Analysis 17.8. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 8 Blood loss.

Study or subgroup	Study or subgroup Misop		prostol Ergometrine plus oxytocin		Mea	an Difference	Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)	Rar	ndom, 95% CI		Random, 95% CI	
Bamigboye 1998b	231	187 (92)	233	183 (68)			18.03%	4[-10.73,18.73]	
Caliskan 2003	388	328 (152)	398	296 (168)			15.65%	32[9.61,54.39]	
Cook 1999	424	279 (300.6)	310	255.1 (338.8)		+	8.66%	23.86[-23.48,71.2]	
El-Refaey 2000	501	256 (137)	499	251 (136.8)			17.37%	5[-11.97,21.97]	
Fararjeh 2003	49	588 (360)	48	387.1 (273.4)			1.89%	200.87[73.82,327.92]	
Harriott 2009	70	180.1 (120)	70	197 (177)	-	•	8.1%	-16.9[-67,33.2]	
Ng 2001	1026	296 (160)	1032	254 (157)			18.32%	42[28.3,55.7]	
Ng 2007	178	289 (178)	177	255 (149)		+	11.96%	34[-0.14,68.14]	
Total ***	2867		2767			•	100%	22.86[4.5,41.22]	
Heterogeneity: Tau ² =430.06;	Chi ² =28.94, df=7	(P=0); I ² =75.81%	6			į			
Test for overall effect: Z=2.44	1(P=0.01)								
			Favou	rs Misoprostol	-100 -50	0 50	100 Favours Erg	ometrine plus oxytocin	

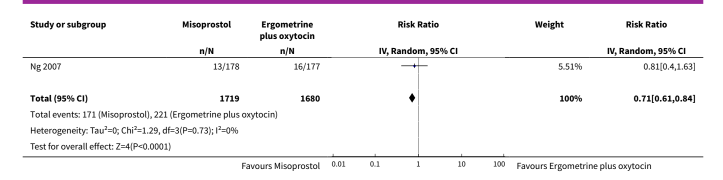
Analysis 17.9. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup	Mis			ometrine oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Bamigboye 1998b	83	2.3 (16)	83	2.8 (19)		5.57%	-0.5[-5.84,4.84]
Caliskan 2002	396	15 (12)	402	15 (12)		13.07%	0[-1.67,1.67]
Caliskan 2003	388	14 (11)	398	15 (12)		13.19%	-1[-2.61,0.61]
Cook 1999	424	6.9 (17.5)	310	0.5 (14.1)		11.58%	6.4[4.11,8.69]
El-Refaey 2000	218	10 (12)	236	10 (16)		10.84%	0[-2.59,2.59]
Fararjeh 2003	49	17.5 (7.3)	48	12.8 (7.3)		10.08%	4.7[1.79,7.61]
Harriott 2009	70	5 (7.7)	70	5.6 (7.2)		11.13%	-0.6[-3.07,1.87]
Ng 2001	1026	13.4 (12.7)	1032	13.4 (12.7)	+	14.23%	0[-1.1,1.1]
Ng 2007	178	17 (14)	177	16 (13)		10.31%	1[-1.81,3.81]
Total ***	2832		2756		•	100%	1.09[-0.49,2.67]
Heterogeneity: Tau ² =4.26; Ch	ni²=39.15, df=8(P	<0.0001); I ² =79.5	57%				
Test for overall effect: Z=1.35	6(P=0.18)						
			Favour	rs Misoprostol -	10 -5 0 5 10	Favours Erg	ometrine plus oxytocin

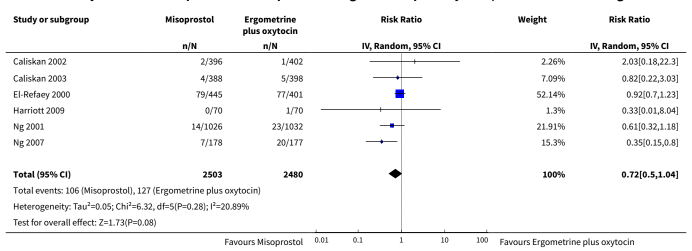
Analysis 17.11. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 11 Nausea.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Rand	om, 95º	% CI			IV, Random, 95% CI
El-Refaey 2000	138/445	175/401		-	F			85.88%	0.71[0.59,0.85]
Harriott 2009	0/70	3/70	\leftarrow	+	+			0.31%	0.14[0.01,2.72]
Ng 2001	20/1026	27/1032		_	+			8.3%	0.75[0.42,1.32]
	Favours Misoprostol			0.1	1	10	100	Favours Ergometrine	e plus oxytocin

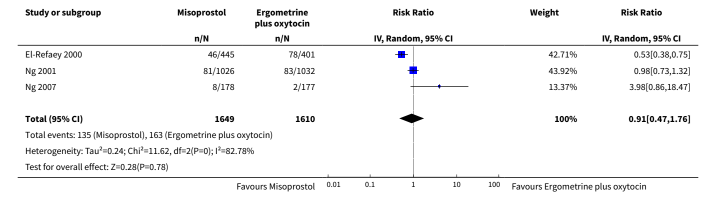




Analysis 17.12. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 12 Vomiting.



Analysis 17.13. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 13 Headache.

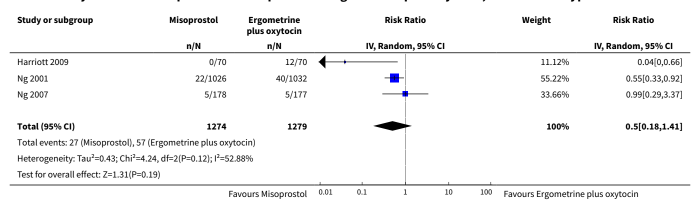




Analysis 17.14. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Misoprostol Ergometrine plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ra	ındom, 95% (CI			IV, Random, 95% CI
El-Refaey 2000	217/445	218/401			+			100%	0.9[0.79,1.02]
Total (95% CI)	445	401			•			100%	0.9[0.79,1.02]
Total events: 217 (Misoprostol), 2	18 (Ergometrine plus ox	(ytocin)							
Heterogeneity: Not applicable									
Test for overall effect: Z=1.63(P=0	.1)								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Ergometrin	e plus oxytocin

Analysis 17.15. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 15 Hypertension.



Analysis 17.16. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 16 Shivering.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI
	n/N	n/N	IV, Random, 95% CI		
Caliskan 2002	47/396	19/402	-	17.06%	2.51[1.5,4.2]
Caliskan 2003	44/388	15/398		15.6%	3.01[1.7,5.32]
El-Refaey 2000	319/445	147/401		27.81%	1.96[1.7,2.25]
Harriott 2009	11/67	6/70	+-	8.67%	1.92[0.75,4.89]
Ng 2001	310/1026	102/1032		26.28%	3.06[2.49,3.76]
Ng 2007	35/178	2/177		4.58%	17.4[4.25,71.25]
Total (95% CI)	2500	2480	•	100%	2.71[1.95,3.76]
Total events: 766 (Misoprosto	ol), 291 (Ergometrine plus o	kytocin)			
Heterogeneity: Tau ² =0.1; Chi ²	² =21.39, df=5(P=0); I ² =76.62 ⁰	%			
Test for overall effect: Z=5.94	(P<0.0001)				
	Fa	avours Misoprostol	0.01 0.1 1 10 10	⁰⁰ Favours Ergometrin	e plus oxytocin



Analysis 17.17. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 17 Fever.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin		Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ındom, 95% (:1			IV, Random, 95% CI	
Caliskan 2002	16/396	6/402			-	-		23.76%	2.71[1.07,6.85]	
Caliskan 2003	17/388	6/398			-	-		23.98%	2.91[1.16,7.29]	
El-Refaey 2000	24/432	1/416			_	•	\rightarrow	8.06%	23.11[3.14,170.06]	
Harriott 2009	0/70	0/70							Not estimable	
Lam 2004	10/30	0/30				+	\rightarrow	4.46%	21[1.29,342.93]	
Ng 2001	87/1026	13/1032			-	-		35.46%	6.73[3.78,11.98]	
Ng 2007	7/178	0/177			+	+	→	4.28%	14.92[0.86,259.21]	
Total (95% CI)	2520	2525			•	>		100%	5.33[2.87,9.88]	
Total events: 161 (Misoprosto	ol), 26 (Ergometrine plus oxy	rtocin)								
Heterogeneity: Tau ² =0.19; Ch	ni ² =7.84, df=5(P=0.17); I ² =36.	19%								
Test for overall effect: Z=5.31	(P<0.0001)					1				
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Ergometrine	e plus oxvtocin	

Analysis 17.18. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% C			IV, Random, 95% CI
Caliskan 2002	11/396	10/402			-		24.42%	1.12[0.48,2.6]
Caliskan 2003	15/388	17/398			-		37.7%	0.91[0.46,1.79]
Cook 1999	1/424	0/310					1.71%	2.2[0.09,53.71]
El-Refaey 2000	17/445	14/401			-		36.17%	1.09[0.55,2.19]
Harriott 2009	0/70	0/70						Not estimable
Ng 2007	0/178	0/177						Not estimable
Total (95% CI)	1901	1758			•		100%	1.04[0.68,1.57]
Total events: 44 (Misoprostol), 41 (Ergometrine plus oxyt	ocin)						
Heterogeneity: Tau ² =0; Chi ² =	0.42, df=3(P=0.94); I ² =0%							
Test for overall effect: Z=0.17	(P=0.87)							
	Fa	vours Misoprostol	0.01	0.1	1	10 100	Favours Ergometrin	e plus oxvtocin

Analysis 17.20. Comparison 17 Misoprostol vs Ergometrine plus oxytocin, Outcome 20 Maternal satisfaction.

Study or subgroup	М	isoprostol		gometrine us oxytocin		Me	an Differe		Mean Difference			
	N	Mean(SD)	N	Mean(SD)	ean(SD) Random, 95% CI				Random, 95% CI			
17.20.1 Woman's satisfaction using an eight item Client Satisfaction Questionna												
Ng 2007	178	178 27 (3.1)		26.4 (2.9)			-	-		0.6[-0.02,1.22]		
			Fa	vours Misoprostol	-5	-2.5	0	2.5	5	Favours Ergometrine		



Comparison 18. Misoprostol vs Misoprostol plus oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.85 [1.03, 3.33]
3 Blood transfusion	2	1589	Risk Ratio (IV, Random, 95% CI)	2.97 [1.40, 6.30]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.93 [0.99, 3.76]
7 Additional uterotonics	3	1689	Risk Ratio (IV, Random, 95% CI)	1.89 [1.26, 2.83]
8 Blood loss	1	792	Mean Difference (IV, Random, 95% CI)	48.0 [24.68, 71.32]
9 Change in haemoglo- bin	3	1689	Mean Difference (IV, Random, 95% CI)	0.27 [-0.42, 0.96]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	100	Risk Ratio (IV, Random, 95% CI)	1.83 [0.73, 4.57]
12 Vomiting	3	1689	Risk Ratio (IV, Random, 95% CI)	1.39 [0.51, 3.82]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	1689	Risk Ratio (IV, Random, 95% CI)	0.94 [0.72, 1.22]
17 Fever	3	1689	Risk Ratio (IV, Random, 95% CI)	0.97 [0.62, 1.53]
18 Diarrhoea	2	1589	Risk Ratio (IV, Random, 95% CI)	1.22 [0.70, 2.13]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Analysis 18.2. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% C	I			IV, Random, 95% CI
Caliskan 2002	17/396	11/401			+			61.69%	1.56[0.74,3.3]
Caliskan 2003	14/388	6/404			-			38.31%	2.43[0.94,6.26]
Total (95% CI)	784	805			•			100%	1.85[1.03,3.33]
Total events: 31 (Misoprostol)), 17 (Misoprostol plus oxyto	cin)							
Heterogeneity: Tau ² =0; Chi ² =0	0.51, df=1(P=0.47); I ² =0%								
Test for overall effect: Z=2.06((P=0.04)					1			
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxvtocin

Analysis 18.3. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin		1	Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95%	6 CI			IV, Random, 95% CI
Caliskan 2002	12/396	4/401			-			44.79%	3.04[0.99,9.34]
Caliskan 2003	14/388	5/404			-	_		55.21%	2.92[1.06,8.02]
Total (95% CI)	784	805			•	-		100%	2.97[1.4,6.3]
Total events: 26 (Misoprostol)), 9 (Misoprostol plus oxytoc	in)							
Heterogeneity: Tau ² =0; Chi ² =0	0, df=1(P=0.96); I ² =0%								
Test for overall effect: Z=2.84((P=0)					1	1		
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin

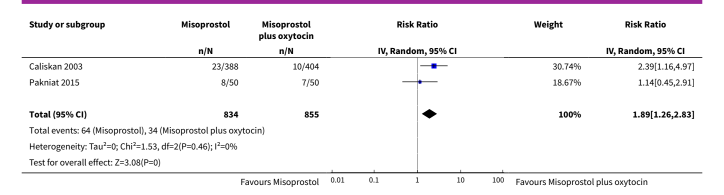
Analysis 18.6. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Caliskan 2002	39/396	28/401			-			54.67%	1.41[0.89,2.25]
Caliskan 2003	35/388	13/404			-			45.33%	2.8[1.51,5.22]
Total (95% CI)	784	805			•			100%	1.93[0.99,3.76]
Total events: 74 (Misoprostol)	, 41 (Misoprostol plus oxyto	cin)							
Heterogeneity: Tau ² =0.16; Chi	i ² =3.01, df=1(P=0.08); l ² =66.7	7%							
Test for overall effect: Z=1.92((P=0.06)								
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin

Analysis 18.7. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Caliskan 2002	33/396	17/401		_ 				50.59%	1.97[1.11,3.47]
	Fav	vours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin





Analysis 18.8. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 8 Blood loss.

Study or subgroup	Misoprostol		Misoprostol plus oxytocin			Mean Difference				Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ndom, 95%	CI			Random, 95% CI
Caliskan 2003	388	328 (152)	404	280 (182)				1		100%	48[24.68,71.32]
Total ***	388		404					~		100%	48[24.68,71.32]
Heterogeneity: Not applicable											
Test for overall effect: Z=4.03(P<0.0	0001)										
			Favour	s Misoprostol	-100	-50	0	50	100	Favours Mis	oprostol plus oxytocin

Analysis 18.9. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol	Misoprostol plus oxytocin			Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95% CI			Random, 95% CI
Caliskan 2002	396	15 (12)	401	15 (13)			+		15.59%	0[-1.74,1.74]
Caliskan 2003	388	14 (11)	404	14 (13)			•		16.77%	0[-1.67,1.67]
Pakniat 2015	50	11.4 (2.9)	50	11 (0.8)					67.64%	0.4[-0.43,1.23]
Total ***	834		855						100%	0.27[-0.42,0.96]
Heterogeneity: Tau ² =0; Chi ² =	0.29, df=2(P=0.8	7); I ² =0%								
Test for overall effect: Z=0.77	(P=0.44)									
			Favour	s Misoprostol	-100	-50	0 50	100	Favours Mis	oprostol plus oxytocin

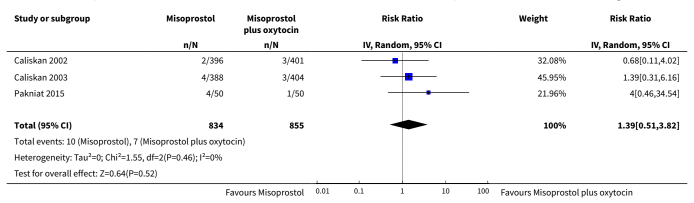
Analysis 18.11. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 11 Nausea.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, F	andom, 95%	6 CI			IV, Random, 95% CI
Pakniat 2015	11/50	6/50			-	-		100%	1.83[0.73,4.57]
Total (95% CI)	50	50			•	-		100%	1.83[0.73,4.57]
Total events: 11 (Misoprostol)	, 6 (Misoprostol plus oxytocii	n)							
	Fav	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin	



Study or subgroup	Misoprostol	Misoprostol Misoprostol plus oxytocin			Risk Ratio)		Weight Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI		IV, Random, 95% CI
Heterogeneity: Not applicable								
Test for overall effect: Z=1.3(P=0.19)								
		Favours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol plus oxytocin

Analysis 18.12. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 12 Vomiting.



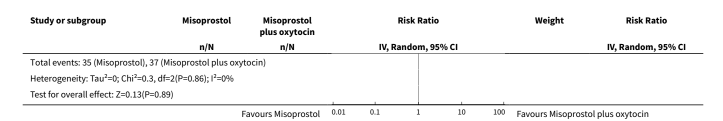
Analysis 18.16. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 16 Shivering.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Caliskan 2002	47/396	52/401			#			51.34%	0.92[0.63,1.32]
Caliskan 2003	44/388	49/404			+			47.85%	0.93[0.64,1.37]
Pakniat 2015	3/50	0/50				•	—	0.81%	7[0.37,132.1]
Total (95% CI)	834	855			•			100%	0.94[0.72,1.22]
Total events: 94 (Misoprostol), 101 (Misoprostol plus oxy	tocin)							
Heterogeneity: Tau ² =0; Chi ² =	1.82, df=2(P=0.4); I ² =0%								
Test for overall effect: Z=0.46	(P=0.65)								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin

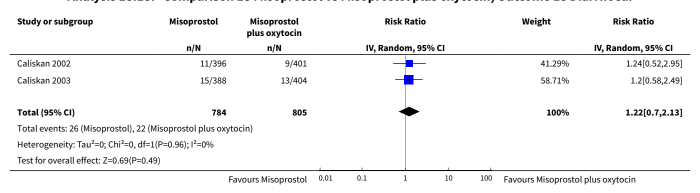
Analysis 18.17. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 17 Fever.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95°	% CI			IV, Random, 95% CI
Caliskan 2002	16/396	19/401			-			48.5%	0.85[0.44,1.63]
Caliskan 2003	17/388	16/404			-			45.93%	1.11[0.57,2.16]
Pakniat 2015	2/50	2/50						5.56%	1[0.15,6.82]
Total (95% CI)	834	855			•			100%	0.97[0.62,1.53]
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Misoprostol	plus oxytocin





Analysis 18.18. Comparison 18 Misoprostol vs Misoprostol plus oxytocin, Outcome 18 Diarrhoea.



Comparison 19. Carbetocin vs Injectable prostaglandins

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

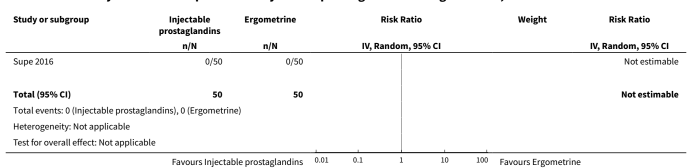
Comparison 20. Injectable prostaglandins vs Ergometrine

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
3 Blood transfusion	4	384	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.58]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	399	Risk Ratio (IV, Random, 95% CI)	1.42 [0.64, 3.17]
7 Additional uterotonics	5	684	Risk Ratio (IV, Random, 95% CI)	0.78 [0.40, 1.51]
8 Blood loss	8	1195	Mean Difference (IV, Random, 95% CI)	-16.08 [-57.17, 25.00]
9 Change in haemoglo- bin	2	300	Mean Difference (IV, Random, 95% CI)	-1.28 [-6.58, 4.01]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	5	684	Risk Ratio (IV, Random, 95% CI)	1.70 [0.98, 2.94]
12 Vomiting	5	684	Risk Ratio (IV, Random, 95% CI)	2.70 [0.97, 7.55]
13 Headache	1	80	Risk Ratio (IV, Random, 95% CI)	2.0 [0.39, 10.31]

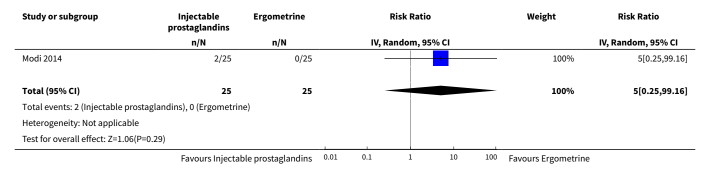


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14 Abdominal pain	3	384	Risk Ratio (IV, Random, 95% CI)	1.70 [0.07, 40.44]
15 Hypertension	2	315	Risk Ratio (IV, Random, 95% CI)	0.16 [0.02, 1.37]
16 Shivering	3	434	Risk Ratio (IV, Random, 95% CI)	0.38 [0.16, 0.91]
17 Fever	4	534	Risk Ratio (IV, Random, 95% CI)	4.52 [0.76, 26.80]
18 Diarrhoea	5	664	Risk Ratio (IV, Random, 95% CI)	10.09 [2.75, 37.00]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 20.1. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 1 Death.



Analysis 20.2. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 2 PPH >= 1000 mL.





Analysis 20.3. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Ergometrine	ometrine Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, Rai	ndom, 9	5% CI			IV, Random, 95% CI
Biswas 2007	0/50	2/50	-	-		_		49.82%	0.2[0.01,4.06]
Modi 2014	2/25	0/25		_	_	-		50.18%	5[0.25,99.16]
Supe 2016	0/50	0/50							Not estimable
Vaid 2009	0/67	0/67							Not estimable
Total (95% CI)	192	192						100%	1.01[0.04,23.58]
Total events: 2 (Injectable pr	ostaglandins), 2 (Ergometrin	e)							
Heterogeneity: Tau ² =2.84; Cl	hi²=2.21, df=1(P=0.14); l²=54.	79%							
Test for overall effect: Z=0(P=	=1)								
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	

Analysis 20.6. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Ergometrine			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% C	1			IV, Random, 95% CI
Kushtagi 2006	7/108	5/107						34.67%	1.39[0.45,4.23]
Modi 2014	6/25	0/25			+	+	\rightarrow	7.5%	13[0.77,219.11]
Vaid 2009	13/67	12/67			-			57.83%	1.08[0.53,2.2]
Total (95% CI)	200	199			•			100%	1.42[0.64,3.17]
Total events: 26 (Injectable p	orostaglandins), 17 (Ergomet	rine)							
Heterogeneity: Tau ² =0.16; Ch	ni²=2.82, df=2(P=0.24); I²=29.	08%							
Test for overall effect: Z=0.86	(P=0.39)								
	Favours Injectal	ble prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	

Analysis 20.7. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ra	atio		Weight	Risk Ratio
	n/N	n/N		IV, Random	, 95% CI			IV, Random, 95% CI
Khurshid 2010	0/100	1/100		+ +			4.25%	0.33[0.01,8.09]
Modi 2014	2/25	0/25		-	+		4.84%	5[0.25,99.16]
Supe 2016	4/50	2/50		-	•		15.71%	2[0.38,10.43]
Tripti 2006	0/100	2/100	\leftarrow	+			4.72%	0.2[0.01,4.11]
Vaid 2009	9/67	14/67		-			70.48%	0.64[0.3,1.38]
Total (95% CI)	342	342		•			100%	0.78[0.4,1.51]
Total events: 15 (Injectable p	rostaglandins), 19 (Ergomet	rine)						
Heterogeneity: Tau ² =0.01; Ch	ni ² =4.03, df=4(P=0.4); I ² =0.76	%						
Test for overall effect: Z=0.74	(P=0.46)							
	Favours Injectal	ole prostaglandins	0.01	0.1 1	10	100	Favours Ergometrine	



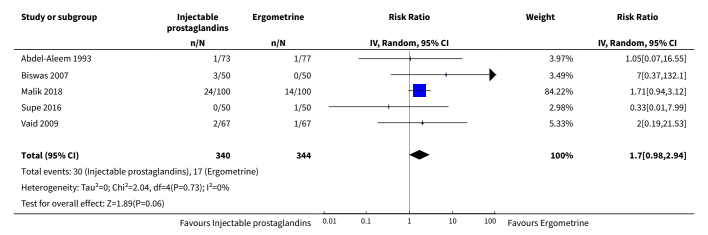
Analysis 20.8. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 8 Blood loss.

Study or subgroup	roup Injectable Ergometrine prostaglandins			Mean Differe	nce		Weight	Mean Difference		
	N	Mean(SD)	N	Mean(SD)		Random, 95%	6 CI			Random, 95% CI
Abdel-Aleem 1993	73	179 (59)	77	319 (52.3)	•				13.04%	-140[-157.88,-122.12]
Khurshid 2010	100	63.6 (10.1)	100	83.6 (14.1)		+			13.35%	-20[-23.4,-16.6]
Kushtagi 2006	108	235.7 (99.3)	107	214.1 (110)		+			12.58%	21.6[-6.42,49.62]
Malik 2018	100	129 (27.3)	100	250 (35.2)	•				13.29%	-121[-129.73,-112.27]
Modi 2014	25	435 (147.6)	25	131 (72)					10.06%	304[239.63,368.37]
Reddy 2001	40	113 (127)	40	202 (84)	←				11.36%	-89[-136.19,-41.81]
Supe 2016	50	153.8 (43.5)	50	152.2 (49.3)					13.02%	1.6[-16.61,19.81]
Tripti 2006	100	74.9 (27.2)	100	93.6 (32.7)		-			13.29%	-18.74[-27.07,-10.41]
Total ***	596		599				-		100%	-16.08[-57.17,25]
Heterogeneity: Tau ² =3287.59	; Chi²=734.67, d	f=7(P<0.0001); I ² :	=99.05%							
Test for overall effect: Z=0.77	(P=0.44)									
		Favours Inj	ectable p	rostaglandins	-100	-50 0	50	100	Favours Erg	ometrine

Analysis 20.9. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 9 Change in haemoglobin.

Study or subgroup	Injectable prostaglandins		Ergometrine			Mean Difference				Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95% CI				Random, 95% CI	
Malik 2018	100	4.7 (3.2)	100	8.7 (3.2)						49.7%	-4[-4.89,-3.11]	
Supe 2016	50	9 (1.2)	50	7.6 (0.2)			•			50.3%	1.4[1.06,1.74]	
Total ***	150		150				•			100%	-1.28[-6.58,4.01]	
Heterogeneity: Tau ² =14.46; C	Chi ² =124.4, df=1(I	P<0.0001); I ² =99.	2%									
Test for overall effect: Z=0.48	(P=0.63)						į					
		Favours Inj	ectable p	rostaglandins	-100	-50	0	50	100	Favours Erg	ometrine	

Analysis 20.11. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 11 Nausea.

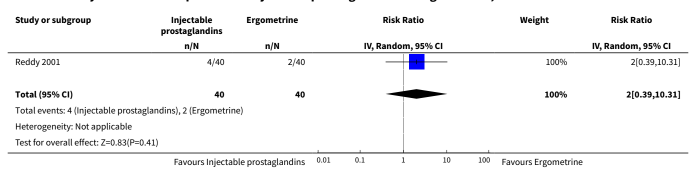




Analysis 20.12. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 12 Vomiting.

Study or subgroup	Injectable prostaglandins	Ergometrine		Ris	sk Ratio	0		Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 9	5% CI			IV, Random, 95% CI
Abdel-Aleem 1993	12/73	1/77			-			17.95%	12.66[1.69,94.91]
Biswas 2007	1/50	2/50			-			14.21%	0.5[0.05,5.34]
Malik 2018	12/100	6/100			+-			38.97%	2[0.78,5.12]
Supe 2016	8/50	1/50			-	•		17.62%	8[1.04,61.62]
Vaid 2009	1/67	1/67						11.25%	1[0.06,15.66]
Total (95% CI)	340	344				>		100%	2.7[0.97,7.55]
Total events: 34 (Injectable p	orostaglandins), 11 (Ergomet	rine)							
Heterogeneity: Tau ² =0.48; Ch	ni ² =6.15, df=4(P=0.19); I ² =34.	99%							
Test for overall effect: Z=1.89	(P=0.06)								
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	

Analysis 20.13. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 13 Headache.



Analysis 20.14. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 14 Abdominal pain.

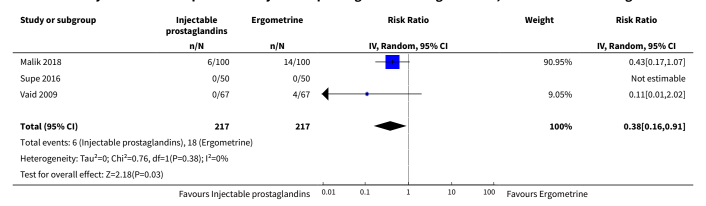
Study or subgroup	Injectable prostaglandins	,			Weight	Risk Ratio
	n/N	n/N	IV, Random, 95	% CI		IV, Random, 95% CI
Abdel-Aleem 1993	6/73	0/77		-	32.05%	13.7[0.79,238.98]
Supe 2016	1/50	9/50			36.82%	0.11[0.01,0.84]
Vaid 2009	2/67	0/67		•	31.13%	5[0.24,102.22]
Total (95% CI)	190	194			100%	1.7[0.07,40.44]
Total events: 9 (Injectable pr	ostaglandins), 9 (Ergometrin	e)				
Heterogeneity: Tau ² =6.03; C	hi²=8.81, df=2(P=0.01); l²=77.	3%				
Test for overall effect: Z=0.33	8(P=0.74)					
	Favours Injectal	ole prostaglandins	0.01 0.1 1	10 100	Favours Ergometrine	



Analysis 20.15. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 15 Hypertension.

Study or subgroup	Injectable prostaglandins	Ergometrine		R	isk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95	% CI			IV, Random, 95% CI
Biswas 2007	0/50	5/50	$\overline{}$	1	-			55.28%	0.09[0.01,1.6]
Kushtagi 2006	0/108	1/107		-				44.72%	0.33[0.01,8.02]
Total (95% CI)	158	157	-					100%	0.16[0.02,1.37]
Total events: 0 (Injectable pr	ostaglandins), 6 (Ergometrin	e)							
Heterogeneity: Tau ² =0; Chi ² =	:0.35, df=1(P=0.56); I ² =0%								
Test for overall effect: Z=1.67	(P=0.09)								
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	

Analysis 20.16. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 16 Shivering.



Analysis 20.17. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 17 Fever.

Study or subgroup	Injectable prostaglandins	Ergometrine			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, F	andom, 95	% CI			IV, Random, 95% CI
Biswas 2007	1/50	0/50						31.36%	3[0.13,71.92]
Malik 2018	4/100	0/100			-	-		37.41%	9[0.49,165]
Supe 2016	0/50	0/50							Not estimable
Vaid 2009	1/67	0/67			-			31.24%	3[0.12,72.35]
Total (95% CI)	267	267					-	100%	4.52[0.76,26.8]
Total events: 6 (Injectable pr	ostaglandins), 0 (Ergometrin	ie)							
Heterogeneity: Tau ² =0; Chi ² =	0.34, df=2(P=0.84); I ² =0%								
Test for overall effect: Z=1.66	(P=0.1)								
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	



Analysis 20.18. Comparison 20 Injectable prostaglandins vs Ergometrine, Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ra	tio		Weight	Risk Ratio
	n/N	n/N		IV, Random,	95% CI			IV, Random, 95% CI
Abdel-Aleem 1993	2/73	0/77		-	-	→	18.53%	5.27[0.26,107.96]
Malik 2018	2/100	0/100		-	+	→	18.48%	5[0.24,102.85]
Reddy 2001	7/40	0/40		+	-	\rightarrow	21.1%	15[0.89,254.13]
Supe 2016	7/50	0/50		+	-	\rightarrow	21%	15[0.88,255.78]
Vaid 2009	7/67	0/67				→	20.9%	15[0.87,257.48]
Total (95% CI)	330	334			-	_	100%	10.09[2.75,37]
Total events: 25 (Injectable p	rostaglandins), 0 (Ergometri	ine)						
Heterogeneity: Tau ² =0; Chi ² =	0.61, df=4(P=0.96); I ² =0%							
Test for overall effect: Z=3.49	(P=0)				1			
	Favours Injectal	ole prostaglandins	0.01	0.1 1	10	100	Favours Ergometrine	

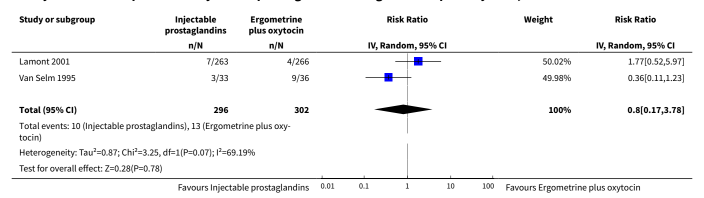
Comparison 21. Injectable prostaglandins vs Ergometrine plus oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	598	Risk Ratio (IV, Random, 95% CI)	0.80 [0.17, 3.78]
3 Blood transfusion	1	69	Risk Ratio (IV, Random, 95% CI)	0.65 [0.17, 2.53]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	598	Risk Ratio (IV, Random, 95% CI)	1.29 [0.92, 1.79]
7 Additional uterotonics	1	112	Risk Ratio (IV, Random, 95% CI)	1.07 [0.16, 7.36]
8 Blood loss	2	598	Mean Difference (IV, Random, 95% CI)	-26.18 [-104.63, 52.27]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	529	Risk Ratio (IV, Random, 95% CI)	5.06 [1.12, 22.86]
12 Vomiting	1	529	Risk Ratio (IV, Random, 95% CI)	0.92 [0.40, 2.13]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

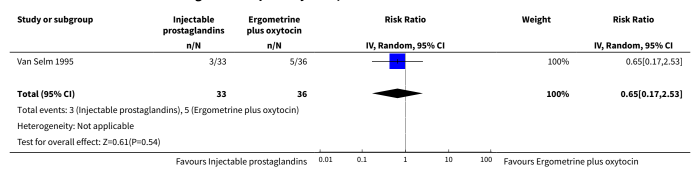


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	641	Risk Ratio (IV, Random, 95% CI)	23.70 [7.55, 74.45]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 21.2. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 2 PPH >= 1000 mL.



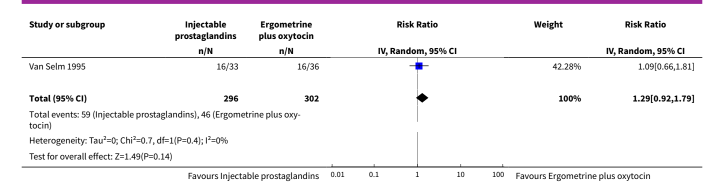
Analysis 21.3. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 3 Blood transfusion.



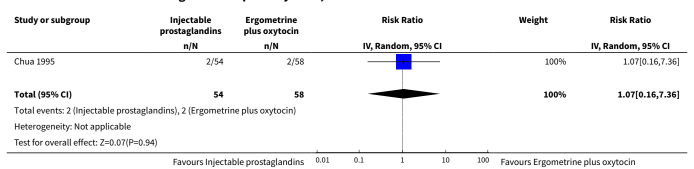
Analysis 21.6. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Lamont 2001	43/263	30/266			-			57.72%	1.45[0.94,2.24]
	Favours Injectab	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	e plus oxytocin

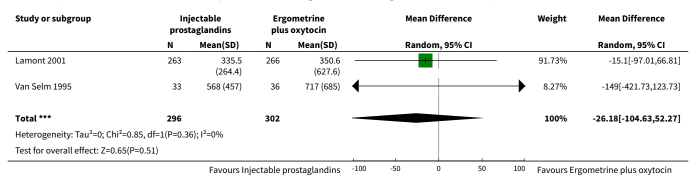




Analysis 21.7. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 7 Additional uterotonics.



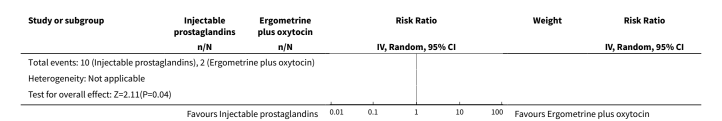
Analysis 21.8. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 8 Blood loss.



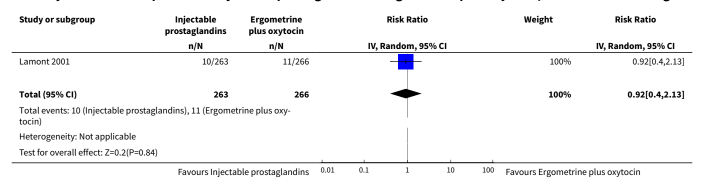
Analysis 21.11. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin			Risk Ratio)		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Lamont 2001	10/263	2/266				1		100%	5.06[1.12,22.86]
Total (95% CI)	263	266					1	100%	5.06[1.12,22.86]
	Favours Injecta	ble prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	plus oxytocin





Analysis 21.12. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 12 Vomiting.



Analysis 21.18. Comparison 21 Injectable prostaglandins vs Ergometrine plus oxytocin, Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rar	ıdom, 95% CI			IV, Random, 95% CI
Chua 1995	16/54	1/58				•	33.21%	17.19[2.36,125.22]
Lamont 2001	55/263	2/266			-		66.79%	27.81[6.86,112.85]
Total (95% CI)	317	324				~	100%	23.7[7.55,74.45]
Total events: 71 (Injectable p	rostaglandins), 3 (Ergometr	ne plus oxytocin)						
Heterogeneity: Tau ² =0; Chi ² =	0.15, df=1(P=0.7); I ² =0%							
Test for overall effect: Z=5.42	(P<0.0001)		1			1	i	
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10 100	Favours Ergometrin	e plus oxytocin

Comparison 22. Misoprostol plus oxytocin vs Injectable prostaglandins

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Comparison 23. Ergometrine vs Carbetocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Comparison 24. Carbetocin vs Ergometrine plus oxytocin

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Death	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	4	1210	Risk Ratio (IV, Random, 95% CI)	0.34 [0.13, 0.86]
3 Blood transfusion	4	1030	Risk Ratio (IV, Random, 95% CI)	1.42 [0.42, 4.82]
4 Severe maternal morbidity: intensive care admissions	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	4	1030	Risk Ratio (IV, Random, 95% CI)	0.96 [0.44, 2.09]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7 Additional uterotonics	6	1530	Risk Ratio (IV, Random, 95% CI)	0.54 [0.30, 1.00]
8 Blood loss	5	1330	Mean Difference (IV, Random, 95% CI)	-44.08 [-82.41, -5.75]
9 Change in haemoglo- bin	5	1160	Mean Difference (IV, Random, 95% CI)	-2.57 [-4.32, -0.82]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	6	1530	Risk Ratio (IV, Random, 95% CI)	0.29 [0.19, 0.45]
12 Vomiting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.27 [0.16, 0.46]
13 Headache	5	1330	Risk Ratio (IV, Random, 95% CI)	1.41 [0.45, 4.38]
14 Abdominal pain	4	930	Risk Ratio (IV, Random, 95% CI)	0.57 [0.36, 0.91]
15 Hypertension	3	660	Risk Ratio (IV, Random, 95% CI)	0.56 [0.22, 1.39]
16 Shivering	5	1290	Risk Ratio (IV, Random, 95% CI)	0.45 [0.26, 0.80]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 24.1. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 1 Death.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		R	isk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rai	ndom, 95% CI			IV, Random, 95% CI
Nirmala 2009	0/60	0/60						Not estimable
Samimi 2013	0/100	0/100						Not estimable
Total (95% CI)	160	160						Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytoci	n)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable						1		
	F	avours Carbetocin	0.01	0.1	1 10	100	Favours Ergometri	ne plus oxvtocinCarbetocin



Analysis 24.2. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Askar 2011	0/120	1/120		8.52%	0.33[0.01,8.1]
Leung 2006	0/150	1/150		8.51%	0.33[0.01,8.12]
Maged 2017	4/150	15/150		74.46%	0.27[0.09,0.78]
Su 2009	1/185	0/185	+	8.5%	3[0.12,73.17]
Total (95% CI)	605	605	•	100%	0.34[0.13,0.86]
Total events: 5 (Carbetocin), 1	7 (Ergometrine plus oxytoc	in)			
Heterogeneity: Tau ² =0; Chi ² =1	.98, df=3(P=0.58); I ² =0%				
Test for overall effect: Z=2.27(F	P=0.02)				
	F	avours Carbetocin	0.01 0.1 1 10	100 Favours Ergometrin	e plus oxytocinCarbetocin

Analysis 24.3. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 3 Blood transfusion.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Ris	sk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI	
Askar 2011	0/120	1/120		+	+		14.57%	0.33[0.01,8.1]	
Leung 2006	5/150	2/150		-			56.23%	2.5[0.49,12.69]	
Nirmala 2009	0/60	1/60					14.66%	0.33[0.01,8.02]	
Su 2009	1/185	0/185			+		14.54%	3[0.12,73.17]	
Total (95% CI)	515	515		-			100%	1.42[0.42,4.82]	
Total events: 6 (Carbetocin), 4	4 (Ergometrine plus oxytoci	n)							
Heterogeneity: Tau ² =0; Chi ² =2	2.27, df=3(P=0.52); I ² =0%								
Test for overall effect: Z=0.57((P=0.57)					1			
	F	avours Carbetocin	0.01	0.1	1 10	100	Favours Fronmetrin	e plus oxytocinCarbetocin	

Analysis 24.4. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	om, 95% CI		IV, Random, 95% CI
Nirmala 2009	0/60	0/60				Not estimable
Samimi 2013	0/100	0/100				Not estimable
Total (95% CI)	160	160				Not estimable
Total events: 0 (Carbetocin), 0 (E	Ergometrine plus oxytoci	n)				
Heterogeneity: Not applicable						
Test for overall effect: Not applic	cable					
	F	avours Carbetocin	0.01 0.1	1 10	100 Favours Frgometr	ine plus oxytocinCarbetocin



Analysis 24.6. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin	Carbetocin Ergometrine plus oxytocin			sk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI	
Askar 2011	2/120	3/120		-	•		19.18%	0.67[0.11,3.92]	
Leung 2006	6/150	2/150					23.88%	3[0.62,14.63]	
Nirmala 2009	3/60	6/60					33.16%	0.5[0.13,1.91]	
Su 2009	3/185	3/185					23.79%	1[0.2,4.89]	
Total (95% CI)	515	515		•	•		100%	0.96[0.44,2.09]	
Total events: 14 (Carbetocin),	14 (Ergometrine plus oxyto	ocin)							
Heterogeneity: Tau ² =0.01; Chi	i ² =3.06, df=3(P=0.38); I ² =2.0	9%							
Test for overall effect: Z=0.11(P=0.91)					1			
	F	avours Carbetocin	0.01	0.1	1 10	100	Favours Ergometrin	e plus oxytocinCarbetocin	

Analysis 24.7. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI	
Askar 2011	18/120	21/120		-		21.97%	0.86[0.48,1.53]	
Leung 2006	13/150	10/150				18.68%	1.3[0.59,2.87]	
Maged 2017	5/150	26/150				16.68%	0.19[0.08,0.49]	
Nirmala 2009	3/60	9/60				12.61%	0.33[0.09,1.17]	
Samimi 2013	1/100	11/100				6.76%	0.09[0.01,0.69]	
Su 2009	25/185	31/185		-		23.31%	0.81[0.5,1.31]	
Total (95% CI)	765	765		•		100%	0.54[0.3,1]	
Total events: 65 (Carbetocin)	, 108 (Ergometrine plus oxyt	cocin)						
Heterogeneity: Tau ² =0.35; Ch	i ² =15.76, df=5(P=0.01); l ² =68	3.28%						
Test for overall effect: Z=1.97	(P=0.05)				1			
	F	avours Carbetocin	0.01	0.1 1 10	100	Favours Ergometrin	e plus oxytocinCarbetocin	

Analysis 24.8. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 8 Blood loss.

Study or subgroup	Ca	Carbetocin		Ergometrine plus oxytocin		Mean Difference		Weight		Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95%	6 CI			Random, 95% CI
Askar 2011	120	224.6 (110.6)	120	306.1 (95.7)	+				21.48%	-81.5[-107.66,-55.34]
Leung 2006	150	232 (122)	150	249 (175)					20.07%	-17[-51.14,17.14]
Maged 2017	150	578 (178)	150	602 (213)					18.07%	-24[-68.42,20.42]
Nirmala 2009	60	244 (114)	60	343 (143)	-	-			17.71%	-99[-145.27,-52.73]
Su 2009	185	217.4 (99.2)	185	223.1 (76.3)					22.67%	-5.7[-23.73,12.33]
Total ***	665		665						100%	-44.08[-82.41,-5.75]
Heterogeneity: Tau ² =1602.31	; Chi²=30.82, df=	4(P<0.0001); I ² =	87.02%							
Test for overall effect: Z=2.25	(P=0.02)									
	Favours Carbetocir				-100	-50 0	50	100	Favours Erg Carbetocin	ometrine plus oxytocin-



Analysis 24.9. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup	Car			ometrine oxytocin					Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95	% CI			Random, 95% CI
Askar 2011	120	8 (2)	120	11 (3)		-			24.41%	-3[-3.65,-2.35]
Leung 2006	150	14 (11)	150	15 (13)		-+-			15.67%	-1[-3.73,1.73]
Maged 2017	150	11 (12)	150	12 (13)		-+			15.21%	-1[-3.83,1.83]
Nirmala 2009	60	3 (2)	60	4 (2)		-			24.23%	-1[-1.72,-0.28]
Samimi 2013	100	4.1 (3.6)	100	10.4 (7.8)		-			20.47%	-6.3[-7.98,-4.62]
Total ***	580		580			•			100%	-2.57[-4.32,-0.82]
Heterogeneity: Tau ² =3.16; Ch	i ² =40.53, df=4(P	<0.0001); I ² =90.1	3%			İ				
Test for overall effect: Z=2.88	(P=0)									
			Favours Carbetocin		-10	-5 0	5	10	Favours Erg Carbetocin	ometrine plus oxytocin-

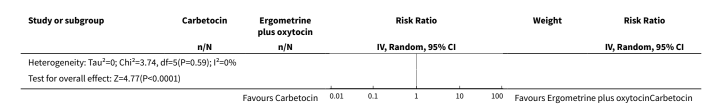
Analysis 24.11. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 11 Nausea.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
Askar 2011	4/120	13/120		16.13%	0.31[0.1,0.92]
Leung 2006	2/150	11/150		8.66%	0.18[0.04,0.81]
Maged 2017	5/150	11/150		18.03%	0.45[0.16,1.28]
Nirmala 2009	0/60	1/60		1.9%	0.33[0.01,8.02]
Samimi 2013	2/100	3/100		6.15%	0.67[0.11,3.9]
Su 2009	11/185	46/185	-	49.13%	0.24[0.13,0.45]
Total (95% CI)	765	765	•	100%	0.29[0.19,0.45]
Total events: 24 (Carbetocin),	85 (Ergometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0; Chi ² =2	2.34, df=5(P=0.8); I ² =0%				
Test for overall effect: Z=5.49((P<0.0001)				
	F	avours Carbetocin	0.01 0.1 1 10	100 Favours Ergometrin	e plus oxytocinCarbetocin

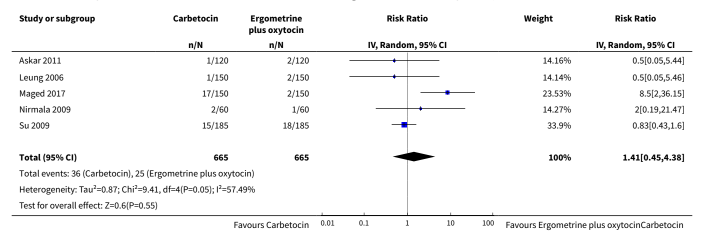
Analysis 24.12. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	CI		IV, Random, 95% CI
Askar 2011	3/120	12/120				18.89%	0.25[0.07,0.86]
Leung 2006	1/150	10/150				6.95%	0.1[0.01,0.77]
Maged 2017	4/150	10/150				22.44%	0.4[0.13,1.25]
Nirmala 2009	0/60	2/60	\leftarrow	+ -		3.19%	0.2[0.01,4.08]
Samimi 2013	1/100	0/100				2.85%	3[0.12,72.77]
Su 2009	7/185	30/185				45.68%	0.23[0.11,0.52]
Total (95% CI)	765	765		•		100%	0.27[0.16,0.46]
Total events: 16 (Carbetocin),	, 64 (Ergometrine plus oxyto	cin)			1		
	F	avours Carbetocin	0.01	0.1 1	10 100	Favours Ergometrine	e plus oxytocinCarbetocin

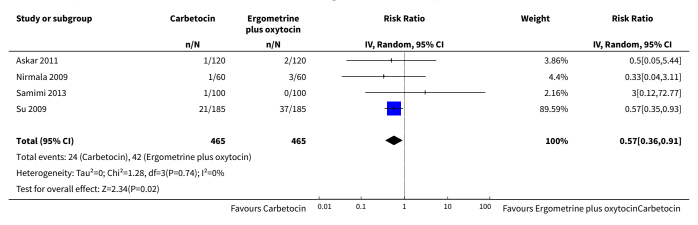




Analysis 24.13. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 13 Headache.



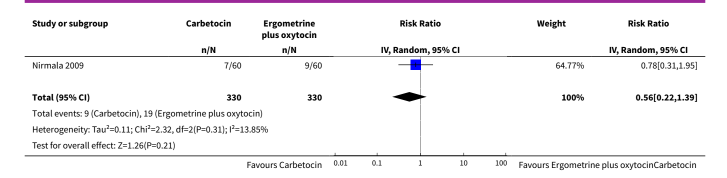
Analysis 24.14. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 14 Abdominal pain.



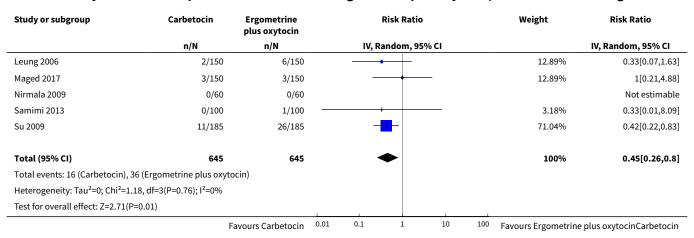
Analysis 24.15. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 15 Hypertension.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Askar 2011	2/120	4/120		-	-			25.6%	0.5[0.09,2.68]
Leung 2006	0/150	6/150	_	•	+	1		9.63%	0.08[0,1.35]
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Ergometrine	plus oxytocinCarbetocin





Analysis 24.16. Comparison 24 Carbetocin vs Ergometrine plus oxytocin, Outcome 16 Shivering.



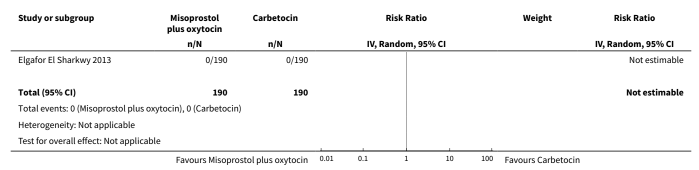
Comparison 25. Misoprostol plus oxytocin vs Carbetocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	380	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	380	Risk Ratio (IV, Random, 95% CI)	4.00 [0.45, 35.46]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	1	380	Risk Ratio (IV, Random, 95% CI)	1.19 [0.74, 1.93]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

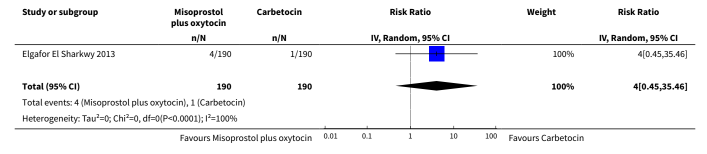


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9 Change in haemoglobin	1	380	Mean Difference (IV, Random, 95% CI)	-1.70 [-3.72, 0.32]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	380	Risk Ratio (IV, Random, 95% CI)	1.21 [0.62, 2.39]
12 Vomiting	1	380	Risk Ratio (IV, Random, 95% CI)	1.33 [0.58, 3.09]
13 Headache	1	380	Risk Ratio (IV, Random, 95% CI)	2.0 [0.37, 10.79]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	380	Risk Ratio (IV, Random, 95% CI)	7.83 [3.43, 17.88]
17 Fever	1	380	Risk Ratio (IV, Random, 95% CI)	8.5 [1.99, 36.28]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 25.1. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 1 Death.



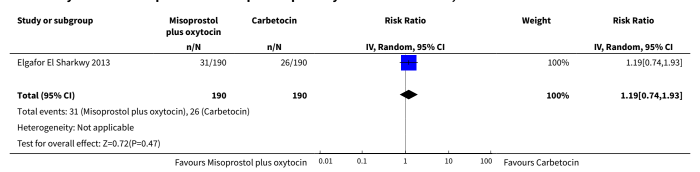
Analysis 25.3. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 3 Blood transfusion.



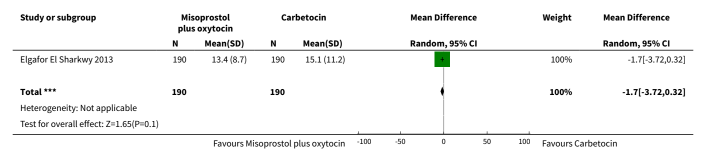


Study or subgroup	Misoprostol plus oxytocin	Carbetocin			Risk Ratio	•		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Test for overall effect: Z=1.25(P=0.21)			_						
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 25.7. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 7 Additional uterotonics.



Analysis 25.9. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 9 Change in haemoglobin.



Analysis 25.11. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 11 Nausea.

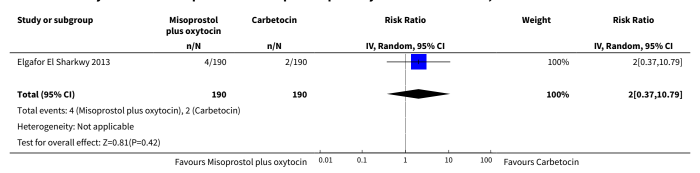
Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio					Weight	Risk Ratio
	n/N	n/N		IV, F	andom, 95% (CI			IV, Random, 95% CI
Elgafor El Sharkwy 2013	17/190	14/190			-			100%	1.21[0.62,2.39]
Total (95% CI)	190	190			•			100%	1.21[0.62,2.39]
Total events: 17 (Misoprostol plus ox	kytocin), 14 (Carbetoc	in)							
Heterogeneity: Not applicable									
Test for overall effect: Z=0.56(P=0.57	·)								
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	



Analysis 25.12. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio					Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Elgafor El Sharkwy 2013	12/190	9/190						100%	1.33[0.58,3.09]
Total (95% CI)	190	190			•			100%	1.33[0.58,3.09]
Total events: 12 (Misoprostol plus oxy	ytocin), 9 (Carbetocin)							
Heterogeneity: Not applicable									
Test for overall effect: Z=0.67(P=0.5)									
	Favours Misopros	tol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 25.13. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 13 Headache.



Analysis 25.16. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 16 Shivering.

Study or subgroup	bgroup Misoprostol Carbetocin Risk Ratio plus oxytocin			Weight	Risk Ratio				
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Elgafor El Sharkwy 2013	47/190	6/190				-		100%	7.83[3.43,17.88]
Total (95% CI)	190	190				•		100%	7.83[3.43,17.88]
Total events: 47 (Misoprostol pl	us oxytocin), 6 (Carbetocii	n)							
Heterogeneity: Not applicable									
Test for overall effect: Z=4.89(P<	<0.0001)					1			
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	

Analysis 25.17. Comparison 25 Misoprostol plus oxytocin vs Carbetocin, Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Elgafor El Sharkwy 2013	17/190	2/190			-	1	_	100%	8.5[1.99,36.28]
Total (95% CI)	190	190			-	~	-	100%	8.5[1.99,36.28]
Total events: 17 (Misoprostol pl	us oxytocin), 2 (Carbetocin)							
	Favours Misopros	tol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	



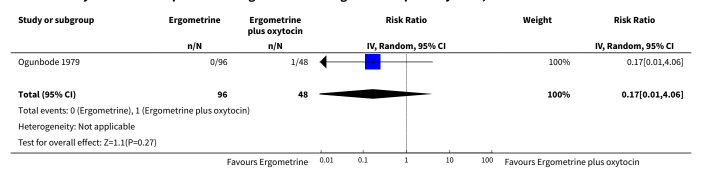
Study or subgroup	Misoprostol plus oxytocin	Carbetocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Heterogeneity: Not applicable									
Test for overall effect: Z=2.89(P=0)									
	Favours Misonro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	

Comparison 26. Ergometrine vs Ergometrine plus oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	144	Risk Ratio (IV, Random, 95% CI)	0.17 [0.01, 4.06]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	144	Mean Difference (IV, Random, 95% CI)	20.10 [5.76, 34.44]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Analysis 26.6. Comparison 26 Ergometrine vs Ergometrine plus oxytocin, Outcome 6 PPH >= 500 mL.



Analysis 26.8. Comparison 26 Ergometrine vs Ergometrine plus oxytocin, Outcome 8 Blood loss.

Study or subgroup	Erge	ometrine	_	ometrine oxytocin		Ме	an Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ndom, 95% CI		Random, 95% CI
Ogunbode 1979	96	96 (54.2)	48	75.9 (33.2)				100%	20.1[5.76,34.44]
Total ***	96		48				•	100%	20.1[5.76,34.44]
Heterogeneity: Not applicable									
Test for overall effect: Z=2.75(P=0.01)					1				
			Favours	Ergometrine	-100	-50	0 50	100 Favours Erg	ometrine plus oxytocin

Comparison 27. Misoprostol plus oxytocin vs Ergometrine

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

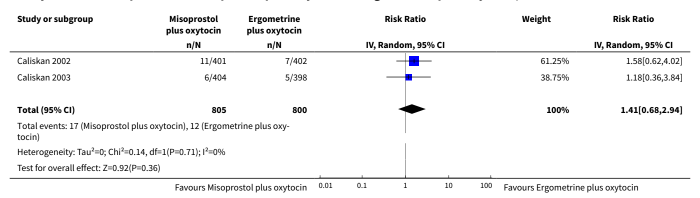
Comparison 28. Misoprostol plus oxytocin vs Ergometrine plus oxytocin

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.41 [0.68, 2.94]
3 Blood transfusion	2	1605	Risk Ratio (IV, Random, 95% CI)	0.89 [0.36, 2.19]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.39 [0.65, 2.99]
7 Additional uterotonics	2	1605	Risk Ratio (IV, Random, 95% CI)	1.48 [0.82, 2.69]
8 Blood loss	1	802	Mean Difference (IV, Random, 95% CI)	-16.0 [-40.24, 8.24]
9 Change in haemoglo- bin	2	1605	Mean Difference (IV, Random, 95% CI)	-0.50 [-1.72, 0.72]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

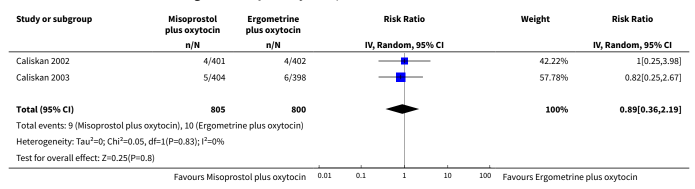


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12 Vomiting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.04 [0.23, 4.77]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	1605	Risk Ratio (IV, Random, 95% CI)	2.95 [2.02, 4.29]
17 Fever	2	1605	Risk Ratio (IV, Random, 95% CI)	2.89 [1.51, 5.54]
18 Diarrhoea	2	1605	Risk Ratio (IV, Random, 95% CI)	0.81 [0.46, 1.41]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

Analysis 28.2. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 2 PPH >= 1000 mL.



Analysis 28.3. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 3 Blood transfusion.

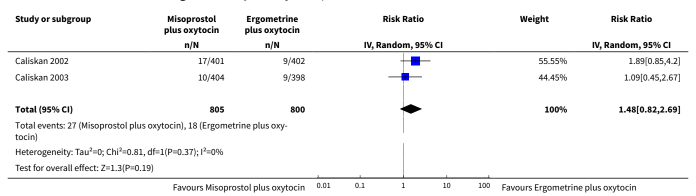




Analysis 28.6. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, F	Random, 95% C	:1			IV, Random, 95% CI
Caliskan 2002	28/401	14/402			-			53.35%	2[1.07,3.75]
Caliskan 2003	13/404	14/398			_			46.65%	0.91[0.44,1.92]
Total (95% CI)	805	800						100%	1.39[0.65,2.99]
Total events: 41 (Misoprostol tocin)	l plus oxytocin), 28 (Ergomet	rine plus oxy-							
Heterogeneity: Tau ² =0.19; Ch	ni ² =2.51, df=1(P=0.11); I ² =60.	13%							
Test for overall effect: Z=0.84	(P=0.4)								
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Ergometrin	e plus oxytocin

Analysis 28.7. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 7 Additional uterotonics.



Analysis 28.8. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 8 Blood loss.

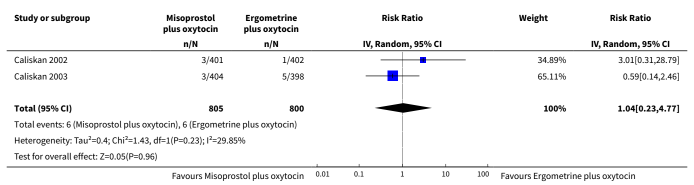
Study or subgroup		oprostol oxytocin		ometrine oxytocin		Mea	an Differei	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	dom, 95%	CI			Random, 95% CI
Caliskan 2003	404	280 (182)	398	296 (168)						100%	-16[-40.24,8.24]
Total ***	404		398			→				100%	-16[-40.24,8.24]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.29(P=0.2)											
		Favours Mis	oprostol	plus oxytocin	-100	-50	0	50	100	Favours Erg	ometrine plus oxytocin



Analysis 28.9. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 9 Change in haemoglobin.

Study or subgroup		oprostol oxytocin		ometrine oxytocin	М	ean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	R	andom, 95% CI			Random, 95% CI
Caliskan 2002	401	15 (13)	402	15 (12)		•		50.02%	0[-1.73,1.73]
Caliskan 2003	404	14 (13)	398	15 (12)				49.98%	-1[-2.73,0.73]
Total ***	805		800					100%	-0.5[-1.72,0.72]
Heterogeneity: Tau ² =0; Chi ² =0	.64, df=1(P=0.42	2); I ² =0%							
Test for overall effect: Z=0.8(P	=0.42)								
		Favours Mis	oprostol	plus oxytocin -10	00 -50	0 50	100	Favours Erg	ometrine plus oxytocin

Analysis 28.12. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 12 Vomiting.



Analysis 28.16. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 16 Shivering.

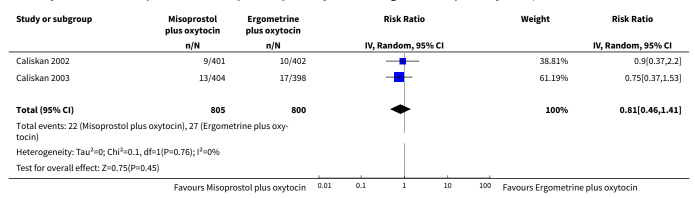
Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Caliskan 2002	52/401	19/402		-		55.1%	2.74[1.65,4.55]
Caliskan 2003	49/404	15/398		-		44.9%	3.22[1.84,5.64]
Total (95% CI)	805	800		•		100%	2.95[2.02,4.29]
Total events: 101 (Misoprostotocin)	ol plus oxytocin), 34 (Ergom	etrine plus oxy-					
Heterogeneity: Tau ² =0; Chi ² =	0.17, df=1(P=0.68); I ² =0%						
Test for overall effect: Z=5.63	(P<0.0001)						
	Favours Misopro	ostol plus oxytocin	0.01 0.	1 1 10	100	Favours Ergometrin	e plus oxytocin



Analysis 28.17. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
Caliskan 2002	19/401	6/402				51.12%	3.17[1.28,7.87]
Caliskan 2003	16/404	6/398		-		48.88%	2.63[1.04,6.64]
Total (95% CI)	805	800		•		100%	2.89[1.51,5.54]
Total events: 35 (Misoprostol tocin)	plus oxytocin), 12 (Ergome	trine plus oxy-					
Heterogeneity: Tau ² =0; Chi ² =	0.08, df=1(P=0.77); I ² =0%						
Test for overall effect: Z=3.21	(P=0)						
	Favours Misopro	ostol plus oxvtocin	0.01	0.1 1 10	100	Favours Ergometrin	e plus oxytocin

Analysis 28.18. Comparison 28 Misoprostol plus oxytocin vs Ergometrine plus oxytocin, Outcome 18 Diarrhoea.



Comparison 29. Oxytocin vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	3	1920	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	3	1920	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	11	8782	Risk Ratio (IV, Random, 95% CI)	0.61 [0.51, 0.72]
2.1 Vaginal birth	10	8731	Risk Ratio (IV, Random, 95% CI)	0.61 [0.51, 0.72]
2.2 Caesarean	1	51	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	8	6717	Risk Ratio (IV, Random, 95% CI)	0.75 [0.51, 1.12]
3.1 Vaginal birth	7	6643	Risk Ratio (IV, Random, 95% CI)	0.75 [0.51, 1.12]
3.2 Caesarean	1	74	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbid- ity: intensive care admis- sions	1	1950	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	1	1950	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidi- ty: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	9	7021	Risk Ratio (IV, Random, 95% CI)	0.61 [0.52, 0.71]
6.1 Vaginal birth	8	6970	Risk Ratio (IV, Random, 95% CI)	0.61 [0.52, 0.72]
6.2 Caesarean	1	51	Risk Ratio (IV, Random, 95% CI)	0.48 [0.17, 1.40]
7 Additional uterotonics	8	5047	Risk Ratio (IV, Random, 95% CI)	0.43 [0.32, 0.58]
7.1 Vaginal birth	6	4922	Risk Ratio (IV, Random, 95% CI)	0.49 [0.35, 0.66]
7.2 Caesarean	2	125	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.44]
8 Blood loss	8	3521	Mean Difference (IV, Random, 95% CI)	-118.52 [-141.40, -95.64]
8.1 Vaginal birth	6	3396	Mean Difference (IV, Random, 95% CI)	-121.22 [-144.50, -97.94]
8.2 Caesarean	2	125	Mean Difference (IV, Random, 95% CI)	-42.70 [-166.12, 80.72]
9 Change in haemoglobin	5	2304	Mean Difference (IV, Random, 95% CI)	-2.68 [-4.47, -0.89]
9.1 Vaginal birth	4	2253	Mean Difference (IV, Random, 95% CI)	-3.19 [-4.98, -1.40]
9.2 Caesarean	1	51	Mean Difference (IV, Random, 95% CI)	-0.20 [-3.51, 3.11]
10 Breastfeeding	1	1540	Risk Ratio (IV, Random, 95% CI)	1.00 [0.95, 1.05]
10.1 Vaginal birth	1	1540	Risk Ratio (IV, Random, 95% CI)	1.00 [0.95, 1.05]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	3	1788	Risk Ratio (IV, Random, 95% CI)	0.82 [0.47, 1.42]
11.1 Vaginal birth	2	1714	Risk Ratio (IV, Random, 95% CI)	0.86 [0.49, 1.51]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
11.2 Caesarean	1	74	Risk Ratio (IV, Random, 95% CI)	0.25 [0.02, 3.83]
12 Vomiting	3	2150	Risk Ratio (IV, Random, 95% CI)	1.40 [0.44, 4.41]
12.1 Vaginal birth	2	2076	Risk Ratio (IV, Random, 95% CI)	1.40 [0.44, 4.41]
12.2 Caesarean	1	74	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	2	1704	Risk Ratio (IV, Random, 95% CI)	1.56 [0.52, 4.74]
13.1 Vaginal birth	1	1653	Risk Ratio (IV, Random, 95% CI)	1.26 [0.63, 2.51]
13.2 Caesarean	1	51	Risk Ratio (IV, Random, 95% CI)	6.74 [0.37, 124.21]
14 Abdominal pain	1	1665	Risk Ratio (IV, Random, 95% CI)	0.89 [0.80, 1.00]
14.1 Vaginal birth	1	1665	Risk Ratio (IV, Random, 95% CI)	0.89 [0.80, 1.00]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Ceasarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal sense of well- being	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth - Women's perceptions of well-being at 3 months postpartum: Less energy than before birth	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Vaginal birth - Women's perceptions of well-being at	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3 months postpartum: Experiencing (some) fatigue				
19.3 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth - Did management influence positively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Vaginal birth - Did management influence negatively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Vaginal birth - Did management make no difference in the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.4 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 29.1. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 1 Death.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.1.1 Vaginal birth					
de Groot 1996	0/78	0/143			Not estimable
Jerbi 2007	0/65	0/65			Not estimable
Stanton 2013	0/682	0/887			Not estimable
Subtotal (95% CI)	825	1095			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
29.1.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	825	1095			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Test for subgroup differences: Not app	olicable				
		Favours Oxytocin (0.01 0.1 1 10	100 Favours Placebo o	no treatment



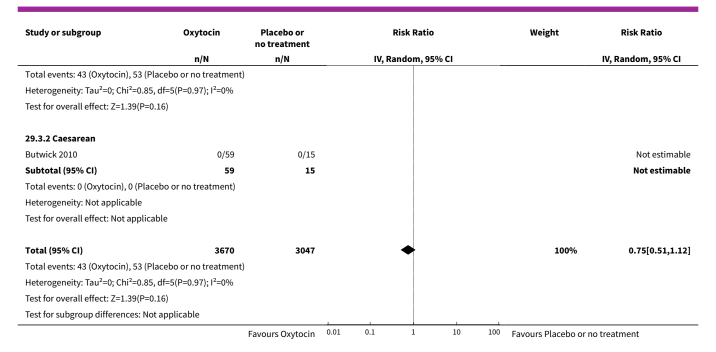
Analysis 29.2. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.2.1 Vaginal birth					
Abdel-Aleem 2010	4/1291	4/659		1.5%	0.51[0.13,2.03]
Al-Sawaf 2013	1/37	6/39		0.67%	0.18[0.02,1.39]
Benchimol 2001	12/196	13/220		4.96%	1.04[0.48,2.22]
de Groot 1996	7/78	16/143		4.02%	0.8[0.34,1.87]
Jangsten 2011	82/810	138/821	-	43.96%	0.6[0.47,0.78]
Jans 2017	54/851	99/835	-	28.45%	0.54[0.39,0.74]
Jerbi 2007	0/65	0/65			Not estimable
Nordstrom 1997	32/513	43/487	-+ 	14.79%	0.71[0.45,1.1]
Poeschmann 1991	2/28	3/24		0.99%	0.57[0.1,3.14]
Stanton 2013	1/682	8/887		0.67%	0.16[0.02,1.3]
Subtotal (95% CI)	4551	4180	•	100%	0.61[0.51,0.72]
Total events: 195 (Oxytocin), 330 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =6.37,	df=8(P=0.61); I ² =0%				
Test for overall effect: Z=5.77(P<0.	.0001)				
29.2.2 Caesarean					
Rosseland 2013	0/26	0/25			Not estimable
Subtotal (95% CI)	26	25			Not estimable
Total events: 0 (Oxytocin), 0 (Place	ebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applica	ble				
Total (95% CI)	4577	4205	•	100%	0.61[0.51,0.72]
Total events: 195 (Oxytocin), 330 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =6.37,	df=8(P=0.61); I ² =0%				
Test for overall effect: Z=5.77(P<0.	.0001)				
Test for subgroup differences: Not	applicable				

Analysis 29.3. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
29.3.1 Vaginal birth							
Abdel-Aleem 2010	8/1257	7/642				15.56%	0.58[0.21,1.6]
Al-Sawaf 2013	0/37	1/39				1.58%	0.35[0.01,8.35]
de Groot 1996	2/78	3/143		+		5.08%	1.22[0.21,7.16]
Jangsten 2011	18/810	23/821		-		42.77%	0.79[0.43,1.46]
Jans 2017	10/851	12/835				22.83%	0.82[0.36,1.88]
Jerbi 2007	0/65	0/65					Not estimable
Nordstrom 1997	5/513	7/487				12.19%	0.68[0.22,2.12]
Subtotal (95% CI)	3611	3032		•		100%	0.75[0.51,1.12]
		Favours Oxytocin	0.01	0.1 1 10	100	Favours Placebo or	no treatment





Analysis 29.4. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Oxytocin	Placebo or no treatment			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
29.4.1 Vaginal birth								
Abdel-Aleem 2010	0/1291	0/659						Not estimable
Subtotal (95% CI)	1291	659						Not estimable
Total events: 0 (Oxytocin), 0 (Placebo o	r no treatment)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
29.4.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Oxytocin), 0 (Placebo o	r no treatment)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total (95% CI)	1291	659						Not estimable
Total events: 0 (Oxytocin), 0 (Placebo o	r no treatment)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Test for subgroup differences: Not appl	icable							
		Favours Oxytocin	0.01	0.1	1 10	100	Favours Placebo o	r no treatment



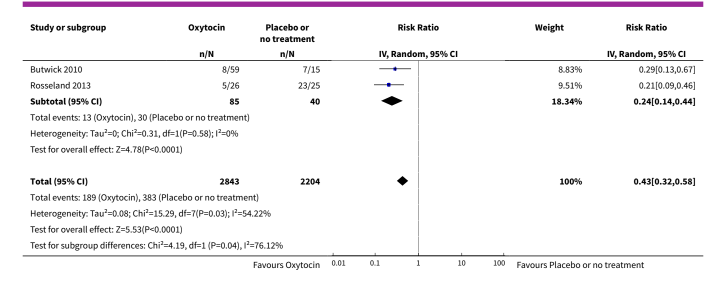
Analysis 29.6. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.6.1 Vaginal birth					
Abdel-Aleem 2010	63/1291	65/659		13.7%	0.49[0.35,0.69]
Al-Sawaf 2013	2/37	8/39		1.12%	0.26[0.06,1.16]
Benchimol 2001	29/196	60/220		10.87%	0.54[0.36,0.81]
de Groot 1996	25/78	55/143	-+	11.46%	0.83[0.57,1.22]
Jans 2017	276/851	369/835	•	28.46%	0.73[0.65,0.83]
Nordstrom 1997	104/513	175/487	+	21.6%	0.56[0.46,0.7]
Poeschmann 1991	7/28	10/24		3.58%	0.6[0.27,1.33]
Stanton 2013	18/682	49/887		7.13%	0.48[0.28,0.81]
Subtotal (95% CI)	3676	3294	•	97.91%	0.61[0.52,0.72]
Total events: 524 (Oxytocin), 791	(Placebo or no treatmen	nt)			
Heterogeneity: Tau ² =0.02; Chi ² =1	3.17, df=7(P=0.07); l ² =46	.84%			
Test for overall effect: Z=5.81(P<0	.0001)				
29.6.2 Caesarean					
Rosseland 2013	4/26	8/25		2.09%	0.48[0.17,1.4]
Subtotal (95% CI)	26	25	•	2.09%	0.48[0.17,1.4]
Total events: 4 (Oxytocin), 8 (Plac	ebo or no treatment)				
Heterogeneity: Tau ² =0; Chi ² =0, df	=0(P<0.0001); I ² =100%				
Test for overall effect: Z=1.34(P=0	.18)				
Total (95% CI)	3702	3319	•	100%	0.61[0.52,0.71]
Total events: 528 (Oxytocin), 799	(Placebo or no treatmen	it)			
Heterogeneity: Tau ² =0.02; Chi ² =1	3.49, df=8(P=0.1); I ² =40.7	71%			
	0001)		İ		
Test for overall effect: Z=6.11(P<0	.0001)				

Analysis 29.7. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI		IV, Random, 95% CI
29.7.1 Vaginal birth						
Abdel-Aleem 2010	41/1260	55/641			19.26%	0.38[0.26,0.56]
Al-Sawaf 2013	2/37	8/39			3.59%	0.26[0.06,1.16]
de Groot 1996	14/78	26/143		+	13.67%	0.99[0.55,1.78]
Jans 2017	79/842	195/830		-	24.2%	0.4[0.31,0.51]
Nordstrom 1997	40/513	67/487			19.97%	0.57[0.39,0.82]
Poeschmann 1991	0/28	2/24	\leftarrow	-	0.97%	0.17[0.01,3.42]
Subtotal (95% CI)	2758	2164		◆	81.66%	0.49[0.35,0.66]
Total events: 176 (Oxytocin), 35	53 (Placebo or no treatme	nt)				
Heterogeneity: Tau ² =0.07; Chi ²	=10.86, df=5(P=0.05); I ² =53	3.95%				
Test for overall effect: Z=4.54(P	2<0.0001)					
29.7.2 Caesarean						
		Favours Oxytocin	0.01	0.1 1 10	100 Favours Placebo or r	no treatment



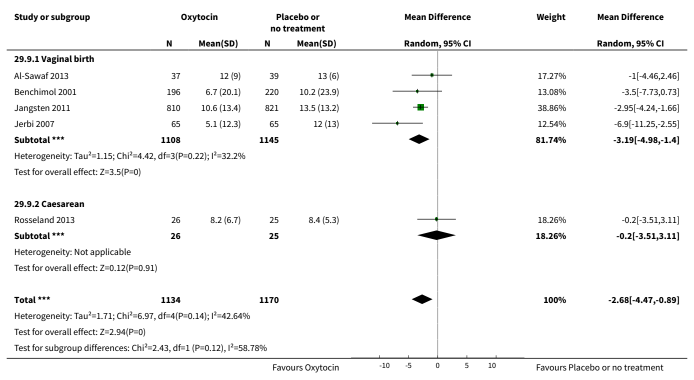


Analysis 29.8. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	o	xytocin		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
29.8.1 Vaginal birth							
Al-Sawaf 2013	37	314.7 (94.6)	39	438.6 (130.2)		20.14%	-123.9[-174.88,-72.92]
Benchimol 2001	196	278 (254)	220	382 (269.5)		20.66%	-104[-154.32,-53.68]
de Groot 1996	78	499 (454)	143	520 (419)	+	3.52%	-21[-142.93,100.93]
Jangsten 2011	810	535 (414.5)	821	680 (486.7)		27.21%	-145[-188.85,-101.15]
Nordstrom 1997	513	409 (345)	487	527 (412)		23.46%	-118[-165.23,-70.77]
Poeschmann 1991	28	374 (279)	24	548 (376)	+ +	1.57%	-174[-356.51,8.51]
Subtotal ***	1662		1734		•	96.56%	-121.22[-144.5,-97.94]
Heterogeneity: Tau ² =0; Chi ² =	4.52, df=5(P=0.4	·8); I ² =0%					
Test for overall effect: Z=10.2	1(P<0.0001)						
29.8.2 Caesarean							
Butwick 2010	59	750.8 (160.9)	15	800 (256)	+	2.83%	-49.22[-185.12,86.68]
Rosseland 2013	26	841 (556)	25	853 (518)	+	0.6%	-12[-306.8,282.8]
	85		40			3.44%	-42.7[-166.12,80.72]
Subtotal ***	65					3.7770	12.1[200:22,00:12]
Subtotal *** Heterogeneity: Tau ² =0; Chi ² =		2); I ² =0%				3.4470	42.7[100.12,00.72]
	0.05, df=1(P=0.8	2); I ² =0%				3.4470	12.11 200122,001121
Heterogeneity: Tau ² =0; Chi ² =	0.05, df=1(P=0.8	2); I ² =0%	1774		•	100%	-118.52[-141.4,-95.64]
Heterogeneity: Tau ² =0; Chi ² = Test for overall effect: Z=0.68	0.05, df=1(P=0.8 (P=0.5)		1774		•		. , ,
Heterogeneity: Tau ² =0; Chi ² = Test for overall effect: Z=0.68 Total ***	0.05, df=1(P=0.8 (P=0.5) 1747 6.08, df=7(P=0.5		1774		•		. , ,



Analysis 29.9. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.

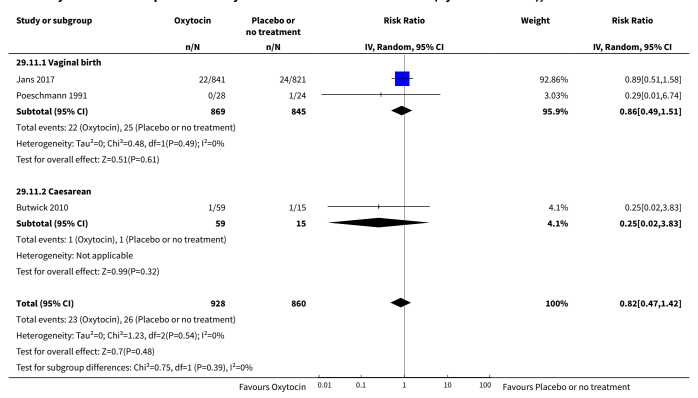


Analysis 29.10. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 10 Breastfeeding.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.10.1 Vaginal birth					
Jans 2017	618/779	606/761		100%	1[0.95,1.05]
Subtotal (95% CI)	779	761	▼	100%	1[0.95,1.05]
Total events: 618 (Oxytocin), 606 (Plac	ebo or no treatme	nt)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.15(P=0.88)					
29.10.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	779	761	•	100%	1[0.95,1.05]
Total events: 618 (Oxytocin), 606 (Plac	ebo or no treatme	nt)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.15(P=0.88)					
Test for subgroup differences: Not app	olicable				
		Favours Oxytocin	1	Favours Placebo or	no treatment



Analysis 29.11. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 11 Nausea.



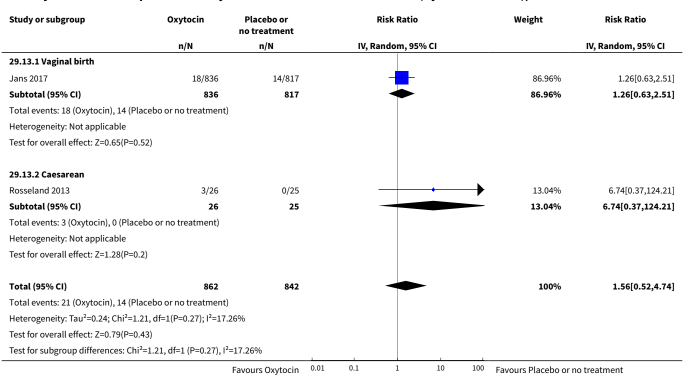
Analysis 29.12. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 12 Vomiting.

Study or subgroup O:	cytocin	Placebo or no treatment		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI
29.12.1 Vaginal birth								
Benchimol 2001	1/196	1/220			•	-	17.23%	1.12[0.07,17.83]
Jans 2017	6/840	4/820		-			82.77%	1.46[0.41,5.17]
Subtotal (95% CI)	1036	1040		-			100%	1.4[0.44,4.41]
Total events: 7 (Oxytocin), 5 (Placebo or no	treatment)							
Heterogeneity: Tau ² =0; Chi ² =0.03, df=1(P=0	.86); I ² =0%							
Test for overall effect: Z=0.57(P=0.57)								
29.12.2 Caesarean								
Butwick 2010	0/59	0/15						Not estimable
Subtotal (95% CI)	59	15						Not estimable
Total events: 0 (Oxytocin), 0 (Placebo or no	treatment)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total (95% CI)	1095	1055					100%	1.4[0.44,4.41]
Total events: 7 (Oxytocin), 5 (Placebo or no	treatment)							
Heterogeneity: Tau ² =0; Chi ² =0.03, df=1(P=0	.86); I ² =0%							
Test for overall effect: Z=0.57(P=0.57)								
		Favours Oxytocin	0.01	0.1	1 10	100	Favours Placebo or	no treatment



Study or subgroup	Oxytocin	Placebo or no treatment			Risk Ratio	•		Weight Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI		IV, Random, 95% CI
Test for subgroup differences: Not applicable			_					
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or no treatment

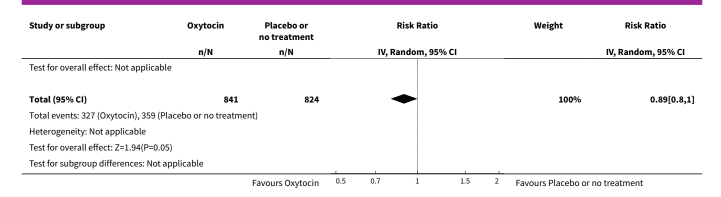
Analysis 29.13. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 13 Headache.



Analysis 29.14. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.14.1 Vaginal birth					
Jans 2017	327/841	359/824	-	100%	0.89[0.8,1]
Subtotal (95% CI)	841	824	•	100%	0.89[0.8,1]
Total events: 327 (Oxytocin), 359 (Pla	acebo or no treatmen	t)			
Heterogeneity: Not applicable					
Test for overall effect: Z=1.94(P=0.05))				
29.14.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	o or no treatment)				
Heterogeneity: Not applicable					
		Favours Oxytocin	0.5 0.7 1 1.5	Favours Placebo or I	no treatment





Analysis 29.16. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
29.16.1 Vaginal birth					
Benchimol 2001	0/196	0/220			Not estimable
Subtotal (95% CI)	196	220			Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	o or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicabl	e				
29.16.2 Ceasarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	o or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	е				
Total (95% CI)	196	220			Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	o or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicabl	e				
Test for subgroup differences: Not a	pplicable				
		Favours Oxytocin (0.01 0.1 1 10	100 Favours Placebo	or no treatment

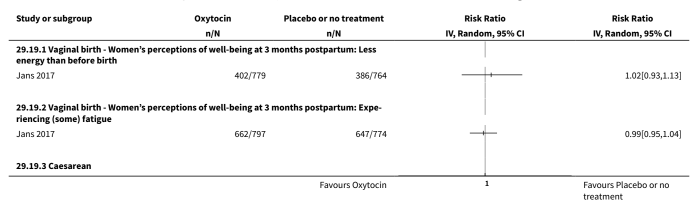
Analysis 29.17. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 17 Fever.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, F	Random, 95% (:1			IV, Random, 95% CI
29.17.1 Vaginal birth									
Benchimol 2001	0/196	0/220							Not estimable
Subtotal (95% CI)	196	220							Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	o or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	2								
				_					
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or no	o treatment

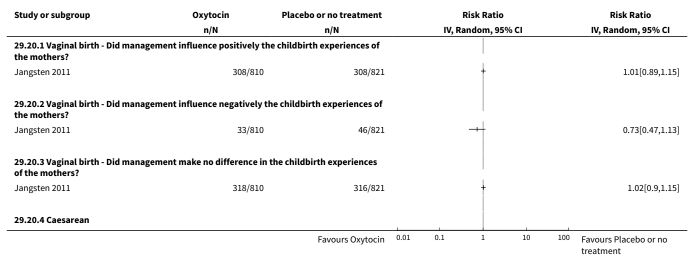


Study or subgroup	Oxytocin	Placebo or no treatment			Risk Ratio	•		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
29.17.2 Caesarean									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Total (95% CI)	196	220							Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Test for subgroup differences: Not app	licable								
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or	no treatment

Analysis 29.19. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 19 Maternal sense of well-being.



Analysis 29.20. Comparison 29 Oxytocin vs Placebo or no treatment (by mode of birth), Outcome 20 Maternal satisfaction.





Comparison 30. Carbetocin vs Placebo or no treatment (by mode of birth)

Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	1	50	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.75 [0.30, 1.85]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Caesarean	1	50	Risk Ratio (IV, Random, 95% CI)	0.75 [0.30, 1.85]
7 Additional uterotonics	2	169	Risk Ratio (IV, Random, 95% CI)	0.19 [0.12, 0.32]
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Caesarean	2	169	Risk Ratio (IV, Random, 95% CI)	0.19 [0.12, 0.32]
8 Blood loss	1	50	Mean Difference (IV, Random, 95% CI)	-274.0 [-591.60, 43.60]
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or sub- No. of studies No. of partici- group title pants			Statistical method	Effect size
8.2 Caesarean	1	50	Mean Difference (IV, Random, 95% CI)	-274.0 [-591.60, 43.60]
9 Change in haemo- globin	1	50	Mean Difference (IV, Random, 95% CI)	-3.40 [-7.23, 0.43]
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	1	50	Mean Difference (IV, Random, 95% CI)	-3.40 [-7.23, 0.43]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



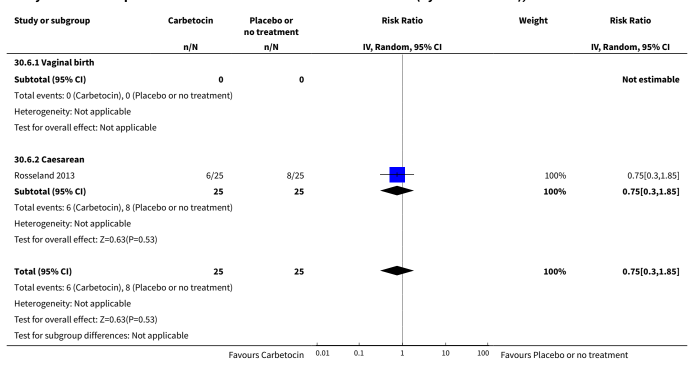
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 30.2. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI		IV, Random, 95% CI
30.2.1 Vaginal birth			,			
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
30.2.2 Caesarean						
Rosseland 2013	0/25	0/25				Not estimable
Subtotal (95% CI)	25	25				Not estimable
Total events: 0 (Carbetocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	25	25				Not estimable
Total events: 0 (Carbetocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not appl	icable					
	F:	avours Carbetocin	0.01 0.	1 1 10	100 Favours Placeho or r	no treatment



Analysis 30.6. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.



Analysis 30.7. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
30.7.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Pla	acebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applica	ble				
30.7.2 Caesarean					
Barton 1996	8/62	41/57		58.55%	0.18[0.09,0.35]
Rosseland 2013	5/25	23/25		41.45%	0.22[0.1,0.48]
Subtotal (95% CI)	87	82	•	100%	0.19[0.12,0.32]
Total events: 13 (Carbetocin), 64 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =0.13,	df=1(P=0.72); I ² =0%				
Test for overall effect: Z=6.29(P<0.	.0001)				
Total (95% CI)	87	82	•	100%	0.19[0.12,0.32]
Total events: 13 (Carbetocin), 64 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =0.13,	df=1(P=0.72); I ² =0%				
Test for overall effect: Z=6.29(P<0.	.0001)				
Test for subgroup differences: Not	applicable				
	ı	Favours Carbetocin 0.01	0.1 1 10	100 Favours Placebo or	no treatment



Analysis 30.8. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Ca	rbetocin	Placebo or no treatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
30.8.1 Vaginal birth							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
30.8.2 Caesarean							
Rosseland 2013	25	579 (623)	25	853 (518)		100%	-274[-591.6,43.6]
Subtotal ***	25		25			100%	-274[-591.6,43.6]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.69(P=0.09)							
Total ***	25		25			100%	-274[-591.6,43.6]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.69(P=0.09)							
Test for subgroup differences: Not ap	plicable						
			Favou	rs Carbetocin	-500 -250 0 250 500	Favours Pla	cebo or no treatment

Analysis 30.9. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Ca	rbetocin	Placebo or no treatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
30.9.1 Vaginal birth							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
30.9.2 Caesarean							
Rosseland 2013	25	5 (8.2)	25	8.4 (5.3)	+	100%	-3.4[-7.23,0.43]
Subtotal ***	25		25		•	100%	-3.4[-7.23,0.43]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.74(P=0.08)							
Total ***	25		25		•	100%	-3.4[-7.23,0.43]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.74(P=0.08)							
Test for subgroup differences: Not app	olicable						
			Favou	rs Carbetocin -100	-50 0 50	100 Favours Plac	ebo or no treatment



Analysis 30.13. Comparison 30 Carbetocin vs Placebo or no treatment (by mode of birth), Outcome 13 Headache.

Study or subgroup	Carbetocin	Placebo or no treatment	Risk Rat	io	Weight	Risk Ratio
	n/N	n/N	IV, Random,	95% CI		IV, Random, 95% CI
30.13.1 Vaginal birth						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Placel	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
30.13.2 Caesarean						
Rosseland 2013	2/25	0/25	-		100%	5[0.25,99.16]
Subtotal (95% CI)	25	25			100%	5[0.25,99.16]
Total events: 2 (Carbetocin), 0 (Placel	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.06(P=0.29)						
Total (95% CI)	25	25			100%	5[0.25,99.16]
Total events: 2 (Carbetocin), 0 (Placel	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.06(P=0.29)						
Test for subgroup differences: Not ap	plicable					
	•	vours Carbetocin	0.01 0.1 1	10 100	Favours Placebo or n	o treatment

Comparison 31. Misoprostol vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3497	Risk Ratio (IV, Random, 95% CI)	1.00 [0.10, 9.59]
1.1 Vaginal birth	4	3497	Risk Ratio (IV, Random, 95% CI)	1.00 [0.10, 9.59]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	8	5467	Risk Ratio (IV, Random, 95% CI)	0.73 [0.56, 0.95]
2.1 Vaginal birth	8	5467	Risk Ratio (IV, Random, 95% CI)	0.73 [0.56, 0.95]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	6	3134	Risk Ratio (IV, Random, 95% CI)	0.46 [0.15, 1.47]
3.1 Vaginal birth	6	3134	Risk Ratio (IV, Random, 95% CI)	0.46 [0.15, 1.47]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	1	1620	Risk Ratio (IV, Random, 95% CI)	1.00 [0.14, 7.05]
4.1 Vaginal birth	1	1620	Risk Ratio (IV, Random, 95% CI)	1.00 [0.14, 7.05]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	7	4047	Risk Ratio (IV, Random, 95% CI)	0.75 [0.59, 0.94]
6.1 Vaginal birth	7	4047	Risk Ratio (IV, Random, 95% CI)	0.75 [0.59, 0.94]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	8	3746	Risk Ratio (IV, Random, 95% CI)	0.67 [0.52, 0.87]
7.1 Vaginal birth	8	3746	Risk Ratio (IV, Random, 95% CI)	0.67 [0.52, 0.87]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	8	4146	Mean Difference (IV, Random, 95% CI)	-42.07 [-52.47, -31.68]
8.1 Vaginal birth	8	4146	Mean Difference (IV, Random, 95% CI)	-42.07 [-52.47, -31.68]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	5	2334	Mean Difference (IV, Random, 95% CI)	-2.53 [-3.53, -1.52]
9.1 Vaginal birth	5	2334	Mean Difference (IV, Random, 95% CI)	-2.53 [-3.53, -1.52]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3997	Risk Ratio (IV, Random, 95% CI)	1.18 [0.78, 1.78]
11.1 Vaginal birth	4	3997	Risk Ratio (IV, Random, 95% CI)	1.18 [0.78, 1.78]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	6	4949	Risk Ratio (IV, Random, 95% CI)	1.41 [0.92, 2.16]
12.1 Vaginal birth	6	4949	Risk Ratio (IV, Random, 95% CI)	1.41 [0.92, 2.16]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	1116	Risk Ratio (IV, Random, 95% CI)	0.94 [0.32, 2.77]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
13.1 Vaginal birth	1	1116	Risk Ratio (IV, Random, 95% CI)	0.94 [0.32, 2.77]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	4	1894	Risk Ratio (IV, Random, 95% CI)	0.98 [0.29, 3.37]
14.1 Vaginal birth	4	1894	Risk Ratio (IV, Random, 95% CI)	0.98 [0.29, 3.37]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	5286	Risk Ratio (IV, Random, 95% CI)	2.94 [2.38, 3.64]
16.1 Vaginal birth	9	5286	Risk Ratio (IV, Random, 95% CI)	2.94 [2.38, 3.64]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	5	4396	Risk Ratio (IV, Random, 95% CI)	4.09 [2.01, 8.32]
17.1 Vaginal birth	5	4396	Risk Ratio (IV, Random, 95% CI)	4.09 [2.01, 8.32]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	6	4791	Risk Ratio (IV, Random, 95% CI)	2.80 [1.21, 6.49]
18.1 Vaginal birth	6	4791	Risk Ratio (IV, Random, 95% CI)	2.80 [1.21, 6.49]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



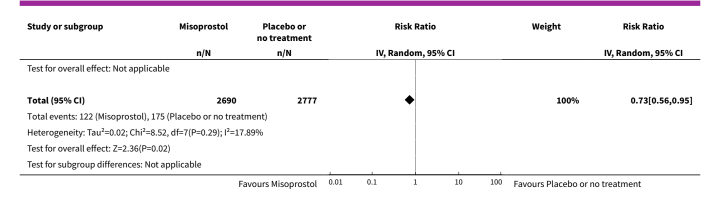
Analysis 31.1. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 1 Death.

Study or subgroup	Misoprostol Placebo or no treatment		Risl	k Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rand	om, 95% CI		IV, Random, 95% CI
31.1.1 Vaginal birth						
Derman 2006	0/812	1/808			49.97%	0.33[0.01,8.13]
Hoj 2005	1/330	0/331	-		- 50.03%	3.01[0.12,73.6]
Mobeen 2011	0/533	0/583				Not estimable
Supe 2016	0/50	0/50				Not estimable
Subtotal (95% CI)	1725	1772			100%	1[0.1,9.59]
Total events: 1 (Misoprostol), 1 (Pla	cebo or no treatment)					
Heterogeneity: Tau ² =0; Chi ² =0.91, d	lf=1(P=0.34); I ² =0%					
Test for overall effect: Z=0(P=1)						
31.1.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Pla	cebo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	le					
Total (95% CI)	1725	1772			100%	1[0.1,9.59]
Total events: 1 (Misoprostol), 1 (Pla	cebo or no treatment)					
Heterogeneity: Tau ² =0; Chi ² =0.91, d	f=1(P=0.34); I ² =0%					
Test for overall effect: Z=0(P=1)						
Test for subgroup differences: Not a	pplicable					
	Fav	ours Misoprostol	0.01 0.1	1 10	100 Favours Placebo or i	no treatment

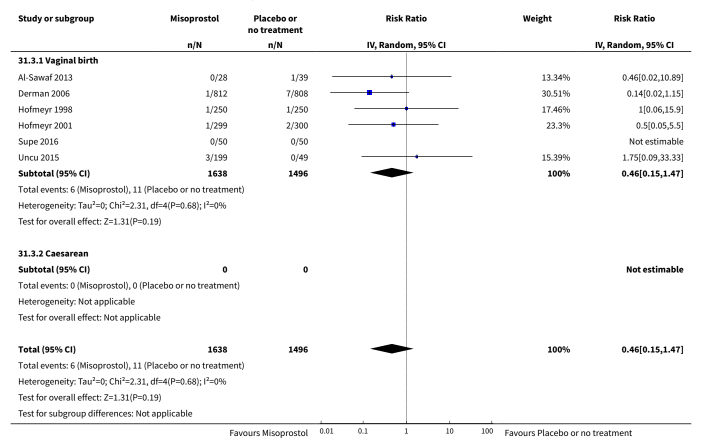
Analysis 31.2. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ra	tio		Weight	Risk Ratio
	n/N	n/N		IV, Random,	95% CI			IV, Random, 95% CI
31.2.1 Vaginal birth								
Al-Sawaf 2013	2/28	6/39			_		2.78%	0.46[0.1,2.13]
Bamigboye 1998a	13/270	19/272		+			11.94%	0.69[0.35,1.37]
Benchimol 2001	16/186	13/220		+			11.37%	1.46[0.72,2.95]
Derman 2006	2/812	10/808	_				2.82%	0.2[0.04,0.91]
Hofmeyr 1998	15/250	23/250		+			13.82%	0.65[0.35,1.22]
Hofmeyr 2001	27/300	29/299		+			19.58%	0.93[0.56,1.53]
Hoj 2005	37/330	56/331		-			27.59%	0.66[0.45,0.98]
Mobeen 2011	10/514	19/558		-+-			10.1%	0.57[0.27,1.22]
Subtotal (95% CI)	2690	2777		•			100%	0.73[0.56,0.95]
Total events: 122 (Misoprostol), 175 (F	lacebo or no treatr	ment)						
Heterogeneity: Tau ² =0.02; Chi ² =8.52, o	df=7(P=0.29); I ² =17.	89%						
Test for overall effect: Z=2.36(P=0.02)								
31.2.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol), 0 (Place	bo or no treatment)						
Heterogeneity: Not applicable								
	Fa	vours Misoprostol	0.01	0.1 1	10	100	Favours Placebo or r	no treatment



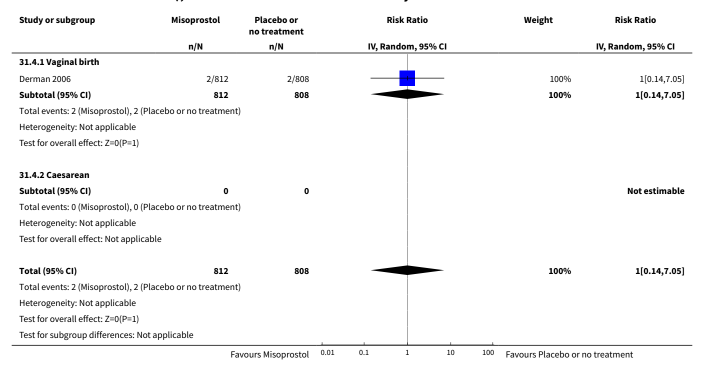


Analysis 31.3. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 3 Blood transfusion.





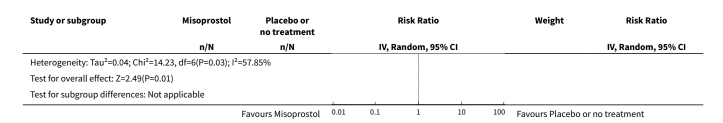
Analysis 31.4. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.



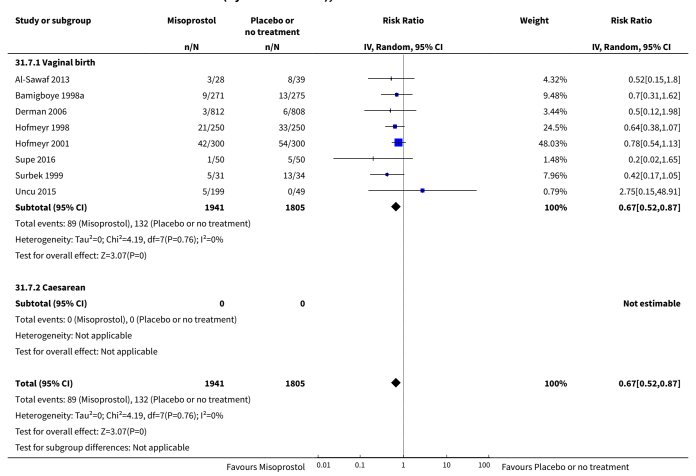
Analysis 31.6. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
31.6.1 Vaginal birth					
Al-Sawaf 2013	3/28	8/39		3.16%	0.52[0.15,1.8]
Benchimol 2001	52/186	60/220	+	20.35%	1.03[0.75,1.41]
Derman 2006	52/812	97/808	-	20.03%	0.53[0.39,0.74]
Fu 2003	2/80	9/76		2.21%	0.21[0.05,0.95]
Hoj 2005	150/330	170/331	-	28.47%	0.89[0.76,1.04]
Mobeen 2011	85/514	122/558		23.74%	0.76[0.59,0.97]
Surbek 1999	2/31	5/34		2.04%	0.44[0.09,2.1]
Subtotal (95% CI)	1981	2066	•	100%	0.75[0.59,0.94]
Total events: 346 (Misoprostol), 4	71 (Placebo or no treati	ment)			
Heterogeneity: Tau ² =0.04; Chi ² =1	4.23, df=6(P=0.03); l ² =57	7.85%			
Test for overall effect: Z=2.49(P=0	.01)				
31.6.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (P	lacebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applica	able				
Total (95% CI)	1981	2066	•	100%	0.75[0.59,0.94]
Total events: 346 (Misoprostol), 4	71 (Placebo or no treatr	ment)			
	Fa	vours Misoprostol 0.0	01 0.1 1 10	100 Favours Placebo or	no treatment





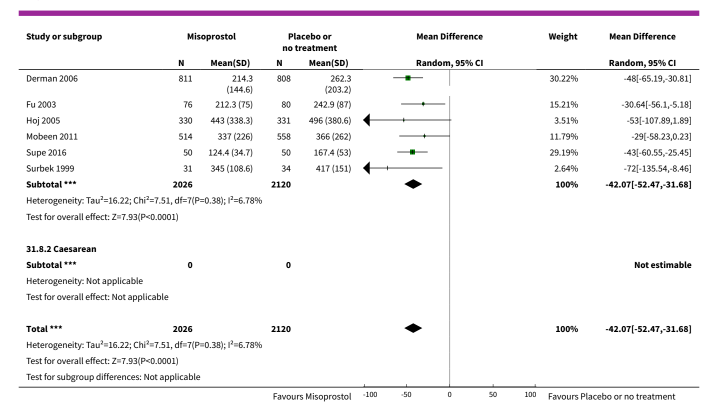
Analysis 31.7. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.



Analysis 31.8. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Mis	oprostol		acebo or reatment		Mea	an Differer	ice		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ran	ndom, 95%	CI			Random, 95% CI
31.8.1 Vaginal birth											
Al-Sawaf 2013	28	348 (112)	39	438.6 (130.2)	←		İ			3.13%	-90.6[-148.83,-32.37]
Benchimol 2001	186	374 (238.4)	220	382 (269.5)			+	—		4.32%	-8[-57.42,41.42]
			Favour	s Misoprostol	-100	-50	0	50	100	Favours Pla	cebo or no treatment





Analysis 31.9. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Misoprostol			acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
31.9.1 Vaginal birth							
Al-Sawaf 2013	28	13 (9)	39	13 (6)		6.13%	0[-3.83,3.83]
Benchimol 2001	186	9.6 (23.5)	220	10.2 (23.9)		4.35%	-0.6[-5.22,4.02]
Hoj 2005	330	4.7 (18.2)	331	7.2 (18.2)		10.63%	-2.5[-5.27,0.27]
Mobeen 2011	528	11 (12)	572	13 (14)		24.13%	-2[-3.54,-0.46]
Supe 2016	50	4.9 (0.5)	50	8.1 (0.2)	+	54.76%	-3.2[-3.35,-3.05]
Subtotal ***	1122		1212		◆	100%	-2.53[-3.53,-1.52]
Heterogeneity: Tau ² =0.47; Chi ² =6	.41, df=4(P=	0.17); I ² =37.59%)				
Test for overall effect: Z=4.93(P<0	.0001)						
31.9.2 Caesarean							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applica	ible						
Total ***	1122		1212		•	100%	-2.53[-3.53,-1.52]
Heterogeneity: Tau ² =0.47; Chi ² =6.	.41, df=4(P=	0.17); I ² =37.59%)				
Test for overall effect: Z=4.93(P<0	.0001)						
Test for subgroup differences: No	t applicable						
			Favou	rs Misoprostol	-10 -5 0 5 10	Favours Pla	cebo or no treatment



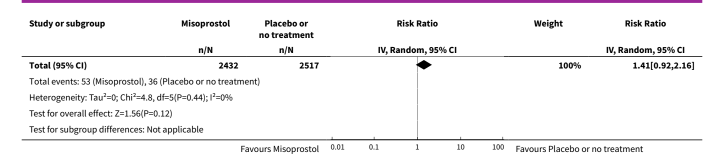
Analysis 31.11. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
31.11.1 Vaginal birth					
Derman 2006	35/812	29/808	 	72.56%	1.2[0.74,1.95]
Hofmeyr 2001	5/300	1/300	+	3.68%	5[0.59,42.54]
Hoj 2005	2/330	4/331		5.91%	0.5[0.09,2.72]
Mobeen 2011	8/533	8/583		17.84%	1.09[0.41,2.89]
Subtotal (95% CI)	1975	2022	*	100%	1.18[0.78,1.78]
Total events: 50 (Misoprostol), 42	(Placebo or no treatmer	nt)			
Heterogeneity: Tau ² =0; Chi ² =2.76	i, df=3(P=0.43); I ² =0%				
Test for overall effect: Z=0.8(P=0.4	42)				
31.11.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (P	Placebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applica	able				
Total (95% CI)	1975	2022	•	100%	1.18[0.78,1.78]
Total events: 50 (Misoprostol), 42	! (Placebo or no treatmer	nt)			
Heterogeneity: Tau ² =0; Chi ² =2.76	i, df=3(P=0.43); I ² =0%				
Test for overall effect: Z=0.8(P=0.4	42)				
Test for subgroup differences: No	t applicable				
	Fav	ours Misoprostol 0.01	0.1 1 10 1	.00 Favours Placebo or	no treatment

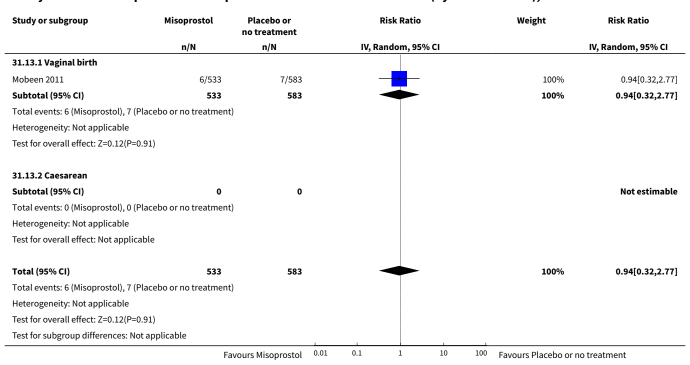
Analysis 31.12. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ra	tio	Weight	Risk Ratio
	n/N	n/N	IV, Random,	95% CI		IV, Random, 95% CI
31.12.1 Vaginal birth						
Bamigboye 1998a	1/271	1/275			2.41%	1.01[0.06,16.14]
Benchimol 2001	7/186	1/220	-	+	4.24%	8.28[1.03,66.68]
Derman 2006	28/812	25/808	-	-	65.64%	1.11[0.66,1.89]
Hofmeyr 2001	4/300	2/300		+	6.47%	2[0.37,10.84]
Hoj 2005	10/330	4/331	+	+	13.98%	2.51[0.79,7.92]
Mobeen 2011	3/533	3/583			7.25%	1.09[0.22,5.4]
Subtotal (95% CI)	2432	2517	•	•	100%	1.41[0.92,2.16]
Total events: 53 (Misoprostol), 36 (Pl	acebo or no treatme	ent)				
Heterogeneity: Tau ² =0; Chi ² =4.8, df=	5(P=0.44); I ² =0%					
Test for overall effect: Z=1.56(P=0.12))					
31.12.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Place	ebo or no treatment)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	•					
	Fa	vours Misoprostol	0.01 0.1 1	10 100	Favours Placebo or r	no treatment





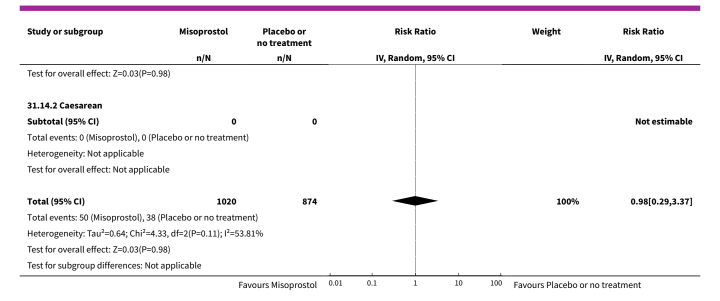
Analysis 31.13. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 13 Headache.



Analysis 31.14. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95% (CI			IV, Random, 95% CI
31.14.1 Vaginal birth									
Bamigboye 1998a	1/271	0/275			+			11.94%	3.04[0.12,74.4]
Hofmeyr 1998	2/250	7/250			-			30.86%	0.29[0.06,1.36]
Hofmeyr 2001	47/300	31/300			-			57.2%	1.52[0.99,2.32]
Uncu 2015	0/199	0/49							Not estimable
Subtotal (95% CI)	1020	874			*			100%	0.98[0.29,3.37]
Total events: 50 (Misoprostol), 38 (Placebo or no treatme	nt)							
Heterogeneity: Tau ² =0.64; Ch	i ² =4.33, df=2(P=0.11); l ² =53.8	81%							
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or i	no treatment



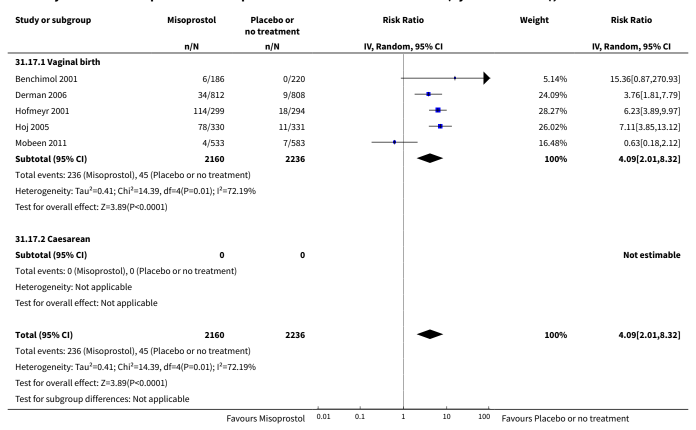


Analysis 31.16. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 16 Shivering.

Misoprostol	Placebo or no treatment			Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
1/34	4/36		0.96%	0.26[0.03,2.25]
5/186	0/220	+	0.53%	13[0.72,233.56]
419/812	140/808	-	29.5%	2.98[2.53,3.51]
48/250	13/250		9.6%	3.69[2.05,6.64]
133/300	33/300	-+-	18.39%	4.03[2.85,5.7]
189/330	78/331	-	26.21%	2.43[1.96,3.01]
50/533	23/583		12.69%	2.38[1.47,3.84]
7/31	1/34		1.06%	7.68[1,58.92]
10/199	1/49		1.06%	2.46[0.32,18.78]
2675	2611	•	100%	2.94[2.38,3.64]
293 (Placebo or no treatn	nent)			
14.21, df=8(P=0.08); l ² =43	.69%			
0.0001)				
0	0			Not estimable
Placebo or no treatment)				
able				
2675	2611	•	100%	2.94[2.38,3.64]
293 (Placebo or no treatn	nent)			
14.21, df=8(P=0.08); l ² =43	.69%			
0.0001)				
ot applicable				
	n/N 1/34 5/186 419/812 48/250 133/300 189/330 50/533 7/31 10/199 2675 293 (Placebo or no treatn 4.2.1, df=8(P=0.08); l²=43 0.0001) 0 Placebo or no treatment) able 2675 293 (Placebo or no treatment) 2675 293 (Placebo or no treatment) 2675 293 (Placebo or no treatment) 2675	1/34 4/36 5/186 0/220 419/812 140/808 48/250 13/250 133/300 33/300 189/330 78/331 50/533 23/583 7/31 1/34 10/199 1/49 2675 2611 293 (Placebo or no treatment) 4.21, df=8(P=0.08); l²=43.69% 0.0001) 1/2675 2611 293 (Placebo or no treatment) 293 (Placebo or no treatment) 293 (Placebo or no treatment) 294 (Placebo or no treatment) 295 (Placebo or no treatment) 297 (Placebo or no treatment) 298 (Placebo or no treatment) 299 (Placebo or no treatment) 299 (Placebo or no treatment)	no treatment n/N n/N n/N 1/34 4/36 5/186 0/220 419/812 140/808 48/250 13/250 133/300 33/300 189/330 78/331 50/533 23/583 7/31 1/34 10/199 1/49 2675 2611 293 (Placebo or no treatment) 4.2.1, df=8(P=0.08); l²=43.69% 0.0001) 1/34 1/34 1/35 1/35 1/35 1/36 1/36 1/36 1/36 1/36 1/36 1/36 1/36	no treatment n/N n/N n/N 1/34 4/36 5/186 0/220 419/812 140/808 48/250 133/300 33/300 189/330 78/331 50/533 23/583 7/31 1/34 10/99 1/49 2675 2611 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0



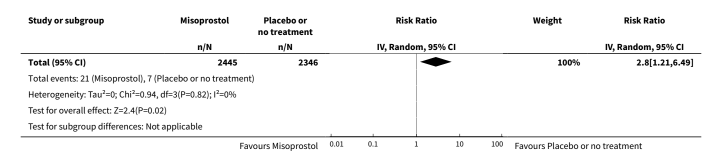
Analysis 31.17. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 17 Fever.



Analysis 31.18. Comparison 31 Misoprostol vs Placebo or no treatment (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
31.18.1 Vaginal birth					
Bamigboye 1998a	0/271	0/275			Not estimable
Derman 2006	9/812	2/808		30.27%	4.48[0.97,20.66]
Hofmeyr 2001	1/300	1/300		9.24%	1[0.06,15.91]
Hoj 2005	10/330	4/331		53.57%	2.51[0.79,7.92]
Mobeen 2011	1/533	0/583	+	6.92%	3.28[0.13,80.36]
Uncu 2015	0/199	0/49			Not estimable
Subtotal (95% CI)	2445	2346	•	100%	2.8[1.21,6.49]
Total events: 21 (Misoprostol), 7 (Plac	cebo or no treatmen	t)			
Heterogeneity: Tau ² =0; Chi ² =0.94, df	=3(P=0.82); I ² =0%				
Test for overall effect: Z=2.4(P=0.02)					
31.18.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Place	ebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
	Fa	vours Misoprostol	0.01 0.1 1 10	100 Favours Placebo or	no treatment





Comparison 32. Injectable prostaglandins vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.04, 3.24]
2.1 Vaginal birth	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.04, 3.24]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 vaginal birth	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.55 [0.22, 1.35]
6.1 Vaginal birth	1	46	Risk Ratio (IV, Random, 95% CI)	0.55 [0.22, 1.35]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size		
7 Additional uterotonics	2	146	Risk Ratio (IV, Random, 95% CI)	0.66 [0.21, 2.09]		
7.1 Vaginal birth	2	146	Risk Ratio (IV, Random, 95% CI)	0.66 [0.21, 2.09]		
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
8 Blood loss	2	146	Mean Difference (IV, Random, 95% CI)	-95.17 [-296.09, 105.75]		
8.1 Vaginal birth	2	146	Mean Difference (IV, Random, 95% CI)	-95.17 [-296.09, 105.75]		
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
9 Change in haemoglo- bin	1	100	Mean Difference (IV, Random, 95% CI)	0.90 [0.56, 1.24]		
9.1 Vaginal birth	1	100	Mean Difference (IV, Random, 95% CI)	0.90 [0.56, 1.24]		
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
11 Nausea	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.02, 8.46]		
11.1 Vaginal birth	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.02, 8.46]		
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]		

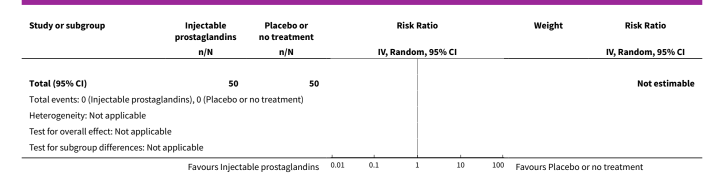


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

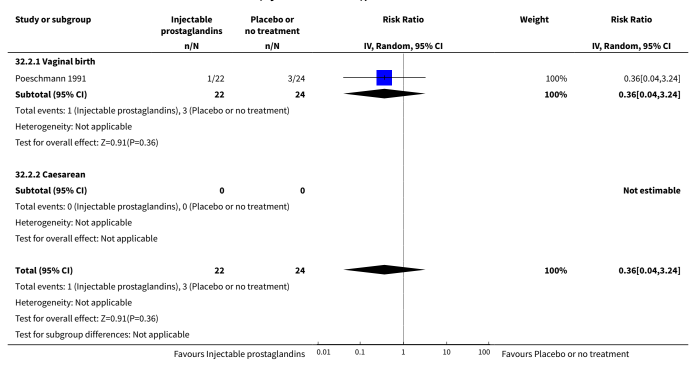
Analysis 32.1. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins				Weight	Risk Ratio		
	n/N	n/N		IV, Ra	ndom, 95% CI			IV, Random, 95% CI
32.1.1 Vaginal birth								
Supe 2016	0/50	0/50						Not estimable
Subtotal (95% CI)	50	50						Not estimable
Total events: 0 (Injectable pro	staglandins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable	2							
Test for overall effect: Not app	olicable							
32.1.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable pro	staglandins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable	e				ĺ			
Test for overall effect: Not app	olicable							
	Favours Injecta	ble prostaglandins	0.01	0.1	1 10	100	Favours Placebo c	r no treatment





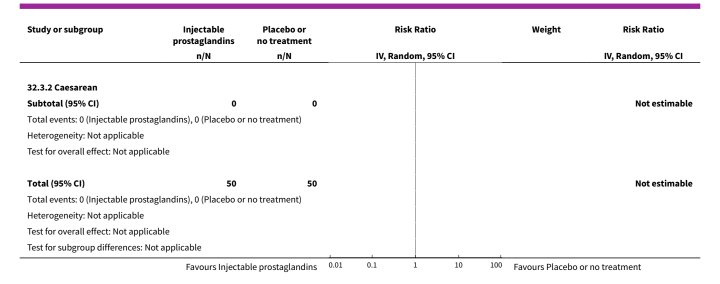
Analysis 32.2. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.



Analysis 32.3. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, F	Random, 95% (:I			IV, Random, 95% CI
32.3.1 vaginal birth									
Supe 2016	0/50	0/50							Not estimable
Subtotal (95% CI)	50	50							Not estimable
Total events: 0 (Injectable pro	staglandins), 0 (Placebo or	no treatment)							
Heterogeneity: Not applicable	e								
Test for overall effect: Not app	olicable								
	Favours Injecta	ole prostaglandins	0.01	0.1	1	10	100	Favours Placebo or I	no treatment



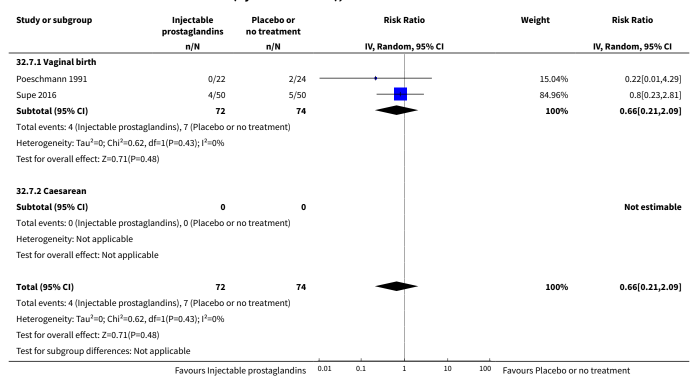


Analysis 32.6. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95	% CI		IV, Random, 95% CI
32.6.1 Vaginal birth							
Poeschmann 1991	5/22	10/24		-		100%	0.55[0.22,1.35]
Subtotal (95% CI)	22	24				100%	0.55[0.22,1.35]
Total events: 5 (Injectable prostag	landins), 10 (Placebo d	or no treatment)					
Heterogeneity: Not applicable							
Test for overall effect: Z=1.31(P=0.	19)						
32.6.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Injectable prostag	landins), 0 (Placebo or	no treatment)					
Heterogeneity: Not applicable							
Test for overall effect: Not applical	ble						
Total (95% CI)	22	24				100%	0.55[0.22,1.35]
Total events: 5 (Injectable prostag	landins), 10 (Placebo d	or no treatment)					
Heterogeneity: Not applicable							
Test for overall effect: Z=1.31(P=0.	19)			ĺ			
Test for subgroup differences: Not	applicable			ĺ			
	Favours Injecta	ble prostaglandins	0.01	0.1 1	10 100	Favours Placebo or i	no treatment



Analysis 32.7. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.

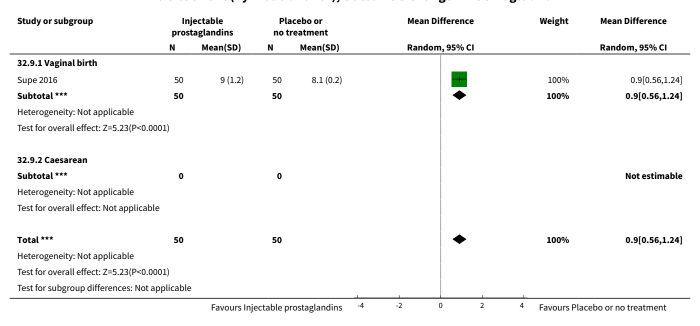


Analysis 32.8. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.

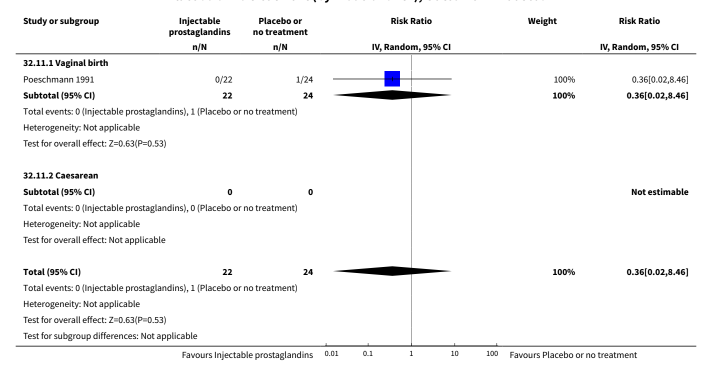
Study or subgroup		jectable taglandins		acebo or reatment		Mean Diffe	erence		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 9	5% CI			Random, 95% CI
32.8.1 Vaginal birth										
Poeschmann 1991	22	324 (302)	24	548 (376)	\leftarrow				38.77%	-224[-420.35,-27.65]
Supe 2016	50	153.8 (43.5)	50	167.4 (53)		-			61.23%	-13.6[-32.59,5.39]
Subtotal ***	72		74						100%	-95.17[-296.09,105.75]
Heterogeneity: Tau ² =17068.99; Chi ²	=4.37, df=	:1(P=0.04); I ² =77.	12%							
Test for overall effect: Z=0.93(P=0.3	5)									
32.8.2 Caesarean										
Subtotal ***	0		0							Not estimable
Heterogeneity: Not applicable										
Test for overall effect: Not applicab	le									
Total ***	72		74						100%	-95.17[-296.09,105.75]
Heterogeneity: Tau ² =17068.99; Chi ²	=4.37, df=	:1(P=0.04); I ² =77.	12%							
Test for overall effect: Z=0.93(P=0.3	5)									
Test for subgroup differences: Not a	pplicable	!								
		Favours Inje	ectable p	rostaglandins	-100	-50 0	50	100	Favours Pl	acebo or no treatment



Analysis 32.9. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.



Analysis 32.11. Comparison 32 Injectable prostaglandins vs Placebo or no treatment (by mode of birth), Outcome 11 Nausea.





Comparison 33. Ergometrine vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.09 [0.01, 0.72]
2.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	0.09 [0.01, 0.72]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1529	Risk Ratio (IV, Random, 95% CI)	0.34 [0.04, 3.28]
3.1 Vaginal birth	2	1529	Risk Ratio (IV, Random, 95% CI)	0.34 [0.04, 3.28]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.42]
6.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.42]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	1529	Risk Ratio (IV, Random, 95% CI)	0.18 [0.09, 0.37]
7.1 Vaginal birth	2	1529	Risk Ratio (IV, Random, 95% CI)	0.18 [0.09, 0.37]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	1529	Mean Difference (IV, Random, 95% CI)	-50.64 [-119.92, 18.65]
8.1 Vaginal birth	2	1529	Mean Difference (IV, Random, 95% CI)	-50.64 [-119.92, 18.65]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	1	100	Mean Difference (IV, Random, 95% CI)	-0.5 [-0.58, -0.42]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
9.1 Vaginal birth	1	100	Mean Difference (IV, Random, 95% CI)	-0.5 [-0.58, -0.42]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	1	1429	Risk Ratio (IV, Random, 95% CI)	42.10 [2.55, 694.80]	
11.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	42.10 [2.55, 694.80]	
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12 Vomiting	1	1429	Risk Ratio (IV, Random, 95% CI)	25.67 [1.52, 432.78]	
12.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	25.67 [1.52, 432.78]	
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13 Headache	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [0.37, 138.91]	
13.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [0.37, 138.91]	
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14 Abdominal pain	1	1429	Risk Ratio (IV, Random, 95% CI)	2.05 [1.04, 4.08]	
14.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	2.05 [1.04, 4.08]	
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15 Hypertension	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [2.83, 18.24]	
15.1 Vaginal birth	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [2.83, 18.24]	
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



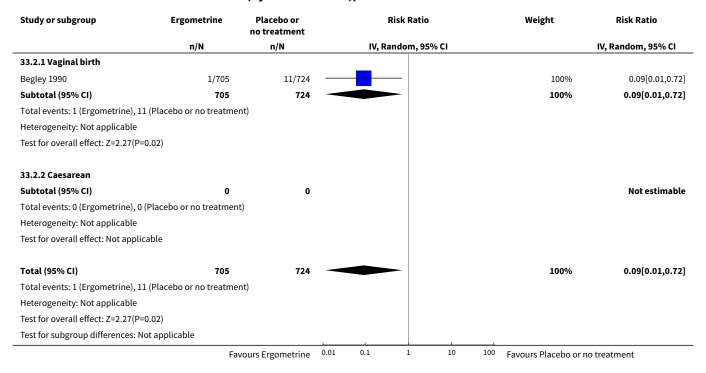
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Casarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 33.1. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 1 Death.

Study or subgroup	Ergometrine	Placebo or no treatment		Risl	k Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rand	om, 95% CI			IV, Random, 95% CI
33.1.1 Vaginal birth								
Supe 2016	0/50	0/50						Not estimable
Subtotal (95% CI)	50	50						Not estimable
Total events: 0 (Ergometrine), 0 (Place	ebo or no treatment	:)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
33.1.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine), 0 (Place	ebo or no treatment	:)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total (95% CI)	50	50						Not estimable
Total events: 0 (Ergometrine), 0 (Place	ebo or no treatment	:)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Test for subgroup differences: Not ap	plicable							
	Fav	ours Ergometrine	0.01	0.1	1 10	100	Favours Placebo or	no treatment



Analysis 33.2. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.

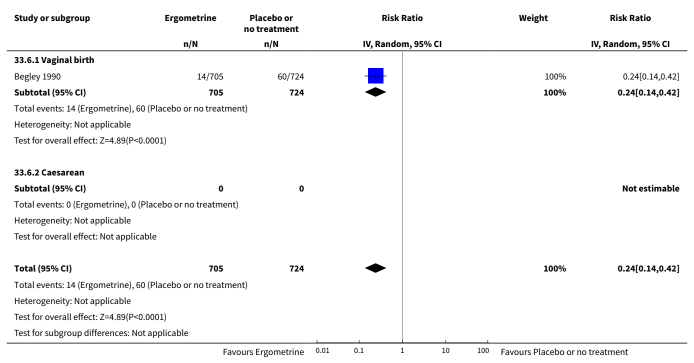


Analysis 33.3. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
33.3.1 Vaginal birth					
Begley 1990	1/705	3/724		100%	0.34[0.04,3.28]
Supe 2016	0/50	0/50	_		Not estimable
Subtotal (95% CI)	755	774		100%	0.34[0.04,3.28]
Total events: 1 (Ergometrine), 3 (Pla	cebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.93(P=0.35	5)				
33.3.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Pla	cebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	755	774		100%	0.34[0.04,3.28]
Total events: 1 (Ergometrine), 3 (Pla	cebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.93(P=0.35	5)				
Test for subgroup differences: Not a	pplicable				
	Favo	ours Ergometrine 0.0	1 0.1 1 10	100 Favours Placebo or r	no treatment



Analysis 33.6. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.



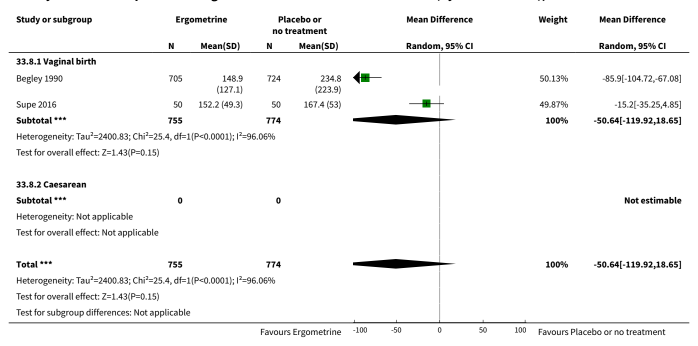
Analysis 33.7. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
33.7.1 Vaginal birth					
Begley 1990	14/705	93/724		82.12%	0.15[0.09,0.27]
Supe 2016	2/50	5/50		17.88%	0.4[0.08,1.97]
Subtotal (95% CI)	755	774	•	100%	0.18[0.09,0.37]
Total events: 16 (Ergometrine), 98	8 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0.08; Chi ² =1	.22, df=1(P=0.27); I ² =18.1	.9%			
Test for overall effect: Z=4.66(P<0	.0001)				
33.7.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (I	Placebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applica	able				
Total (95% CI)	755	774	•	100%	0.18[0.09,0.37]
Total events: 16 (Ergometrine), 98	8 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0.08; Chi ² =1	.22, df=1(P=0.27); I ² =18.1	.9%			
Test for overall effect: Z=4.66(P<0	.0001)				
	Fav	ours Ergometrine 0.01	0.1 1 10	LOO Favours Placebo or	no treatment



Study or subgroup	Ergometrine	netrine Placebo or no treatment		Risk Ratio				Weight Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI		IV, Random, 95% CI
Test for subgroup differences	: Not applicable		_				_	
		Favours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or no treatment

Analysis 33.8. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.



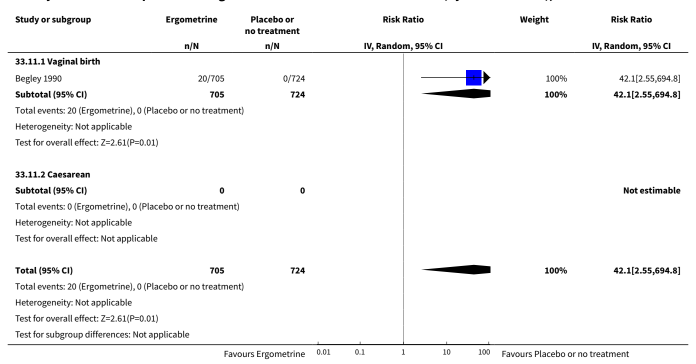
Analysis 33.9. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Erg	ometrine		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
33.9.1 Vaginal birth							
Supe 2016	50	7.6 (0.2)	50	8.1 (0.2)		100%	-0.5[-0.58,-0.42]
Subtotal ***	50		50		T	100%	-0.5[-0.58,-0.42]
Heterogeneity: Not applicable							
Test for overall effect: Z=12.5(P<0.0	001)						
33.9.2 Caesarean							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	le						
Total ***	50		50			100%	-0.5[-0.58,-0.42]
Heterogeneity: Not applicable							
Test for overall effect: Z=12.5(P<0.0	001)						
			Favour	s Ergometrine -100	-50 0 50	100 Favours Pla	cebo or no treatment



Study or subgroup	Erg			lacebo or treatment	Mean Difference			nce		Weight Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95%	6 CI		Random, 95% CI
Test for subgroup differences:	Not applicable				_					
			Favou	rs Ergometrine	-100	-50	0	50	100	Favours Placebo or no treatment

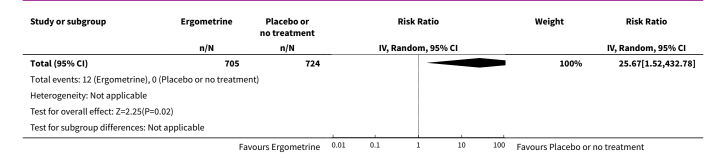
Analysis 33.11. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 11 Nausea.



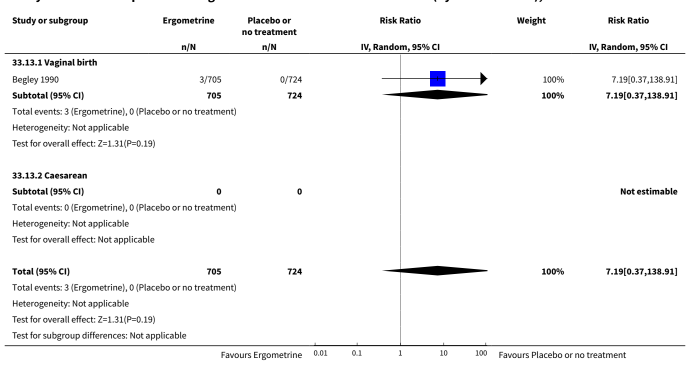
Analysis 33.12. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI		IV, Random, 95% CI
33.12.1 Vaginal birth						
Begley 1990	12/705	0/724			100%	25.67[1.52,432.78]
Subtotal (95% CI)	705	724			100%	25.67[1.52,432.78]
Total events: 12 (Ergometrine), 0 (Pla	acebo or no treatmen	t)				
Heterogeneity: Not applicable						
Test for overall effect: Z=2.25(P=0.02))					
33.12.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Ergometrine), 0 (Plac	cebo or no treatment)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	•					
	Fav	ours Ergometrine	0.01 0	.1 1 10 1	00 Favours Placebo or i	no treatment





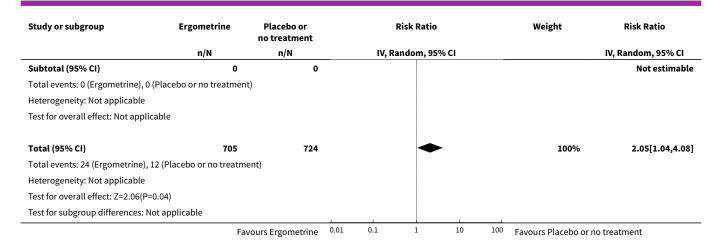
Analysis 33.13. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 13 Headache.



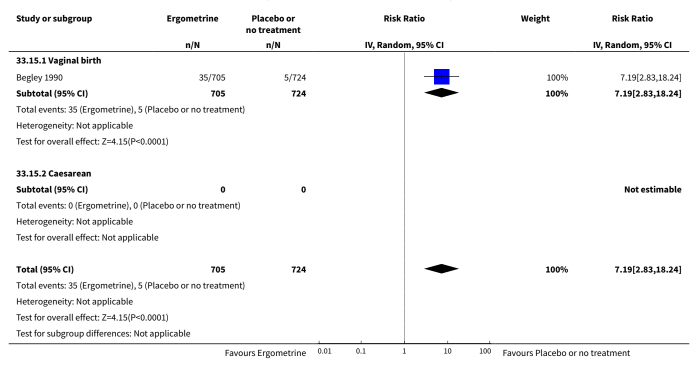
Analysis 33.14. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
33.14.1 Vaginal birth									
Begley 1990	24/705	12/724			-			100%	2.05[1.04,4.08]
Subtotal (95% CI)	705	724			•			100%	2.05[1.04,4.08]
Total events: 24 (Ergometrine)	, 12 (Placebo or no treatme	ent)							
Heterogeneity: Not applicable									
Test for overall effect: Z=2.06(F	P=0.04)								
33.14.2 Caesarean									
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Placebo or	no treatment





Analysis 33.15. Comparison 33 Ergometrine vs Placebo or no treatment (by mode of birth), Outcome 15 Hypertension.



Comparison 34. Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2 PPH >= 1000 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.44 [0.18, 1.05]
2.1 Vaginal birth	2	3207	Risk Ratio (IV, Random, 95% CI)	0.44 [0.18, 1.05]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	3	3400	Risk Ratio (IV, Random, 95% CI)	0.34 [0.18, 0.66]
3.1 Vaginal birth	3	3400	Risk Ratio (IV, Random, 95% CI)	0.34 [0.18, 0.66]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.37 [0.30, 0.46]
6.1 Vaginal birth	2	3207	Risk Ratio (IV, Random, 95% CI)	0.37 [0.30, 0.46]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	3	3400	Risk Ratio (IV, Random, 95% CI)	0.19 [0.15, 0.24]
7.1 Vaginal birth	3	3400	Risk Ratio (IV, Random, 95% CI)	0.19 [0.15, 0.24]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	1705	Mean Difference (IV, Random, 95% CI)	-35.02 [-101.63, 31.59]
8.1 Vaginal birth	2	1705	Mean Difference (IV, Random, 95% CI)	-35.02 [-101.63, 31.59]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	1454	Mean Difference (IV, Random, 95% CI)	-3.57 [-6.50, -0.63]
9.1 Vaginal birth	2	1454	Mean Difference (IV, Random, 95% CI)	-3.57 [-6.50, -0.63]



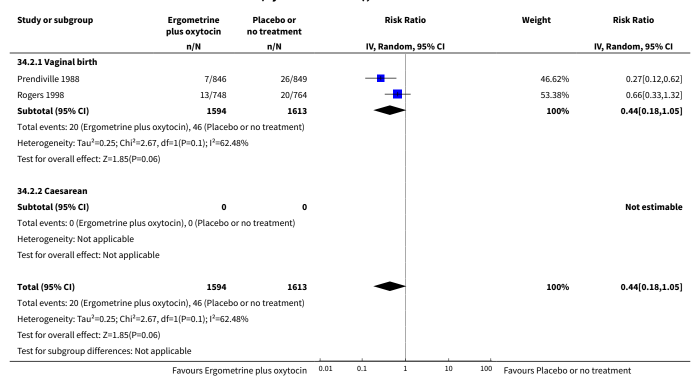
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	2	3207	Risk Ratio (IV, Random, 95% CI)	1.03 [0.99, 1.07]
10.1 Vaginal birth	2	3207	Risk Ratio (IV, Random, 95% CI)	1.03 [0.99, 1.07]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	1512	Risk Ratio (IV, Random, 95% CI)	1.95 [1.38, 2.76]
11.1 Vaginal birth	1	1512	Risk Ratio (IV, Random, 95% CI)	1.95 [1.38, 2.76]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	3207	Risk Ratio (IV, Random, 95% CI)	2.15 [1.46, 3.18]
12.1 Vaginal delivery	2	3207	Risk Ratio (IV, Random, 95% CI)	2.15 [1.46, 3.18]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	2	3207	Risk Ratio (IV, Random, 95% CI)	1.65 [0.78, 3.48]
13.1 Vaginal birth	2	3207	Risk Ratio (IV, Random, 95% CI)	1.65 [0.78, 3.48]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth - General health at 6 weeks postpar- tum (Worse than prepregnan- cy)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Vaginal birth - General health at 6 weeks postpar- tum (Exhausted since birth)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.3 Vaginal birth - General health at 6 weeks postpar- tum (Exhausted at 6 weeks)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.4 Vaginal birth - General health at 6 weeks postpar- tum (Blues)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.5 Vaginal birth - General health at 6 weeks postpar- tum (Depressed)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.6 Vaginal birth - General health at 6 weeks postpar- tum (Help for depression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.7 Vaginal birth - General health at 6 weeks postpar- tum (Admission to hospital for depression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.8 Vaginal birth - General health at 6 weeks postpar- tum (No health problems re- ported)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.9 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth - Satisfied with third-stage management	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Vaginal birth - Felt in control during third stage	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 34.2. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 2 PPH >= 1000 mL.



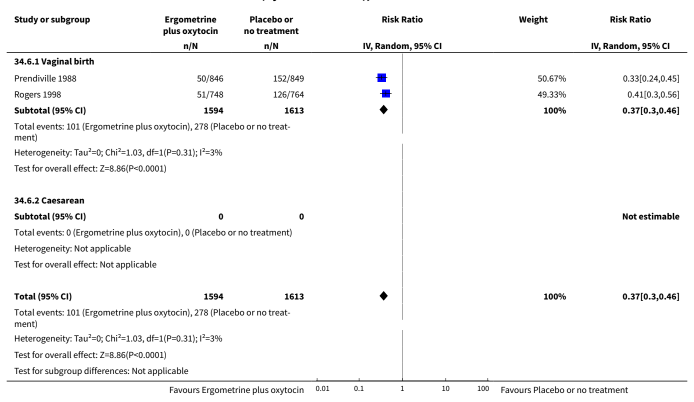
Analysis 34.3. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		Risk Ra	atio		Weight	Risk Ratio
	n/N	n/N		IV, Random	, 95% CI			IV, Random, 95% CI
34.3.1 Vaginal birth								
Prendiville 1988	18/846	48/849		-			66.88%	0.38[0.22,0.64]
Rogers 1998	4/748	20/764					28.96%	0.2[0.07,0.59]
Thilaganathan 1993	1/103	0/90			+		4.16%	2.63[0.11,63.64]
Subtotal (95% CI)	1697	1703		•			100%	0.34[0.18,0.66]
Total events: 23 (Ergometrine pl	us oxytocin), 68 (Placebo	o or no treatment)						
Heterogeneity: Tau ² =0.1; Chi ² =2.	.58, df=2(P=0.28); I ² =22.4	18%						
Test for overall effect: Z=3.18(P=	0)							
34.3.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine plu	ıs oxytocin), 0 (Placebo c	or no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applic	cable							
Total (95% CI)	1697	1703		•			100%	0.34[0.18,0.66]
Total events: 23 (Ergometrine pl	us oxytocin), 68 (Placeb	o or no treatment)						
Heterogeneity: Tau ² =0.1; Chi ² =2.	.58, df=2(P=0.28); I ² =22.4	18%						
Test for overall effect: Z=3.18(P=	(0)							
	Favours Ergome	trine plus oxytocin	0.01	0.1 1	10	100	Favours Placebo or r	no treatment



Study or subgroup	Ergometrine plus oxytocin	3			Risk Ratio	•		Weight Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI		IV, Random, 95% CI
Test for subgroup differences	s: Not applicable		_					
	Favours Fronme	atrine nlus avvtacin	0.01	0.1	1	10	100	Favours Placeho or no treatment

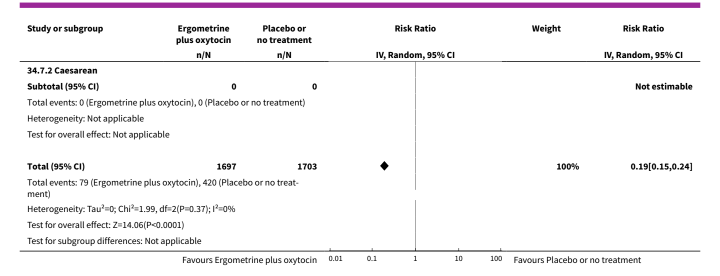
Analysis 34.6. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 6 PPH >= 500 mL.



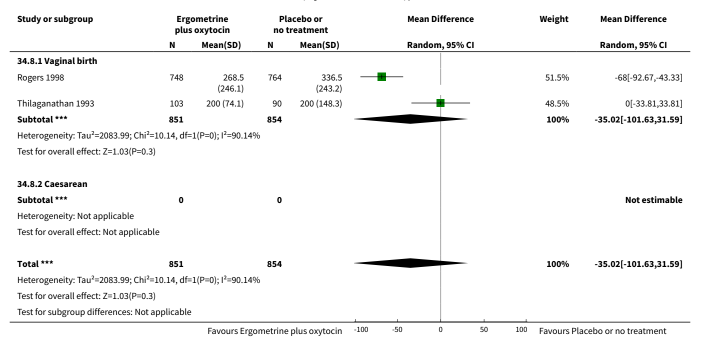
Analysis 34.7. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
34.7.1 Vaginal birth									
Prendiville 1988	54/846	252/849		-				68.36%	0.22[0.16,0.28]
Rogers 1998	24/748	161/764		-				30.42%	0.15[0.1,0.23]
Thilaganathan 1993	1/103	7/90						1.23%	0.12[0.02,1]
Subtotal (95% CI)	1697	1703		•				100%	0.19[0.15,0.24]
Total events: 79 (Ergometrine pment)	olus oxytocin), 420 (Placeb	oo or no treat-							
Heterogeneity: Tau ² =0; Chi ² =1.	99, df=2(P=0.37); I ² =0%								
Test for overall effect: Z=14.06(P<0.0001)								
				1		1	1		
	Favours Ergome	trine plus oxytocin	0.01	0.1	1	10	100	Favours Placebo or	no treatment





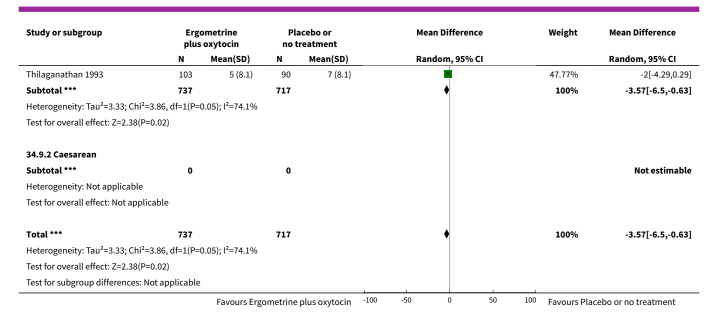
Analysis 34.8. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 8 Blood loss.



Analysis 34.9. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup		ometrine oxytocin		cebo or reatment		М	ean Differe	ence		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ndom, 95	% CI			Random, 95% CI
34.9.1 Vaginal birth											
Prendiville 1988	634	1 (21)	627	6 (13)			+			52.23%	-5[-6.93,-3.07]
		Favours Ergometrine plus oxytocin			-100	-50	0	50	100	Favours Pla	cebo or no treatment



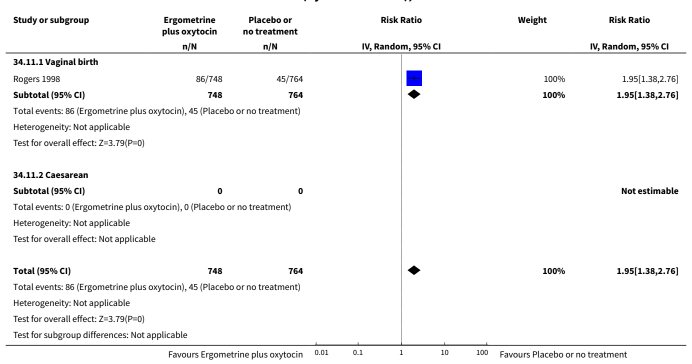


Analysis 34.10. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 10 Breastfeeding.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
34.10.1 Vaginal birth					
Prendiville 1988	637/846	632/849	•	57.43%	1.01[0.96,1.07]
Rogers 1998	546/748	531/764	•	42.57%	1.05[0.99,1.12]
Subtotal (95% CI)	1594	1613	•	100%	1.03[0.99,1.07]
Total events: 1183 (Ergometrine plus ment)	oxytocin), 1163 (Pla	acebo or no treat-			
Heterogeneity: Tau ² =0; Chi ² =0.76, df	=1(P=0.38); I ² =0%				
Test for overall effect: Z=1.29(P=0.2)					
34.10.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus ox	ytocin), 0 (Placebo d	or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	•				
Total (95% CI)	1594	1613	!	100%	1.03[0.99,1.07]
Total events: 1183 (Ergometrine plus ment)	oxytocin), 1163 (Pla	acebo or no treat-			
Heterogeneity: Tau ² =0; Chi ² =0.76, df	=1(P=0.38); I ² =0%				
Test for overall effect: Z=1.29(P=0.2)					
Test for subgroup differences: Not ap	plicable				
	Favours Ergome	trine plus oxytocin 0.0	1 0.1 1 10 1	100 Favours Placebo or	no treatment



Analysis 34.11. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 11 Nausea.

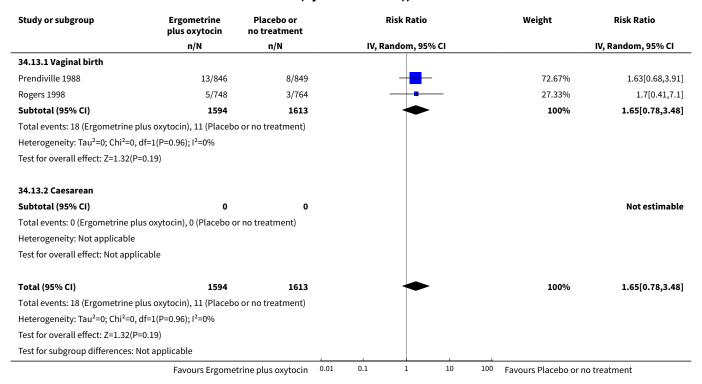


Analysis 34.12. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		ı	Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95% CI			IV, Random, 95% CI
34.12.1 Vaginal delivery								
Prendiville 1988	102/846	55/849			-		64.91%	1.86[1.36,2.55]
Rogers 1998	47/748	17/764			-		35.09%	2.82[1.64,4.87]
Subtotal (95% CI)	1594	1613			•		100%	2.15[1.46,3.18]
Total events: 149 (Ergometrine plus ment)	oxytocin), 72 (Placel	oo or no treat-						
Heterogeneity: Tau ² =0.04; Chi ² =1.69	df=1(P=0.19); I ² =40	.7%						
Test for overall effect: Z=3.86(P=0)								
34.12.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine plus ox	ytocin), 0 (Placebo o	r no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
Total (95% CI)	1594	1613			•		100%	2.15[1.46,3.18]
Total events: 149 (Ergometrine plus ment)	oxytocin), 72 (Placeb	oo or no treat-						
Heterogeneity: Tau ² =0.04; Chi ² =1.69	df=1(P=0.19); I ² =40	.7%						
Test for overall effect: Z=3.86(P=0)								
Test for subgroup differences: Not ap	plicable							
	Favours Ergome	trine plus oxytocin	0.01	0.1	1 1	.0 100	Favours Placebo or i	no treatment



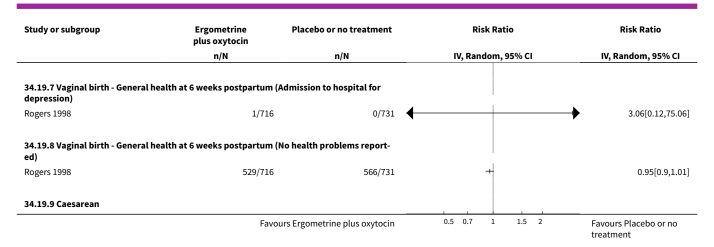
Analysis 34.13. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 13 Headache.



Analysis 34.19. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 19 Maternal well-being.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI	
34.19.1 Vaginal birth - Genera	l health at 6 weeks postpartum (Worse than prepregnancy)			
Rogers 1998	64/716	66/731		0.99[0.71,1.37]	
34.19.2 Vaginal birth - Genera	l health at 6 weeks postpartum (Exhausted since birth)			
Rogers 1998	163/716	175/731		0.95[0.79,1.15]	
34.19.3 Vaginal birth - Genera	l health at 6 weeks postpartum (Exhausted at 6 weeks)			
Rogers 1998	105/716	113/731		0.95[0.74,1.21]	
34.19.4 Vaginal birth - Genera	l health at 6 weeks postpartum (Blues)			
Rogers 1998	313/716	343/731	+	0.93[0.83,1.04]	
34.19.5 Vaginal birth - Genera	l health at 6 weeks postpartum (Depressed)			
Rogers 1998	55/716	46/731	+	1.22[0.84,1.78]	
34.19.6 Vaginal birth - Genera	l health at 6 weeks postpartum (Help for depression)			
Rogers 1998	107/716	104/731		1.05[0.82,1.35]	
	Favo	urs Ergometrine plus oxytocin	0.5 0.7 1 1.5 2	Favours Placebo or no treatment	





Analysis 34.20. Comparison 34 Ergometrine plus oxytocin vs Placebo or no treatment (by mode of birth), Outcome 20 Maternal satisfaction.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio	io Risk Ratio	
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI	
34.20.1 Vaginal birth - Satisfie	ed with third-stage management				
Rogers 1998	721/745	718/762	-	1.03[1,1.05]	
34.20.2 Vaginal birth - Felt in o	control during third stage				
Rogers 1998	621/745	667/762		0.95[0.91,0.99]	
34.20.3 Caesarean					
	Favou	urs Ergometrine plus oxytocin	1	Favours Placebo or no treatment	

Comparison 35. Misoprostol plus oxytocin vs Placebo or no treatment (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.1 Vaginl birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI) 0.0 [0.0, 0		
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Comparison 36. Misoprostol vs Oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	24	28520	Risk Ratio (IV, Random, 95% CI)	0.62 [0.14, 2.74]
1.1 Vaginal birth	21	27955	Risk Ratio (IV, Random, 95% CI)	0.74 [0.14, 3.95]
1.2 Caesarean	3	565	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.09]
2 PPH >= 1000 mL	38	34261	Risk Ratio (IV, Random, 95% CI)	1.26 [1.11, 1.43]
2.1 Vaginal birth	31	33496	Risk Ratio (IV, Random, 95% CI)	1.31 [1.15, 1.49]
2.2 Caesarean	7	765	Risk Ratio (IV, Random, 95% CI)	0.83 [0.54, 1.26]
3 Blood transfusion	42	35467	Risk Ratio (IV, Random, 95% CI)	0.81 [0.65, 1.00]
3.1 Vaginal birth	34	34417	Risk Ratio (IV, Random, 95% CI)	0.83 [0.67, 1.03]
3.2 Caesarean	8	1050	Risk Ratio (IV, Random, 95% CI)	0.48 [0.19, 1.21]
4 Severe maternal morbidity: intensive care admissions	10	21698	Risk Ratio (IV, Random, 95% CI)	1.16 [0.55, 2.43]
4.1 Vaginal birth	9	21508	Risk Ratio (IV, Random, 95% CI)	1.16 [0.55, 2.43]
4.2 Caesarean	1	190	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	44	36920	Risk Ratio (IV, Random, 95% CI)	1.08 [0.94, 1.24]
6.1 Vaginal birth	38	36215	Risk Ratio (IV, Random, 95% CI)	1.08 [0.90, 1.31]
6.2 Caesarean	6	705	Risk Ratio (IV, Random, 95% CI)	1.07 [0.92, 1.25]
7 Additional uterotonics	47	35981	Risk Ratio (IV, Random, 95% CI)	1.01 [0.85, 1.20]
7.1 Vaginal birth	35	34521	Risk Ratio (IV, Random, 95% CI)	1.06 [0.86, 1.30]
7.2 Caesarean	12	1460	Risk Ratio (IV, Random, 95% CI)	0.89 [0.69, 1.16]
8 Blood loss	43	35239	Mean Difference (IV, Random, 95% CI)	-8.90 [-23.45, 5.65]
8.1 Vaginal birth	34	34339	Mean Difference (IV, Random, 95% CI)	-2.08 [-16.92, 12.77]
8.2 Caesarean	9	900	Mean Difference (IV, Random, 95% CI)	-59.79 [-89.04, -30.54]
9 Change in haemoglo- bin	31	12028	Mean Difference (IV, Random, 95% CI)	-0.14 [-0.74, 0.47]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.1 Vaginal birth	24	11240	Mean Difference (IV, Random, 95% CI)	-0.00 [-0.62, 0.62]
9.2 Caesarean	7	788	Mean Difference (IV, Random, 95% CI)	-0.76 [-2.26, 0.73]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	33	29732	Risk Ratio (IV, Random, 95% CI)	1.22 [0.93, 1.60]
11.1 Vaginal birth	26	28907	Risk Ratio (IV, Random, 95% CI)	1.30 [0.94, 1.79]
11.2 Caesarean	7	825	Risk Ratio (IV, Random, 95% CI)	0.97 [0.61, 1.54]
12 Vomiting	41	32687	Risk Ratio (IV, Random, 95% CI)	1.51 [1.19, 1.91]
12.1 Vaginal birth	30	31402	Risk Ratio (IV, Random, 95% CI)	1.82 [1.43, 2.33]
12.2 Caesarean	11	1285	Risk Ratio (IV, Random, 95% CI)	0.89 [0.55, 1.45]
13 Headache	10	4079	Risk Ratio (IV, Random, 95% CI)	0.88 [0.54, 1.42]
13.1 Vaginal birth	5	3494	Risk Ratio (IV, Random, 95% CI)	1.42 [0.99, 2.04]
13.2 Caesarean	5	585	Risk Ratio (IV, Random, 95% CI)	0.47 [0.19, 1.15]
14 Abdominal pain	8	3382	Risk Ratio (IV, Random, 95% CI)	0.91 [0.79, 1.06]
14.1 Vaginal birth	7	3207	Risk Ratio (IV, Random, 95% CI)	0.87 [0.73, 1.04]
14.2 Caesarean	1	175	Risk Ratio (IV, Random, 95% CI)	1.03 [0.78, 1.37]
15 Hypertension	3	1028	Risk Ratio (IV, Random, 95% CI)	3.64 [0.60, 22.27]
15.1 Vaginal birth	2	828	Risk Ratio (IV, Random, 95% CI)	4.00 [0.44, 36.03]
15.2 Caesarean	1	200	Risk Ratio (IV, Random, 95% CI)	3.0 [0.12, 72.77]
16 Shivering	49	34865	Risk Ratio (IV, Random, 95% CI)	4.02 [3.23, 4.99]
16.1 Vaginal birth	36	33355	Risk Ratio (IV, Random, 95% CI)	4.16 [3.27, 5.29]
16.2 Caesarean	13	1510	Risk Ratio (IV, Random, 95% CI)	3.58 [2.06, 6.21]
17 Fever	41	33008	Risk Ratio (IV, Random, 95% CI)	3.75 [2.73, 5.15]
17.1 Vaginal birth	32	31858	Risk Ratio (IV, Random, 95% CI)	4.62 [3.33, 6.42]
17.2 Caesarean	9	1150	Risk Ratio (IV, Random, 95% CI)	1.52 [0.80, 2.87]
18 Diarrhoea	26	30733	Risk Ratio (IV, Random, 95% CI)	2.13 [1.55, 2.93]

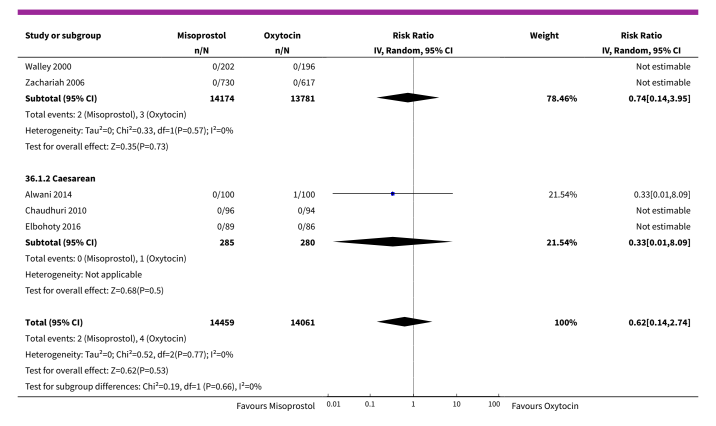


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
18.1 Vaginal birth	26	30733	Risk Ratio (IV, Random, 95% CI)	2.13 [1.55, 2.93]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Satisfied or very satisfied with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Complaints about or problems with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Would take drug again after subsequent deliveries	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.4 Would recommend drug to a friend	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 36.1. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.1.1 Vaginal birth					
Afolabi 2010	0/100	0/100			Not estimable
Amin 2014	0/100	0/100			Not estimable
Atukunda 2014	0/570	0/570			Not estimable
Baskett 2007	0/311	0/311			Not estimable
Bellad 2012	0/321	0/331			Not estimable
Bhatti 2014	0/60	0/60			Not estimable
Chaudhuri 2012	0/265	0/265			Not estimable
Diop 2016	0/252	0/141			Not estimable
Gulmezoglu 2001	2/9225	2/9230		57.02%	1[0.14,7.1]
Kundodyiwa 2001	0/243	0/256			Not estimable
Lumbiganon 1999	0/397	0/200			Not estimable
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	0/257	0/257			Not estimable
Oboro 2003	0/247	0/249			Not estimable
Parsons 2006	0/225	0/225			Not estimable
Parsons 2007	0/224	1/226	•	21.45%	0.34[0.01,8.21]
Perez-Rumbos 2017	0/195	0/197			Not estimable
Shrestha 2011	0/100	0/100			Not estimable
Tewatia 2014	0/50	0/50			Not estimable
Tematia 2011	·	ours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

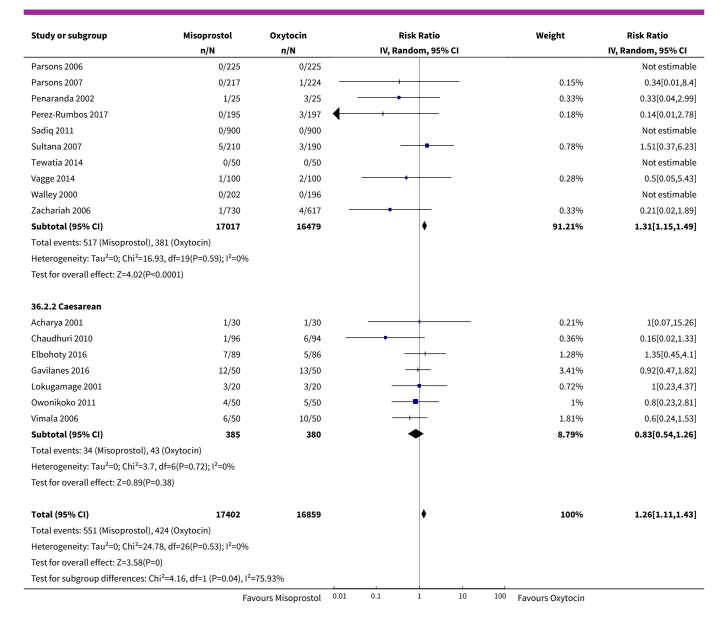




Analysis 36.2. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	n/N IV, Random, 95% CI			IV, Random, 95% CI
36.2.1 Vaginal birth						
Afolabi 2010	0/100	0/100				Not estimable
Al-Sawaf 2013	2/28	1/37		+	0.29%	2.64[0.25,27.7]
Atukunda 2014	18/570	14/570		+-	3.32%	1.29[0.65,2.56]
Baskett 2007	14/311	7/311		+	1.97%	2[0.82,4.89]
Bellad 2012	0/321	0/331				Not estimable
Benchimol 2001	16/186	12/196		+	3.03%	1.41[0.68,2.89]
Bhatti 2014	0/60	0/60				Not estimable
Bugalho 2001	0/323	1/339	-	+	0.15%	0.35[0.01,8.56]
Caliskan 2002	17/396	14/407			3.27%	1.25[0.62,2.5]
Caliskan 2003	14/388	15/384			3.08%	0.92[0.45,1.89]
Chaudhuri 2012	1/265	2/265		+	0.27%	0.5[0.05,5.48]
Cook 1999	13/424	0/129		+	0.2%	8.26[0.49,137.98]
Diallo 2017	2/154	4/150		-+-	0.56%	0.49[0.09,2.62]
Gerstenfeld 2001	15/154	14/161			3.27%	1.12[0.56,2.24]
Gulmezoglu 2001	366/9214	263/9228		+	64.84%	1.39[1.19,1.63]
Gupta 2006	0/100	0/100				Not estimable
Kundodyiwa 2001	9/243	5/256			1.35%	1.9[0.64,5.58]
Lumbiganon 1999	22/397	13/200			3.57%	0.85[0.44,1.66]
Modi 2014	0/25	0/25		ĺ		Not estimable
Nasr 2009	0/257	0/257		į		Not estimable
Oboro 2003	0/247	0/249		ĺ		Not estimable
	Fav	ours Misoprostol	0.01 0.1	1 10	100 Favours Oxytocin	

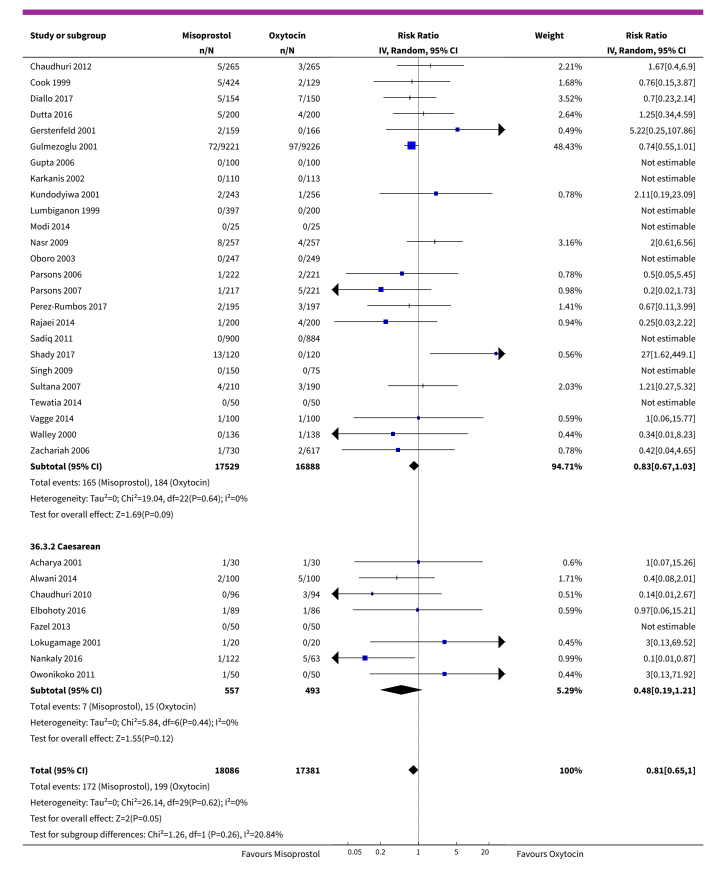




Analysis 36.3. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

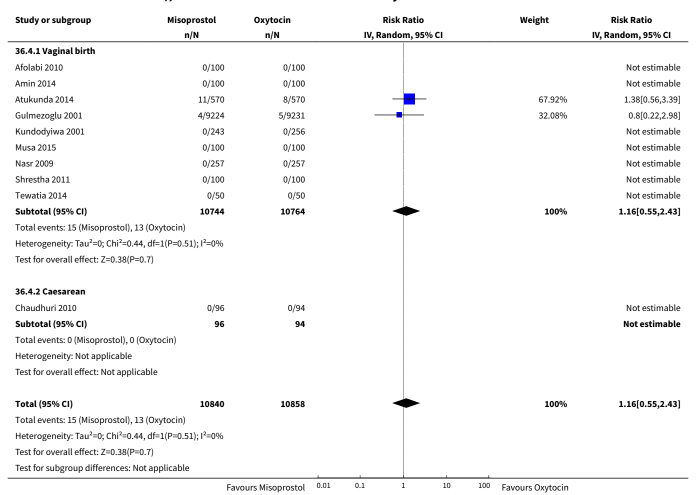
Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.3.1 Vaginal birth					
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	0/28	0/37			Not estimable
Atukunda 2014	7/570	16/570		5.75%	0.44[0.18,1.06]
Baskett 2007	0/311	0/311			Not estimable
Bellad 2012	1/321	1/331		0.58%	1.03[0.06,16.42]
Bhatti 2014	1/60	1/60		0.59%	1[0.06,15.62]
Bugalho 2001	2/323	1/339	-	0.78%	2.1[0.19,23.04]
Caliskan 2002	12/396	13/407		7.48%	0.95[0.44,2.05]
Caliskan 2003	14/388	13/384	· · · · · · · · · · · · · · · · · · ·	8.11%	1.07[0.51,2.24]
	Fav	ours Misoprostol	0.05 0.2 1 5 20	Favours Oxytocin	







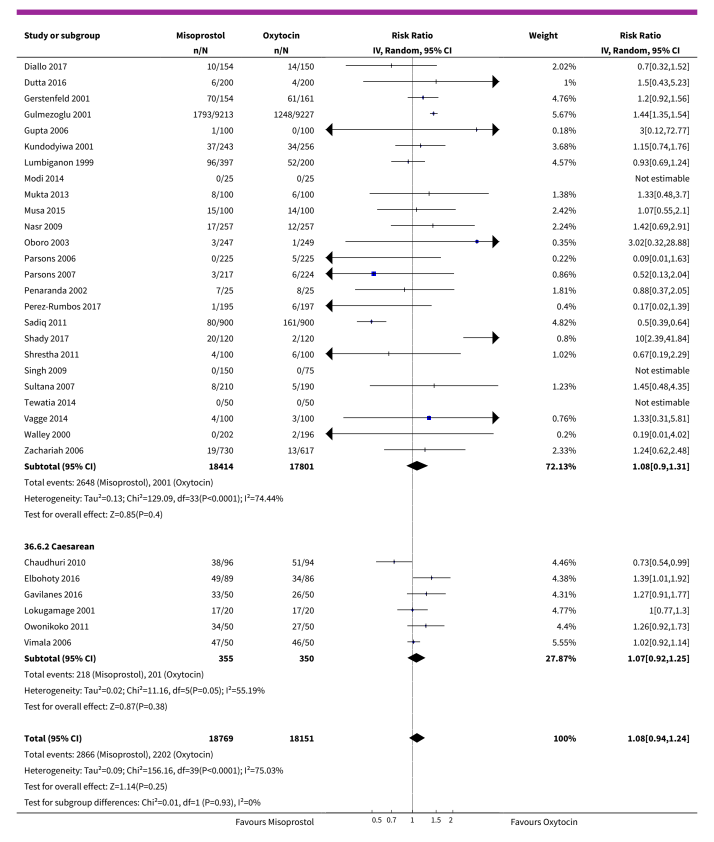
Analysis 36.4. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.



Analysis 36.6. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
36.6.1 Vaginal birth						
Afolabi 2010	0/100	0/100			Not estimable	
Al-Sawaf 2013	3/28	2/37		0.58%	1.98[0.35,11.08]	
Amin 2014	4/100	3/100		0.76%	1.33[0.31,5.81]	
Asmat 2017	123/839	120/839		4.94%	1.02[0.81,1.29]	
Atukunda 2014	89/570	57/570		4.44%	1.56[1.14,2.13]	
Bellad 2012	10/321	30/331	——	2.32%	0.34[0.17,0.69]	
Benchimol 2001	52/186	29/196		3.83%	1.89[1.26,2.84]	
Bhatti 2014	2/60	5/60	+ -	0.66%	0.4[0.08,1.98]	
Bugalho 2001	10/323	15/339		2.01%	0.7[0.32,1.53]	
Caliskan 2002	39/396	33/407		3.61%	1.21[0.78,1.89]	
Caliskan 2003	35/388	28/384		3.41%	1.24[0.77,1.99]	
Chaudhuri 2012	16/265	15/265		2.38%	1.07[0.54,2.11]	
Cook 1999	63/424	1/129		0.45%	19.17[2.68,136.84]	
	Fav	ours Misoprostol	0.5 0.7 1 1.5 2	Favours Oxytocin		



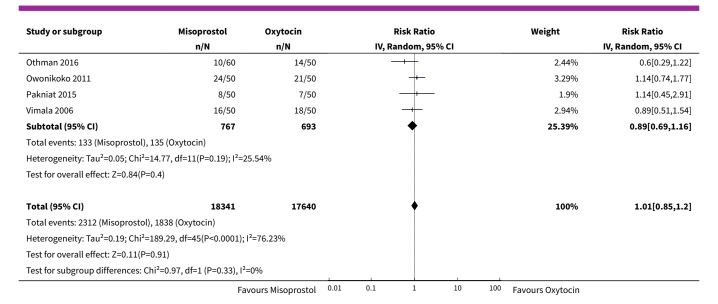




Analysis 36.7. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.7.1 Vaginal birth					
Afolabi 2010	3/100	4/100		1.06%	0.75[0.17,3.27]
Al-Sawaf 2013	3/28	2/37		0.83%	1.98[0.35,11.08]
Atukunda 2014	47/570	31/570	 	3.28%	1.52[0.98,2.35]
Baskett 2007	159/311	126/311	+	3.97%	1.26[1.06,1.5]
Bellad 2012	1/321	8/331	+	0.61%	0.13[0.02,1.02]
Bhatti 2014	1/60	3/60		0.54%	0.33[0.04,3.11]
Bugalho 2001	7/323	7/339		1.69%	1.05[0.37,2.96]
Caliskan 2002	33/396	26/407	+-	3.11%	1.3[0.8,2.14]
Caliskan 2003	23/388	26/384	-	2.96%	0.88[0.51,1.51]
Chaudhuri 2012	20/265	23/265	 -	2.86%	0.87[0.49,1.54]
Cook 1999	95/424	6/129		2.22%	4.82[2.16,10.73]
Diallo 2017	7/154	6/150		1.63%	1.14[0.39,3.3]
Gerstenfeld 2001	36/159	18/166		3.02%	2.09[1.24,3.52]
Gulmezoglu 2001	1398/9225	1002/9228	+	4.1%	1.4[1.29,1.51]
Gupta 2006	5/100	1/100		0.58%	5[0.59,42.04]
Karkanis 2002	28/110	20/113		3.06%	1.44[0.86,2.4]
Kundodyiwa 2001	13/243	7/256	<u> </u>	1.97%	1.96[0.79,4.82]
Lumbiganon 1999	41/397	28/200		3.25%	0.74[0.47,1.16]
Modi 2014	0/25	0/25		3.2370	Not estimable
Mukta 2013	22/100	16/100		2.84%	1.38[0.77,2.46]
Musa 2015				2.84%	
Nasr 2009	20/100	19/100		1.33%	1.05[0.6,1.85]
Oboro 2003	6/257	4/257			1.5[0.43,5.25]
	31/247	27/249		3.14%	1.16[0.71,1.88]
Parsons 2006	16/225	21/225		2.71%	0.76[0.41,1.42]
Parsons 2007	9/223	19/224		2.3%	0.48[0.22,1.03]
Perez-Rumbos 2017	7/195	22/197		2.15%	0.32[0.14,0.74]
Rajaei 2014	9/200	21/200	.	2.34%	0.43[0.2,0.91]
Sadiq 2011	32/900	148/900	-	3.49%	0.22[0.15,0.31]
Shady 2017	20/120	2/120		1.1%	10[2.39,41.84]
Singh 2009	2/150	2/75		0.68%	0.5[0.07,3.48]
Sultana 2007	5/210	6/190		1.45%	0.75[0.23,2.43]
Tewatia 2014	7/50	3/50	+-	1.27%	2.33[0.64,8.51]
Vagge 2014	4/100	3/100	- 	1.06%	1.33[0.31,5.81]
Walley 2000	6/168	8/172		1.69%	0.77[0.27,2.17]
Zachariah 2006	63/730	38/617	 	3.43%	1.4[0.95,2.07]
Subtotal (95% CI)	17574	16947	†	74.61%	1.06[0.86,1.3]
Total events: 2179 (Misoprostol)	, 1703 (Oxytocin)				
Heterogeneity: Tau ² =0.21; Chi ² =	166.61, df=33(P<0.0001); I	2=80.19%			
Test for overall effect: Z=0.52(P=	0.6)				
36.7.2 Caesarean					
Acharya 2001	2/30	3/30		0.83%	0.67[0.12,3.71]
Alwani 2014	4/100	9/100		1.5%	0.44[0.14,1.4]
Chaudhuri 2010	11/96	14/94	+	2.39%	0.77[0.37,1.61]
Eftekhari 2009	7/50	16/50		2.23%	0.44[0.2,0.97]
Elbohoty 2016	20/89	11/86	+-	2.57%	1.76[0.9,3.45]
Gavilanes 2016	10/50	12/50	 -	2.37%	0.83[0.4,1.75]
Lokugamage 2001	6/20	1/20	+	0.64%	6[0.79,45.42]
Nankaly 2016	15/122	9/63	 	2.3%	0.86[0.4,1.86]
	Fav	ours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

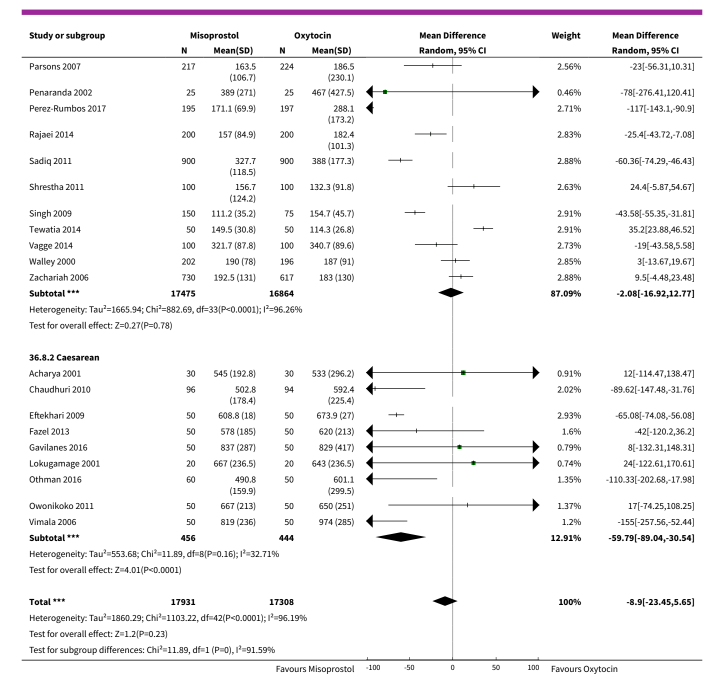




Analysis 36.8. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Mis	oprostol	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
36.8.1 Vaginal birth							
Afolabi 2010	100	153.2 (58)	100	155.6 (58)		2.86%	-2.4[-18.47,13.67]
Al-Sawaf 2013	28	348 (112)	37	314.7 (94.6)		2.16%	33.3[-18.18,84.78]
Amin 2014	100	300 (262.8)	100	250 (262.8)		1.7%	50[-22.83,122.83]
Asmat 2017	839	322 (199.9)	839	337 (211.4)	- + 	2.81%	-15[-34.69,4.69]
Atukunda 2014	570	341.5 (206.2)	570	304.2 (190.8)		2.76%	37.3[14.24,60.36]
Bellad 2012	321	192 (124)	331	366 (135.9)		2.81%	-174[-193.96,-154.04]
Benchimol 2001	186	374 (238.4)	196	278 (254)		2.21%	96[46.63,145.37]
Bhatti 2014	60	200 (125)	60	360 (136)		2.27%	-160[-206.74,-113.26]
Bugalho 2001	323	155 (122)	339	157.3 (138.7)		2.81%	-2.3[-22.17,17.57]
Caliskan 2003	388	328 (152)	384	312 (176)	+	2.76%	16[-7.21,39.21]
Chaudhuri 2012	265	153.2 (143.5)	265	146.9 (158.5)		2.71%	6.3[-19.45,32.05]
Cook 1999	424	279 (300.6)	129	98 (71.6)		2.61%	181[149.83,212.17]
Dasuki 2002	98	238.7 (94.5)	98	225.9 (94.5)	+-	2.7%	12.86[-13.61,39.33]
Diallo 2017	154	196.5 (210)	150	208.4 (324)		1.94%	-11.9[-73.45,49.65]
Dutta 2016	200	185.7 (84.4)	200	168.5 (68.4)		2.87%	17.2[2.14,32.26]
Gulmezoglu 2001	9213	332.8 (274.6)	9227	289.7 (262.1)	+	2.94%	43.1[35.35,50.85]
Gupta 2006	100	161.7 (76.8)	100	151 (69.1)	+-	2.8%	10.7[-9.56,30.96]
Kundodyiwa 2001	243	354 (99.3)	256	348 (99.3)	+-	2.84%	6[-11.42,23.42]
Lumbiganon 1999	397	355.9 (310.8)	200	353 (310)		2.13%	2.86[-49.87,55.59]
Modi 2014	25	255.8 (102.2)	25	223.2 (122.5)		1.92%	32.6[-29.94,95.14]
Musa 2015	100	325.9 (164.7)	100	304 (163.3)		2.3%	21.9[-23.56,67.36]
Oboro 2003	247	341 (19.3)	249	339 (18.9)	+	2.96%	2[-1.36,5.36]
Parsons 2006	225	150 (74.1)	225	150 (74.1)		2.89%	0[-13.69,13.69]

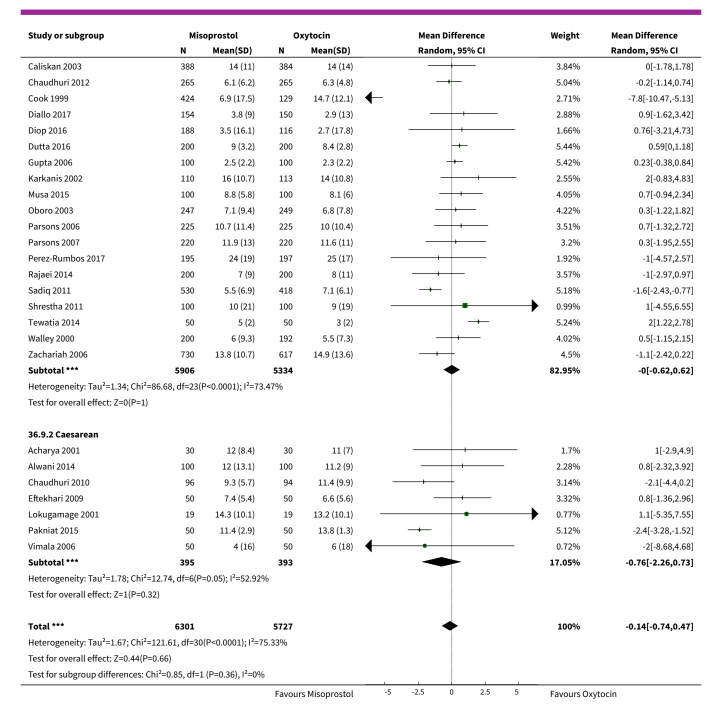




Analysis 36.9. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol	0:	xytocin	Mean Di	ifference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Randon	1, 95% CI		Random, 95% CI
36.9.1 Vaginal birth								
Afolabi 2010	100	3 (10.2)	100	4 (10.2)			2.55%	-1[-3.83,1.83]
Al-Sawaf 2013	28	13 (9)	37	12 (9)		-	1.42%	1[-3.42,5.42]
Atukunda 2014	570	9.4 (14.3)	569	9.6 (16.7)		 	3.8%	-0.2[-2.01,1.61]
Benchimol 2001	186	9.6 (23.5)	196	6.7 (20.1)		-	1.43%	2.9[-1.5,7.3]
Caliskan 2002	396	15 (12)	407	14 (14)		 	3.81%	1[-0.8,2.8]
			Favour	s Misoprostol	-5 -2.5	0 2.5	5 Favours Oxy	tocin

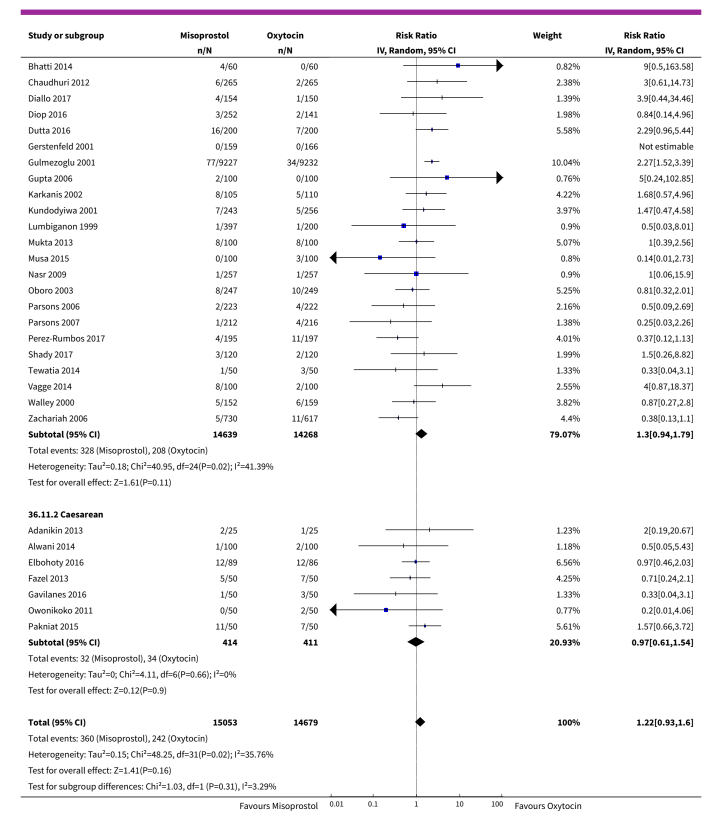




Analysis 36.11. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Misoprostol	Oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Randon	n, 95% CI			IV, Random, 95% CI
36.11.1 Vaginal birth								
Afolabi 2010	4/100	0/100		-		-	0.82%	9[0.49,165]
Atukunda 2014	138/570	86/570			+		11.67%	1.6[1.26,2.05]
Bellad 2012	12/321	0/331					0.87%	25.78[1.53,433.55]
	Fav	ours Misoprostol	0.01 0	.1 1	10	100	Favours Oxytocin	



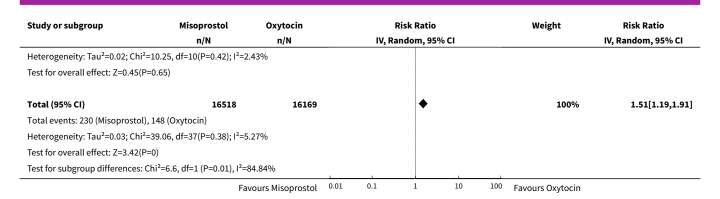




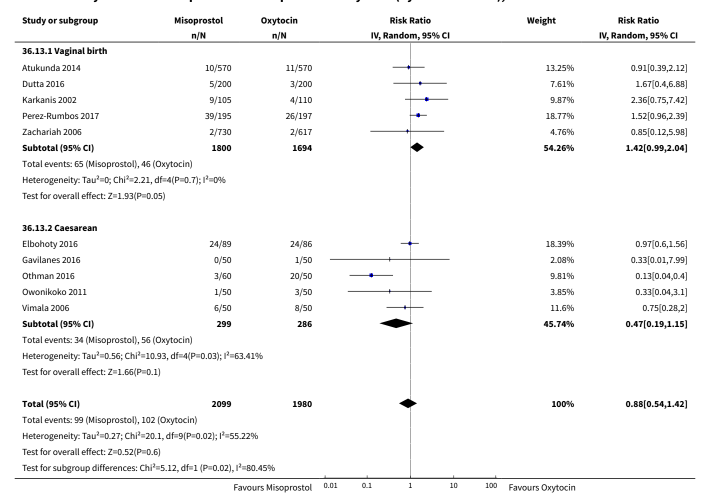
Analysis 36.12. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.12.1 Vaginal birth	- /	- /			
Afolabi 2010	0/100	0/100			Not estimable
Amin 2014	12/100	2/100		2.44%	6[1.38,26.12]
Atukunda 2014	35/569	19/570	_	13.68%	1.85[1.07,3.19]
Bellad 2012	5/321	0/331	 	0.65%	11.34[0.63,204.28]
Benchimol 2001	7/186	1/196	 	1.24%	7.38[0.92,59.38]
Bhatti 2014	3/60	1/60	+	1.09%	3[0.32,28.03]
Bugalho 2001	2/323	1/337	-	0.95%	2.09[0.19,22.9]
Caliskan 2002	2/396	2/407		1.41%	1.03[0.15,7.26]
Caliskan 2003	4/388	3/384	+	2.38%	1.32[0.3,5.86]
Chaudhuri 2012	5/265	2/265	+	2%	2.5[0.49,12.77]
Diallo 2017	2/154	1/150		0.95%	1.95[0.18,21.26]
Diop 2016	2/252	0/141	•	0.6%	2.81[0.14,58.05]
Dutta 2016	4/200	3/200	- +	2.4%	1.33[0.3,5.88]
Gerstenfeld 2001	0/159	0/166			Not estimable
Gulmezoglu 2001	66/9227	25/9232	-	17.44%	2.64[1.67,4.18]
Karkanis 2002	6/105	4/110	+	3.38%	1.57[0.46,5.41]
Kundodyiwa 2001	2/243	2/256		1.41%	1.05[0.15,7.42]
umbiganon 1999	0/397	1/200	•	0.54%	0.17[0.01,4.11]
Mukta 2013	4/100	2/100		1.9%	2[0.37,10.67]
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	6/257	5/257		3.73%	1.2[0.37,3.88]
Oboro 2003	12/247	9/249		6.73%	1.34[0.58,3.13]
Parsons 2006	1/221	4/224	+	1.14%	0.25[0.03,2.25]
Parsons 2007	1/214	2/213		0.95%	0.5[0.05,5.45]
Penaranda 2002	0/25	1/25 —		0.55%	0.33[0.01,7.81]
Perez-Rumbos 2017	2/195	5/197		2.01%	0.4[0.08,2.06]
Shady 2017	3/120	2/120		1.71%	1.5[0.26,8.82]
Tewatia 2014	1/50	1/50		0.72%	1[0.06,15.55]
Walley 2000	5/164	4/177		3.09%	1.35[0.37,4.94]
Zachariah 2006	6/730	2/617	+	2.08%	2.54[0.51,12.52]
Subtotal (95% CI)	15868	15534	•	77.17%	1.82[1.43,2.33]
Γotal events: 198 (Misoprostol),	, 104 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =22	· · · · · · · · · · · · · · · · · · ·				
Fest for overall effect: Z=4.84(P	<0.0001)				
36.12.2 Caesarean					
Acharya 2001	2/30	3/30		1.81%	0.67[0.12,3.71]
Adanikin 2013	2/25	2/25		1.52%	1[0.15,6.55]
llwani 2014	1/100	1/100		0.72%	1[0.06,15.77]
Chaudhuri 2010	2/96	3/94		1.72%	0.65[0.11,3.82]
Elbohoty 2016	9/89	7/86		5.57%	1.24[0.48,3.19]
Fazel 2013	2/50	3/50		1.76%	0.67[0.12,3.82]
Gavilanes 2016	1/50	1/50		0.72%	1[0.06,15.55]
Othman 2016	1/60	15/50		1.36%	0.06[0.01,0.41]
Owonikoko 2011	0/50	1/50 —		0.54%	0.33[0.01,7.99]
Pakniat 2015	4/50	2/50		1.95%	2[0.38,10.43]
'imala 2006	8/50	6/50	<u> </u>	5.16%	1.33[0.5,3.56]
Subtotal (95% CI)	650	635		22.83%	0.89[0.55,1.45]
abiolal (3370 CI)	650	033	_	22.83%	0.05[0.55,1.45]





Analysis 36.13. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 13 Headache.





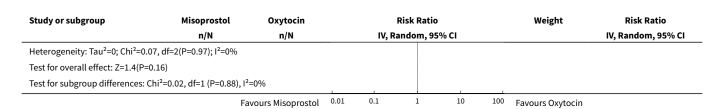
Analysis 36.14. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.14.1 Vaginal birth					
Atukunda 2014	132/570	163/570	<u></u>	56.18%	0.81[0.66,0.99]
Bellad 2012	0/321	0/331			Not estimable
Dutta 2016	2/200	0/200	+	0.24%	5[0.24,103.49]
Karkanis 2002	12/105	13/110		4.05%	0.97[0.46,2.02]
Mukta 2013	26/100	22/100	-	8.97%	1.18[0.72,1.94]
Shrestha 2011	7/100	7/100		2.16%	1[0.36,2.75]
Sultana 2007	5/210	4/190		1.3%	1.13[0.31,4.15]
Subtotal (95% CI)	1606	1601	♦	72.89%	0.87[0.73,1.04]
Total events: 184 (Misoprostol), 20	9 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =3.56,	df=5(P=0.61); I ² =0%				
Test for overall effect: Z=1.54(P=0.3	12)				
36.14.2 Caesarean					
Elbohoty 2016	47/89	44/86	+	27.11%	1.03[0.78,1.37]
Subtotal (95% CI)	89	86	*	27.11%	1.03[0.78,1.37]
Total events: 47 (Misoprostol), 44 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.22(P=0.8	33)				
Total (95% CI)	1695	1687	†	100%	0.91[0.79,1.06]
Total events: 231 (Misoprostol), 25	3 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =4.53,	df=6(P=0.61); I ² =0%				
Test for overall effect: Z=1.2(P=0.23	3)				
Test for subgroup differences: Chi ²	=0.97, df=1 (P=0.32), I ² =	:0%			
	Fav	ours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

Analysis 36.15. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 15 Hypertension.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.15.1 Vaginal birth					
Parsons 2006	1/208	0/215		32.1%	3.1[0.13,75.68]
Parsons 2007	2/202	0/203		35.68%	5.02[0.24,104.01]
Subtotal (95% CI)	410	418		67.78%	4[0.44,36.03]
Total events: 3 (Misoprostol), 0 (C	Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.05	, df=1(P=0.83); I ² =0%				
Test for overall effect: Z=1.24(P=0	0.22)				
36.15.2 Caesarean					
Alwani 2014	1/100	0/100		32.22%	3[0.12,72.77]
Subtotal (95% CI)	100	100		32.22%	3[0.12,72.77]
Total events: 1 (Misoprostol), 0 (C	Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.68(P=0	0.5)				
Total (95% CI)	510	518		100%	3.64[0.6,22.27]
Total events: 4 (Misoprostol), 0 (C	Oxytocin)				
	Fav	vours Misoprostol	0.01 0.1 1 10	100 Favours Oxytocin	

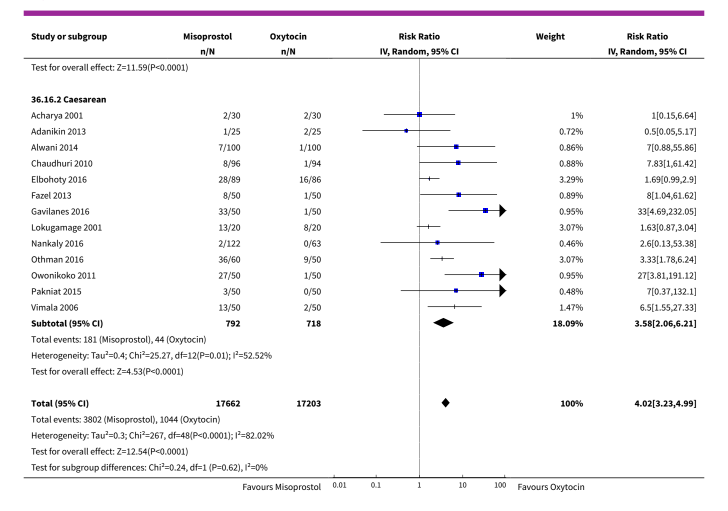




Analysis 36.16. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.16.1 Vaginal birth					
Afolabi 2010	4/100	2/100	- 	1.2%	2[0.37,10.67]
Amin 2014	25/100	4/100		2.16%	6.25[2.26,17.31]
Atukunda 2014	321/570	168/570	+	4.04%	1.91[1.65,2.21]
Baskett 2007	21/311	0/311	-	0.53%	43[2.62,706.74]
Bellad 2012	173/321	14/331	-	3.33%	12.74[7.56,21.49]
Benchimol 2001	5/186	0/196	+	0.5%	11.59[0.65,208.12]
Bhatti 2014	32/60	3/60		1.95%	10.67[3.45,32.96]
Bugalho 2001	123/323	51/337	+	3.84%	2.52[1.89,3.36]
Caliskan 2002	47/396	16/407		3.26%	3.02[1.74,5.23]
Caliskan 2003	44/388	19/384		3.34%	2.29[1.36,3.85]
Chaudhuri 2012	51/265	2/265		1.52%	25.5[6.27,103.67]
Dasuki 2002	13/98	2/98		1.44%	6.5[1.51,28.05]
Diallo 2017	11/154	3/150		1.73%	3.57[1.02,12.55]
Diop 2016	154/191	7/140		2.82%	16.13[7.81,33.31]
Dutta 2016	41/200	4/200		2.18%	10.25[3.74,28.08]
Gerstenfeld 2001	7/159	7/166		2.15%	1.04[0.37,2.91]
Gulmezoglu 2001	1620/9227	466/9232	+	4.09%	3.48[3.15,3.84]
Gupta 2006	16/100	13/100	- 	2.94%	1.23[0.63,2.42]
Karkanis 2002	26/105	15/110	<u> </u>	3.19%	1.82[1.02,3.23]
Kundodyiwa 2001	106/243	78/256	 +	3.93%	1.43[1.13,1.81]
Lumbiganon 1999	94/397	25/200	-	3.6%	1.89[1.26,2.85]
Mukta 2013	50/100	8/100		2.9%	6.25[3.13,12.5]
Musa 2015	42/100	15/100		3.33%	2.8[1.66,4.71]
Nasr 2009	80/257	0/257		0.53%	161[10.04,2582.33]
Oboro 2003	141/247	35/249	 	3.77%	4.06[2.93,5.62]
Parsons 2006	180/223	8/223	_ 	2.93%	22.5[11.36,44.56]
Parsons 2007	16/213	2/213	· · · · · · · · · · · · · · · · · · ·	1.44%	8[1.86,34.37]
Penaranda 2002	1/25	0/25		0.43%	3[0.13,70.3]
Perez-Rumbos 2017	11/195	2/197		1.4%	5.56[1.25,24.74]
Rajaei 2014	2/200	2/200		0.95%	1[0.14,7.03]
Singh 2009	19/150	0/75	-	0.53%	19.63[1.2,320.72]
Sultana 2007	13/210	2/190		1.42%	5.88[1.34,25.72]
Tewatia 2014	10/50	0/50	· · · · · · · · · · · · · · · · · · ·	0.52%	21[1.26,348.93]
Vagge 2014	15/100	3/100		1.81%	5[1.49,16.74]
Walley 2000	39/176	10/176		2.98%	3.9[2.01,7.57]
Zachariah 2006	68/730	14/617	—	3.22%	4.11[2.33,7.22]
Subtotal (95% CI)	16870	16485	•	81.91%	4.16[3.27,5.29]
Total events: 3621 (Misoprosto					- ,
Heterogeneity: Tau ² =0.3; Chi ² =	· · ·	=85.48%			

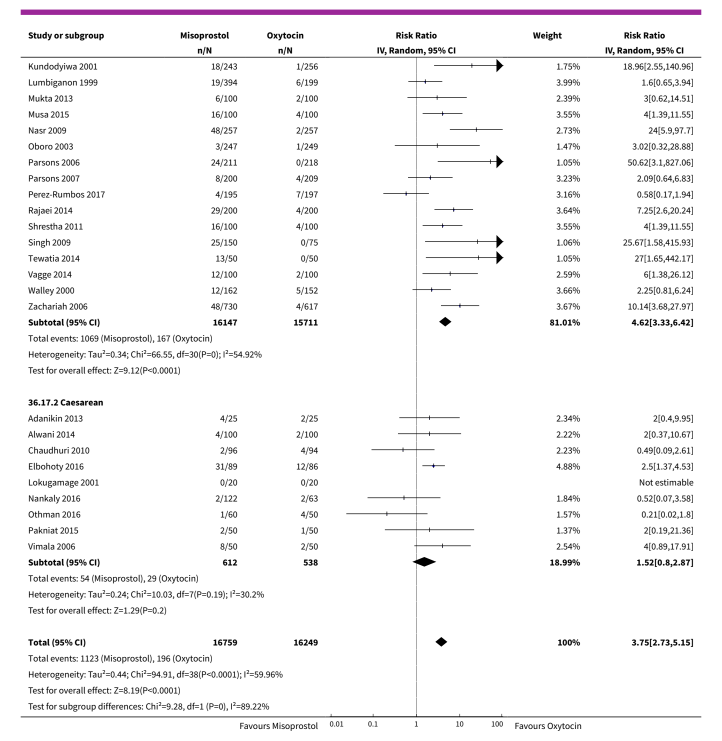




Analysis 36.17. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
36.17.1 Vaginal birth					
Afolabi 2010	0/100	0/100			Not estimable
Amin 2014	15/100	3/100		3.17%	5[1.49,16.74]
Atukunda 2014	53/570	12/570		4.82%	4.42[2.39,8.18]
Baskett 2007	39/311	1/311	_	1.78%	39[5.39,282.1]
Bellad 2012	4/321	0/331	-	0.98%	9.28[0.5,171.66]
Benchimol 2001	6/186	0/196	+	1.01%	13.7[0.78,241.41]
Bhatti 2014	2/60	0/60	-	0.93%	5[0.25,102]
Caliskan 2002	16/396	6/407		3.91%	2.74[1.08,6.93]
Caliskan 2003	17/388	5/384		3.75%	3.36[1.25,9.03]
Chaudhuri 2012	6/265	0/265	 	1.01%	13[0.74,229.61]
Diallo 2017	4/154	1/150		1.55%	3.9[0.44,34.46]
Diop 2016	8/252	2/141		2.47%	2.24[0.48,10.4]
Dutta 2016	17/200	1/200	_	1.75%	17[2.28,126.53]
Gulmezoglu 2001	559/9198	78/9205	+	5.71%	7.17[5.67,9.07]
Gupta 2006	2/100	0/100	-	0.92%	5[0.24,102.85]
Karkanis 2002	20/107	12/112	· · · · · · · · · · · · · · · · · · ·	4.68%	1.74[0.9,3.39]
	Fav	ours Misoprostol	0.01 0.1 1 10 100	Favours Oxytocin	

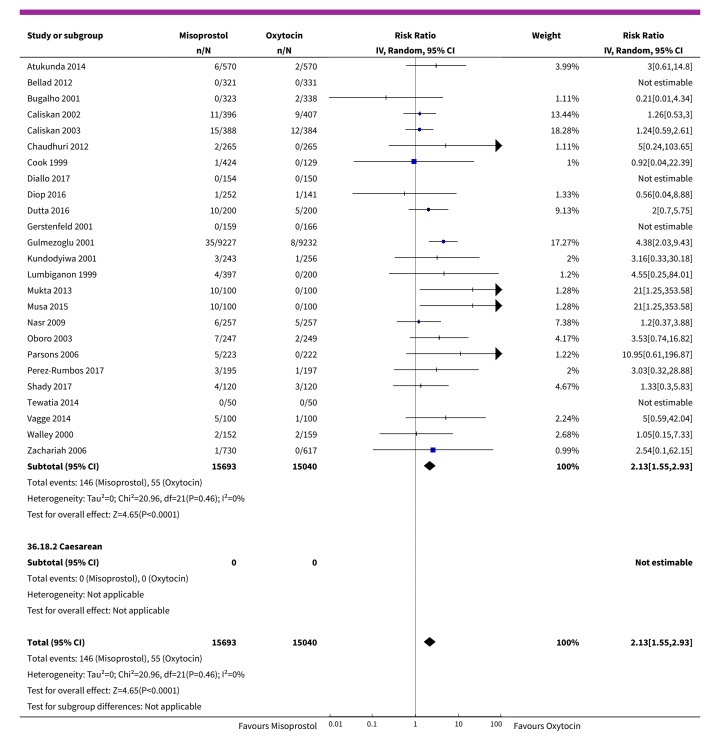




Analysis 36.18. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol	Oxytocin			Risk Ratio	,		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
36.18.1 Vaginal birth									
Amin 2014	5/100	1/100				+ ,	_ ,	2.24%	5[0.59,42.04]
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Oxytocin	

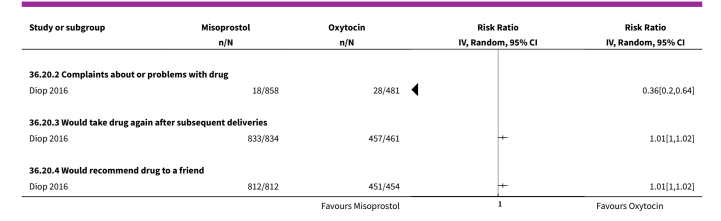




Analysis 36.20. Comparison 36 Misoprostol vs Oxytocin (by mode of birth), Outcome 20 Maternal satisfaction.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI	
36.20.1 Satisfied or very satisf	fied with drug				
Diop 2016	854/854	480/484	+	1.01[1,1.02]	
		Favours Misoprostol	1	Favours Oxytocin	





Comparison 37. Injectable prostaglandins vs Oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	100	Risk Ratio (IV, Random, 95% CI)	1.43 [0.20, 10.31]
2.1 Vaginal birth	2	100	Risk Ratio (IV, Random, 95% CI)	1.43 [0.20, 10.31]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	250	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.65]
3.1 Vaginal birth	2	250	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.65]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	4	500	Risk Ratio (IV, Random, 95% CI)	0.84 [0.26, 2.71]
6.1 Vaginal birth	4	500	Risk Ratio (IV, Random, 95% CI)	0.84 [0.26, 2.71]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	4	500	Risk Ratio (IV, Random, 95% CI)	0.29 [0.09, 0.94]
7.1 Vaginal birth	4	500	Risk Ratio (IV, Random, 95% CI)	0.29 [0.09, 0.94]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	4	500	Mean Difference (IV, Random, 95% CI)	-15.83 [-152.28, 120.62]
8.1 Vaginal birth	4	500	Mean Difference (IV, Random, 95% CI)	-15.83 [-152.28, 120.62]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	3	450	Risk Ratio (IV, Random, 95% CI)	1.17 [0.40, 3.41]
11.1 Vaginal birth	3	450	Risk Ratio (IV, Random, 95% CI)	1.17 [0.40, 3.41]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	400	Risk Ratio (IV, Random, 95% CI)	2.48 [0.57, 10.73]
12.1 Vaginal birth	2	400	Risk Ratio (IV, Random, 95% CI)	2.48 [0.57, 10.73]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	200	Risk Ratio (IV, Random, 95% CI)	0.20 [0.01, 4.11]
13.1 Vaginal birth	1	200	Risk Ratio (IV, Random, 95% CI)	0.20 [0.01, 4.11]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

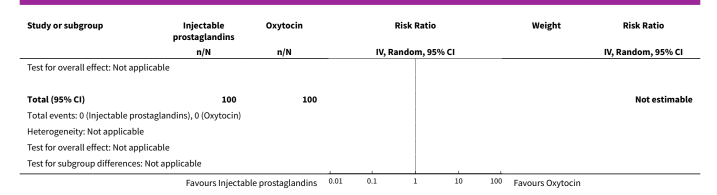


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	400	Risk Ratio (IV, Random, 95% CI)	0.91 [0.11, 7.73]
16.1 Vaginal birth	2	400	Risk Ratio (IV, Random, 95% CI)	0.91 [0.11, 7.73]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	200	Risk Ratio (IV, Random, 95% CI)	2.0 [0.18, 21.71]
17.1 Vaginal birth	1	200	Risk Ratio (IV, Random, 95% CI)	2.0 [0.18, 21.71]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	400	Risk Ratio (IV, Random, 95% CI)	10.38 [1.96, 54.98]
18.1 Vaginal birth	2	400	Risk Ratio (IV, Random, 95% CI)	10.38 [1.96, 54.98]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

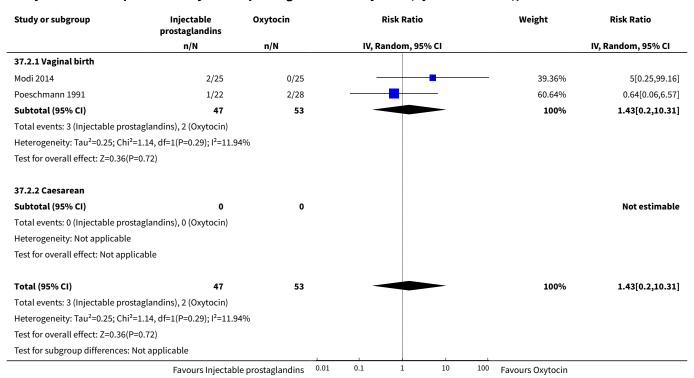
Analysis 37.1. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 959	% CI			IV, Random, 95% CI
37.1.1 Vaginal birth									
Kumar 2016	0/100	0/100							Not estimable
Subtotal (95% CI)	100	100							Not estimable
Total events: 0 (Injectable prostag	glandins), 0 (Oxytocin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applica	ble								
37.1.2 Caesarean									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Injectable prostag	glandins), 0 (Oxytocin)								
Heterogeneity: Not applicable									
	Favours Injectabl	e prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	





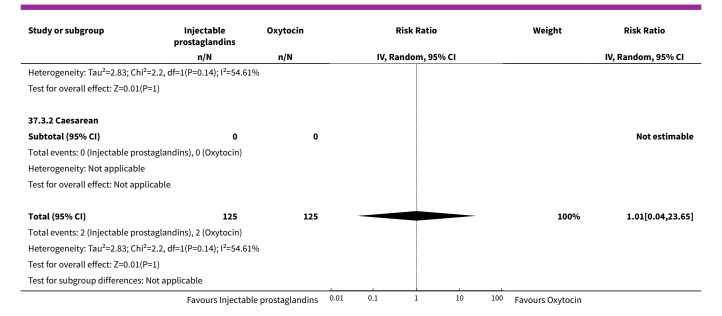
Analysis 37.2. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.



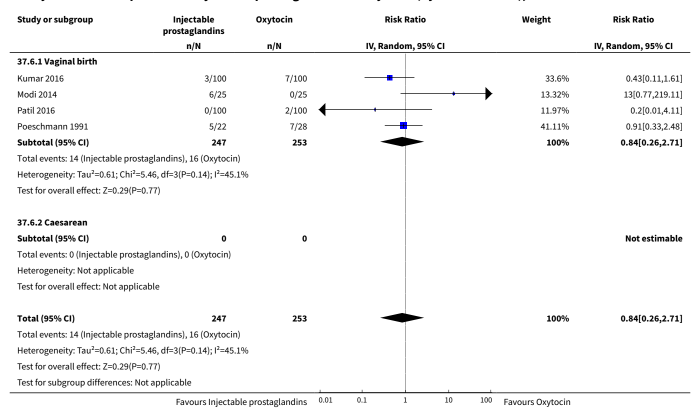
Analysis 37.3. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Oxytocin	Risk Ratio IV, Random, 95% CI					Weight	Risk Ratio
	n/N	n/N							IV, Random, 95% CI
37.3.1 Vaginal birth									
Kumar 2016	0/100	2/100	\leftarrow	-	-	_		49.72%	0.2[0.01,4.11]
Modi 2014	2/25	0/25		_	-	1		50.28%	5[0.25,99.16]
Subtotal (95% CI)	125	125						100%	1.01[0.04,23.65]
Total events: 2 (Injectable pr	ostaglandins), 2 (Oxytocin)								
	Favours Injectabl	e prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	



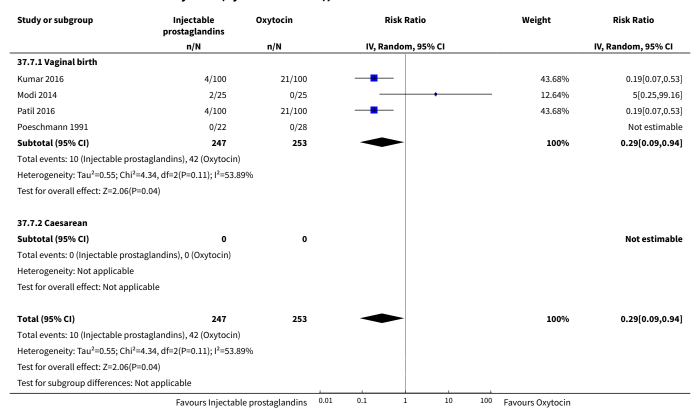


Analysis 37.6. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.





Analysis 37.7. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.



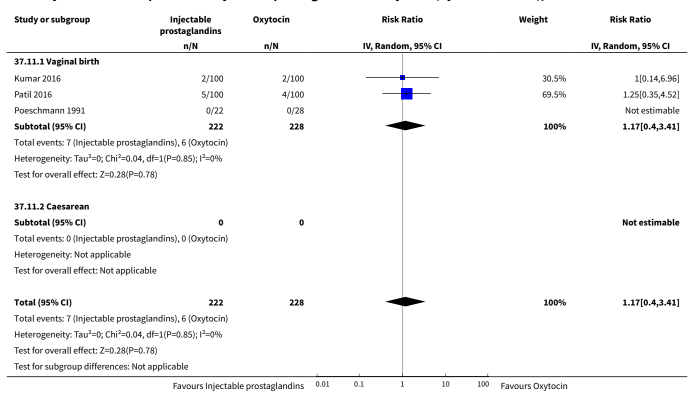
Analysis 37.8. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup		jectable taglandins	0	xytocin		Mean Differe	ice	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95%	CI		Random, 95% CI
37.8.1 Vaginal birth									
Kumar 2016	100	170.2 (197.4)	100	281.1 (197.4)	•	_		26.67%	-110.85[-165.57,-56.13]
Modi 2014	25	435 (147.6)	25	223.2 (122.5)			•	25.69%	211.8[136.61,286.99]
Patil 2016	100	170.2 (50.2)	100	281.1 (84.8)	◀			27.71%	-110.85[-130.17,-91.53]
Poeschmann 1991	22	324 (302)	28	374 (279)	\leftarrow	-		19.93%	-50[-213.11,113.11]
Subtotal ***	247		253					100%	-15.83[-152.28,120.62]
Heterogeneity: Tau ² =17392.8; Chi ² =	=67.08, df=	=3(P<0.0001); I ² =9	95.53%			İ			
Test for overall effect: Z=0.23(P=0.8	2)								
37.8.2 Caesarean									
Subtotal ***	0		0			İ			Not estimable
Heterogeneity: Not applicable						İ			
Test for overall effect: Not applicab	le								
Total ***	247		253					- 100%	-15.83[-152.28,120.62]
Heterogeneity: Tau ² =17392.8; Chi ² =	=67.08, df=	=3(P<0.0001); I ² =9	95.53%			į			
Test for overall effect: Z=0.23(P=0.8	2)					ĺ			
		Favours Inj	ectable p	rostaglandins	-100	-50 0	50 100	Favours O	kytocin



Study or subgroup	dy or subgroup Injectable prostagland					Mean Difference				Weight Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95%	CI		Random, 95% CI	
Test for subgroup differences:	Not applicable				_						
		Favours Ini	ectable	nrostaglandins	-100	-50	0	50	100	Favours Ovytocin	

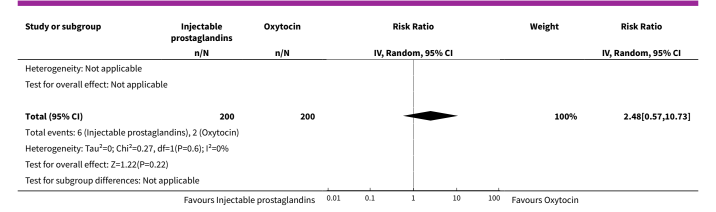
Analysis 37.11. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 11 Nausea.



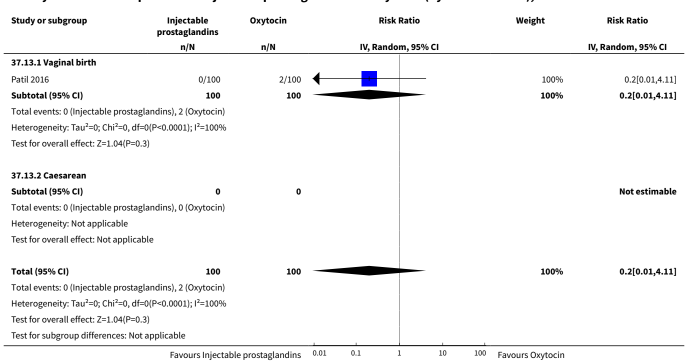
Analysis 37.12. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Injectable prostaglandins	Oxytocin			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI				IV, Random, 95% CI
37.12.1 Vaginal birth								
Kumar 2016	2/100	0/100		_	-	\longrightarrow	23.47%	5[0.24,102.85]
Patil 2016	4/100	2/100			-		76.53%	2[0.37,10.67]
Subtotal (95% CI)	200	200					100%	2.48[0.57,10.73]
Total events: 6 (Injectable pro	ostaglandins), 2 (Oxytocin)							
Heterogeneity: Tau ² =0; Chi ² =	0.27, df=1(P=0.6); I ² =0%							
Test for overall effect: Z=1.22	(P=0.22)							
37.12.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable pro	ostaglandins), 0 (Oxytocin)							
	Favours Injectab	le prostaglandins	0.01	0.1	1 10	100	Favours Oxytocin	





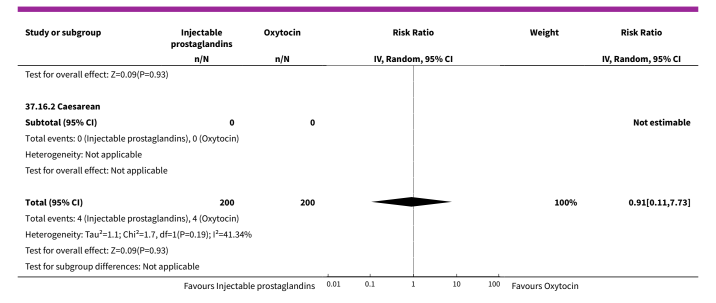
Analysis 37.13. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 13 Headache.



Analysis 37.16. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Injectable prostaglandins	Oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI					IV, Random, 95% CI	
37.16.1 Vaginal birth									
Kumar 2016	4/100	2/100			-			65.56%	2[0.37,10.67]
Patil 2016	0/100	2/100	\leftarrow	-		_		34.44%	0.2[0.01,4.11]
Subtotal (95% CI)	200	200						100%	0.91[0.11,7.73]
Total events: 4 (Injectable pro	ostaglandins), 4 (Oxytocin)								
Heterogeneity: Tau ² =1.1; Chi	² =1.7, df=1(P=0.19); I ² =41.34%								
	Favours Injectabl	e prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	





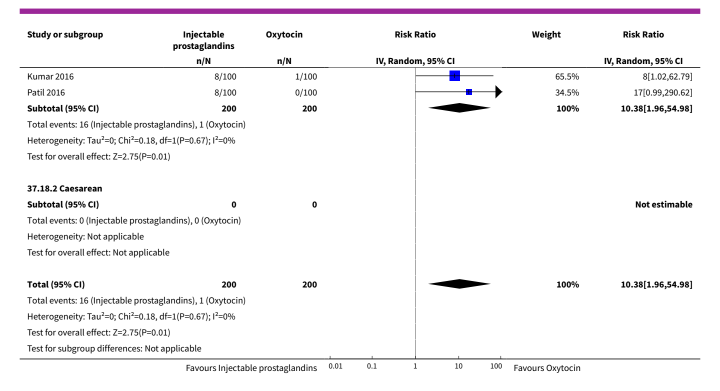
Analysis 37.17. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 17 Fever.

prostaglandins	Oxytocin	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
2/100	1/100		100%	2[0.18,21.71]
100	100		100%	2[0.18,21.71]
glandins), 1 (Oxytocin)				
.57)				
0	0			Not estimable
glandins), 0 (Oxytocin)				
ble				
100	100		100%	2[0.18,21.71]
glandins), 1 (Oxytocin)				
.57)				
t applicable				
	n/N 2/100 100 glandins), 1 (Oxytocin) .57) 0 glandins), 0 (Oxytocin)	2/100 1/100 100 100 glandins), 1 (Oxytocin) .57) 0 0 glandins), 0 (Oxytocin) ble 100 100 glandins), 1 (Oxytocin)	n/N	2/100 1/100 100% 100 100 100% glandins), 1 (Oxytocin) 0 0 glandins), 0 (Oxytocin) blee 100 100 100 glandins), 1 (Oxytocin)

Analysis 37.18. Comparison 37 Injectable prostaglandins vs Oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Oxytocin	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
37.18.1 Vaginal birth						1			
	Favours Injectab	le prostaglandins	0.01	0.1	1	10	100	Favours Oxytocin	





Comparison 38. Carbetocin vs Oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	5	30327	Risk Ratio (IV, Random, 95% CI)	2.00 [0.37, 10.92]
1.1 Vaginal birth	1	29539	Risk Ratio (IV, Random, 95% CI)	2.00 [0.37, 10.92]
1.2 Caesarean	4	788	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	10	30633	Risk Ratio (IV, Random, 95% CI)	0.73 [0.45, 1.19]
2.1 Vaginal birth	4	29682	Risk Ratio (IV, Random, 95% CI)	0.68 [0.21, 2.20]
2.2 Caesarean	6	951	Risk Ratio (IV, Random, 95% CI)	0.62 [0.31, 1.23]
3 Blood transfusion	16	32115	Risk Ratio (IV, Random, 95% CI)	0.68 [0.38, 1.22]
3.1 Vaginal birth	5	30028	Risk Ratio (IV, Random, 95% CI)	1.04 [0.52, 2.10]
3.2 Caesarean	10	2032	Risk Ratio (IV, Random, 95% CI)	0.50 [0.23, 1.10]
3.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	0.16 [0.01, 2.93]
4 Severe maternal morbidity: intensive care admissions	2	29847	Risk Ratio (IV, Random, 95% CI)	1.16 [0.67, 2.02]
4.1 Vaginal birth	1	29470	Risk Ratio (IV, Random, 95% CI)	1.13 [0.65, 1.98]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.2 Caesarean	1	377	Risk Ratio (IV, Random, 95% CI)	3.02 [0.12, 73.56]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	30633	Risk Ratio (IV, Random, 95% CI)	0.75 [0.58, 0.98]
6.1 Vaginal birth	5	29955	Risk Ratio (IV, Random, 95% CI)	0.67 [0.34, 1.30]
6.2 Caesarean	6	678	Risk Ratio (IV, Random, 95% CI)	0.71 [0.47, 1.07]
7 Additional uterotonics	22	32699	Risk Ratio (IV, Random, 95% CI)	0.48 [0.34, 0.68]
7.1 Vaginal birth	6	30187	Risk Ratio (IV, Random, 95% CI)	0.54 [0.30, 0.99]
7.2 Caesarean	15	2457	Risk Ratio (IV, Random, 95% CI)	0.45 [0.27, 0.74]
7.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	0.37 [0.02, 8.71]
8 Blood loss	16	2115	Mean Difference (IV, Random, 95% CI)	-92.73 [-154.97, -30.49]
8.1 Vaginal birth	4	485	Mean Difference (IV, Random, 95% CI)	-68.42 [-143.52, 6.68]
8.2 Caesarean	12	1630	Mean Difference (IV, Random, 95% CI)	-101.54 [-178.53, -24.55]
9 Change in haemoglo- bin	12	2096	Mean Difference (IV, Random, 95% CI)	-1.66 [-3.81, 0.50]
9.1 Vaginal birth	3	415	Mean Difference (IV, Random, 95% CI)	-4.21 [-5.34, -3.07]
9.2 Caesarean	8	1626	Mean Difference (IV, Random, 95% CI)	-0.63 [-3.48, 2.22]
9.3 Both caesarean and vaginal birth	1	55	Mean Difference (IV, Random, 95% CI)	-1.70 [-6.97, 3.57]
10 Breastfeeding	2	190	Risk Ratio (IV, Random, 95% CI)	0.94 [0.86, 1.03]
10.1 Vaginal birth	1	135	Risk Ratio (IV, Random, 95% CI)	0.93 [0.85, 1.03]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	0.98 [0.79, 1.22]
11 Nausea	12	2288	Risk Ratio (IV, Random, 95% CI)	1.11 [0.78, 1.56]
11.1 Vaginal birth	4	555	Risk Ratio (IV, Random, 95% CI)	1.30 [0.37, 4.60]

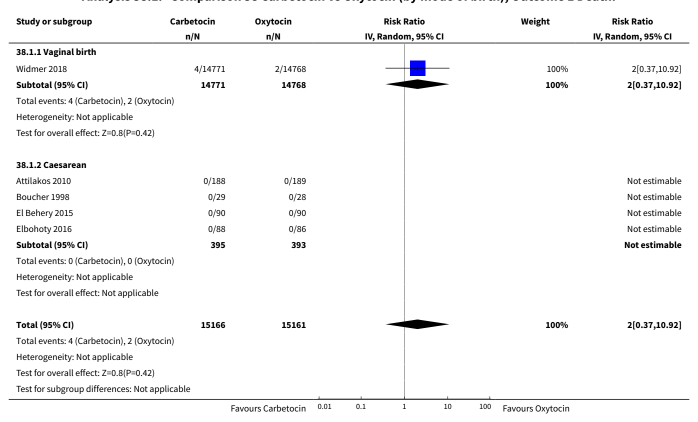


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
11.2 Caesarean	8	1733	Risk Ratio (IV, Random, 95% CI)	1.10 [0.75, 1.61]
12 Vomiting	13	31833	Risk Ratio (IV, Random, 95% CI)	0.90 [0.53, 1.50]
12.1 Vaginal birth	5	30055	Risk Ratio (IV, Random, 95% CI)	0.85 [0.14, 5.25]
12.2 Caesarean	7	1723	Risk Ratio (IV, Random, 95% CI)	0.83 [0.44, 1.60]
12.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	3.33 [0.14, 78.42]
13 Headache	15	2620	Risk Ratio (IV, Random, 95% CI)	0.84 [0.63, 1.12]
13.1 Vaginal birth	4	555	Risk Ratio (IV, Random, 95% CI)	1.15 [0.41, 3.26]
13.2 Caesarean	10	2010	Risk Ratio (IV, Random, 95% CI)	0.81 [0.61, 1.08]
13.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	7.78 [0.42, 143.81]
14 Abdominal pain	10	31293	Risk Ratio (IV, Random, 95% CI)	1.18 [0.97, 1.44]
14.1 Vaginal birth	5	29946	Risk Ratio (IV, Random, 95% CI)	1.09 [0.79, 1.49]
14.2 Caesarean	5	1347	Risk Ratio (IV, Random, 95% CI)	1.21 [0.90, 1.62]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	1998	Risk Ratio (IV, Random, 95% CI)	0.78 [0.49, 1.23]
16.1 Vaginal birth	3	495	Risk Ratio (IV, Random, 95% CI)	1.22 [0.49, 3.07]
16.2 Caesarean	6	1503	Risk Ratio (IV, Random, 95% CI)	0.70 [0.40, 1.20]
17 Fever	4	466	Risk Ratio (IV, Random, 95% CI)	1.58 [0.27, 9.35]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	3	411	Risk Ratio (IV, Random, 95% CI)	2.90 [0.19, 43.87]
17.3 Both caesarean and vaginal birth	1	55	Risk Ratio (IV, Random, 95% CI)	0.37 [0.02, 8.71]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

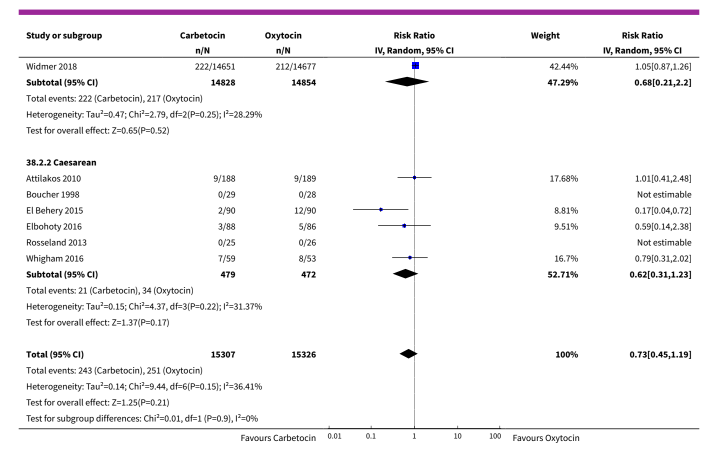
Analysis 38.1. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 1 Death.



Analysis 38.2. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin Oxytocin		Risk Ratio					Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI						IV, Random, 95% CI
38.2.1 Vaginal birth									
Fenix 2012	0/30	0/30							Not estimable
Kabir 2015	0/47	4/47	\leftarrow					2.65%	0.11[0.01,2.01]
Maged 2016	0/100	1/100	.—					2.21%	0.33[0.01,8.09]
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

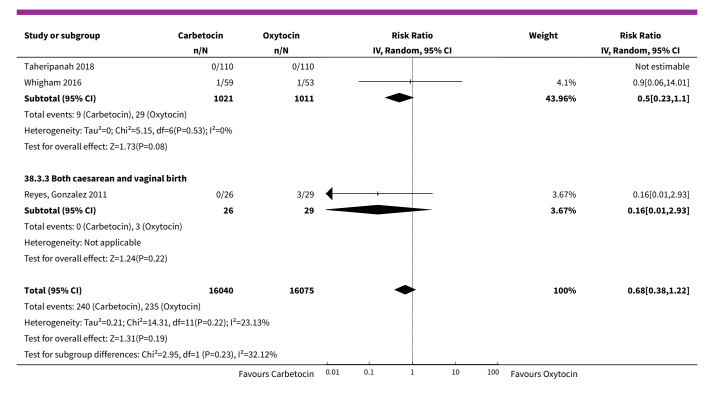




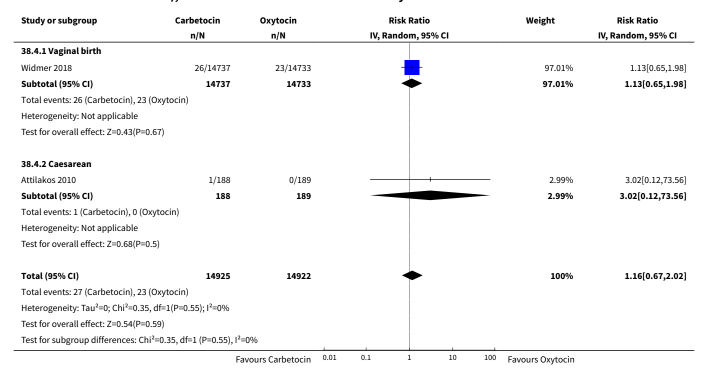
Analysis 38.3. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI
	n/N	n/N	IV, Random, 95% CI		
38.3.1 Vaginal birth					
Fenix 2012	0/30	0/30			Not estimable
Kabir 2015	0/47	3/47	+	3.63%	0.14[0.01,2.69]
Maged 2016	1/100	2/100		5.27%	0.5[0.05,5.43]
Reyes 2011	1/45	0/90	+	3.13%	5.93[0.25,142.84]
Widmer 2018	229/14771	198/14768	-	40.33%	1.16[0.96,1.4]
Subtotal (95% CI)	14993	15035	*	52.36%	1.04[0.52,2.1]
Total events: 231 (Carbetocin)	, 203 (Oxytocin)				
Heterogeneity: Tau ² =0.15; Chi	² =3.43, df=3(P=0.33); I ² =12.5	9%			
Test for overall effect: Z=0.12(F	P=0.9)				
38.3.2 Caesarean					
Attilakos 2010	4/188	5/189		13.69%	0.8[0.22,2.95]
Boucher 1998	0/29	0/28			Not estimable
Carrillo-Gaucin 2016	1/60	2/57	+	5.31%	0.48[0.04,5.1]
Dansereau 1999	2/317	2/318		7.4%	1[0.14,7.08]
El Behery 2015	0/90	14/90	— ——	3.95%	0.03[0,0.57]
Elbohoty 2016	0/88	1/86	+	3.12%	0.33[0.01,7.89]
Fahmy 2015	0/50	0/50			Not estimable
Fahmy 2016	1/30	4/30		6.39%	0.25[0.03,2.11]
	Fa	vours Carbetocin	0.01 0.1 1 10 10	00 Favours Oxytocin	



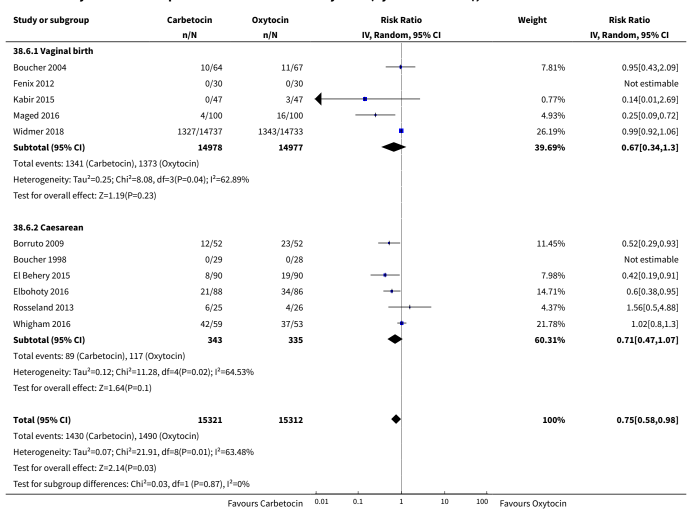


Analysis 38.4. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.





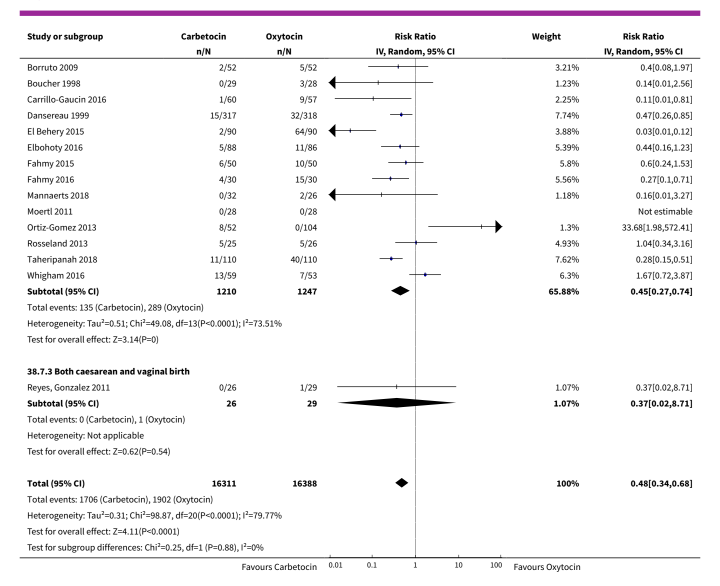
Analysis 38.6. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.



Analysis 38.7. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
38.7.1 Vaginal birth					
Boucher 2004	12/83	12/77		6.89%	0.93[0.44,1.94]
Fenix 2012	3/30	27/30		5.07%	0.11[0.04,0.33]
Kabir 2015	0/47	5/47		1.27%	0.09[0.01,1.6]
Maged 2016	23/100	37/100	-	8.62%	0.62[0.4,0.97]
Reyes 2011	0/45	3/90	+ +	1.22%	0.28[0.01,5.36]
Widmer 2018	1533/14770	1528/14768	+	9.98%	1[0.94,1.07]
Subtotal (95% CI)	15075	15112	•	33.05%	0.54[0.3,0.99]
Total events: 1571 (Carbetocin),	, 1612 (Oxytocin)				
Heterogeneity: Tau ² =0.31; Chi ² =	23.52, df=5(P=0); I ² =78.74%	6			
Test for overall effect: Z=2(P=0.0	04)				
38.7.2 Caesarean					
Attilakos 2010	63/188	86/189	. +	9.5%	0.74[0.57,0.95]
	Fav	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	

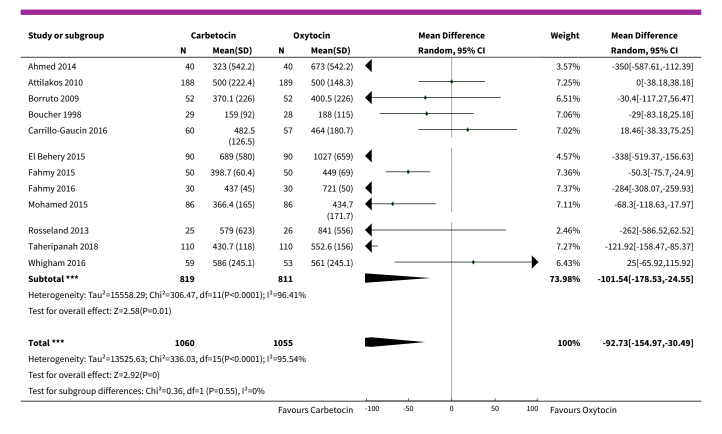




Analysis 38.8. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Caı	rbetocin	0:	xytocin		Me	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
38.8.1 Vaginal birth										
Boucher 2004	64	413.3 (197.5)	67	410.4 (194.1)			+	_	6.86%	2.9[-64.19,69.99]
Fenix 2012	30	296 (183.3)	30	493.3 (183.3)	•				6.4%	-197.3[-290.04,-104.56]
Kabir 2015	47	325 (306)	47	389 (366)	\leftarrow	+			5.49%	-64[-200.39,72.39]
Maged 2016	100	337.7 (118.8)	100	378 (143.2)	-	+			7.27%	-40.27[-76.73,-3.81]
Subtotal ***	241		244						26.02%	-68.42[-143.52,6.68]
Heterogeneity: Tau ² =4124.92; C	hi²=12.39, df=	3(P=0.01); I ² =75.	78%							
Test for overall effect: Z=1.79(P=	=0.07)									
38.8.2 Caesarean										
			Favou	rs Carbetocin	-100	-50	0 50	100	Favours Ox	ytocin

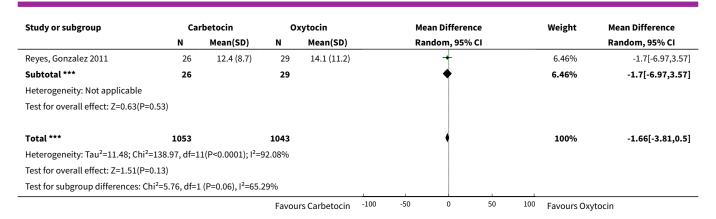




Analysis 38.9. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Ca	rbetocin	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
38.9.1 Vaginal birth							
Boucher 2004	82	12.8 (10.8)	73	15.9 (11.6)	+	8.2%	-3.1[-6.64,0.44]
Fenix 2012	30	6 (4.6)	30	11 (4.6)	*	9.37%	-5[-7.35,-2.65]
Maged 2016	100	5.5 (3.5)	100	9.6 (6.2)	+	10.09%	-4.1[-5.5,-2.7]
Subtotal ***	212		203		•	27.66%	-4.21[-5.34,-3.07]
Heterogeneity: Tau ² =0; Chi ² =0	0.84, df=2(P=0.6	6); I ² =0%					
Test for overall effect: Z=7.26(P<0.0001)						
38.9.2 Caesarean							
Attilakos 2010	183	16 (2.2)	185	16 (2.2)	†	10.49%	0[-0.45,0.45]
Dansereau 1999	317	7.5 (10)	318	8.3 (10)	•	9.99%	-0.8[-2.36,0.76]
El Behery 2015	90	17.4 (8.7)	90	9.4 (6.7)		9.44%	8[5.73,10.27]
Mannaerts 2018	32	14.5 (11)	26	15 (9)	+	6.58%	-0.5[-5.65,4.65]
Moertl 2011	28	11 (9.9)	28	11.4 (7.6)	+	7.1%	-0.4[-5.02,4.22]
Rosseland 2013	25	5 (8.2)	26	8.2 (6.7)	+	7.61%	-3.2[-7.32,0.92]
Taheripanah 2018	110	11 (10.1)	110	20.5 (12)	*	8.82%	-9.5[-12.43,-6.57]
Whigham 2016	30	22 (11.5)	28	21 (11.5)	+	5.86%	1[-4.94,6.94]
Subtotal ***	815		811		♦	65.88%	-0.63[-3.48,2.22]
Heterogeneity: Tau ² =13.62; Ch	ni²=91.63, df=7(P<0.0001); I ² =92.	36%				
Test for overall effect: Z=0.43(P=0.66)						
38.9.3 Both caesarean and v	aginal birth						
			Favou	ırs Carbetocin -100	-50 0 50	100 Favours Ox	ytocin





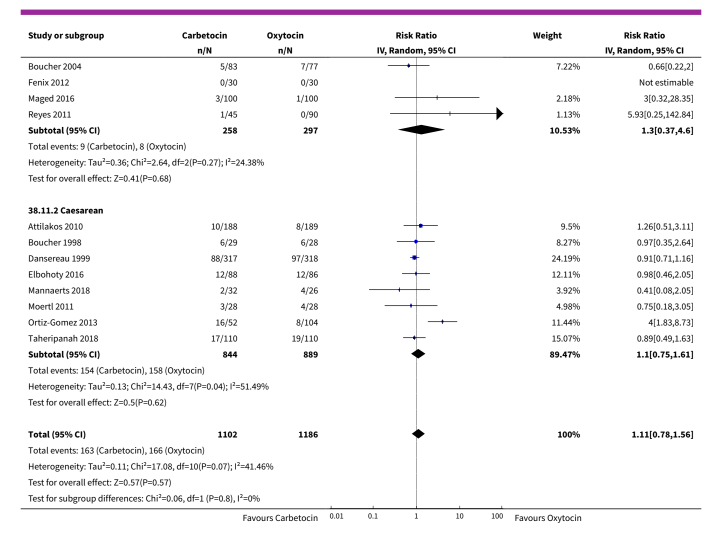
Analysis 38.10. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 10 Breastfeeding.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
38.10.1 Vaginal birth					
Reyes 2011	41/45	88/90	+	83.79%	0.93[0.85,1.03]
Subtotal (95% CI)	45	90		83.79%	0.93[0.85,1.03]
Total events: 41 (Carbetocin), 88 (Oxy	rtocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.44(P=0.15)					
38.10.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Oxyto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
38.10.3 Both caesarean and vaginal	birth				
Reyes, Gonzalez 2011	22/26	25/29	+	16.21%	0.98[0.79,1.22]
Subtotal (95% CI)	26	29	\	16.21%	0.98[0.79,1.22]
Total events: 22 (Carbetocin), 25 (Oxy	rtocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.17(P=0.87)					
Total (95% CI)	71	119		100%	0.94[0.86,1.03]
Total events: 63 (Carbetocin), 113 (Ox	ytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.18, df=	1(P=0.67); I ² =0%		ĺ		
Test for overall effect: Z=1.38(P=0.17)					
Test for subgroup differences: Chi ² =0.	.18, df=1 (P=0.67), I ² =	=0%	ĺ		
	Fa	vours Carbetocin 0.01	0.1 1 10	100 Favours Oxytocin	

Analysis 38.11. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Carbetocin	Oxytocin	tocin Risk Ratio					Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
38.11.1 Vaginal birth				1		1			
	F	avours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

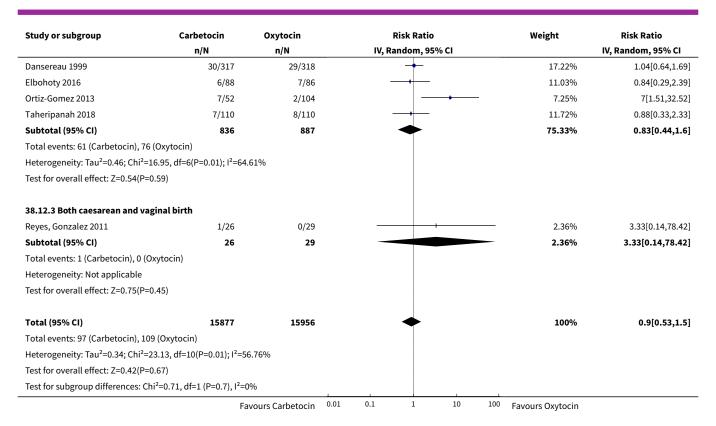




Analysis 38.12. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Oxytocin	ı	Risk Ratio		Weight	Risk Ratio
	n/N	n/N	n/N IV, Random, 95% CI				IV, Random, 95% CI
38.12.1 Vaginal birth							
Boucher 2004	0/83	6/77	+ +			2.8%	0.07[0,1.25]
Fenix 2012	0/30	0/30					Not estimable
Maged 2016	2/100	0/100	_			2.54%	5[0.24,102.85]
Reyes 2011	0/45	0/90					Not estimable
Widmer 2018	33/14757	27/14743		+-		16.98%	1.22[0.73,2.03]
Subtotal (95% CI)	15015	15040				22.32%	0.85[0.14,5.25]
Total events: 35 (Carbetocin),	33 (Oxytocin)						
Heterogeneity: Tau ² =1.52; Chi	² =4.59, df=2(P=0.1); l ² =56.38	%					
Test for overall effect: Z=0.17(F	P=0.86)						
38.12.2 Caesarean							
Attilakos 2010	5/188	5/189	_			9.48%	1.01[0.3,3.42]
Borruto 2009	4/52	20/52		_		11.49%	0.2[0.07,0.54]
Boucher 1998	2/29	5/28				7.13%	0.39[0.08,1.83]
	Fa	vours Carbetocin	0.01 0.1	1 10	100	Favours Oxytocin	

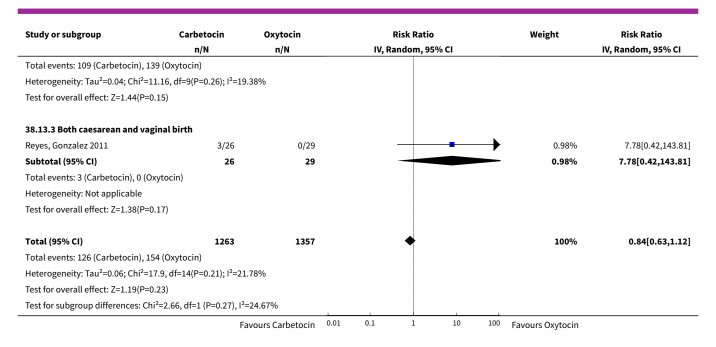




Analysis 38.13. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 13 Headache.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
38.13.1 Vaginal birth					
Boucher 2004	6/83	11/77		7.6%	0.51[0.2,1.3]
Fenix 2012	1/30	1/30		1.12%	1[0.07,15.26]
Maged 2016	5/100	2/100	+	3%	2.5[0.5,12.59]
Reyes 2011	2/45	1/90	+	1.46%	4[0.37,42.95]
Subtotal (95% CI)	258	297		13.17%	1.15[0.41,3.26]
Total events: 14 (Carbetocin), 15 ((Oxytocin)				
Heterogeneity: Tau ² =0.38; Chi ² =4.	.48, df=3(P=0.21); I ² =33.0	17%			
Test for overall effect: Z=0.27(P=0	.79)				
38.13.2 Caesarean					
Attilakos 2010	0/188	1/189 -	•	0.82%	0.34[0.01,8.17]
Borruto 2009	7/52	15/52		9.62%	0.47[0.21,1.05]
Boucher 1998	1/29	0/28	-	0.84%	2.9[0.12,68.33]
	46/317	43/318	L	22.49%	1.07[0.73,1.58]
Dansereau 1999	40/311	43/316	<u> </u>	22.49%	1.07[0.75,1.50]
Dansereau 1999 El Behery 2015	23/90	30/90	-	19.37%	0.77[0.48,1.21]
	•	•	-		
El Behery 2015	23/90	30/90		19.37%	0.77[0.48,1.21]
El Behery 2015 Elbohoty 2016	23/90 20/88	30/90 24/86		19.37% 17.26%	0.77[0.48,1.21] 0.81[0.49,1.36]
El Behery 2015 Elbohoty 2016 Moertl 2011	23/90 20/88 2/28	30/90 24/86 2/28		19.37% 17.26% 2.25%	0.77[0.48,1.21] 0.81[0.49,1.36] 1[0.15,6.61]
El Behery 2015 Elbohoty 2016 Moertl 2011 Ortiz-Gomez 2013	23/90 20/88 2/28 6/52	30/90 24/86 2/28 8/104		19.37% 17.26% 2.25% 6.89%	0.77[0.48,1.21] 0.81[0.49,1.36] 1[0.15,6.61] 1.5[0.55,4.1]





Analysis 38.14. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
38.14.1 Vaginal birth					
Boucher 2004	5/83	0/77	 	0.47%	10.21[0.57,181.71]
Fenix 2012	7/30	8/30		4.62%	0.88[0.36,2.11]
Kabir 2015	4/47	5/47		2.4%	0.8[0.23,2.8]
Reyes 2011	0/45	1/90		0.39%	0.66[0.03,15.87]
Widmer 2018	63/14754	56/14743	-	18.28%	1.12[0.78,1.61]
Subtotal (95% CI)	14959	14987	*	26.16%	1.09[0.79,1.49]
Total events: 79 (Carbetocin), 70	(Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =2.9	2, df=4(P=0.57); I ² =0%				
Test for overall effect: Z=0.52(P=	0.6)				
38.14.2 Caesarean					
Attilakos 2010	1/188	1/189		0.51%	1.01[0.06,15.95]
Borruto 2009	21/52	20/52	-	12.59%	1.05[0.65,1.69]
Boucher 1998	0/29	1/28 —	•	0.39%	0.32[0.01,7.59]
Dansereau 1999	131/317	127/318	+	32.1%	1.03[0.86,1.25]
Elbohoty 2016	72/88	44/86	-	28.24%	1.6[1.27,2.01]
Subtotal (95% CI)	674	673	•	73.84%	1.21[0.9,1.62]
Total events: 225 (Carbetocin), 1	193 (Oxytocin)				
Heterogeneity: Tau ² =0.05; Chi ² =	9.4, df=4(P=0.05); l ² =57.43	%			
Test for overall effect: Z=1.24(P=	0.21)				
Total (95% CI)	15633	15660	*	100%	1.18[0.97,1.44]
Total events: 304 (Carbetocin), 2	263 (Oxytocin)		į		
Heterogeneity: Tau ² =0.02; Chi ² =	12.7, df=9(P=0.18); I ² =29.1	1%	į		
Test for overall effect: Z=1.62(P=	0.11)				
	hi ² =0.22, df=1 (P=0.64), I ² =		İ		



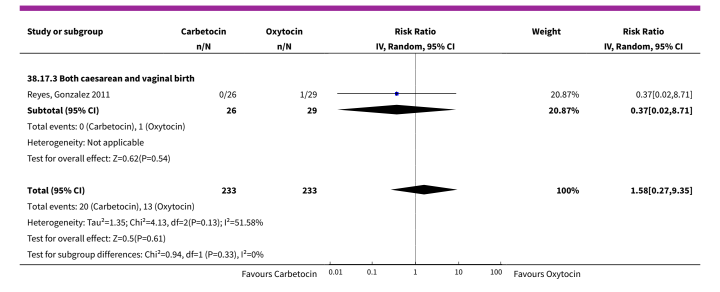
Analysis 38.16. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 16 Shivering.

8/83 2/100 0/45 228 0.0.34); I ² =0%	n/N 7/77 0/100 0/90 267	IV, Random, 95% CI	16.6% 2.19% 18.79%	1.06[0.4,2.79] 5[0.24,102.85] Not estimable 1.22[0.49,3.07]
2/100 0/45 228	0/100 0/90	•	2.19%	5[0.24,102.85] Not estimable
2/100 0/45 228	0/100 0/90	•	2.19%	5[0.24,102.85] Not estimable
0/45 228	0/90		•	Not estimable
228	•		18.79%	
)	267		18.79%	1.22[0.49,3.07]
0.34); I ² =0%				
		I I		
2/188	4/189		6.57%	0.5[0.09,2.71]
6/52	0/52	+	2.45%	13[0.75,225]
1/29	0/28		2.01%	2.9[0.12,68.33]
37/317	49/318	-	43.78%	0.76[0.51,1.13]
8/88	16/86		21.88%	0.49[0.22,1.08]
1/52	7/104		4.51%	0.29[0.04,2.26]
726	777	•	81.21%	0.7[0.4,1.2]
n)				
(P=0.25); I ² =24.38	8%			
954	1044	•	100%	0.78[0.49,1.23]
n)				
(P=0.27); I ² =19.7	5%			
df=1 (P=0.3), I ² =6.	.77%			
(6/52 1/29 37/317 8/88 1/52 726 n) (P=0.25); I ² =24.3 954 n) (P=0.27); I ² =19.7	6/52 0/52 1/29 0/28 37/317 49/318 8/88 16/86 1/52 7/104 726 777 n) (P=0.25); l ² =24.38%	6/52 0/52 1/29 0/28 37/317 49/318 8/88 16/86 1/52 7/104 726 777 1) (P=0.25); l²=24.38% 954 1044 1) (P=0.27); l²=19.75%	6/52 0/52 2.45% 1/29 0/28 2.01% 37/317 49/318 43.78% 8/88 16/86 21.88% 1/52 7/104 4.51% 726 777 81.21% 1) (P=0.25); l²=24.38% 41044 100% 1) (P=0.27); l²=19.75%

Analysis 38.17. Comparison 38 Carbetocin vs Oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
38.17.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Oxytoo	in)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
38.17.2 Caesarean					
Boucher 1998	0/29	0/28			Not estimable
El Behery 2015	8/90	0/90	•	23.91%	17[1,290.19]
Elbohoty 2016	12/88	12/86		55.22%	0.98[0.46,2.05]
Subtotal (95% CI)	207	204		79.13%	2.9[0.19,43.87]
Total events: 20 (Carbetocin), 12 (Oxyt	cocin)				
Heterogeneity: Tau ² =2.96; Chi ² =3.64, c	If=1(P=0.06); I ² =72.5	55%			
Test for overall effect: Z=0.77(P=0.44)					
	Fa	vours Carbetocin 0.	01 0.1 1 10 100	Favours Oxytocin	





Comparison 39. Ergometrine vs Oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	1293	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	1293	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	6	2591	Risk Ratio (IV, Random, 95% CI)	1.30 [0.52, 3.27]
2.1 Vaginal birth	6	2591	Risk Ratio (IV, Random, 95% CI)	1.30 [0.52, 3.27]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	5	1893	Risk Ratio (IV, Random, 95% CI)	1.44 [0.20, 10.23]
3.1 Vaginal birth	5	1893	Risk Ratio (IV, Random, 95% CI)	1.44 [0.20, 10.23]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6 PPH >= 500 mL	10	3221	Risk Ratio (IV, Random, 95% CI)	1.31 [0.86, 1.99]
6.1 Vaginal birth	10	3221	Risk Ratio (IV, Random, 95% CI)	1.31 [0.86, 1.99]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	6	2493	Risk Ratio (IV, Random, 95% CI)	1.46 [0.61, 3.48]
7.1 Vaginal birth	6	2493	Risk Ratio (IV, Random, 95% CI)	1.46 [0.61, 3.48]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	10	3221	Mean Difference (IV, Random, 95% CI)	8.09 [-17.83, 34.00]
8.1 Vaginal birth	10	3221	Mean Difference (IV, Random, 95% CI)	8.09 [-17.83, 34.00]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	2	1893	Mean Difference (IV, Random, 95% CI)	0.42 [-0.30, 1.13]
9.1 Vaginal birth	2	1893	Mean Difference (IV, Random, 95% CI)	0.42 [-0.30, 1.13]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	6	2529	Risk Ratio (IV, Random, 95% CI)	4.56 [1.13, 18.44]
11.1 Vaginal birth	6	2529	Risk Ratio (IV, Random, 95% CI)	4.56 [1.13, 18.44]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	5	2343	Risk Ratio (IV, Random, 95% CI)	3.83 [1.10, 13.28]
12.1 Vaginal birth	5	2343	Risk Ratio (IV, Random, 95% CI)	3.83 [1.10, 13.28]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	4	2293	Risk Ratio (IV, Random, 95% CI)	5.63 [0.93, 33.96]
13.1 Vaginal birth	4	2293	Risk Ratio (IV, Random, 95% CI)	5.63 [0.93, 33.96]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

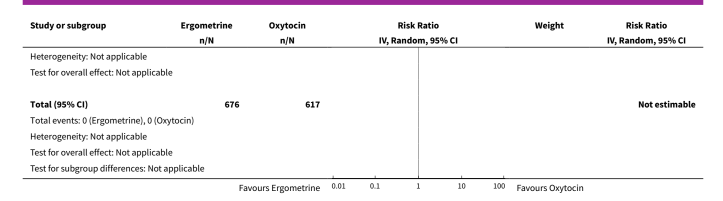


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15 Hypertension	3	1410	Risk Ratio (IV, Random, 95% CI)	13.39 [2.01, 89.44]
15.1 Vaginal birth	3	1410	Risk Ratio (IV, Random, 95% CI)	13.39 [2.01, 89.44]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	1493	Risk Ratio (IV, Random, 95% CI)	1.73 [0.93, 3.25]
16.1 Vaginal birth	3	1493	Risk Ratio (IV, Random, 95% CI)	1.73 [0.93, 3.25]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1443	Risk Ratio (IV, Random, 95% CI)	2.97 [0.97, 9.05]
17.1 Vaginal birth	2	1443	Risk Ratio (IV, Random, 95% CI)	2.97 [0.97, 9.05]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	1393	Risk Ratio (IV, Random, 95% CI)	3.74 [0.42, 33.53]
18.1 Vaginal birth	2	1393	Risk Ratio (IV, Random, 95% CI)	3.74 [0.42, 33.53]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

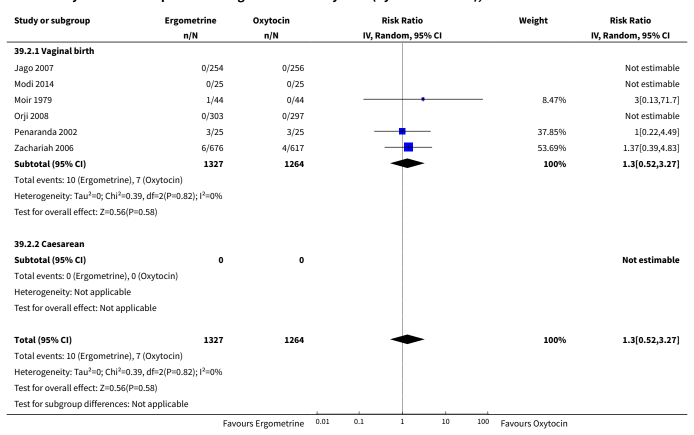
Analysis 39.1. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Ergometrine	Oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N n/N		IV, F	Random, 95	% CI			IV, Random, 95% CI
39.1.1 Vaginal birth									
Zachariah 2006	0/676	0/617							Not estimable
Subtotal (95% CI)	676	617							Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
39.1.2 Caesarean									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)								
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	





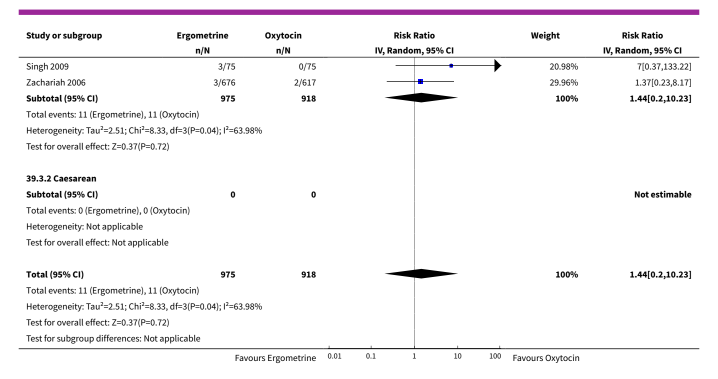
Analysis 39.2. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.



Analysis 39.3. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Ergometrine Oxytocin			Risk Ratio)		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI						IV, Random, 95% CI	
39.3.1 Vaginal birth										
Dhananjaya 2014	4/50	0/50			-		\rightarrow	21.32%	9[0.5,162.89]	
Ezeama 2014	1/149	9/151						27.74%	0.11[0.01,0.88]	
Modi 2014	0/25	0/25							Not estimable	
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin		

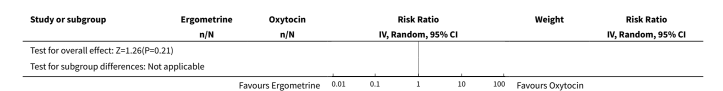




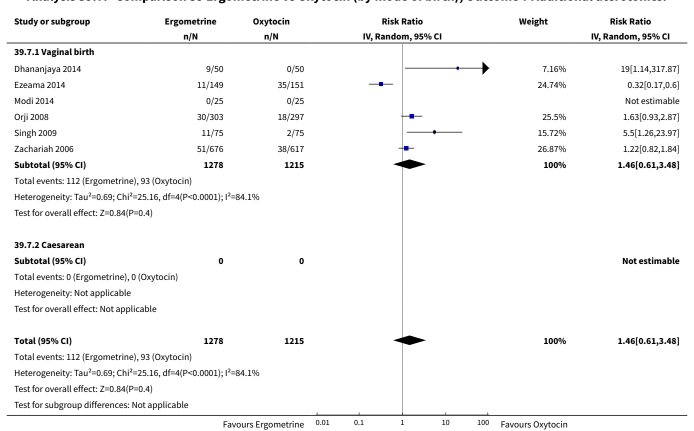
Analysis 39.6. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
39.6.1 Vaginal birth					
Dhananjaya 2014	6/50	1/50	+	3.72%	6[0.75,48.05]
Ezeama 2014	3/149	12/151		9.07%	0.25[0.07,0.88]
Jago 2007	0/254	0/256			Not estimable
Modi 2014	0/25	0/25			Not estimable
Moir 1979	2/44	2/44		4.32%	1[0.15,6.79]
Moodie 1976	14/40	10/40	+	20.72%	1.4[0.71,2.77]
Orji 2008	18/303	12/297	 • -	19.75%	1.47[0.72,3]
Penaranda 2002	12/25	8/25		20.09%	1.5[0.74,3.03]
Singh 2009	2/75	0/75		1.84%	5[0.24,102.42]
Zachariah 2006	20/676	13/617	-	20.49%	1.4[0.7,2.8]
Subtotal (95% CI)	1641	1580	•	100%	1.31[0.86,1.99]
Total events: 77 (Ergometrine), 5	8 (Oxytocin)				
Heterogeneity: Tau ² =0.1; Chi ² =9.	89, df=7(P=0.19); l ² =29.22	%			
Test for overall effect: Z=1.26(P=	0.21)				
39.6.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 ((Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applic	able				
Total (95% CI)	1641	1580	•	100%	1.31[0.86,1.99]
Total events: 77 (Ergometrine), 5	8 (Oxytocin)				
Heterogeneity: Tau ² =0.1; Chi ² =9.	89 df=7(P=0.19)·1 ² =29.22	%			





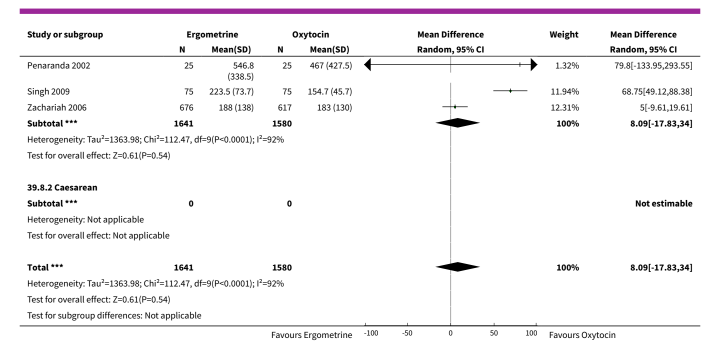
Analysis 39.7. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.



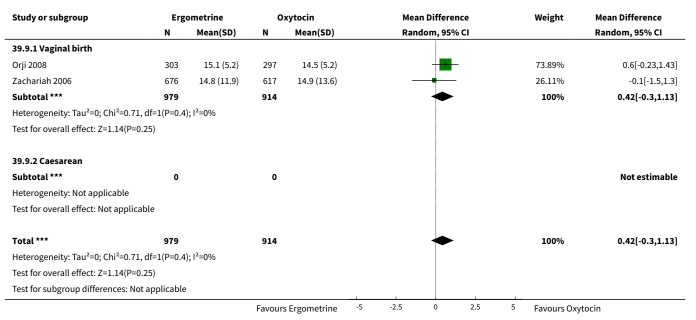
Analysis 39.8. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	0	xytocin		М	ean Differenc	:e		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	andom, 95% (CI			Random, 95% CI
39.8.1 Vaginal birth											
Dhananjaya 2014	50	345 (109.5)	50	219 (86.3)					→	9.97%	126[87.35,164.65]
Ezeama 2014	149	287.1 (84.4)	151	301.8 (109.2)		_	+			11.73%	-14.7[-36.77,7.37]
Jago 2007	254	150.2 (63.6)	256	171.9 (81.6)		-	→			12.43%	-21.7[-34.39,-9.01]
Modi 2014	25	131 (72)	25	223.2 (122.5)	4					8.05%	-92.2[-147.92,-36.48]
Moir 1979	44	201 (50)	44	208 (58)			+			11.68%	-7[-29.63,15.63]
Moodie 1976	40	369 (118)	40	391 (129)			•			8.21%	-22[-76.18,32.18]
Orji 2008	303	246.6 (95.4)	297	245.7 (77.6)						12.36%	0.92[-12.98,14.82]
			Favour	s Ergometrine	-100	-50	0	50	100	Favours Oxyto	cin





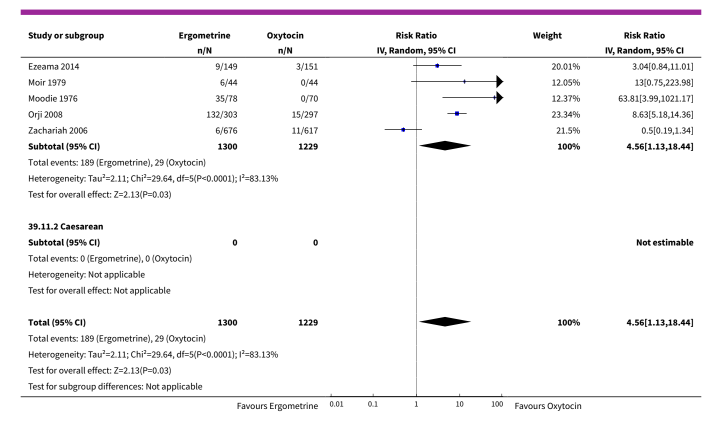
Analysis 39.9. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.



Analysis 39.11. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Ergometrine	Oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
39.11.1 Vaginal birth									
Dhananjaya 2014	1/50	0/50				+		10.73%	3[0.13,71.92]
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	



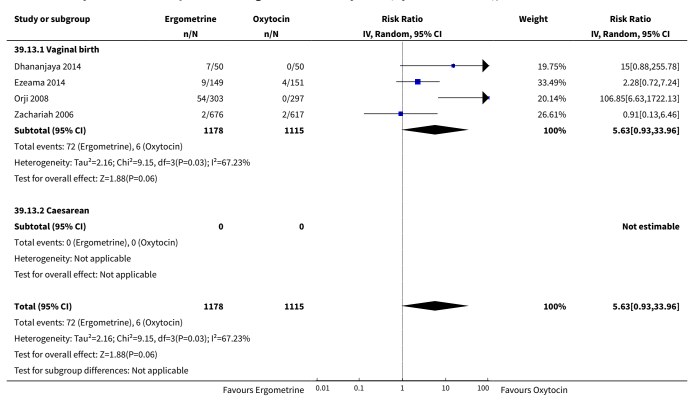


Analysis 39.12. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
39.12.1 Vaginal birth					
Dhananjaya 2014	4/50	0/50		12.59%	9[0.5,162.89]
Ezeama 2014	4/149	1/151	-	17.87%	4.05[0.46,35.85]
Orji 2008	132/303	12/297	-	36.56%	10.78[6.1,19.04]
Penaranda 2002	0/25	1/25	+	11.17%	0.33[0.01,7.81]
Zachariah 2006	3/676	2/617		21.8%	1.37[0.23,8.17]
Subtotal (95% CI)	1203	1140		100%	3.83[1.1,13.28]
Total events: 143 (Ergometrine), 16 (Oxytocin)				
Heterogeneity: Tau ² =1.02; Chi ² =9.08,	df=4(P=0.06); I ² =55.9	7%			
Test for overall effect: Z=2.11(P=0.03))				
39.12.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Oxy	tocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	1203	1140		100%	3.83[1.1,13.28]
Total events: 143 (Ergometrine), 16 (Oxytocin)				
Heterogeneity: Tau ² =1.02; Chi ² =9.08,	df=4(P=0.06); I ² =55.9	7%			
Test for overall effect: Z=2.11(P=0.03))				
Test for subgroup differences: Not ap	plicable				
	Favo	ours Ergometrine	0.01 0.1 1 10 100	Favours Oxytocin	



Analysis 39.13. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 13 Headache.



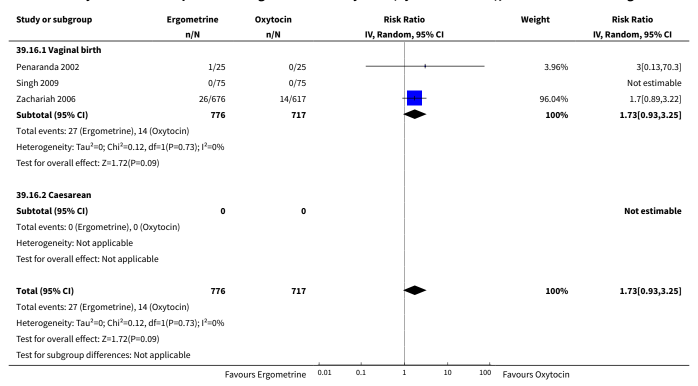
Analysis 39.15. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
39.15.1 Vaginal birth							
Ezeama 2014	7/149	0/151		-		23.98%	15.2[0.88,263.78]
Jago 2007	134/254	29/256		-		51.35%	4.66[3.24,6.69]
Orji 2008	54/303	0/297			—	24.66%	106.85[6.63,1722.13]
Subtotal (95% CI)	706	704				100%	13.39[2.01,89.44]
Total events: 195 (Ergometrine), 29 (O	kytocin)						
Heterogeneity: Tau ² =1.79; Chi ² =5.39, d	f=2(P=0.07); I ² =62.9	1%					
Test for overall effect: Z=2.68(P=0.01)							
39.15.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)						
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Total (95% CI)	706	704				100%	13.39[2.01,89.44]
Total events: 195 (Ergometrine), 29 (O	kytocin)						
Heterogeneity: Tau ² =1.79; Chi ² =5.39, d	f=2(P=0.07); I ² =62.9	1%					
Test for overall effect: Z=2.68(P=0.01)							
	Fave	ours Ergometrine	0.01	0.1 1 10	100	Favours Oxytocin	



Study or subgroup	Ergometrine n/N	Oxytocin n/N			Risk Ratio andom, 95			Weight	Risk Ratio IV, Random, 95% CI
Test for subgroup differences	: Not applicable					1			
		Favours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	

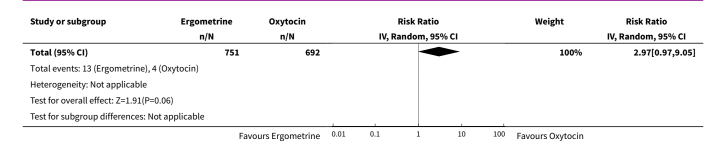
Analysis 39.16. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 16 Shivering.



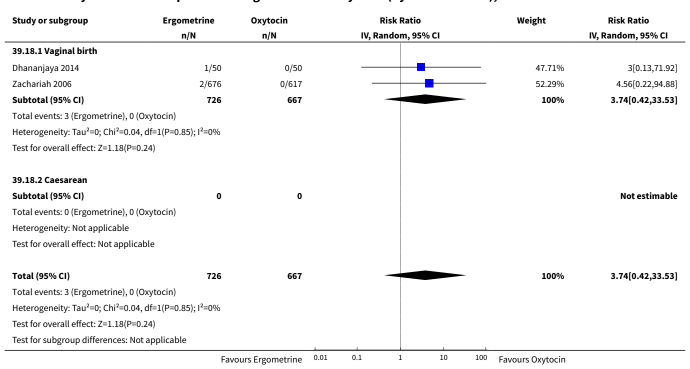
Analysis 39.17. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Ergometrine	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	CI		IV, Random, 95% CI
39.17.1 Vaginal birth							
Singh 2009	0/75	0/75					Not estimable
Zachariah 2006	13/676	4/617		-		100%	2.97[0.97,9.05]
Subtotal (95% CI)	751	692			-	100%	2.97[0.97,9.05]
Total events: 13 (Ergometrine), 4 (Oxyt	ocin)						
Heterogeneity: Not applicable							
Test for overall effect: Z=1.91(P=0.06)							
39.17.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)						
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
	Fav	ours Ergometrine	0.01).1 1	10 100	Favours Oxytocin	





Analysis 39.18. Comparison 39 Ergometrine vs Oxytocin (by mode of birth), Outcome 18 Diarrhoea.



Comparison 40. Ergometine plus oxytocin vs Oxytocin (by mode of birth)

Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	2	1314	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	2	1314	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	10	10990	Risk Ratio (IV, Random, 95% CI)	0.73 [0.57, 0.93]
2.1 Vaginal birth	10	10990	Risk Ratio (IV, Random, 95% CI)	0.73 [0.57, 0.93]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
3 Blood transfusion	11	10985	Risk Ratio (IV, Random, 95% CI)	0.88 [0.53, 1.44]
3.1 Vaginal birth	9	10521	Risk Ratio (IV, Random, 95% CI)	1.00 [0.61, 1.64]
3.2 Caesarean	2	464	Risk Ratio (IV, Random, 95% CI)	0.39 [0.17, 0.91]
4 Severe maternal morbidity: intensive care admissions	1	991	Risk Ratio (IV, Random, 95% CI)	2.99 [0.12, 73.32]
4.1 Vaginal birth	1	991	Risk Ratio (IV, Random, 95% CI)	2.99 [0.12, 73.32]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	11090	Risk Ratio (IV, Random, 95% CI)	0.72 [0.57, 0.91]
6.1 Vaginal birth	11	11090	Risk Ratio (IV, Random, 95% CI)	0.72 [0.57, 0.91]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	10	8968	Risk Ratio (IV, Random, 95% CI)	0.79 [0.59, 1.07]
7.1 Vaginal birth	8	8504	Risk Ratio (IV, Random, 95% CI)	0.80 [0.58, 1.10]
7.2 Caesarean	2	464	Risk Ratio (IV, Random, 95% CI)	0.74 [0.22, 2.48]
8 Blood loss	10	4248	Mean Difference (IV, Random, 95% CI)	-10.31 [-40.32, 19.70]
8.1 Vaginal birth	7	3729	Mean Difference (IV, Random, 95% CI)	-0.12 [-33.85, 33.60]
8.2 Caesarean	3	519	Mean Difference (IV, Random, 95% CI)	-49.54 [-88.07, -11.01]
9 Change in haemo- globin	6	4324	Mean Difference (IV, Random, 95% CI)	-2.23 [-5.24, 0.77]
9.1 Vaginal birth	6	4324	Mean Difference (IV, Random, 95% CI)	-2.23 [-5.24, 0.77]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	1	3483	Risk Ratio (IV, Random, 95% CI)	0.99 [0.96, 1.01]
10.1 Vaginal birth	1	3483	Risk Ratio (IV, Random, 95% CI)	0.99 [0.96, 1.01]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	7	6931	Risk Ratio (IV, Random, 95% CI)	1.72 [0.84, 3.53]



Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
11.1 Vaginal birth	5	6467	Risk Ratio (IV, Random, 95% CI)	1.58 [0.65, 3.86]
11.2 Caesarean	2	464	Risk Ratio (IV, Random, 95% CI)	2.09 [0.53, 8.22]
12 Vomiting	9	10227	Risk Ratio (IV, Random, 95% CI)	3.05 [1.76, 5.29]
12.1 Vaginal birth	7	9763	Risk Ratio (IV, Random, 95% CI)	2.93 [1.50, 5.71]
12.2 Caesarean	2	464	Risk Ratio (IV, Random, 95% CI)	2.83 [1.12, 7.15]
13 Headache	5	5105	Risk Ratio (IV, Random, 95% CI)	1.26 [0.79, 1.99]
13.1 Vaginal birth	4	4689	Risk Ratio (IV, Random, 95% CI)	1.74 [0.67, 4.55]
13.2 Caesarean	1	416	Risk Ratio (IV, Random, 95% CI)	1.06 [0.59, 1.91]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	1362	Risk Ratio (IV, Random, 95% CI)	2.00 [0.29, 13.97]
15.1 Vaginal birth	2	1314	Risk Ratio (IV, Random, 95% CI)	4.57 [0.65, 32.04]
15.2 Caesarean	1	48	Risk Ratio (IV, Random, 95% CI)	0.25 [0.03, 2.08]
16 Shivering	2	1591	Risk Ratio (IV, Random, 95% CI)	0.96 [0.60, 1.53]
16.1 Vaginal birth	2	1591	Risk Ratio (IV, Random, 95% CI)	0.96 [0.60, 1.53]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1591	Risk Ratio (IV, Random, 95% CI)	1.08 [0.48, 2.43]
17.1 Vaginal birth	2	1591	Risk Ratio (IV, Random, 95% CI)	1.08 [0.48, 2.43]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	3	2030	Risk Ratio (IV, Random, 95% CI)	1.26 [0.72, 2.22]
18.1 Vaginal birth	3	2030	Risk Ratio (IV, Random, 95% CI)	1.26 [0.72, 2.22]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0	,	Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfac- tion	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Outcome or sub- group title	No. of studies No. of participants		Statistical method	Effect size
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

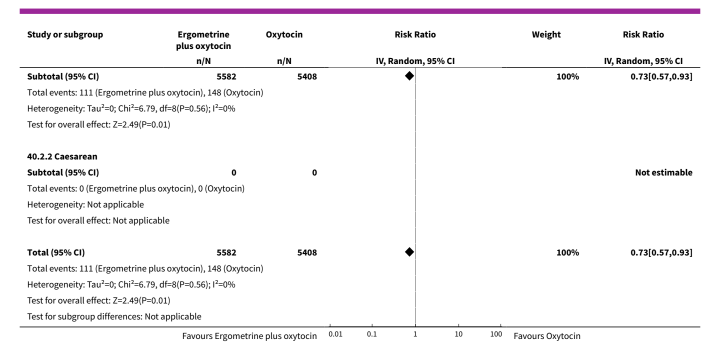
Analysis 40.1. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
40.1.1 Vaginal birth					
Nuamsiri 2016	0/162	0/161			Not estimable
Yuen 1995	0/496	0/495			Not estimable
Subtotal (95% CI)	658	656			Not estimable
Total events: 0 (Ergometrine plus oxy	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	•				
40.1.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus oxy	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	•				
Total (95% CI)	658	656			Not estimable
Total events: 0 (Ergometrine plus oxy	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	•				
Test for subgroup differences: Not ap	plicable				
	Favours Ergometri	ne plus oxytocin 0.01	0.1 1 10	100 Favours Oxytocin	

Analysis 40.2. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% C	CI CO	IV, Random, 95% CI
40.2.1 Vaginal birth					
Caliskan 2002	7/402	14/407		7.6%	0.51[0.21,1.24]
Caliskan 2003	5/398	15/384		6.09%	0.32[0.12,0.88]
Choy 2002	3/500	6/491		3.21%	0.49[0.12,1.95]
Cook 1999	7/310	0/129	-	0.75%	6.27[0.36,108.98]
Khan 1995	9/1016	11/1012	-+	7.96%	0.81[0.34,1.96]
McDonald 1993	68/1730	83/1753		62.17%	0.83[0.61,1.14]
Mitchell 1993	0/228	1/230	+	- 0.6%	0.34[0.01,8.21]
Nuamsiri 2016	0/162	0/161			Not estimable
Rashid 2009	6/340	8/346		5.57%	0.76[0.27,2.18]
Yuen 1995	6/496	10/495	+-	6.06%	0.6[0.22,1.63]
	Favours Frgometi	rine plus oxytocin	0.01 0.1 1	10 100 Favours Oxytocin	

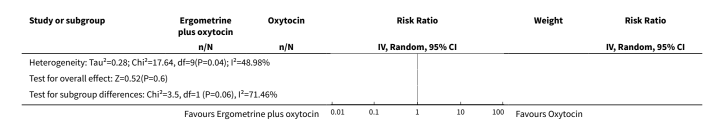




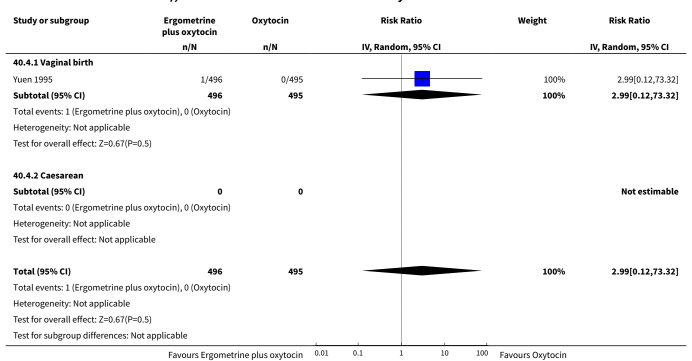
Analysis 40.3. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Risk Ratio Weight	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
40.3.1 Vaginal birth					
Caliskan 2002	4/402	13/407		10.62%	0.31[0.1,0.95]
Caliskan 2003	6/398	13/384	-+-	12.33%	0.45[0.17,1.16]
Choy 2002	13/493	7/487	 • 	12.9%	1.83[0.74,4.56]
Cook 1999	3/310	2/129		5.79%	0.62[0.11,3.69]
Khan 1995	2/1016	1/1012		3.59%	1.99[0.18,21.93]
McDonald 1993	24/1730	16/1753	+-	16.71%	1.52[0.81,2.85]
Nuamsiri 2016	2/162	1/161		3.61%	1.99[0.18,21.7]
Rashid 2009	6/340	2/346	+	6.79%	3.05[0.62,15.02]
Yuen 1995	10/496	12/495		13.93%	0.83[0.36,1.91]
Subtotal (95% CI)	5347	5174	•	86.27%	1[0.61,1.64]
Total events: 70 (Ergometrine ¡	olus oxytocin), 67 (Oxytocin)			
Heterogeneity: Tau ² =0.21; Chi ²	=13.33, df=8(P=0.1); I ² =40%				
Test for overall effect: Z=0.02(P	2=0.99)				
40.3.2 Caesarean					
Balki 2008	0/24	0/24			Not estimable
Koen 2016	7/202	19/214		13.73%	0.39[0.17,0.91]
Subtotal (95% CI)	226	238	•	13.73%	0.39[0.17,0.91]
Total events: 7 (Ergometrine p	lus oxytocin), 19 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2.18(P	2=0.03)				
Total (95% CI)	5573	5412	•	100%	0.88[0.53,1.44]
Total events: 77 (Ergometrine)	olus oxvtocin). 86 (Oxvtocin)			





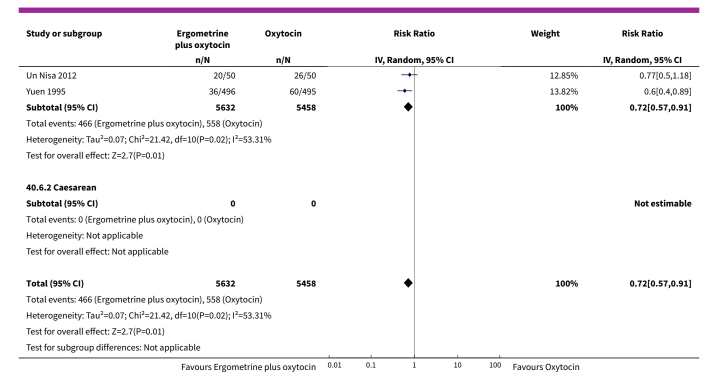
Analysis 40.4. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.



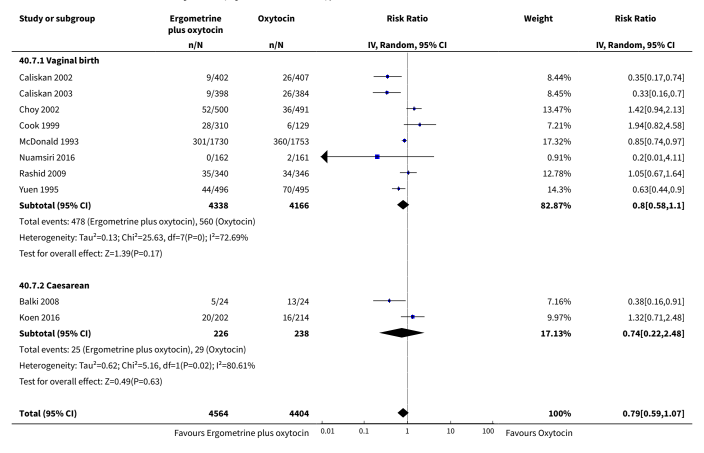
Analysis 40.6. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk	Risk Ratio			Risk Ratio
	n/N	n/N	IV, Rando	om, 95% CI			IV, Random, 95% CI
40.6.1 Vaginal birth							
Caliskan 2002	14/402	33/407				9.06%	0.43[0.23,0.79]
Caliskan 2003	14/398	28/384		-		8.78%	0.48[0.26,0.9]
Choy 2002	23/500	26/491	_	+		10.24%	0.87[0.5,1.5]
Cook 1999	23/310	1/129				1.35%	9.57[1.31,70.13]
Khan 1995	36/1016	41/1012	_	+		12.66%	0.87[0.56,1.36]
McDonald 1993	286/1730	316/1753		+		20.46%	0.92[0.79,1.06]
Mitchell 1993	6/228	17/230				5.23%	0.36[0.14,0.89]
Nuamsiri 2016	0/162	1/161				0.55%	0.33[0.01,8.07]
Rashid 2009	8/340	9/346		 	1	4.99%	0.9[0.35,2.32]
	Favours Ergometi	ine plus oxytocin	0.01 0.1	1 10	100	Favours Oxytocin	

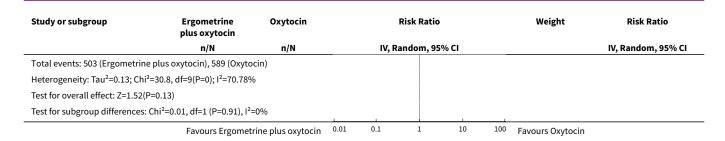




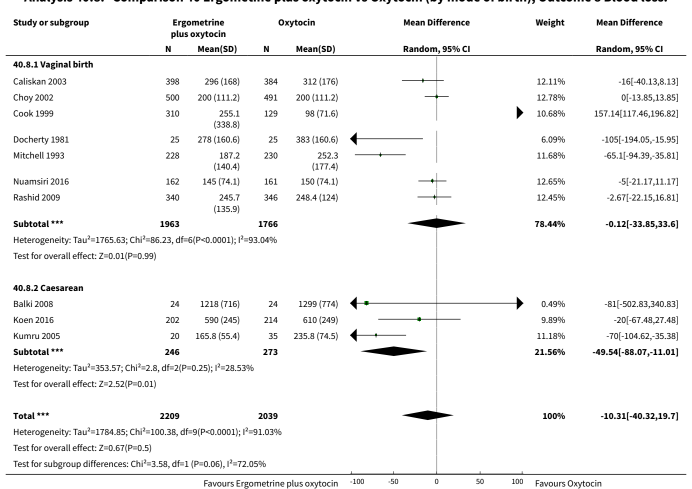
Analysis 40.7. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.







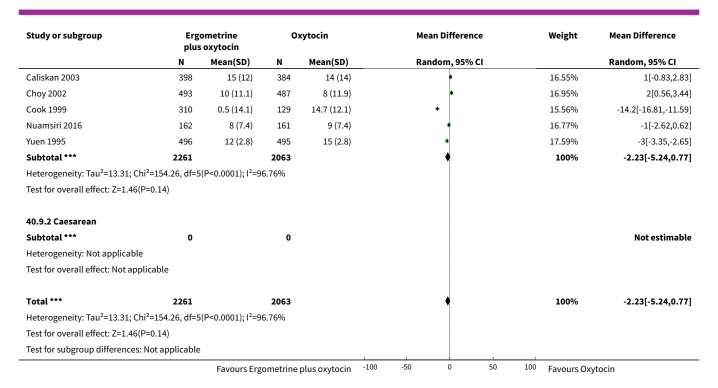
Analysis 40.8. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 8 Blood loss.



Analysis 40.9. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	U	ometrine oxytocin	O	cytocin		Меа	an Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95%	6 CI			Random, 95% CI
40.9.1 Vaginal birth											
Caliskan 2002	402	15 (12)	407	14 (14)			•			16.59%	1[-0.8,2.8]
		Favours Erg	ometrine	plus oxytocin	-100	-50	0	50	100	Favours Oxytoc	in



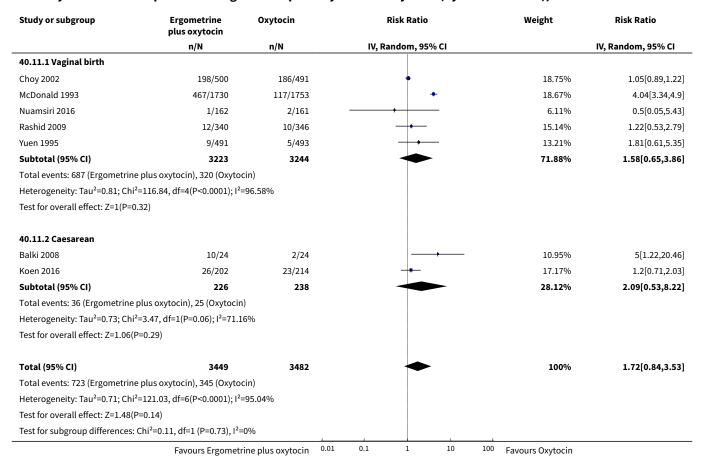


Analysis 40.10. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 10 Breastfeeding.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	ı	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Ra	ndom, 95% CI		IV, Random, 95% CI
40.10.1 Vaginal birth						
McDonald 1993	1478/1730	1518/1753		•	100%	0.99[0.96,1.01]
Subtotal (95% CI)	1730	1753			100%	0.99[0.96,1.01]
Total events: 1478 (Ergometrine plus	s oxytocin), 1518 (Oxyt	tocin)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.99(P=0.32)					
40.10.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Ergometrine plus ox	ytocin), 0 (Oxytocin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	2					
Total (95% CI)	1730	1753			100%	0.99[0.96,1.01]
Total events: 1478 (Ergometrine plus	s oxytocin), 1518 (Oxyt	tocin)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.99(P=0.32)					
Test for subgroup differences: Not ap	oplicable					
	Favours Ergometr	ine plus oxytocin	0.01 0.1	1 10	¹⁰⁰ Favours Oxytocin	



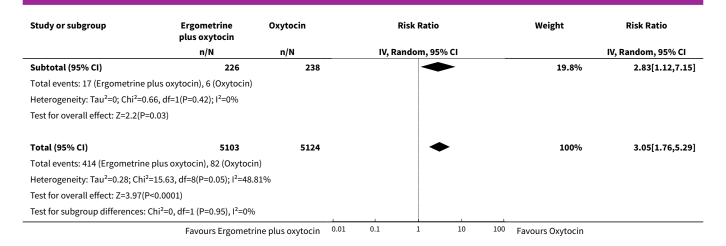
Analysis 40.11. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 11 Nausea.



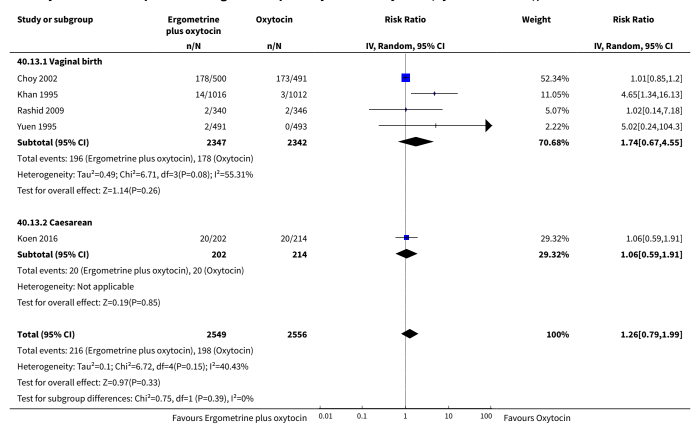
Analysis 40.12. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Ris	sk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Ran	dom, 95% CI		IV, Random, 95% CI	
40.12.1 Vaginal birth							
Caliskan 2002	1/402	2/407			4.44%	0.51[0.05,5.56]	
Caliskan 2003	5/398	3/384	_		9.75%	1.61[0.39,6.68]	
Choy 2002	8/500	2/491		+	8.74%	3.93[0.84,18.4]	
Khan 1995	14/1016	4/1012			13.13%	3.49[1.15,10.55]	
McDonald 1993	358/1730	59/1753		-	26.25%	6.15[4.71,8.03]	
Rashid 2009	4/340	1/346	-	+	5.17%	4.07[0.46,36.23]	
Yuen 1995	7/491	5/493	-	+	12.72%	1.41[0.45,4.4]	
Subtotal (95% CI)	4877	4886		•	80.2%	2.93[1.5,5.71]	
Total events: 397 (Ergometrine	e plus oxytocin), 76 (Oxytoc	n)					
Heterogeneity: Tau ² =0.38; Chi ²	² =13.47, df=6(P=0.04); l ² =55	45%					
Test for overall effect: Z=3.15(F	P=0)						
40.12.2 Caesarean							
Balki 2008	6/24	1/24		+	5.78%	6[0.78,46.14]	
Koen 2016	11/202	5/214		 • .	14.01%	2.33[0.82,6.59]	
	Favours Ergomet	ine plus oxytocin	0.01 0.1	1 10	100 Favours Oxytocin		





Analysis 40.13. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 13 Headache.





Analysis 40.15. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 15 Hypertension.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
40.15.1 Vaginal birth					
Choy 2002	17/500	7/491	-	44.78%	2.38[1,5.7]
Nuamsiri 2016	10/162	0/161	-	24.12%	20.87[1.23,353.21]
Subtotal (95% CI)	662	652		68.91%	4.57[0.65,32.04]
Total events: 27 (Ergometrine plus	s oxytocin), 7 (Oxytocin)				
Heterogeneity: Tau ² =1.21; Chi ² =2.	06, df=1(P=0.15); I ² =51.5	3%			
Test for overall effect: Z=1.53(P=0.	13)				
40.15.2 Caesarean					
Balki 2008	1/24	4/24		31.09%	0.25[0.03,2.08]
Subtotal (95% CI)	24	24		31.09%	0.25[0.03,2.08]
Total events: 1 (Ergometrine plus	oxytocin), 4 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.28(P=0.	2)				
Total (95% CI)	686	676		100%	2[0.29,13.97]
Total events: 28 (Ergometrine plus	s oxytocin), 11 (Oxytocin)			
Heterogeneity: Tau ² =2; Chi ² =6.5, d	ff=2(P=0.04); I ² =69.21%				
Test for overall effect: Z=0.7(P=0.4	9)				
Test for subgroup differences: Chi	² =3.92, df=1 (P=0.05), I ² =	74.48%			
	Favours Froometr	ine plus ovytocin 0.01	0.1 1 10 1	.00 Favours Ovytocin	

Analysis 40.16. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
40.16.1 Vaginal birth					
Caliskan 2002	19/402	16/407	- -	50.89%	1.2[0.63,2.3]
Caliskan 2003	15/398	19/384	 -	49.11%	0.76[0.39,1.48]
Subtotal (95% CI)	800	791	*	100%	0.96[0.6,1.53]
Total events: 34 (Ergometrine plus	oxytocin), 35 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.93,	df=1(P=0.34); I ² =0%				
Test for overall effect: Z=0.17(P=0.8	37)				
40.16.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus o	oxytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	ole				
Total (95% CI)	800	791	*	100%	0.96[0.6,1.53]
Total events: 34 (Ergometrine plus	oxytocin), 35 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.93,	df=1(P=0.34); I ² =0%				
Test for overall effect: Z=0.17(P=0.8	37)				
Test for subgroup differences: Not	applicable				
	Favours Ergometri	ne plus oxytocin 0.01	0.1 1 10 1	00 Favours Oxytocin	



Analysis 40.17. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
40.17.1 Vaginal birth					
Caliskan 2002	6/402	6/407		52.4%	1.01[0.33,3.11]
Caliskan 2003	6/398	5/384		47.6%	1.16[0.36,3.76]
Subtotal (95% CI)	800	791	*	100%	1.08[0.48,2.43]
Total events: 12 (Ergometrine plus o	xytocin), 11 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.03, df	=1(P=0.87); I ² =0%				
Test for overall effect: Z=0.18(P=0.85))				
40.17.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus oxy	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	•				
Total (95% CI)	800	791	•	100%	1.08[0.48,2.43]
Total events: 12 (Ergometrine plus or	xytocin), 11 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.03, df	=1(P=0.87); I ² =0%				
Test for overall effect: Z=0.18(P=0.85))				
Test for subgroup differences: Not ap	plicable				
	Favours Ergometri	ne plus oxytocin 0.01	0.1 1 10 1	.00 Favours Oxytocin	

Analysis 40.18. Comparison 40 Ergometine plus oxytocin vs Oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	•	Weight	Risk Ratio
	n/N	n/N	IV, Random, 9	5% CI		IV, Random, 95% CI
40.18.1 Vaginal birth						
Caliskan 2002	10/402	9/407	_	-	39.93%	1.12[0.46,2.74]
Caliskan 2003	17/398	12/384	-	-	60.07%	1.37[0.66,2.82]
Cook 1999	0/310	0/129				Not estimable
Subtotal (95% CI)	1110	920	•		100%	1.26[0.72,2.22]
Total events: 27 (Ergometrine plus	oxytocin), 21 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0.11, c	ff=1(P=0.74); I ² =0%					
Test for overall effect: Z=0.82(P=0.4	1)					
40.18.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Ergometrine plus o	xytocin), 0 (Oxytocin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicab	le					
Total (95% CI)	1110	920	•		100%	1.26[0.72,2.22]
Total events: 27 (Ergometrine plus	oxytocin), 21 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0.11, c	df=1(P=0.74); I ² =0%		İ			
Test for overall effect: Z=0.82(P=0.4	1)					
	Favours Ergometri	ne plus oxytocin 0.0	01 0.1 1	10 100	Favours Oxytocin	



Study or subgroup	Ergometrine plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
Test for subgroup difference	s: Not applicable								
	Favours Fronmet	rina nlus avvtacin	0.01	0.1	1	10	100	Favours Oxytocin	

Comparison 41. Misoprostol plus oxytocin vs Oxytocin (by mode of birth)

Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	9	4737	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	5	3802	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	4	935	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	17	8514	Risk Ratio (IV, Random, 95% CI)	0.87 [0.69, 1.09]
2.1 Vaginal birth	7	6241	Risk Ratio (IV, Random, 95% CI)	0.77 [0.52, 1.14]
2.2 Caesarean	10	2273	Risk Ratio (IV, Random, 95% CI)	0.93 [0.70, 1.24]
3 Blood transfusion	19	8742	Risk Ratio (IV, Random, 95% CI)	0.50 [0.37, 0.67]
3.1 Vaginal birth	7	5898	Risk Ratio (IV, Random, 95% CI)	0.39 [0.25, 0.61]
3.2 Caesarean	12	2844	Risk Ratio (IV, Random, 95% CI)	0.59 [0.36, 0.96]
4 Severe maternal morbidity: intensive care admissions	3	1886	Risk Ratio (IV, Random, 95% CI)	0.50 [0.05, 5.47]
4.1 Vaginal birth	1	1400	Risk Ratio (IV, Random, 95% CI)	0.50 [0.05, 5.47]
4.2 Caesarean	2	486	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	14	8148	Risk Ratio (IV, Random, 95% CI)	0.71 [0.59, 0.85]
6.1 Vaginal birth	8	6997	Risk Ratio (IV, Random, 95% CI)	0.71 [0.55, 0.92]
6.2 Caesarean	6	1151	Risk Ratio (IV, Random, 95% CI)	0.69 [0.51, 0.92]
7 Additional uterotonics	18	8391	Risk Ratio (IV, Random, 95% CI)	0.54 [0.44, 0.67]
7.1 Vaginal birth	7	5898	Risk Ratio (IV, Random, 95% CI)	0.63 [0.49, 0.79]



Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
7.2 Caesarean	11	2493	Risk Ratio (IV, Random, 95% CI)	0.49 [0.36, 0.66]
8 Blood loss	17	8690	Mean Difference (IV, Random, 95% CI)	-87.26 [-157.83, -16.69
8.1 Vaginal birth	7	6179	Mean Difference (IV, Random, 95% CI)	-12.23 [-26.51, 2.06]
8.2 Caesarean	10	2511	Mean Difference (IV, Random, 95% CI)	-134.79 [-276.45, 6.88]
9 Change in haemo- globin	15	7929	Mean Difference (IV, Random, 95% CI)	-2.59 [-3.70, -1.48]
9.1 Vaginal birth	6	5643	Mean Difference (IV, Random, 95% CI)	-0.89 [-1.61, -0.17]
9.2 Caesarean	9	2286	Mean Difference (IV, Random, 95% CI)	-2.00 [-5.60, -2.39]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	7	3798	Risk Ratio (IV, Random, 95% CI)	2.21 [1.19, 4.10]
11.1 Vaginal birth	2	3003	Risk Ratio (IV, Random, 95% CI)	3.52 [1.55, 7.99]
11.2 Caesarean	5	795	Risk Ratio (IV, Random, 95% CI)	1.71 [0.76, 3.84]
12 Vomiting	11	6718	Risk Ratio (IV, Random, 95% CI)	2.24 [1.52, 3.31]
12.1 Vaginal birth	6	5610	Risk Ratio (IV, Random, 95% CI)	3.32 [2.03, 5.44]
12.2 Caesarean	5	1108	Risk Ratio (IV, Random, 95% CI)	1.51 [0.96, 2.36]
13 Headache	2	303	Risk Ratio (IV, Random, 95% CI)	1.26 [0.26, 6.23]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	2	303	Risk Ratio (IV, Random, 95% CI)	1.26 [0.26, 6.23]
14 Abdominal pain	1	366	Risk Ratio (IV, Random, 95% CI)	1.93 [1.01, 3.67]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	1	366	Risk Ratio (IV, Random, 95% CI)	1.93 [1.01, 3.67]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	19	9458	Risk Ratio (IV, Random, 95% CI)	3.38 [2.50, 4.57]
16.1 Vaginal birth	8	7007	Risk Ratio (IV, Random, 95% CI)	3.68 [2.41, 5.60]

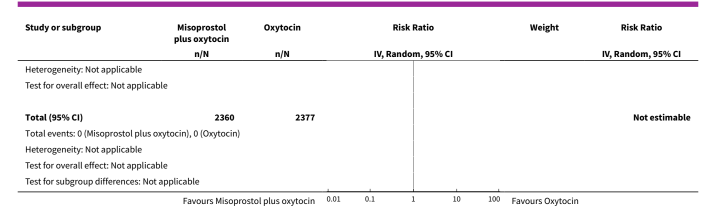


Outcome or sub- group title	No. of studies	No. of participants	Statistical method	Effect size
16.2 Caesarean	11	2451	Risk Ratio (IV, Random, 95% CI)	3.04 [2.00, 4.61]
17 Fever	17	8607	Risk Ratio (IV, Random, 95% CI)	2.99 [2.00, 4.45]
17.1 Vaginal birth	7	6209	Risk Ratio (IV, Random, 95% CI)	4.30 [2.57, 7.21]
17.2 Caesarean	10	2398	Risk Ratio (IV, Random, 95% CI)	1.85 [1.28, 2.67]
18 Diarrhoea	7	5649	Risk Ratio (IV, Random, 95% CI)	2.08 [0.99, 4.38]
18.1 Vaginal birth	5	4887	Risk Ratio (IV, Random, 95% CI)	1.89 [0.82, 4.36]
18.2 Caesarean	2	762	Risk Ratio (IV, Random, 95% CI)	6.07 [0.73, 50.35]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 41.1. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
41.1.1 Vaginal birth					
Badejoko 2012	0/126	0/129			Not estimable
Bhullar 2004	0/377	0/379			Not estimable
Carbonell 2009	0/702	0/698			Not estimable
Chaudhuri 2016	0/144	0/144			Not estimable
Hofmeyr 2011	0/546	0/557			Not estimable
Subtotal (95% CI)	1895	1907			Not estimable
Total events: 0 (Misoprostol plu	s oxytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not appli	cable				
41.1.2 Caesarean					
Chaudhuri 2015	0/198	0/198			Not estimable
El Tahan 2012	0/179	0/187			Not estimable
Lapaire 2006	0/28	0/25			Not estimable
Ugwu 2014	0/60	0/60			Not estimable
Subtotal (95% CI)	465	470			Not estimable
Total events: 0 (Misoprostol plu	s oxytocin), 0 (Oxytocin)				





Analysis 41.2. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
41.2.1 Vaginal birth						
Badejoko 2012	3/126	5/129		2.55%	0.61[0.15,2.52]	
Caliskan 2002	11/401	14/407	-+ -	8.39%	0.8[0.37,1.74]	
Caliskan 2003	6/404	15/384		5.78%	0.38[0.15,0.97]	
Carbonell 2009	13/702	11/698		8%	1.18[0.53,2.6	
Chaudhuri 2016	2/144	4/144		1.79%	0.5[0.09,2.69	
Hofmeyr 2011	5/546	1/553	+	1.1%	5.06[0.59,43.2	
Quibel 2016	13/806	17/797		9.9%	0.76[0.37,1.55	
Subtotal (95% CI)	3129	3112	•	37.52%	0.77[0.52,1.14]	
Total events: 53 (Misoprostol p	olus oxytocin), 67 (Oxytocin)					
Heterogeneity: Tau²=0.03; Chi²	² =6.59, df=6(P=0.36); I ² =8.98%	6				
Test for overall effect: Z=1.29(F	P=0.2)					
41.2.2 Caesarean						
Adanikin 2012	0/109	0/109			Not estimable	
Chaudhuri 2015	5/198	3/198		2.52%	1.67[0.4,6.88	
Elsedeek 2012	0/200	0/200			Not estimable	
Fekih 2009	19/125	24/125	-+ 	16.86%	0.79[0.46,1.37	
Hamm 2005	24/173	22/179	-	17.42%	1.13[0.66,1.94	
Hernandez-Castro 2016	3/60	7/60		2.98%	0.43[0.12,1.58	
Lapaire 2006	13/24	11/19	-	17.95%	0.94[0.55,1.59	
Sitaula 2017	0/100	1/100 —		0.5%	0.33[0.01,8.09	
Sood 2012	6/90	4/84		3.35%	1.4[0.41,4.79	
Ugwu 2014	1/60	2/60		0.9%	0.5[0.05,5.37	
Subtotal (95% CI)	1139	1134	*	62.48%	0.93[0.7,1.24	
Total events: 71 (Misoprostol p	olus oxytocin), 74 (Oxytocin)					
Heterogeneity: Tau²=0; Chi²=3	.92, df=7(P=0.79); I ² =0%					
Test for overall effect: Z=0.47(F	P=0.64)					
Total (95% CI)	4268	4246	•	100%	0.87[0.69,1.09]	
Total events: 124 (Misoprostol	plus oxytocin), 141 (Oxytocir	1)				
Heterogeneity: Tau²=0; Chi²=1	1.14, df=14(P=0.67); I ² =0%					
Test for overall effect: Z=1.22(F	P=0.22)					
Test for subgroup differences:	Chi ² =0.59, df=1 (P=0.44), I ² =0	%				

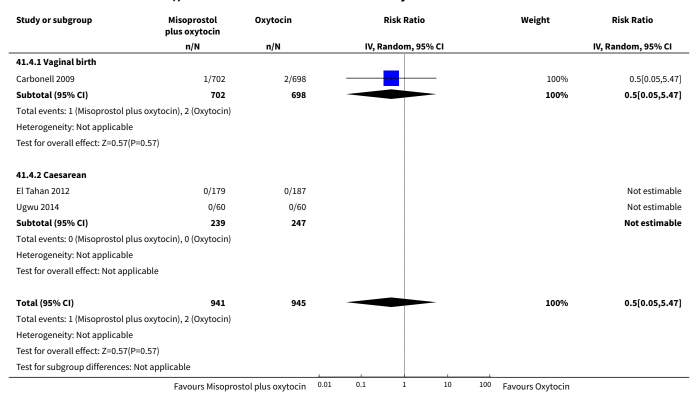


Analysis 41.3. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
41.3.1 Vaginal birth					
Badejoko 2012	1/126	6/129	+	1.89%	0.17[0.02,1.4]
Bhullar 2004	3/377	6/379		4.41%	0.5[0.13,2]
Caliskan 2002	4/401	13/407		6.77%	0.31[0.1,0.95]
Caliskan 2003	5/404	13/384		8.02%	0.37[0.13,1.02]
Carbonell 2009	5/702	13/698		7.96%	0.38[0.14,1.07]
Chaudhuri 2016	5/144	12/144		8.09%	0.42[0.15,1.15]
Quibel 2016	5/806	9/797		7.07%	0.55[0.18,1.63]
Subtotal (95% CI)	2960	2938	•	44.22%	0.39[0.25,0.61]
Total events: 28 (Misoprostol	plus oxytocin), 72 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =1	1.29, df=6(P=0.97); I ² =0%				
Test for overall effect: Z=4.21(P<0.0001)				
41.3.2 Caesarean					
Chaudhuri 2015	10/198	15/198	-++	13.93%	0.67[0.31,1.45]
El Tahan 2012	0/179	11/187		1.05%	0.05[0,0.76]
Elsedeek 2012	0/200	0/200			Not estimable
Fekih 2009	0/125	4/125		0.99%	0.11[0.01,2.04]
Hamm 2005	3/173	3/179		3.33%	1.03[0.21,5.06]
Hernandez-Castro 2016	0/60	5/60	+	1.01%	0.09[0.01,1.61]
Hong 2007	11/96	13/118		14.64%	1.04[0.49,2.22]
Lapaire 2006	0/28	0/25			Not estimable
Nayak 2017	9/100	23/100		16.2%	0.39[0.19,0.8]
Sitaula 2017	0/100	1/100 —	•	0.82%	0.33[0.01,8.09]
Sood 2012	3/90	2/84		2.69%	1.4[0.24,8.17]
Ugwu 2014	1/60	1/59		1.11%	0.98[0.06,15.36]
Subtotal (95% CI)	1409	1435	•	55.78%	0.59[0.36,0.96]
Total events: 37 (Misoprostol	plus oxytocin), 78 (Oxytocin)				
Heterogeneity: Tau ² =0.11; Chi					
Test for overall effect: Z=2.14(
Total (95% CI)	4369	4373	•	100%	0.5[0.37,0.67]
Total events: 65 (Misoprostol			•		
Heterogeneity: Tau ² =0; Chi ² =1		-1			
Test for overall effect: Z=4.7(P					
Test for subgroup differences:		31 91%			
sasp. sap amerences.	Favours Misopros		0.1 1 10	100 Favours Oxytocin	



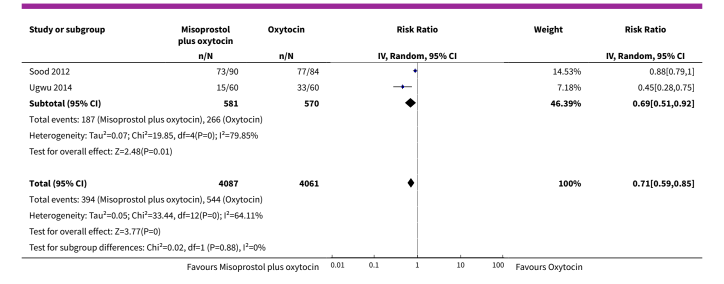
Analysis 41.4. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.



Analysis 41.6. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
41.6.1 Vaginal birth						
Badejoko 2012	28/126	27/129	-	7.6%	1.06[0.66,1.7]	
Bhullar 2004	13/377	20/379	-+ 	4.82%	0.65[0.33,1.29]	
Caliskan 2002	28/401	33/407	-	7.34%	0.86[0.53,1.4]	
Caliskan 2003	13/404	28/384		5.24%	0.44[0.23,0.84]	
Carbonell 2009	28/702	50/698		7.9%	0.56[0.35,0.87]	
Chaudhuri 2016	7/144	19/144		3.6%	0.37[0.16,0.85]	
Hofmeyr 2011	22/546	35/553	-+ 	6.79%	0.64[0.38,1.07]	
Quibel 2016	68/806	66/797	+	10.34%	1.02[0.74,1.41]	
Subtotal (95% CI)	3506	3491	•	53.61%	0.71[0.55,0.92]	
Total events: 207 (Misoprosto	ol plus oxytocin), 278 (Oxytoc	in)				
Heterogeneity: Tau ² =0.06; Ch	ni ² =13.45, df=7(P=0.06); l ² =47	96%				
Test for overall effect: Z=2.63	(P=0.01)					
41.6.2 Caesarean						
Adanikin 2012	0/109	0/109			Not estimable	
Chaudhuri 2015	79/198	132/198	+	13.09%	0.6[0.49,0.73]	
Lapaire 2006	18/24	15/19	+	10.27%	0.95[0.68,1.32]	
Nayak 2017	2/100	9/100		1.32%	0.22[0.05,1]	
	Favours Misopro	stol plus oxytocin	0.01 0.1 1 10	100 Favours Oxytocin		

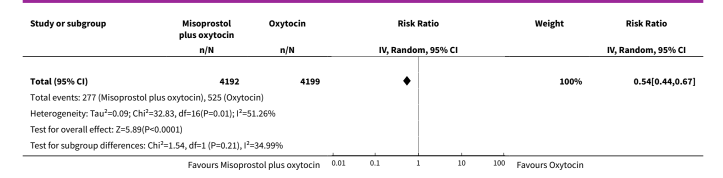




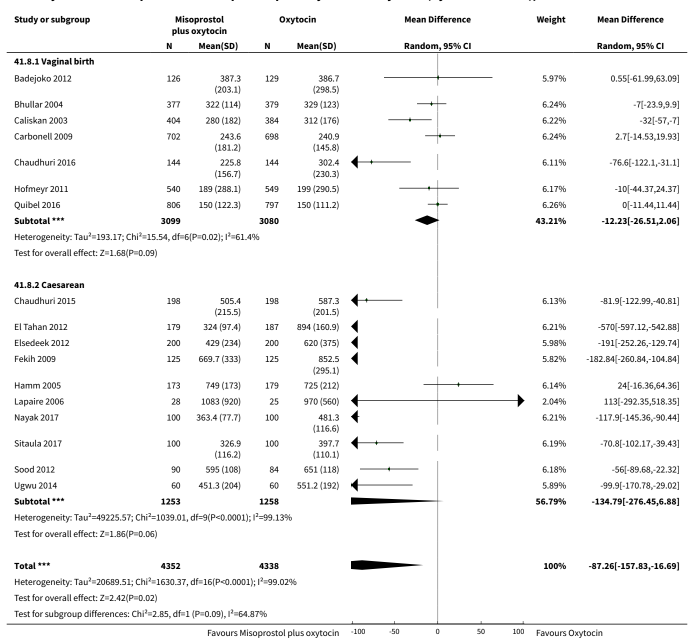
Analysis 41.7. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI	
	n/N	n/N	IV, Random, 95% CI			
41.7.1 Vaginal birth						
Badejoko 2012	6/126	5/129		2.48%	1.23[0.38,3.92]	
Bhullar 2004	10/377	13/379		4.2%	0.77[0.34,1.74]	
Caliskan 2002	17/401	26/407	+	6.08%	0.66[0.37,1.2]	
Caliskan 2003	10/404	26/384		4.94%	0.37[0.18,0.75]	
Carbonell 2009	33/702	54/698		8.23%	0.61[0.4,0.92]	
Chaudhuri 2016	12/144	22/144		5.39%	0.55[0.28,1.06]	
Quibel 2016	19/806	25/797	-+	6.16%	0.75[0.42,1.35]	
Subtotal (95% CI)	2960	2938	•	37.47%	0.63[0.49,0.79]	
Total events: 107 (Misoprostol p	olus oxytocin), 171 (Oxytoc	in)				
Heterogeneity: Tau²=0; Chi²=4.3	32, df=6(P=0.63); I ² =0%					
Test for overall effect: Z=3.89(P=	=0)					
41.7.2 Caesarean						
Chaudhuri 2015	18/198	45/198		7.06%	0.4[0.24,0.67]	
El Tahan 2012	12/179	52/187		6.1%	0.24[0.13,0.44]	
Elsedeek 2012	14/200	36/200		6.19%	0.39[0.22,0.7]	
Hamm 2005	45/173	76/179		9.88%	0.61[0.45,0.83]	
Hernandez-Castro 2016	6/60	24/60		4.15%	0.25[0.11,0.57]	
Hong 2007	28/96	31/118	-	8.04%	1.11[0.72,1.71]	
Lapaire 2006	0/28	0/25			Not estimable	
Nayak 2017	4/100	7/100		2.36%	0.57[0.17,1.89]	
Pakniat 2015	7/50	7/50		3.26%	1[0.38,2.64]	
Sood 2012	20/90	36/84	→	7.71%	0.52[0.33,0.82]	
Ugwu 2014	16/58	40/60	 -	7.78%	0.41[0.26,0.65]	
Subtotal (95% CI)	1232	1261	◆	62.53%	0.49[0.36,0.66	
Total events: 170 (Misoprostol p	olus oxytocin), 354 (Oxytoc	in)				
Heterogeneity: Tau²=0.14; Chi²=	27.05, df=9(P=0); I ² =66.72	%				
Test for overall effect: Z=4.63(P<	<0.0001)		ĺ			



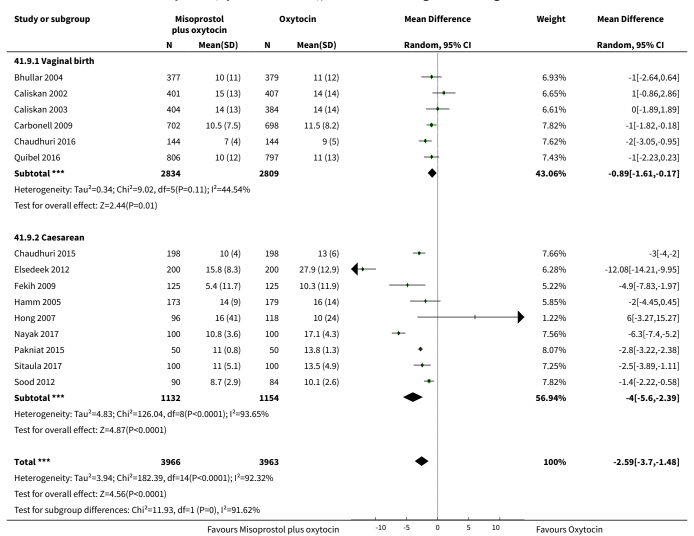


Analysis 41.8. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 8 Blood loss.





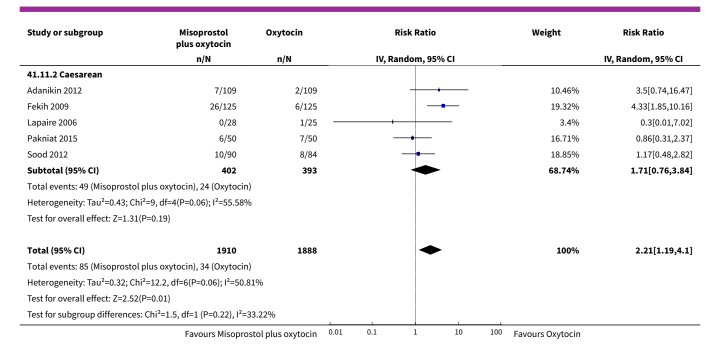
Analysis 41.9. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.



Analysis 41.11. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	ytocin Risk Ratio		Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
41.11.1 Vaginal birth					
Carbonell 2009	14/702	2/698		11.11%	6.96[1.59,30.51]
Quibel 2016	22/806	8/797		20.15%	2.72[1.22,6.07]
Subtotal (95% CI)	1508	1495	-	31.26%	3.52[1.55,7.99]
Total events: 36 (Misoprostol	plus oxytocin), 10 (Oxytocin)	ı			
Heterogeneity: Tau ² =0.07; Ch	i ² =1.2, df=1(P=0.27); I ² =16.61	%			
Test for overall effect: Z=3(P=	0)				
	Favours Misopros	stol plus oxytocin 0.0	1 0.1 1 10	100 Favours Oxytocin	





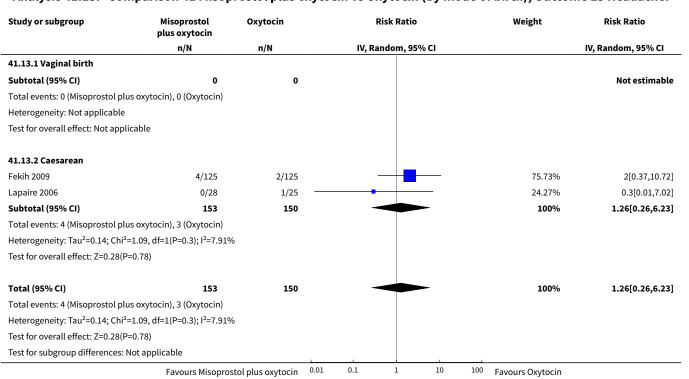
Analysis 41.12. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
41.12.1 Vaginal birth						
Badejoko 2012	29/126	7/129		14.9%	4.24[1.93,9.33]	
Bhullar 2004	5/377	2/379	+	4.94%	2.51[0.49,12.87]	
Caliskan 2002	3/401	2/407		4.23%	1.52[0.26,9.06]	
Caliskan 2003	3/404	3/384		5.16%	0.95[0.19,4.68]	
Carbonell 2009	28/702	4/698		10.22%	6.96[2.45,19.74]	
Quibel 2016	18/806	6/797		12.22%	2.97[1.18,7.43]	
Subtotal (95% CI)	2816	2794	•	51.67%	3.32[2.03,5.44]	
Total events: 86 (Misoprostol p	lus oxytocin), 24 (Oxytocin)					
Heterogeneity: Tau ² =0.04; Chi ² =	=5.57, df=5(P=0.35); I ² =10.2	%				
Test for overall effect: Z=4.78(P-	<0.0001)					
41.12.2 Caesarean						
Adanikin 2012	8/109	3/109	+	7.26%	2.67[0.73,9.79]	
El Tahan 2012	16/179	12/187	+-	16.57%	1.39[0.68,2.86]	
Fekih 2009	15/125	10/125	 •	15.54%	1.5[0.7,3.21]	
Pakniat 2015	1/50	2/50		2.52%	0.5[0.05,5.34]	
Sood 2012	5/90	3/84		6.43%	1.56[0.38,6.31]	
Subtotal (95% CI)	553	555	•	48.33%	1.51[0.96,2.36]	
Total events: 45 (Misoprostol p	lus oxytocin), 30 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =1.6	62, df=4(P=0.8); I ² =0%					
Test for overall effect: Z=1.78(P	=0.07)					
Total (95% CI)	3369	3349	•	100%	2.24[1.52,3.31]	
Total events: 131 (Misoprostol)	olus oxytocin), 54 (Oxytoci	1)				
Heterogeneity: Tau ² =0.1; Chi ² =	13.34, df=10(P=0.21); l ² =25.	06%				



Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, Ra	andom, 95	% CI			IV, Random, 95% CI
Test for overall effect: Z=4.06	6(P<0.0001)								
Test for subgroup difference	s: Chi ² =5.41, df=1 (P=0.02), I ² =	=81.51%				1			
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	

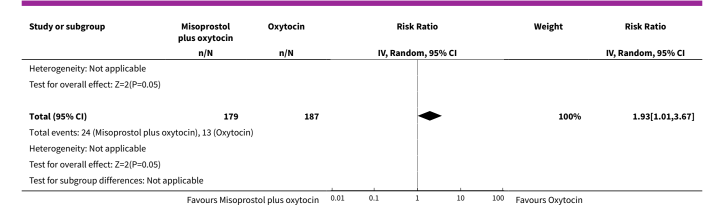
Analysis 41.13. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 13 Headache.



Analysis 41.14. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
41.14.1 Vaginal birth								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol p	olus oxytocin), 0 (Oxytocin)							
Heterogeneity: Not applicable	e							
Test for overall effect: Not app	plicable							
41.14.2 Caesarean								
El Tahan 2012	24/179	13/187			-		100%	1.93[1.01,3.67]
Subtotal (95% CI)	179	187			•		100%	1.93[1.01,3.67]
Total events: 24 (Misoprostol	plus oxytocin), 13 (Oxytocin)					1		
	Favours Misoprost	ol plus oxytocin	0.01	0.1	1 10	100	Favours Oxytocin	





Analysis 41.16. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
41.16.1 Vaginal birth						
Badejoko 2012	34/126	17/129		8.46%	2.05[1.21,3.47]	
Bhullar 2004	2/377	1/379		1.39%	2.01[0.18,22.08]	
Caliskan 2002	52/401	16/407		8.33%	3.3[1.92,5.68]	
Caliskan 2003	49/404	19/384		8.6%	2.45[1.47,4.09]	
Carbonell 2009	390/702	81/698	+	10.8%	4.79[3.86,5.94]	
Chaudhuri 2016	39/144	6/144		6.13%	6.5[2.84,14.88]	
Hofmeyr 2011	142/544	67/556	+	10.5%	2.17[1.66,2.82]	
Quibel 2016	87/806	5/806	_ 	5.68%	17.4[7.1,42.63]	
Subtotal (95% CI)	3504	3503	•	59.89%	3.68[2.41,5.6]	
Total events: 795 (Misoprosto	ol plus oxytocin), 212 (Oxytoc	in)				
Heterogeneity: Tau²=0.25; Ch	ni ² =41.19, df=7(P<0.0001); l ² =	83.01%				
Test for overall effect: Z=6.07	(P<0.0001)					
41.16.2 Caesarean						
Adanikin 2012	11/109	8/109		5.84%	1.38[0.58,3.29]	
Cayan 2010	6/120	2/40		2.81%	1[0.21,4.76]	
Chaudhuri 2015	42/198	11/198		7.59%	3.82[2.03,7.2]	
El Tahan 2012	17/179	6/187		5.61%	2.96[1.19,7.34]	
Elsedeek 2012	0/200	0/200			Not estimable	
Fekih 2009	7/125	1/125	 	1.77%	7[0.87,56.06]	
Hong 2007	1/96	0/118		- 0.83%	3.68[0.15,89.33]	
Lapaire 2006	10/28	2/25		3.23%	4.46[1.08,18.45]	
Pakniat 2015	0/50	0/50			Not estimable	
Sood 2012	19/90	8/84	—	6.54%	2.22[1.03,4.79]	
Ugwu 2014	37/60	5/60		5.9%	7.4[3.12,17.53]	
Subtotal (95% CI)	1255	1196	•	40.11%	3.04[2,4.61]	
Total events: 150 (Misoprosto	ol plus oxytocin), 43 (Oxytoci	n)				
Heterogeneity: Tau²=0.11; Ch	ni ² =11.28, df=8(P=0.19); l ² =29	.07%				
Test for overall effect: Z=5.23	(P<0.0001)					
Total (95% CI)	4759	4699	•	100%	3.38[2.5,4.57]	
Total events: 945 (Misoprosto	ol plus oxytocin), 255 (Oxytoc	in)				
Heterogeneity: Tau ² =0.21: Ch	ni ² =52.87, df=16(P<0.0001); I ²	=69.74%				



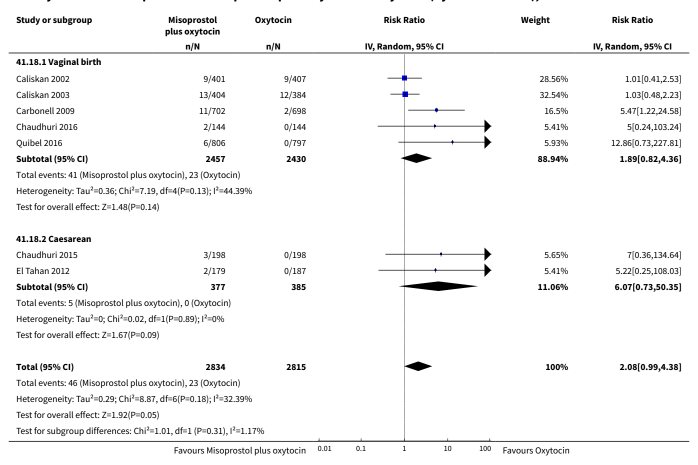
Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio		Weight		Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Test for overall effect: Z=7.93	(P<0.0001)								
Test for subgroup differences	s: Chi ² =0.4, df=1 (P=0.53), I ² =0	0%							
	Favours Misopro	stol plus oxvtocin	0.01	0.1	1	10	100	Favours Oxvtocin	

Analysis 41.17. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI	
	n/N	n/N	IV, Random, 95% CI			
41.17.1 Vaginal birth						
Badejoko 2012	28/126	7/129		7.07%	4.1[1.86,9.03]	
Caliskan 2002	19/401	6/407		6.5%	3.21[1.3,7.96]	
Caliskan 2003	16/404	5/384		6.09%	3.04[1.13,8.22]	
Carbonell 2009	272/702	23/698	-	8.86%	11.76[7.78,17.76]	
Chaudhuri 2016	8/144	2/144	 	4.01%	4[0.86,18.51]	
Hofmeyr 2011	61/522	28/536	-	8.78%	2.24[1.45,3.44]	
Quibel 2016	245/806	50/806	-+-	9.31%	4.9[3.67,6.54]	
Subtotal (95% CI)	3105	3104	•	50.62%	4.3[2.57,7.21]	
Total events: 649 (Misoprosto	ol plus oxytocin), 121 (Oxytoc	cin)				
Heterogeneity: Tau ² =0.34; Ch	ni ² =31.97, df=6(P<0.0001); l ² =	81.23%				
Test for overall effect: Z=5.54	(P<0.0001)					
41.17.2 Caesarean						
Adanikin 2012	8/109	6/109		5.95%	1.33[0.48,3.71]	
Cayan 2010	6/120	1/40		2.66%	2[0.25,16.11]	
Chaudhuri 2015	13/198	4/198		5.6%	3.25[1.08,9.79]	
El Tahan 2012	16/179	8/187	 • •	6.91%	2.09[0.92,4.76]	
Elsedeek 2012	11/200	13/200		7.13%	0.85[0.39,1.84]	
Fekih 2009	9/125	2/125	<u> </u>	4.07%	4.5[0.99,20.41]	
Hong 2007	10/96	5/118	 • •	5.88%	2.46[0.87,6.95]	
Pakniat 2015	2/50	1/50		2.2%	2[0.19,21.36]	
Sood 2012	10/90	6/84		6.21%	1.56[0.59,4.09]	
Ugwu 2014	9/60	1/60	ļ ——	- 2.76%	9[1.18,68.85]	
Subtotal (95% CI)	1227	1171	•	49.38%	1.85[1.28,2.67]	
Total events: 94 (Misoprostol	plus oxytocin), 47 (Oxytocin)				
Heterogeneity: Tau ² =0.02; Ch	ni ² =9.42, df=9(P=0.4); I ² =4.499	%				
Test for overall effect: Z=3.3(F						
Total (95% CI)	4332	4275	•	100%	2.99[2,4.45]	
Total events: 743 (Misoprosto	ol plus oxytocin), 168 (Oxvtoc	cin)			- , -	
Heterogeneity: Tau ² =0.42; Ch						
Test for overall effect: Z=5.37						
Test for subgroup differences	•	=85.39%				



Analysis 41.18. Comparison 41 Misoprostol plus oxytocin vs Oxytocin (by mode of birth), Outcome 18 Diarrhoea.



Comparison 42. Injectable prostaglandins vs Misoprostol (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	170	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.1 Vaginal birth	2	170	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	4	403	Risk Ratio (IV, Random, 95% CI)	0.88 [0.14, 5.39]
3.1 Vaginal birth	4	403	Risk Ratio (IV, Random, 95% CI)	0.88 [0.14, 5.39]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	303	Risk Ratio (IV, Random, 95% CI)	1.60 [0.57, 4.52]
6.1 Vaginal birth	3	303	Risk Ratio (IV, Random, 95% CI)	1.60 [0.57, 4.52]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	4	403	Risk Ratio (IV, Random, 95% CI)	1.02 [0.28, 3.69]
7.1 Vaginal birth	4	403	Risk Ratio (IV, Random, 95% CI)	1.02 [0.28, 3.69]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	3	270	Mean Difference (IV, Random, 95% CI)	52.99 [-39.74, 145.71]
8.1 Vaginal birth	3	270	Mean Difference (IV, Random, 95% CI)	52.99 [-39.74, 145.71]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	2	220	Mean Difference (IV, Random, 95% CI)	2.28 [-1.35, 5.90]
9.1 Vaginal birth	2	220	Mean Difference (IV, Random, 95% CI)	2.28 [-1.35, 5.90]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	2	233	Risk Ratio (IV, Random, 95% CI)	0.23 [0.06, 0.92]
11.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	0.23 [0.06, 0.92]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	233	Risk Ratio (IV, Random, 95% CI)	1.31 [0.01, 163.15]



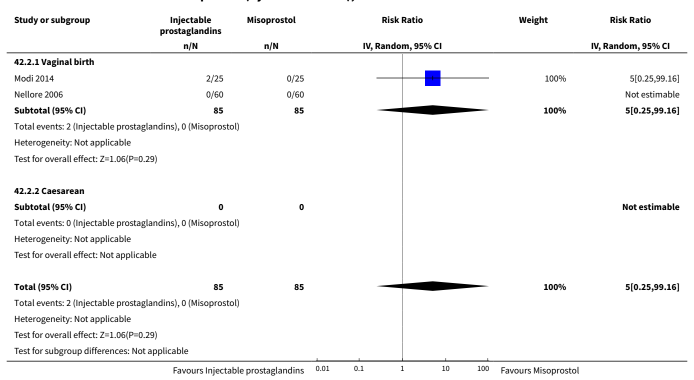
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	1.31 [0.01, 163.15]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	0.99 [0.20, 4.80]
14.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	0.99 [0.20, 4.80]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	353	Risk Ratio (IV, Random, 95% CI)	0.04 [0.01, 0.21]
16.1 Vaginal birth	3	353	Risk Ratio (IV, Random, 95% CI)	0.04 [0.01, 0.21]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	233	Risk Ratio (IV, Random, 95% CI)	0.08 [0.01, 0.39]
17.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	0.08 [0.01, 0.39]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	233	Risk Ratio (IV, Random, 95% CI)	9.03 [1.70, 48.00]
18.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	9.03 [1.70, 48.00]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 42.1. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 1 Death.

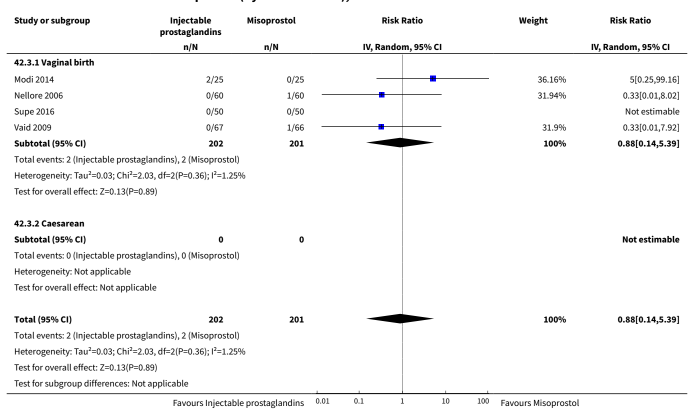
Study or subgroup	Injectable prostaglandins	Misoprostol		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 9	95% CI			IV, Random, 95% CI
42.1.1 Vaginal birth								
Supe 2016	0/50	0/50						Not estimable
Subtotal (95% CI)	50	50						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Misoprostol))						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
42.1.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Misoprostol))						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
Total (95% CI)	50	50						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Misoprostol))						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
Test for subgroup differences: Not ap	oplicable							
	Favours Injectabl	le prostaglandins	0.01	0.1 1	10	100	Favours Misoprostol	

Analysis 42.2. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 2 PPH >= 1000 mL.





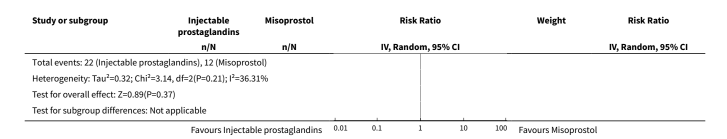
Analysis 42.3. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 3 Blood transfusion.



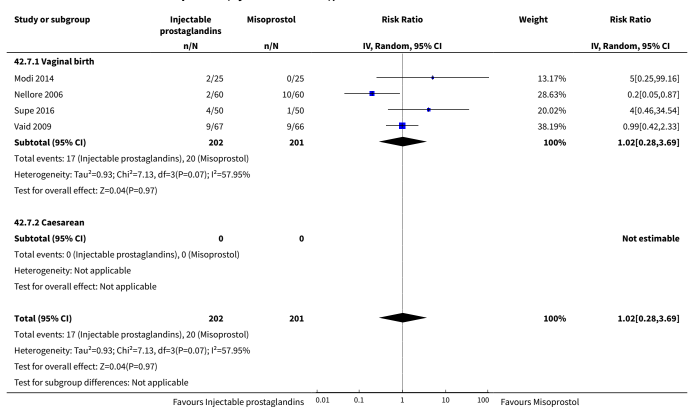
Analysis 42.6. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N IV, Random, 95% CI			IV, Random, 95% CI
42.6.1 Vaginal birth					
Modi 2014	6/25	0/25	+	11.65%	13[0.77,219.11]
Nellore 2006	3/60	4/60		31.99%	0.75[0.18,3.21]
Vaid 2009	13/67	8/66	 	56.36%	1.6[0.71,3.61]
Subtotal (95% CI)	152	151		100%	1.6[0.57,4.52]
Total events: 22 (Injectable pr	ostaglandins), 12 (Misopros	tol)			
Heterogeneity: Tau ² =0.32; Chi	² =3.14, df=2(P=0.21); I ² =36.3	1%			
Test for overall effect: Z=0.89(I	P=0.37)				
42.6.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable pro	staglandins), 0 (Misoprostol)			
Heterogeneity: Not applicable	2				
Test for overall effect: Not app	olicable				
Total (95% CI)	152	151	•	100%	1.6[0.57,4.52]
	Favours Injectab	le prostaglandins	0.01 0.1 1 10 100	Favours Misoprostol	





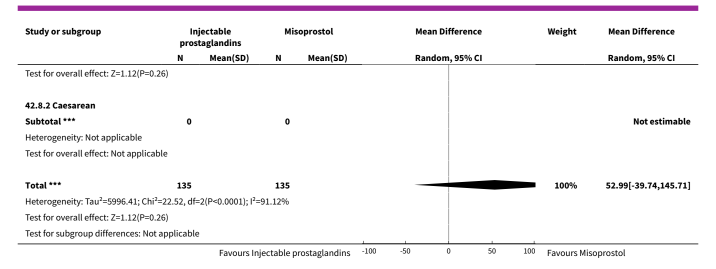
Analysis 42.7. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 7 Additional uterotonics.



Analysis 42.8. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 8 Blood loss.

Study or subgroup		jectable taglandins	Mis	soprostol		Mean D	ifference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Randon	n, 95% CI			Random, 95% CI
42.8.1 Vaginal birth										
Modi 2014	25	435 (147.6)	25	255.8 (102.2)				•	30.72%	179.2[108.84,249.56]
Nellore 2006	60	205 (175)	60	245 (158)		-			32.33%	-40[-99.66,19.66]
Supe 2016	50	153.8 (43.5)	50	124.4 (34.7)			_		36.95%	29.4[13.98,44.82]
Subtotal ***	135		135						100%	52.99[-39.74,145.71]
Heterogeneity: Tau ² =5996.41;	Chi ² =22.52, df=	2(P<0.0001); I ² =9	91.12%							
		Favours Inj	ectable p	rostaglandins	-100	-50	0 50	100	Favours Mis	soprostol





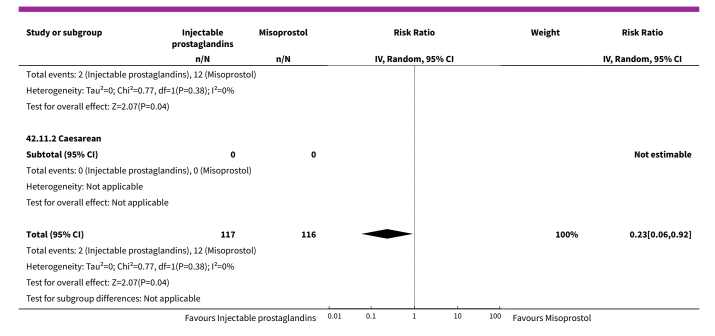
Analysis 42.9. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup		Injectable ostaglandins		oprostol	Mean Differ	ence	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95	% CI		Random, 95% CI
42.9.1 Vaginal birth								
Nellore 2006	60	6.2 (2.8)	60	5.8 (2.3)	•		49.32%	0.4[-0.52,1.32]
Supe 2016	50	9 (1.2)	50	4.9 (0.5)			50.68%	4.1[3.74,4.46]
Subtotal ***	110		110		•		100%	2.28[-1.35,5.9]
Heterogeneity: Tau ² =6.72; Chi ² =54.	19, df=1(P	<0.0001); I ² =98.1	5%					
Test for overall effect: Z=1.23(P=0.2	2)							
42.9.2 Caesarean								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
Test for overall effect: Not applicab	le							
Total ***	110		110		*		100%	2.28[-1.35,5.9]
Heterogeneity: Tau ² =6.72; Chi ² =54.	19, df=1(P	<0.0001); I ² =98.1	5%					
Test for overall effect: Z=1.23(P=0.2	2)							
Test for subgroup differences: Not a	applicable							
		Favours Inj	ectable p	rostaglandins -100	-50 0	50 100	Favours Mis	oprostol

Analysis 42.11. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Misoprostol		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95	5% CI			IV, Random, 95% CI
42.11.1 Vaginal birth									
Supe 2016	0/50	6/50	\leftarrow	-				23.14%	0.08[0,1.33]
Vaid 2009	2/67	6/66			-			76.86%	0.33[0.07,1.57]
Subtotal (95% CI)	117	116			-	1		100%	0.23[0.06,0.92]
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	



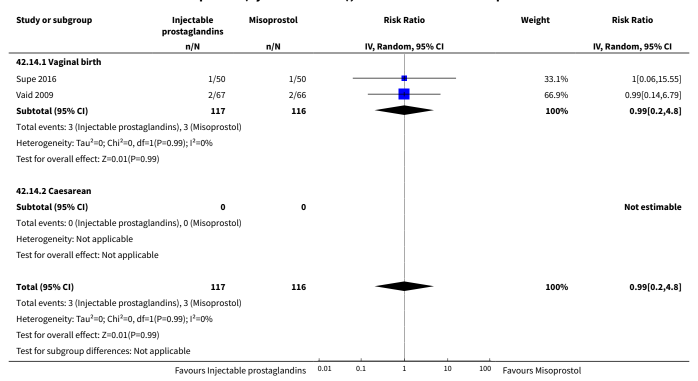


Analysis 42.12. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
42.12.1 Vaginal birth					
Supe 2016	8/50	0/50		47.97%	17[1.01,286.82]
Vaid 2009	1/67	8/66		52.03%	0.12[0.02,0.96]
Subtotal (95% CI)	117	116		100%	1.31[0.01,163.15]
Total events: 9 (Injectable prostagl	andins), 8 (Misoprostol)				
Heterogeneity: Tau ² =10.55; Chi ² =7.	65, df=1(P=0.01); l ² =86.9	93%			
Test for overall effect: Z=0.11(P=0.9	1)				
42.12.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagl	andins), 0 (Misoprostol)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	117	116		- 100%	1.31[0.01,163.15]
Total events: 9 (Injectable prostagl	andins), 8 (Misoprostol)				
Heterogeneity: Tau ² =10.55; Chi ² =7.	65, df=1(P=0.01); I ² =86.9	93%			
Test for overall effect: Z=0.11(P=0.9	1)				
Test for subgroup differences: Not a	applicable				
	Favours Injectable	e prostaglandins 0.	.01 0.1 1 10 10	⁰⁰ Favours Misoprostol	_



Analysis 42.14. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 14 Abdominal pain.

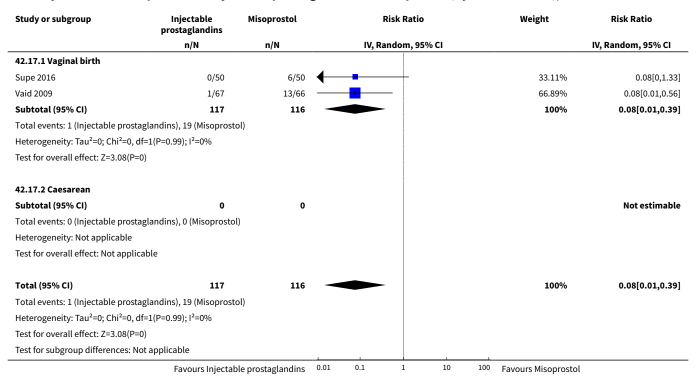


Analysis 42.16. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95%	CI	IV, Random, 95% CI
42.16.1 Vaginal birth					
Nellore 2006	0/60	5/60	-	32.08%	0.09[0.01,1.61]
Supe 2016	0/50	10/50	-	33.53%	0.05[0,0.79]
Vaid 2009	0/67	29/66	-	34.39%	0.02[0,0.27]
Subtotal (95% CI)	177	176		100%	0.04[0.01,0.21]
Total events: 0 (Injectable prostagl	andins), 44 (Misoprosto	ol)			
Heterogeneity: Tau ² =0; Chi ² =0.71, o	df=2(P=0.7); I ² =0%				
Test for overall effect: Z=3.85(P=0)					
42.16.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagl	andins), 0 (Misoprostol)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	177	176		100%	0.04[0.01,0.21]
Total events: 0 (Injectable prostagl	andins), 44 (Misoprosto	ol)			
Heterogeneity: Tau ² =0; Chi ² =0.71, o	df=2(P=0.7); I ² =0%				
Test for overall effect: Z=3.85(P=0)					
Test for subgroup differences: Not	applicable				
	Favours Injectab	le prostaglandins	0.01 0.1 1	10 100 Favours Misoprosto	ıl



Analysis 42.17. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 17 Fever.



Analysis 42.18. Comparison 42 Injectable prostaglandins vs Misoprostol (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Misoprostol	Ris	sk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rand	dom, 95% CI		IV, Random, 95% CI
42.18.1 Vaginal birth						
Supe 2016	7/50	0/50		-	34.7%	15[0.88,255.78]
Vaid 2009	7/67	1/66		-	65.3%	6.9[0.87,54.51]
Subtotal (95% CI)	117	116			100%	9.03[1.7,48]
Total events: 14 (Injectable prostag	landins), 1 (Misoprosto	ol)				
Heterogeneity: Tau ² =0; Chi ² =0.19, d	f=1(P=0.66); I ² =0%					
Test for overall effect: Z=2.58(P=0.0	1)					
42.18.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Injectable prostagla	indins), 0 (Misoprosto)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	е					
Total (95% CI)	117	116			100%	9.03[1.7,48]
Total events: 14 (Injectable prostag	landins), 1 (Misoprosto	ol)				
Heterogeneity: Tau ² =0; Chi ² =0.19, d	f=1(P=0.66); I ² =0%					
Test for overall effect: Z=2.58(P=0.0	1)					
Test for subgroup differences: Not a	pplicable					
	Favours Injectab	le prostaglandins	0.01 0.1	1 10 1	Favours Misoprostol	



Comparison 43. Misoprostol vs Carbetocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	177	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [0.62, 8.64]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [0.62, 8.64]
3 Blood transfusion	1	177	Risk Ratio (IV, Random, 95% CI)	2.97 [0.12, 71.85]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	2.97 [0.12, 71.85]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [1.52, 3.50]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [1.52, 3.50]
7 Additional uterotonics	1	177	Risk Ratio (IV, Random, 95% CI)	3.96 [1.55, 10.07]
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	3.96 [1.55, 10.07]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	0.0 [0.0, 0.0]	
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)		
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	1	177	Risk Ratio (IV, Random, 95% CI)	0.99 [0.47, 2.08]	
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	0.99 [0.47, 2.08]	
12 Vomiting	1	177	Risk Ratio (IV, Random, 95% CI)	1.48 [0.55, 3.99]	
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	1.48 [0.55, 3.99]	
13 Headache	1	177	Risk Ratio (IV, Random, 95% CI)	1.19 [0.71, 1.99]	
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	1.19 [0.71, 1.99]	
14 Abdominal pain	1	177	Risk Ratio (IV, Random, 95% CI)	0.65 [0.52, 0.80]	
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	0.65 [0.52, 0.80]	
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16 Shivering	1	177	Risk Ratio (IV, Random, 95% CI)	3.46 [1.67, 7.17]	
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	3.46 [1.67, 7.17]	
17 Fever	1	177	Risk Ratio (IV, Random, 95% CI)	2.55 [1.41, 4.64]	
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
17.2 Caesarean	1	177	Risk Ratio (IV, Random, 95% CI)	2.55 [1.41, 4.64]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 43.1. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Misoprostol	Carbetocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
43.1.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
43.1.2 Caesarean					
Elbohoty 2016	0/89	0/88			Not estimable
Subtotal (95% CI)	89	88			Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	ocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	89	88			Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	ocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Test for subgroup differences: Not appli	cable				
• • • • • • • • • • • • • • • • • • • •		vours Misoprostol	0.01 0.1 1 1		in



Analysis 43.2. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup N	Iisoprostol	Carbetocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
43.2.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
43.2.2 Caesarean					
Elbohoty 2016	7/89	3/88	- - 	100%	2.31[0.62,8.64]
Subtotal (95% CI)	89	88		100%	2.31[0.62,8.64]
Total events: 7 (Misoprostol), 3 (Carbeto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.24(P=0.21)					
Total (95% CI)	89	88		100%	2.31[0.62,8.64]
Total events: 7 (Misoprostol), 3 (Carbeto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.24(P=0.21)					
Test for subgroup differences: Not applic	able				
	Fa	vours Misoprostol	0.01 0.1 1 10	100 Favours Carbetocin	

Analysis 43.3. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Carbetocin	Carbetocin Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Rando	IV, Random, 95% CI		IV, Random, 95% CI
43.3.1 Vaginal birth						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
43.3.2 Caesarean						
Elbohoty 2016	1/89	0/88			- 100%	2.97[0.12,71.85]
Subtotal (95% CI)	89	88			- 100 %	2.97[0.12,71.85]
·		88			100%	2.91[0.12,11.85]
Total events: 1 (Misoprostol), 0 (Carbeto	icin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=0.67(P=0.5)						
Total (95% CI)	89	88			100%	2.97[0.12,71.85]
Total events: 1 (Misoprostol), 0 (Carbeto	ocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=0.67(P=0.5)						
Test for subgroup differences: Not appli	cable					
	Fa	vours Misoprostol	0.01 0.1	1 10 1	.00 Favours Carbetocin	



Analysis 43.6. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 6 PPH >= 500 mL.

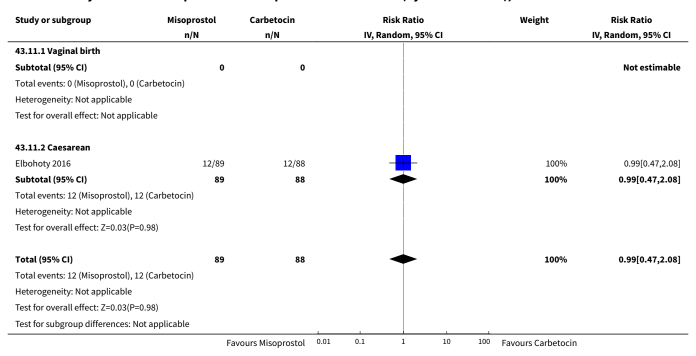
Misoprostol	Carbetocin	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
0	0			Not estimable
arbetocin)				
ble				
49/89	21/88		100%	2.31[1.52,3.5]
89	88	▼	100%	2.31[1.52,3.5]
(Carbetocin)				
.0001)				
89	88	•	100%	2.31[1.52,3.5]
(Carbetocin)				
.0001)				
t applicable				
	n/N 0 arbetocin) arbetocin) 49/89 89 (Carbetocin)	n/N n/N 0 0 arbetocin) bble 49/89 21/88 89 88 (Carbetocin) .0001) 89 88 (Carbetocin)	n/N	n/N

Analysis 43.7. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Carbetocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N IV, Random, 95% CI				IV, Random, 95% CI
43.7.1 Vaginal birth						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
43.7.2 Caesarean						
Elbohoty 2016	20/89	5/88		- 	100%	3.96[1.55,10.07]
Subtotal (95% CI)	89	88		-	100%	3.96[1.55,10.07]
Total events: 20 (Misoprostol), 5 (Carbet	ocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=2.88(P=0)						
Total (95% CI)	89	88		•	100%	3.96[1.55,10.07]
Total events: 20 (Misoprostol), 5 (Carbet	ocin)					. , .
Heterogeneity: Not applicable						
Test for overall effect: Z=2.88(P=0)						
Test for subgroup differences: Not appli	cable					
	Fa	vours Misoprostol	0.01 0.1	1 10 10	DO Favours Carbetocin	



Analysis 43.11. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 11 Nausea.

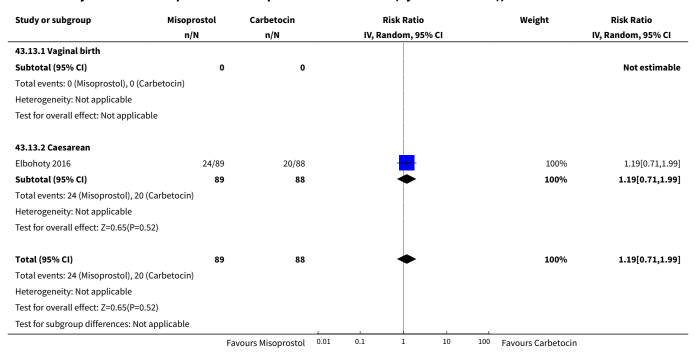


Analysis 43.12. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup N	lisoprostol	Carbetocin			Risk Ratio			Weight	Risk Ratio
	n/N	l n/N		IV, Random, 95% CI				IV, Random, 95% CI	
43.12.1 Vaginal birth									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
43.12.2 Caesarean									
Elbohoty 2016	9/89	6/88						100%	1.48[0.55,3.99]
Subtotal (95% CI)	89	88						100%	1.48[0.55,3.99]
Total events: 9 (Misoprostol), 6 (Carbeto	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.78(P=0.44)									
Total (95% CI)	89	88						100%	1.48[0.55,3.99]
Total events: 9 (Misoprostol), 6 (Carbeto	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.78(P=0.44)									
Test for subgroup differences: Not applic	able								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	



Analysis 43.13. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 13 Headache.



Analysis 43.14. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Carbetocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
43.14.1 Vaginal birth						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
43.14.2 Caesarean						
Elbohoty 2016	47/89	72/88	+		100%	0.65[0.52,0.8]
Subtotal (95% CI)	89	88	♦		100%	0.65[0.52,0.8]
Total events: 47 (Misoprostol), 72 (Carbe	etocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=3.91(P<0.0001)						
Total (050/, CI)	89	88	•		100%	0.65[0.52.0.9]
Total (95% CI)		88	•		100%	0.65[0.52,0.8]
Total events: 47 (Misoprostol), 72 (Carbe	etocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=3.91(P<0.0001)						
Test for subgroup differences: Not applie	cable					
	Fa	vours Misoprostol	0.01 0.1	10 100	Favours Carbetocin	



Analysis 43.16. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 16 Shivering.

Misoprostol	Carbetocin	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
0	0			Not estimable
Carbetocin)				
able				
28/89	8/88	-	100%	3.46[1.67,7.17]
89	88	•	100%	3.46[1.67,7.17]
(Carbetocin)		ĺ		
f=0(P<0.0001); I ² =100%		ĺ		
0)				
89	88	•	100%	3.46[1.67,7.17]
(Carbetocin)				
f=0(P<0.0001); I ² =100%				
0)				
ot applicable				
	n/N 0 Carbetocin) able 28/89 89 (Carbetocin) f=0(P<0.0001); l²=100% 0) 89 (Carbetocin) f=0(P<0.0001); l²=100%	n/N n/N 0 0 Carbetocin) able 28/89 8/88 89 88 (Carbetocin) f=0(P<0.0001); I²=100% 0) 89 88 (Carbetocin) f=0(P<0.0001); I²=100% 0)	n/N n/N IV, Random, 95% CI 0 0 Carbetocin) able 28/89 8/88 89 88 (Carbetocin) f=0(P<0.0001); l²=100% 0) 89 88 (Carbetocin) f=0(P<0.0001); l²=100% 0)	n/N n/N IV, Random, 95% CI 0 0 Carbetocin) able 28/89 8/88 89 88 (Carbetocin) f=0(P<0.0001); l²=100% 0) 89 88 (Carbetocin) f=0(P<0.0001); l²=100% 0)

Analysis 43.17. Comparison 43 Misoprostol vs Carbetocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol	Carbetocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N IV, Random, 95% CI				IV, Random, 95% CI
43.17.1 Vaginal birth						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	cin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
43.17.2 Caesarean						
Elbohoty 2016	31/89	12/88		-	100%	2.55[1.41,4.64]
Subtotal (95% CI)	89	88		•	100%	2.55[1.41,4.64]
Total events: 31 (Misoprostol), 12 (Carbo	etocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=3.08(P=0)						
T-1-1 (050/ CI)	89	88			100%	2 55(1 41 4 64)
Total (95% CI)		88			100%	2.55[1.41,4.64]
Total events: 31 (Misoprostol), 12 (Carbe	etocin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=3.08(P=0)						
Test for subgroup differences: Not appli	cable				1	
	Fa	vours Misoprostol	0.01 0.1	1 10	100 Favours Carbetocin	



Comparison 44. Ergometrine vs Misoprostol (by mode of birth)

Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	7	5254	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	7	5254	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	11	5562	Risk Ratio (IV, Random, 95% CI)	1.25 [0.45, 3.48]
2.1 Vaginal birth	11	5562	Risk Ratio (IV, Random, 95% CI)	1.25 [0.45, 3.48]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	13	5308	Risk Ratio (IV, Random, 95% CI)	1.71 [0.65, 4.50]
3.1 Vaginal birth	13	5308	Risk Ratio (IV, Random, 95% CI)	1.71 [0.65, 4.50]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	1	864	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.16]
4.1 Vaginal birth	1	864	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.16]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	16	6459	Risk Ratio (IV, Random, 95% CI)	1.18 [0.79, 1.75]
6.1 Vaginal birth	16	6459	Risk Ratio (IV, Random, 95% CI)	1.18 [0.79, 1.75]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	16	6565	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.60]
7.1 Vaginal birth	16	6565	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.60]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	15	6337	Mean Difference (IV, Random, 95% CI)	11.66 [-9.85, 33.17]
8.1 Vaginal birth	15	6337	Mean Difference (IV, Random, 95% CI)	11.66 [-9.85, 33.17]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
9 Change in haemo- globin	8	4438	Mean Difference (IV, Random, 95% CI)	0.91 [-0.43, 2.26]
9.1 Vaginal birth	8	4438	Mean Difference (IV, Random, 95% CI)	0.91 [-0.43, 2.26]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	13	5202	Risk Ratio (IV, Random, 95% CI)	1.53 [1.15, 2.04]
11.1 Vaginal birth	13	5202	Risk Ratio (IV, Random, 95% CI)	1.53 [1.15, 2.04]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	14	6136	Risk Ratio (IV, Random, 95% CI)	1.22 [0.81, 1.83]
12.1 Vaginal birth	14	6136	Risk Ratio (IV, Random, 95% CI)	1.22 [0.81, 1.83]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	8	3369	Risk Ratio (IV, Random, 95% CI)	1.70 [0.70, 4.12]
13.1 Vaginal birth	8	3369	Risk Ratio (IV, Random, 95% CI)	1.70 [0.70, 4.12]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	1.58 [0.04, 65.80]
14.1 Vaginal birth	2	233	Risk Ratio (IV, Random, 95% CI)	1.58 [0.04, 65.80]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	2	300	Risk Ratio (IV, Random, 95% CI)	7.55 [0.94, 60.53]
15.1 Vaginal birth	2	300	Risk Ratio (IV, Random, 95% CI)	7.55 [0.94, 60.53]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	17	6860	Risk Ratio (IV, Random, 95% CI)	0.34 [0.26, 0.44]
16.1 Vaginal birth	17	6860	Risk Ratio (IV, Random, 95% CI)	0.34 [0.26, 0.44]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	14	6330	Risk Ratio (IV, Random, 95% CI)	0.24 [0.17, 0.33]
17.1 Vaginal birth	14	6330	Risk Ratio (IV, Random, 95% CI)	0.24 [0.17, 0.33]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



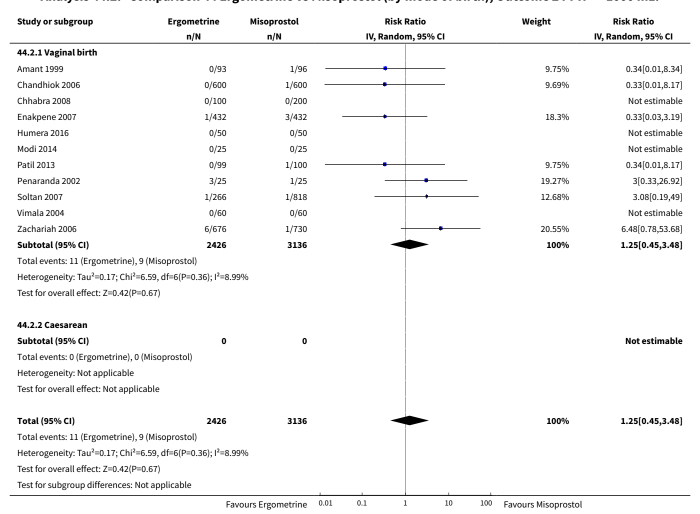
Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
18 Diarrhoea	8	4165	Risk Ratio (IV, Random, 95% CI)	1.04 [0.46, 2.34]
18.1 Vaginal birth	8	4165	Risk Ratio (IV, Random, 95% CI)	1.04 [0.46, 2.34]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 44.1. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 1 Death.

Study or subgroup E	rgometrine	Misoprostol	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
44.1.1 Vaginal birth						
Chandhiok 2006	0/600	0/600				Not estimable
Chhabra 2008	0/100	0/200				Not estimable
Enakpene 2007	0/432	0/432				Not estimable
Fawzy 2012	0/100	0/200				Not estimable
Soltan 2007	0/266	0/818				Not estimable
Supe 2016	0/50	0/50				Not estimable
Zachariah 2006	0/676	0/730				Not estimable
Subtotal (95% CI)	2224	3030				Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
44.1.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	2224	3030				Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not applie	cable					
	Fav	vours Ergometrine	0.01 0.1	10 100	Favours Misoprostol	



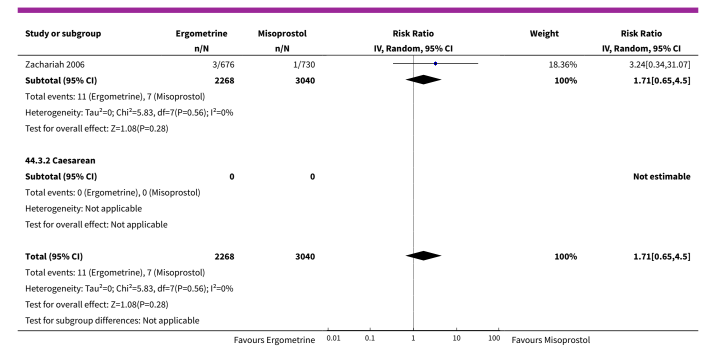
Analysis 44.2. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 2 PPH >= 1000 mL.



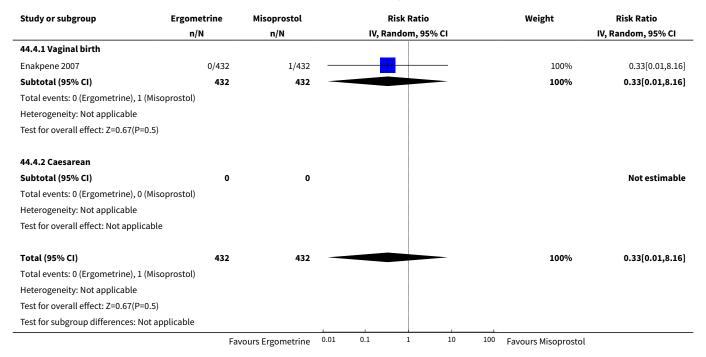
Analysis 44.3. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Misoprostol		Risk	Ratio			Weight	Risk Ratio
	n/N	n/N	IV,	Rando	m, 95% CI				IV, Random, 95% CI
44.3.1 Vaginal birth									
Amant 1999	1/100	1/100						12.34%	1[0.06,15.77]
Chandhiok 2006	0/600	1/600						9.17%	0.33[0.01,8.17]
Chhabra 2008	0/100	0/200							Not estimable
Humera 2016	0/50	0/50							Not estimable
Modi 2014	0/25	0/25							Not estimable
Patil 2013	0/99	1/100						9.23%	0.34[0.01,8.17]
Ray 2001	3/100	1/100			+			18.6%	3[0.32,28.35]
Singh 2009	3/75	0/150		_		+	\rightarrow	10.78%	13.91[0.73,265.81]
Soltan 2007	1/266	1/809	-		 		_	12.25%	3.04[0.19,48.46]
Supe 2016	0/50	0/50							Not estimable
Vaid 2009	0/67	1/66		•				9.26%	0.33[0.01,7.92]
Vimala 2004	0/60	0/60							Not estimable
	Fav	ours Ergometrine	0.01 0.1	1	1 1	10	100	Favours Misoprostol	



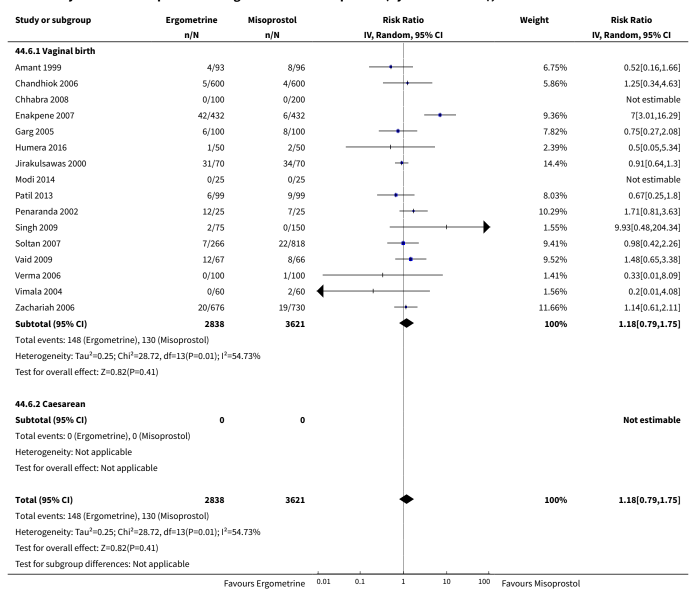


Analysis 44.4. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.





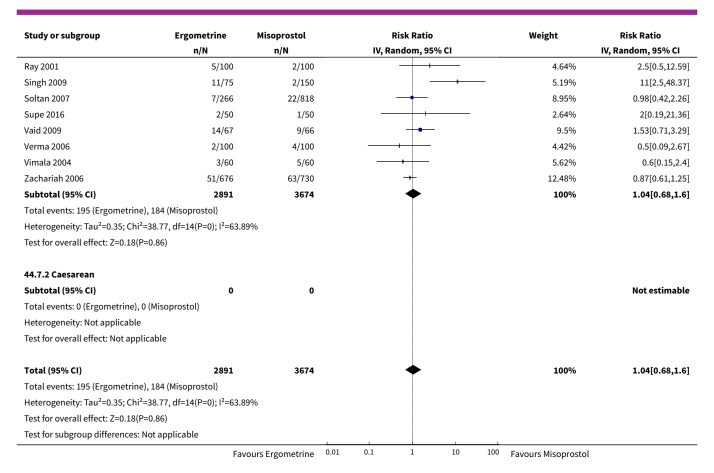
Analysis 44.6. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 6 PPH >= 500 mL.



Analysis 44.7. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Ergometrine	Misoprostol	oprostol Risk Ratio n/N IV, Random, 95% CI			Weight	Risk Ratio	
	n/N	n/N					IV, Random, 95% CI	
44.7.1 Vaginal birth								
Amant 1999	4/91	12/94		-+-			7.22%	0.34[0.12,1.03]
Chandhiok 2006	3/600	4/600					5.14%	0.75[0.17,3.34]
Chhabra 2008	3/100	9/200					6.13%	0.67[0.18,2.41]
Enakpene 2007	80/432	33/432		+-			12.3%	2.42[1.65,3.56]
Garg 2005	7/100	10/100		-+ -			8.34%	0.7[0.28,1.77]
Humera 2016	1/50	2/50					2.64%	0.5[0.05,5.34]
Modi 2014	0/25	0/25						Not estimable
Patil 2013	2/99	6/99					4.79%	0.33[0.07,1.61]
	Fa	vours Ergometrine	0.01	0.1 1	10	100	Favours Misoprostol	

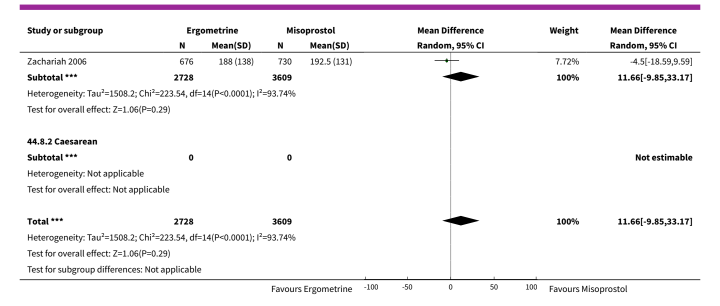




Analysis 44.8. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	Mis	soprostol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
44.8.1 Vaginal birth							
Chandhiok 2006	600	211 (83.4)	600	139.7 (100.4)		7.84%	71.3[60.86,81.74]
Chhabra 2008	100	150 (52)	200	150 (50)	+	7.78%	0[-12.32,12.32]
Enakpene 2007	432	246 (175.5)	432	191.6 (134.5)	 	7.43%	54.4[33.55,75.25]
Fawzy 2012	100	275.8 (165.5)	200	233.5 (132.9)		6.44%	42.22[4.92,79.52]
Humera 2016	50	172.8 (79.7)	50	195.1 (94.3)		6.64%	-22.3[-56.5,11.9]
Jirakulsawas 2000	70	484.7 (120.1)	70	490.5 (109.8)		6.38%	-5.79[-43.91,32.33]
Modi 2014	25	131 (72)	25	255.8 (102.2)	←	5.65%	-124.8[-173.8,-75.8]
Patil 2013	99	178 (137)	99	211 (172)	+++	6.03%	-33[-76.32,10.32]
Penaranda 2002	25	546.8 (338.5)	25	389 (271)	_	1.33%	157.8[-12.17,327.77]
Singh 2009	75	223.5 (162)	150	111.2 (70.4)	→	6.37%	112.33[73.99,150.67]
Soltan 2007	266	149.3 (104)	818	134 (89.8)	├	7.73%	15.28[1.35,29.21]
Supe 2016	50	152.2 (49.3)	50	124.4 (34.7)	_ 	7.62%	27.8[11.09,44.51]
Verma 2006	100	125.8 (72.8)	100	137.6 (72.8)		7.46%	-11.78[-31.96,8.4]
Vimala 2004	60	170 (42)	60	185 (56)	· · · · · · · · · · · · · · · · · · ·	7.58%	-15[-32.71,2.71]
			Favour	s Ergometrine	-100 -50 0 50 100	Favours Mis	oprostol



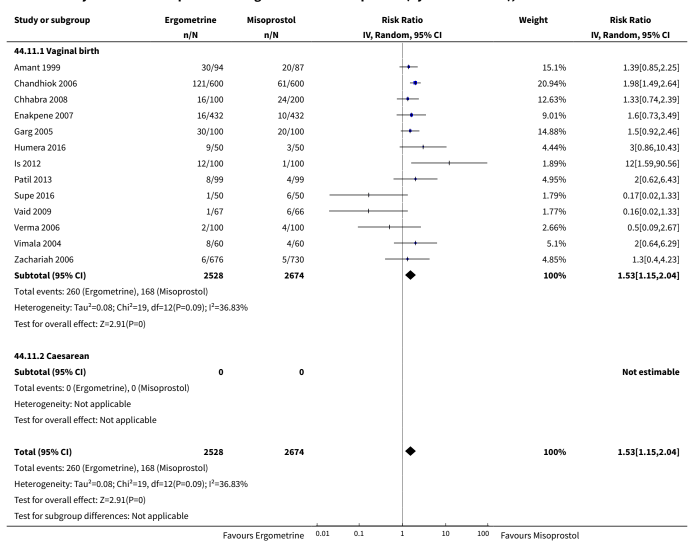


Analysis 44.9. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Erg	ometrine	Mis	oprostol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
44.9.1 Vaginal birth							
Chhabra 2008	100	8 (6.3)	200	7.8 (2)	<u> </u>	11.87%	0.2[-1.07,1.47]
Enakpene 2007	432	4.1 (5.1)	432	1.4 (2)	•	13%	2.72[2.2,3.24]
Gore 2017	182	9.4 (0.7)	182	9.7 (0.8)	+	13.24%	-0.3[-0.45,-0.15]
Soltan 2007	266	4.6 (4)	818	3.6 (2.9)	•	13%	1[0.48,1.52]
Supe 2016	50	7.6 (0.2)	50	4.9 (0.5)	•	13.24%	2.7[2.55,2.85]
Verma 2006	100	2.5 (3)	100	3.1 (3)	•	12.62%	-0.6[-1.43,0.23]
Vimala 2004	60	8 (6.3)	60	7.6 (2)	 	11.01%	0.4[-1.27,2.07]
Zachariah 2006	676	14.8 (11.9)	730	13.8 (10.7)	•	12.02%	1[-0.19,2.19]
Subtotal ***	1866		2572		•	100%	0.91[-0.43,2.26]
Heterogeneity: Tau ² =3.57; Chi ² =81	6.27, df=7(P<0.0001); I ² =99.	14%				
Test for overall effect: Z=1.33(P=0.3	L8)						
44.9.2 Caesarean							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	ole						
Total ***	1866		2572)	100%	0.91[-0.43,2.26]
Heterogeneity: Tau ² =3.57; Chi ² =81	6.27, df=7(P<0.0001); I ² =99.	14%				
Test for overall effect: Z=1.33(P=0.3	L8)						
Test for subgroup differences: Not	applicable						
			Favours	Ergometrine -100	-50 0 50	100 Favours Mis	soprostol



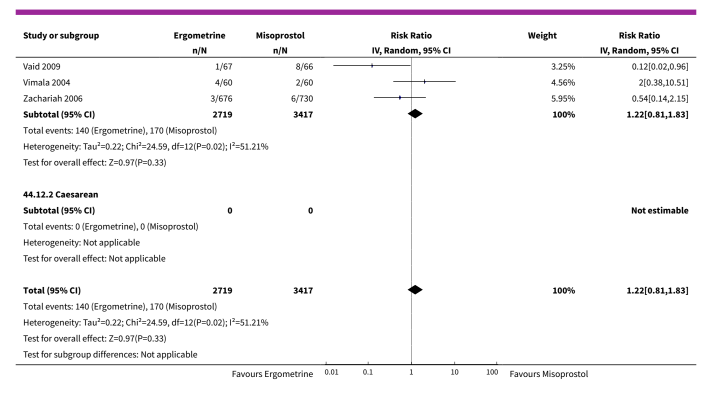
Analysis 44.11. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 11 Nausea.



Analysis 44.12. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
44.12.1 Vaginal birth					
Amant 1999	18/94	13/87	-	12.83%	1.28[0.67,2.46]
Chandhiok 2006	34/600	30/600	+	15.13%	1.13[0.7,1.83]
Chhabra 2008	8/100	12/200		10.28%	1.33[0.56,3.16]
Enakpene 2007	12/432	1/432		3.29%	12[1.57,91.89]
Garg 2005	30/100	19/100	+	14.79%	1.58[0.95,2.61]
Humera 2016	4/50	2/50		4.59%	2[0.38,10.43]
Is 2012	10/100	2/100		5.34%	5[1.12,22.24]
Patil 2013	3/99	3/99		4.93%	1[0.21,4.83]
Penaranda 2002	0/25	0/25			Not estimable
Soltan 2007	12/266	72/818		13.56%	0.51[0.28,0.93]
Supe 2016	1/50	0/50		1.5%	3[0.13,71.92]
	Far	ours Ergometrine	0.01 0.1 1 10	100 Favours Misoprostol	





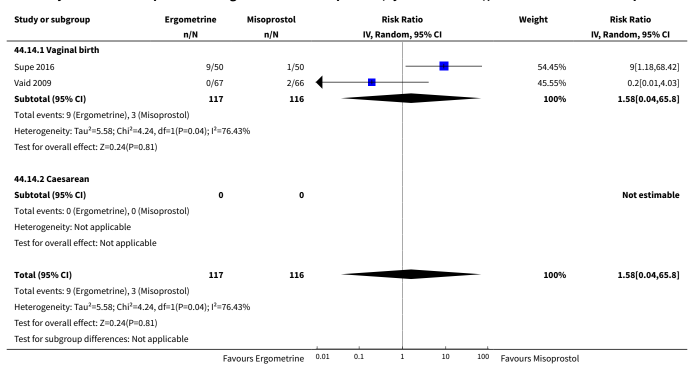
Analysis 44.13. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 13 Headache.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
44.13.1 Vaginal birth					
Amant 1999	12/94	10/87	-	22.73%	1.11[0.51,2.44]
Chhabra 2008	8/100	14/200	-	22.23%	1.14[0.5,2.63]
Enakpene 2007	54/432	1/432	──	11.67%	54[7.5,388.59]
Garg 2005	3/100	4/100		15.71%	0.75[0.17,3.27]
Humera 2016	0/50	0/50			Not estimable
Patil 2013	0/99	0/99			Not estimable
Vimala 2004	4/60	3/60		15.87%	1.33[0.31,5.7]
Zachariah 2006	2/676	2/730		11.78%	1.08[0.15,7.64]
Subtotal (95% CI)	1611	1758	•	100%	1.7[0.7,4.12]
Total events: 83 (Ergometrine), 34 (M	isoprostol)				
Heterogeneity: Tau ² =0.74; Chi ² =14.47	7, df=5(P=0.01); l ² =65	5.45%			
Test for overall effect: Z=1.17(P=0.24))				
44.13.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Misc	_	Ū			Not estimable
Heterogeneity: Not applicable	oprostot)				
Test for overall effect: Not applicable	!				
Total (95% CI)	1611	1758		100%	1.7[0.7,4.12]
Total events: 83 (Ergometrine), 34 (M	isoprostol)				
Heterogeneity: Tau ² =0.74; Chi ² =14.47	7, df=5(P=0.01); l ² =65	5.45%			
Test for overall effect: Z=1.17(P=0.24))				
	Fav	ours Ergometrine 0	0.01 0.1 1 10 100	Favours Misoprosto	1



Study or subgroup	Ergometrine n/N	Misoprostol n/N			Risk Ratio			Weight	Risk Ratio IV, Random, 95% CI
Test for subgroup differences: Not applicable				1		1			
		Favours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	

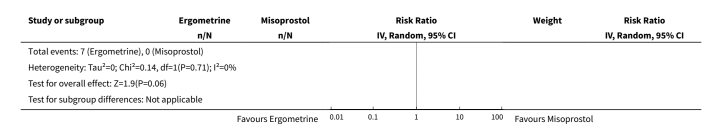
Analysis 44.14. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 14 Abdominal pain.



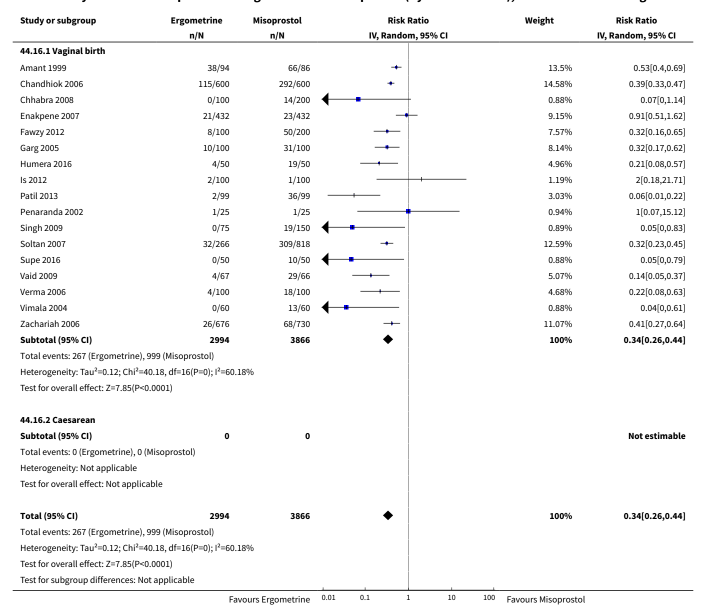
Analysis 44.15. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Misoprostol		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI
44.15.1 Vaginal birth								
Humera 2016	2/50	0/50			-		47.8%	5[0.25,101.58]
Ray 2001	5/100	0/100			+		52.2%	11[0.62,196.33]
Subtotal (95% CI)	150	150					100%	7.55[0.94,60.53]
Total events: 7 (Ergometrine), 0 (M	Misoprostol)							
Heterogeneity: Tau ² =0; Chi ² =0.14,	df=1(P=0.71); I ² =0%							
Test for overall effect: Z=1.9(P=0.0	96)							
44.15.2 Caesarean								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine), 0 (M	Misoprostol)							
Heterogeneity: Not applicable								
Test for overall effect: Not applica	ble							
Total (95% CI)	150	150				_	100%	7.55[0.94,60.53]
	Fav	ours Ergometrine	0.01	0.1	1 10	100	Favours Misoprostol	





Analysis 44.16. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 16 Shivering.





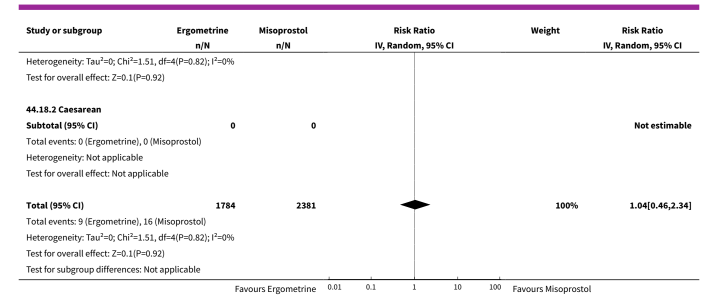
Analysis 44.17. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 17 Fever.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
44.17.1 Vaginal birth					
Amant 1999	3/100	34/100	+	6.89%	0.09[0.03,0.28]
Chandhiok 2006	26/600	58/600		26.22%	0.45[0.29,0.7]
Chhabra 2008	0/100	10/200	+	1.27%	0.09[0.01,1.6]
Enakpene 2007	7/432	31/432		12.25%	0.23[0.1,0.51]
Garg 2005	7/100	29/100		13.02%	0.24[0.11,0.53]
Humera 2016	0/50	2/50	—	1.13%	0.2[0.01,4.06]
Patil 2013	0/99	2/99	—	1.12%	0.2[0.01,4.11]
Singh 2009	0/75	25/150	— ——	1.31%	0.04[0,0.63]
Soltan 2007	6/266	95/818		12.16%	0.19[0.09,0.44]
Supe 2016	0/50	6/50	+	1.25%	0.08[0,1.33]
Vaid 2009	0/67	13/66	——	1.3%	0.04[0,0.6]
Verma 2006	1/100	6/100		2.27%	0.17[0.02,1.36]
Vimala 2004	0/60	4/60		1.21%	0.11[0.01,2.02]
Zachariah 2006	13/676	48/730		18.59%	0.29[0.16,0.53]
Subtotal (95% CI)	2775	3555	•	100%	0.24[0.17,0.33]
Total events: 63 (Ergometrine),	, 363 (Misoprostol)				
Heterogeneity: Tau ² =0.05; Chi ²	=15.24, df=13(P=0.29); I ² =1	14.72%			
Test for overall effect: Z=8.64(P	<0.0001)				
44.17.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0	(Misoprostol)				
Heterogeneity: Not applicable					
Test for overall effect: Not appl	icable				
Total (95% CI)	2775	3555	•	100%	0.24[0.17,0.33]
Total events: 63 (Ergometrine),	, 363 (Misoprostol)				
Heterogeneity: Tau ² =0.05; Chi ²	=15.24, df=13(P=0.29); I ² =1	14.72%			
Test for overall effect: Z=8.64(P	<0.0001)				
Test for subgroup differences: I	Not applicable				

Analysis 44.18. Comparison 44 Ergometrine vs Misoprostol (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine	Misoprostol			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
44.18.1 Vaginal birth									
Amant 1999	0/94	1/86		•				6.44%	0.31[0.01,7.39]
Enakpene 2007	0/432	0/432							Not estimable
Garg 2005	3/100	3/100		_	+	_		26.34%	1[0.21,4.84]
Patil 2013	0/99	0/99							Not estimable
Soltan 2007	4/266	10/818			-	-		49.38%	1.23[0.39,3.89]
Supe 2016	0/50	0/50							Not estimable
Vaid 2009	0/67	1/66	-					6.46%	0.33[0.01,7.92]
Zachariah 2006	2/676	1/730		_	+			11.38%	2.16[0.2,23.76]
Subtotal (95% CI)	1784	2381			*			100%	1.04[0.46,2.34]
Total events: 9 (Ergometrine),	16 (Misoprostol)								
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	





Comparison 45. Misoprostol vs Ergometrine plus oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3258	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	4	3258	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	9	6028	Risk Ratio (IV, Random, 95% CI)	1.61 [1.08, 2.40]
2.1 Vaginal birth	9	6028	Risk Ratio (IV, Random, 95% CI)	1.61 [1.08, 2.40]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	8	6335	Risk Ratio (IV, Random, 95% CI)	1.41 [0.91, 2.21]
3.1 Vaginal birth	8	6335	Risk Ratio (IV, Random, 95% CI)	1.41 [0.91, 2.21]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	10	6492	Risk Ratio (IV, Random, 95% CI)	1.75 [1.35, 2.26]
6.1 Vaginal birth	10	6492	Risk Ratio (IV, Random, 95% CI)	1.75 [1.35, 2.26]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	9	6395	Risk Ratio (IV, Random, 95% CI)	1.87 [1.49, 2.36]
7.1 Vaginal birth	9	6395	Risk Ratio (IV, Random, 95% CI)	1.87 [1.49, 2.36]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	8	5634	Mean Difference (IV, Random, 95% CI)	22.86 [4.50, 41.22]
8.1 Vaginal birth	8	5634	Mean Difference (IV, Random, 95% CI)	22.86 [4.50, 41.22]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	9	5588	Mean Difference (IV, Random, 95% CI)	1.09 [-0.49, 2.67]
9.1 Vaginal birth	9	5588	Mean Difference (IV, Random, 95% CI)	1.09 [-0.49, 2.67]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3399	Risk Ratio (IV, Random, 95% CI)	0.71 [0.61, 0.84]
11.1 Vaginal birth	4	3399	Risk Ratio (IV, Random, 95% CI)	0.71 [0.61, 0.84]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	6	4983	Risk Ratio (IV, Random, 95% CI)	0.72 [0.50, 1.04]
12.1 Vaginal birth	6	4983	Risk Ratio (IV, Random, 95% CI)	0.72 [0.50, 1.04]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	3	3259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.47, 1.76]
13.1 Vaginal birth	3	3259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.47, 1.76]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	1	846	Risk Ratio (IV, Random, 95% CI)	0.90 [0.79, 1.02]
14.1 Vaginal birth	1	846	Risk Ratio (IV, Random, 95% CI)	0.90 [0.79, 1.02]

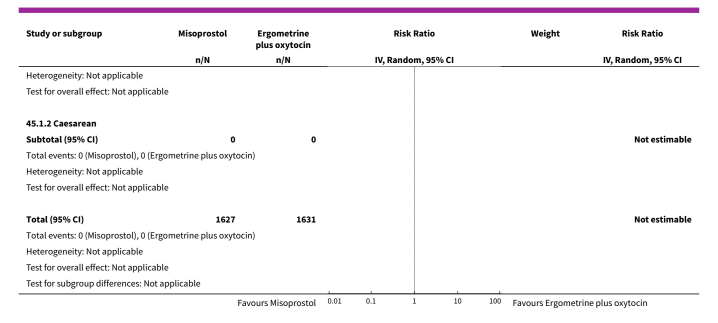


Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	2553	Risk Ratio (IV, Random, 95% CI)	0.50 [0.18, 1.41]
15.1 Vaginal birth	3	2553	Risk Ratio (IV, Random, 95% CI)	0.50 [0.18, 1.41]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	6	4980	Risk Ratio (IV, Random, 95% CI)	2.71 [1.95, 3.76]
16.1 Vaginal birth	6	4980	Risk Ratio (IV, Random, 95% CI)	2.71 [1.95, 3.76]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	7	5045	Risk Ratio (IV, Random, 95% CI)	5.33 [2.87, 9.88]
17.1 Vaginal birth	7	5045	Risk Ratio (IV, Random, 95% CI)	5.33 [2.87, 9.88]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	6	3659	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.57]
18.1 Vaginal birth	6	3659	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.57]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
20.1 Vaginal birth	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]

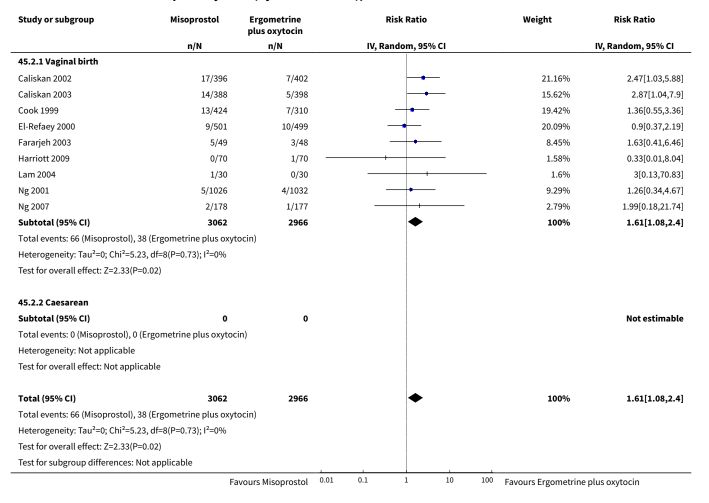
Analysis 45.1. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
45.1.1 Vaginal birth									
El-Refaey 2000	0/501	0/499							Not estimable
Harriott 2009	0/70	0/70							Not estimable
Lam 2004	0/30	0/30							Not estimable
Ng 2001	0/1026	0/1032							Not estimable
Subtotal (95% CI)	1627	1631							Not estimable
Total events: 0 (Misoprostol),	0 (Ergometrine plus oxytoci	n)							
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Ergometrine	plus oxytocin



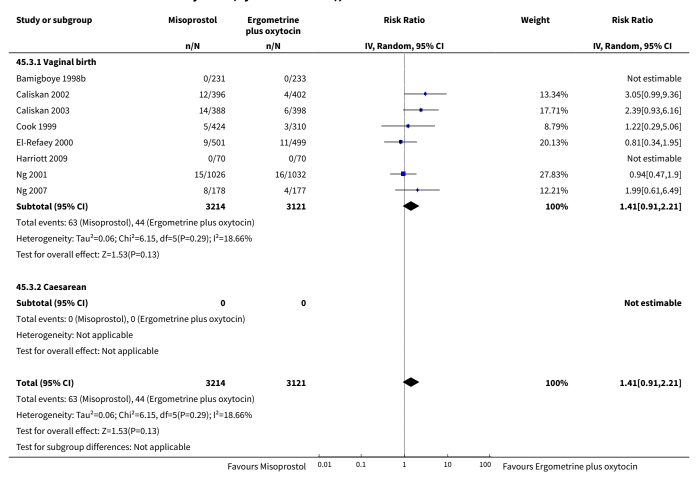


Analysis 45.2. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.





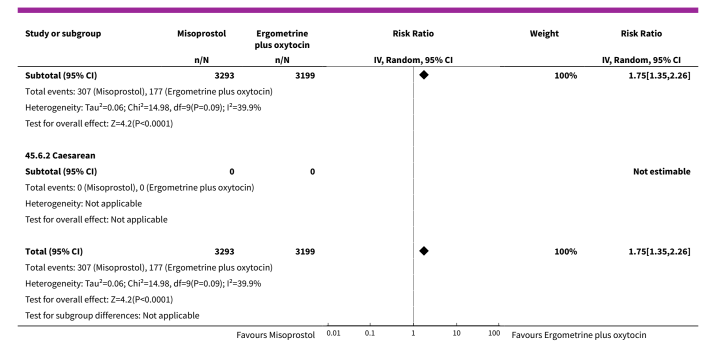
Analysis 45.3. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 3 Blood transfusion.



Analysis 45.6. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
45.6.1 Vaginal birth					
Bamigboye 1998b	2/231	1/233		1.13%	2.02[0.18,22.09]
Caliskan 2002	39/396	14/402		11.53%	2.83[1.56,5.13]
Caliskan 2003	35/388	14/398		11.32%	2.56[1.4,4.69]
Cook 1999	63/424	23/310		15.39%	2[1.27,3.15]
El-Refaey 2000	62/501	56/499	+	19.44%	1.1[0.79,1.55]
Fararjeh 2003	23/49	11/48		11.45%	2.05[1.13,3.73]
Harriott 2009	1/70	3/70		1.29%	0.33[0.04,3.13]
Lam 2004	4/30	2/30	- +	2.36%	2[0.4,10.11]
Ng 2001	60/1026	44/1032	 • -	17.96%	1.37[0.94,2]
Ng 2007	18/178	9/177		8.14%	1.99[0.92,4.31]





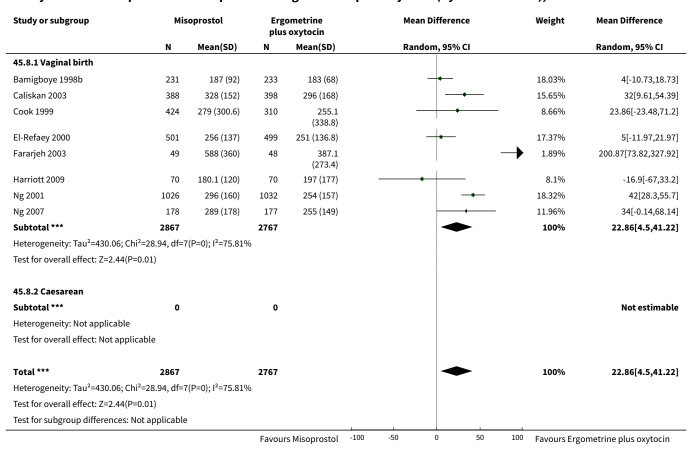
Analysis 45.7. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
45.7.1 Vaginal birth					
Bamigboye 1998b	4/231	1/233		1.09%	4.03[0.45,35.83]
Caliskan 2002	33/396	9/402		7.89%	3.72[1.8,7.68]
Caliskan 2003	23/388	9/398		7.35%	2.62[1.23,5.59]
Cook 1999	95/424	28/310	-	17.13%	2.48[1.67,3.68]
El-Refaey 2000	68/501	50/499	+-	19.5%	1.35[0.96,1.91]
Harriott 2009	6/70	6/70		4.05%	1[0.34,2.95]
Lam 2004	3/30	0/30	+	0.62%	7[0.38,129.93]
Ng 2001	232/1026	144/1032	-	27.76%	1.62[1.34,1.96]
Ng 2007	41/178	24/177	-	14.6%	1.7[1.07,2.69]
Subtotal (95% CI)	3244	3151	•	100%	1.87[1.49,2.36]
Total events: 505 (Misoprostol), 273	1 (Ergometrine plus ox	ytocin)			
Heterogeneity: Tau ² =0.04; Chi ² =13.	5, df=8(P=0.1); I ² =40.7	4%			
Test for overall effect: Z=5.31(P<0.0	0001)				
45.7.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Erg	gometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	3244	3151	•	100%	1.87[1.49,2.36]
Total events: 505 (Misoprostol), 273	1 (Ergometrine plus ox	ytocin)			
Heterogeneity: Tau ² =0.04; Chi ² =13.	5, df=8(P=0.1); I ² =40.7	4%			
Test for overall effect: Z=5.31(P<0.0	0001)				
	Fa	vours Misoprostol 0.01	0.1 1 10 1	00 Favours Ergometrin	e plus oxytocin



Study or subgroup	y or subgroup Misoprostol			Risk Ratio				Weight Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI		IV, Random, 95% CI
Test for subgroup differences	s: Not applicable		_					
		Favours Misoprostol	0.01	0.1	1	10	100	Favours Ergometrine plus oxytocin

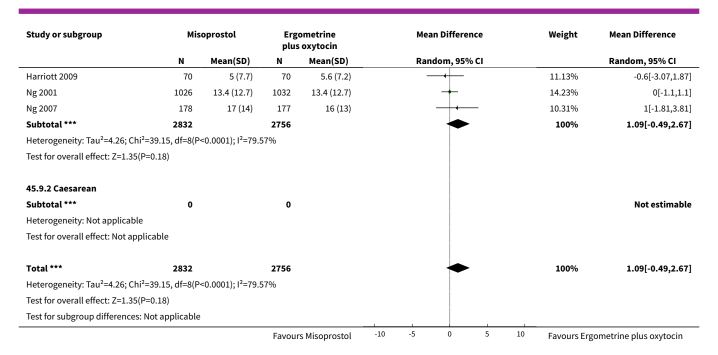
Analysis 45.8. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 8 Blood loss.



Analysis 45.9. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol	U	ometrine oxytocin		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI		Random, 95% CI
45.9.1 Vaginal birth								
Bamigboye 1998b	83	2.3 (16)	83	2.8 (19)		+	5.57%	-0.5[-5.84,4.84]
Caliskan 2002	396	15 (12)	402	15 (12)		-	13.07%	0[-1.67,1.67]
Caliskan 2003	388	14 (11)	398	15 (12)		-+	13.19%	-1[-2.61,0.61]
Cook 1999	424	6.9 (17.5)	310	0.5 (14.1)		_ 	11.58%	6.4[4.11,8.69]
El-Refaey 2000	218	10 (12)	236	10 (16)			10.84%	0[-2.59,2.59]
Fararjeh 2003	49	17.5 (7.3)	48	12.8 (7.3)			10.08%	4.7[1.79,7.61]
			Favour	s Misoprostol	-10	-5 0 5 10	Favours Erg	ometrine plus oxvtocin



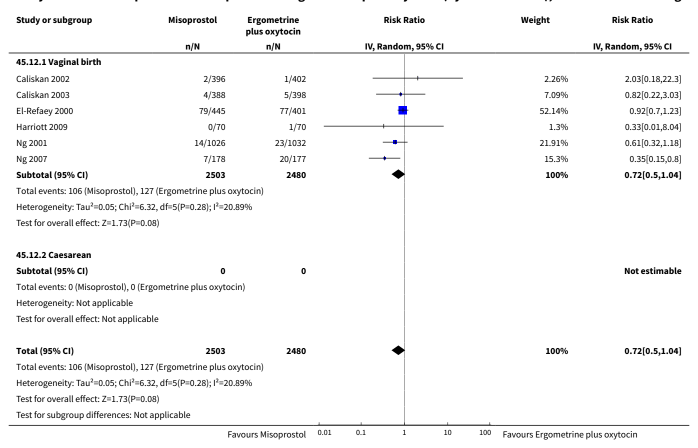


Analysis 45.11. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
45.11.1 Vaginal birth					
El-Refaey 2000	138/445	175/401	+	85.88%	0.71[0.59,0.85]
Harriott 2009	0/70	3/70		0.31%	0.14[0.01,2.72]
Ng 2001	20/1026	27/1032	-+-	8.3%	0.75[0.42,1.32]
Ng 2007	13/178	16/177		5.51%	0.81[0.4,1.63]
Subtotal (95% CI)	1719	1680	◆	100%	0.71[0.61,0.84]
Total events: 171 (Misoprostol), 221	(Ergometrine plus ox	kytocin)			
Heterogeneity: Tau ² =0; Chi ² =1.29, c	f=3(P=0.73); I ² =0%				
Test for overall effect: Z=4(P<0.000	1)				
45.11.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Erg	gometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	1719	1680	•	100%	0.71[0.61,0.84]
Total events: 171 (Misoprostol), 221	(Ergometrine plus ox	(ytocin)			
Heterogeneity: Tau ² =0; Chi ² =1.29, c	df=3(P=0.73); I ² =0%		ĺ		
Test for overall effect: Z=4(P<0.000)	1)				
Test for subgroup differences: Not a	applicable				
	Fa	vours Misoprostol 0.	01 0.1 1 10	100 Favours Ergometrin	e plus oxytocin



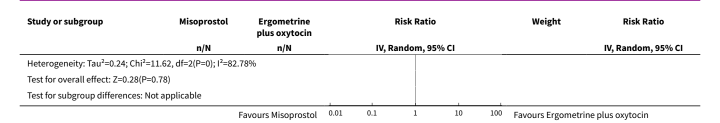
Analysis 45.12. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 12 Vomiting.



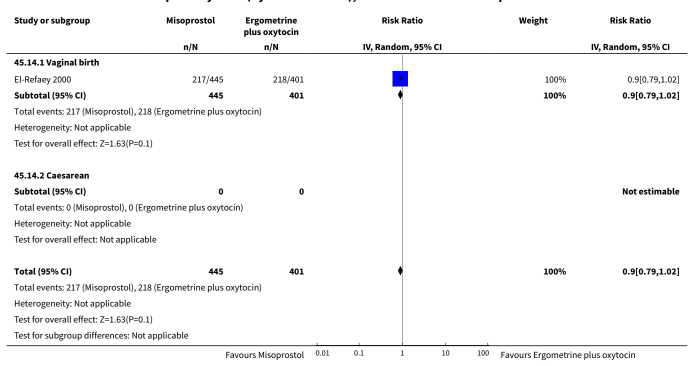
Analysis 45.13. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 13 Headache.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IN.	, Random, 95% CI			IV, Random, 95% CI
45.13.1 Vaginal birth							
El-Refaey 2000	46/445	78/401		-		42.71%	0.53[0.38,0.75]
Ng 2001	81/1026	83/1032		-		43.92%	0.98[0.73,1.32]
Ng 2007	8/178	2/177		+		13.37%	3.98[0.86,18.47]
Subtotal (95% CI)	1649	1610		•		100%	0.91[0.47,1.76]
Total events: 135 (Misoprostol), 163 (Ergometrine plus ox	ytocin)					
Heterogeneity: Tau ² =0.24; Chi ²	² =11.62, df=2(P=0); l ² =82.78	3%					
Test for overall effect: Z=0.28(F	P=0.78)						
45.13.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Misoprostol), 0	(Ergometrine plus oxytoc	in)					
Heterogeneity: Not applicable							
Test for overall effect: Not app	licable						
Total (95% CI)	1649	1610		•		100%	0.91[0.47,1.76]
Total events: 135 (Misoprostol), 163 (Ergometrine plus ox	xytocin)					
	Fa	vours Misoprostol	0.01 0.1	1 10	100	Favours Ergometrin	e plus oxytocin

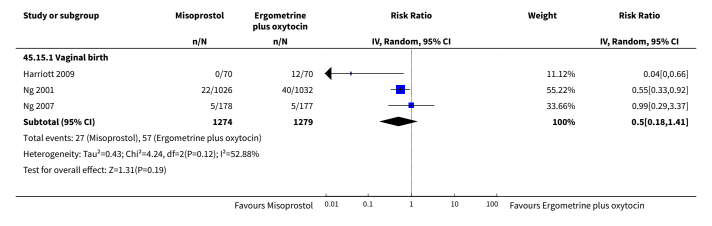




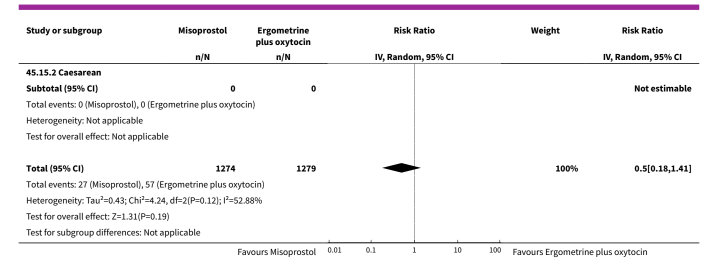
Analysis 45.14. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 14 Abdominal pain.



Analysis 45.15. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 15 Hypertension.





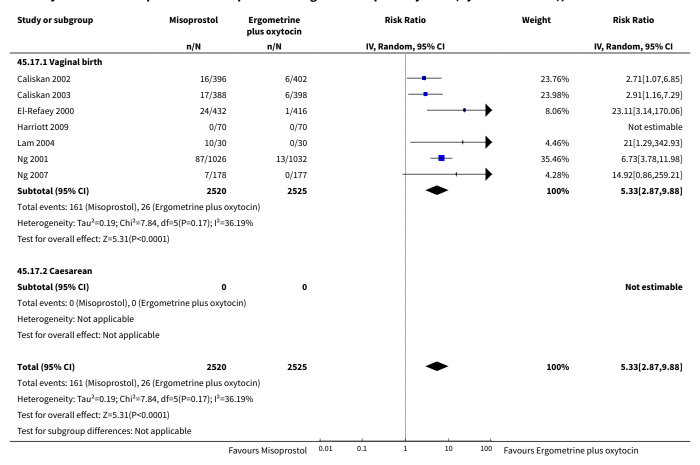


Analysis 45.16. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
45.16.1 Vaginal birth					
Caliskan 2002	47/396	19/402		17.06%	2.51[1.5,4.2]
Caliskan 2003	44/388	15/398		15.6%	3.01[1.7,5.32]
El-Refaey 2000	319/445	147/401		27.81%	1.96[1.7,2.25]
Harriott 2009	11/67	6/70	 • 	8.67%	1.92[0.75,4.89]
Ng 2001	310/1026	102/1032	-	26.28%	3.06[2.49,3.76]
Ng 2007	35/178	2/177		4.58%	17.4[4.25,71.25]
Subtotal (95% CI)	2500	2480	•	100%	2.71[1.95,3.76]
Total events: 766 (Misoprostol), 29	1 (Ergometrine plus o	kytocin)			
Heterogeneity: Tau ² =0.1; Chi ² =21.3	39, df=5(P=0); I ² =76.62	%			
Test for overall effect: Z=5.94(P<0.0	0001)				
45.16.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Erg	gometrine plus oxytoo	in)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	ole				
Total (95% CI)	2500	2480	•	100%	2.71[1.95,3.76]
Total events: 766 (Misoprostol), 29	1 (Ergometrine plus o	xytocin)			
Heterogeneity: Tau ² =0.1; Chi ² =21.3	39, df=5(P=0); I ² =76.62	%			
Test for overall effect: Z=5.94(P<0.0	0001)				
Test for subgroup differences: Not	applicable				
	Fa	avours Misoprostol 0.01	0.1 1 10 1	00 Favours Ergometrin	e plus oxytocin



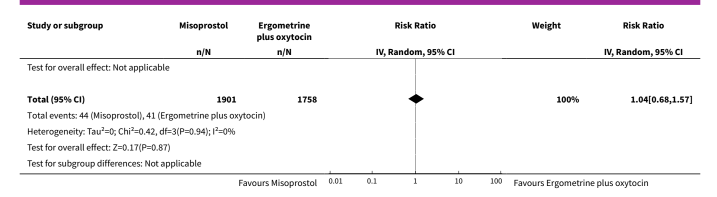
Analysis 45.17. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 17 Fever.



Analysis 45.18. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	ľ	V, Random, 95% CI		IV, Random, 95% CI
45.18.1 Vaginal birth						
Caliskan 2002	11/396	10/402			24.42%	1.12[0.48,2.6]
Caliskan 2003	15/388	17/398		-	37.7%	0.91[0.46,1.79]
Cook 1999	1/424	0/310	_		1.71%	2.2[0.09,53.71]
El-Refaey 2000	17/445	14/401		-	36.17%	1.09[0.55,2.19]
Harriott 2009	0/70	0/70				Not estimable
Ng 2007	0/178	0/177				Not estimable
Subtotal (95% CI)	1901	1758		*	100%	1.04[0.68,1.57]
Total events: 44 (Misoprostol), 41 (Erg	gometrine plus oxyt	ocin)				
Heterogeneity: Tau ² =0; Chi ² =0.42, df=	3(P=0.94); I ² =0%					
Test for overall effect: Z=0.17(P=0.87)						
45.18.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Misoprostol), 0 (Ergor	metrine plus oxytoc	in)				
Heterogeneity: Not applicable						
	Fa	vours Misoprostol	0.01 0.1	1 10	100 Favours Ergometrin	e plus oxytocin





Analysis 45.20. Comparison 45 Misoprostol vs Ergometrine plus oxytocin (by mode of birth), Outcome 20 Maternal satisfaction.

Study or subgroup	М	Misoprostol		Ergometrine plus oxytocin		Mean Difference				Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	xed, 95% CI	l		Fixed, 95% CI
45.20.1 Vaginal birth										
Ng 2007	178	27 (3.1)	177	26.4 (2.9)			-			0.6[-0.02,1.22]
45.20.2 Caesarean										
			Fa	vours Misoprostol	-5	-2.5	0	2.5	5	Favours Ergometrine plus oxytocin

Comparison 46. Misoprostol vs Misoprostol plus oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.85 [1.03, 3.33]
2.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	1.85 [1.03, 3.33]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1589	Risk Ratio (IV, Random, 95% CI)	2.97 [1.40, 6.30]
3.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	2.97 [1.40, 6.30]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



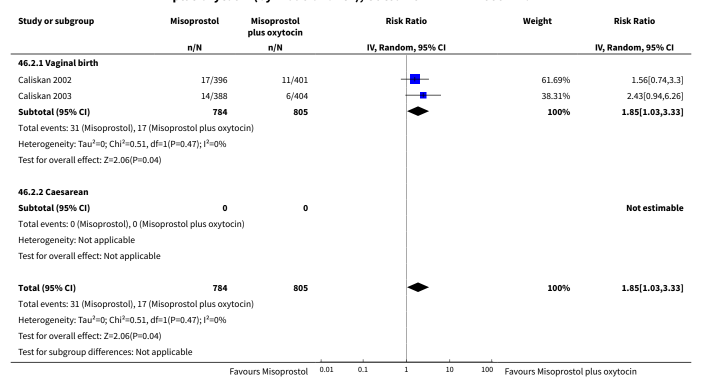
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity:shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.93 [0.99, 3.76]
6.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	1.93 [0.99, 3.76]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	3	1689	Risk Ratio (IV, Random, 95% CI)	1.89 [1.26, 2.83]
7.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	2.12 [1.35, 3.32]
7.2 Caesarean	1	100	Risk Ratio (IV, Random, 95% CI)	1.14 [0.45, 2.91]
8 Blood loss	1	792	Mean Difference (IV, Random, 95% CI)	48.0 [24.68, 71.32]
8.1 Vaginal birth	1	792	Mean Difference (IV, Random, 95% CI)	48.0 [24.68, 71.32]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	3	1689	Mean Difference (IV, Random, 95% CI)	0.27 [-0.42, 0.96]
9.1 Vaginal birth	2	1589	Mean Difference (IV, Random, 95% CI)	0.0 [-1.21, 1.21]
9.2 Caesarean	1	100	Mean Difference (IV, Random, 95% CI)	0.40 [-0.43, 1.23]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	100	Risk Ratio (IV, Random, 95% CI)	1.83 [0.73, 4.57]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	1	100	Risk Ratio (IV, Random, 95% CI)	1.83 [0.73, 4.57]
12 Vomiting	3	1689	Risk Ratio (IV, Random, 95% CI)	1.39 [0.51, 3.82]
12.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	1.03 [0.33, 3.24]
12.2 Caesarean	1	100	Risk Ratio (IV, Random, 95% CI)	4.0 [0.46, 34.54]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	1689	Risk Ratio (IV, Random, 95% CI)	0.94 [0.72, 1.22]
16.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	0.92 [0.71, 1.21]
16.2 Caesarean	1	100	Risk Ratio (IV, Random, 95% CI)	7.0 [0.37, 132.10]
17 Fever	3	1689	Risk Ratio (IV, Random, 95% CI)	0.97 [0.62, 1.53]
17.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	0.97 [0.61, 1.54]
17.2 Caesarean	1	100	Risk Ratio (IV, Random, 95% CI)	1.0 [0.15, 6.82]
18 Diarrhoea	2	1589	Risk Ratio (IV, Random, 95% CI)	1.22 [0.70, 2.13]
18.1 Vaginal birth	2	1589	Risk Ratio (IV, Random, 95% CI)	1.22 [0.70, 2.13]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 46.2. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

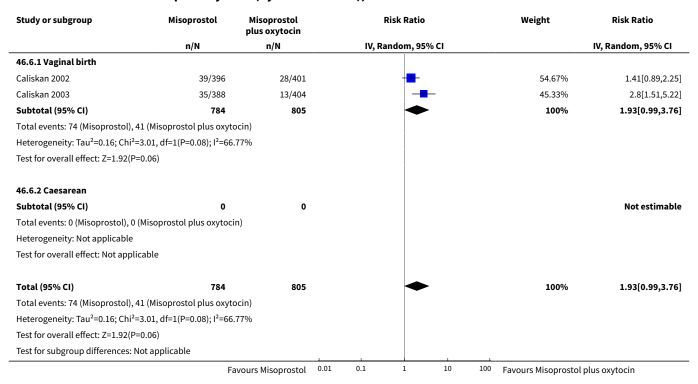


Analysis 46.3. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
46.3.1 Vaginal birth					
Caliskan 2002	12/396	4/401		44.79%	3.04[0.99,9.34]
Caliskan 2003	14/388	5/404		55.21%	2.92[1.06,8.02]
Subtotal (95% CI)	784	805	•	100%	2.97[1.4,6.3]
Total events: 26 (Misoprostol), 9 (Mis	soprostol plus oxytoc	in)			
Heterogeneity: Tau ² =0; Chi ² =0, df=1	P=0.96); I ² =0%				
Test for overall effect: Z=2.84(P=0)					
46.3.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Miso	prostol plus oxytocir	1)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	784	805	•	100%	2.97[1.4,6.3]
Total events: 26 (Misoprostol), 9 (Mis	soprostol plus oxytoc	in)			
Heterogeneity: Tau ² =0; Chi ² =0, df=1	P=0.96); I ² =0%				
Test for overall effect: Z=2.84(P=0)					
Test for subgroup differences: Not a	oplicable				
	Far	vours Misoprostol 0.03	1 0.1 1 10	100 Favours Misoprosto	plus oxytocin



Analysis 46.6. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.



Analysis 46.7. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin	Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Random, 95	% CI		IV, Random, 95% CI
46.7.1 Vaginal birth						
Caliskan 2002	33/396	17/401	-	_	50.59%	1.97[1.11,3.47]
Caliskan 2003	23/388	10/404	-		30.74%	2.39[1.16,4.97]
Subtotal (95% CI)	784	805	•	•	81.33%	2.12[1.35,3.32]
Total events: 56 (Misoprostol), 2	27 (Misoprostol plus oxyto	ocin)				
Heterogeneity: Tau ² =0; Chi ² =0.1	18, df=1(P=0.68); I ² =0%					
Test for overall effect: Z=3.28(P	=0)					
46.7.2 Caesarean						
Pakniat 2015	8/50	7/50	-		18.67%	1.14[0.45,2.91]
Subtotal (95% CI)	50	50	•		18.67%	1.14[0.45,2.91]
Total events: 8 (Misoprostol), 7	(Misoprostol plus oxytoci	n)				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.28(P	=0.78)					
Total (95% CI)	834	855	•		100%	1.89[1.26,2.83]
Total events: 64 (Misoprostol), 3	34 (Misoprostol plus oxyto	ocin)				
	Fa	vours Misoprostol	0.01 0.1 1	10 100 Favo	urs Misoprostol	l plus oxytocin



Study or subgroup	Misoprostol	Misoprostol Misoprostol plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
Heterogeneity: Tau ² =0; Chi ² =	1.53, df=2(P=0.46); I ² =0%								
Test for overall effect: Z=3.08	(P=0)								
Test for subgroup differences	s: Chi ² =1.36, df=1 (P=0.24), I	²=26.39%							
	F	avours Misoprostol	0.01	0.1	1	10	100	Favours Misoprosto	ol plus oxytocin

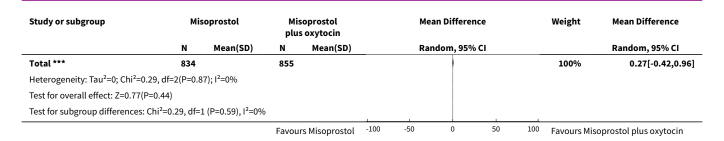
Analysis 46.8. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Mis	oprostol		oprostol oxytocin	Mean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Ra	ndom, 95% CI		Random, 95% CI
46.8.1 Vaginal birth								
Caliskan 2003	388	328 (152)	404	280 (182)			100%	48[24.68,71.32]
Subtotal ***	388		404				100%	48[24.68,71.32]
Heterogeneity: Not applicable								
Test for overall effect: Z=4.03(P<0.0	0001)							
46.8.2 Caesarean								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
Test for overall effect: Not applicab	ole							
Total ***	388		404			•	100%	48[24.68,71.32]
Heterogeneity: Not applicable								
Test for overall effect: Z=4.03(P<0.0	0001)							
Test for subgroup differences: Not	applicable	!						
			Favour	rs Misoprostol -1	00 -50	0 50	100 Favours Mis	oprostol plus oxytocin

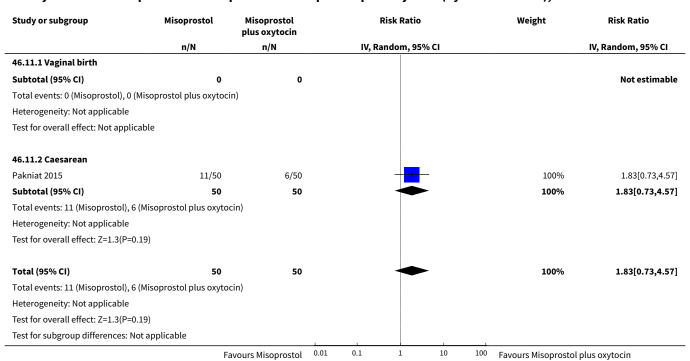
Analysis 46.9. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	Misoprostol		Misoprostol plus oxytocin		Mean Difference		Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
46.9.1 Vaginal birth										
Caliskan 2002	396	15 (12)	401	15 (13)			+		15.59%	0[-1.74,1.74]
Caliskan 2003	388	14 (11)	404	14 (13)			+		16.77%	0[-1.67,1.67]
Subtotal ***	784		805				•		32.36%	0[-1.21,1.21]
Heterogeneity: Tau ² =0; Chi ² =	0, df=1(P=1); I ² =0	0%								
Test for overall effect: Not ap	plicable									
46.9.2 Caesarean										
Pakniat 2015	50	11.4 (2.9)	50	11 (0.8)					67.64%	0.4[-0.43,1.23]
Subtotal ***	50		50						67.64%	0.4[-0.43,1.23]
Heterogeneity: Tau ² =0; Chi ² =	0, df=0(P<0.0001	.); I ² =100%								
Test for overall effect: Z=0.94	(P=0.35)									
							ĺ			
			Favou	rs Misoprostol	-100	-50	0 50	100	Favours Mis	oprostol plus oxytocin





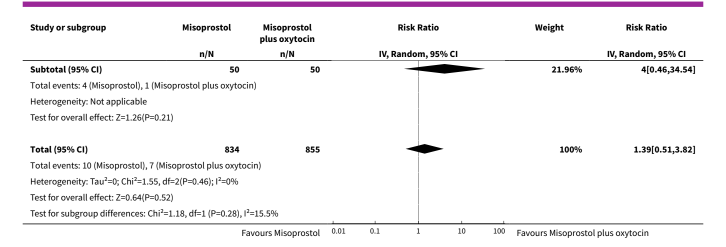
Analysis 46.11. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 11 Nausea.



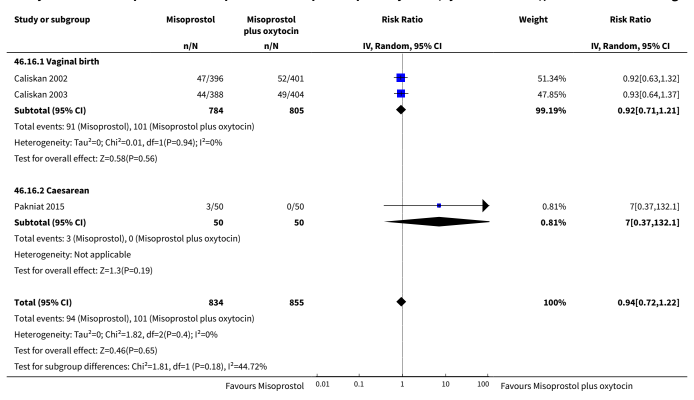
Analysis 46.12. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
46.12.1 Vaginal birth								
Caliskan 2002	2/396	3/401					32.08%	0.68[0.11,4.02]
Caliskan 2003	4/388	3/404			-		45.95%	1.39[0.31,6.16]
Subtotal (95% CI)	784	805		-	>		78.04%	1.03[0.33,3.24]
Total events: 6 (Misoprostol),	6 (Misoprostol plus oxytoci	n)						
Heterogeneity: Tau ² =0; Chi ² =0	0.37, df=1(P=0.54); I ² =0%							
Test for overall effect: Z=0.05(P=0.96)							
46.12.2 Caesarean								
Pakniat 2015	4/50	1/50		_			21.96%	4[0.46,34.54]
	Fa	vours Misoprostol	0.01	0.1	10	100	Favours Misoprosto	l plus oxytocin





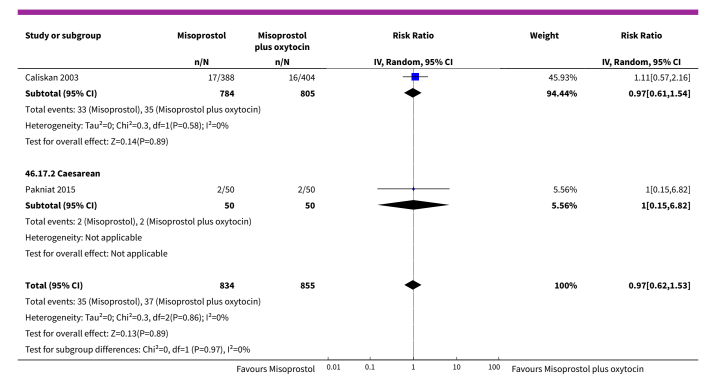
Analysis 46.16. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 16 Shivering.



Analysis 46.17. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Ra	andom, 95	% CI			IV, Random, 95% CI
46.17.1 Vaginal birth									
Caliskan 2002	16/396	19/401			-			48.5%	0.85[0.44,1.63]
	Fav	Favours Misoprostol		0.1	1	10	100	Favours Misoprostol	plus oxytocin





Analysis 46.18. Comparison 46 Misoprostol vs Misoprostol plus oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
46.18.1 Vaginal birth					
Caliskan 2002	11/396	9/401	- -	41.29%	1.24[0.52,2.95]
Caliskan 2003	15/388	13/404		58.71%	1.2[0.58,2.49]
Subtotal (95% CI)	784	805	*	100%	1.22[0.7,2.13]
Total events: 26 (Misoprostol), 22 (Mi	soprostol plus oxyto	ocin)			
Heterogeneity: Tau ² =0; Chi ² =0, df=1(I	P=0.96); I ² =0%				
Test for overall effect: Z=0.69(P=0.49)					
46.18.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Miso	prostol plus oxytocii	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	784	805	•	100%	1.22[0.7,2.13]
Total events: 26 (Misoprostol), 22 (Mi	soprostol plus oxyto	ocin)			
Heterogeneity: Tau ² =0; Chi ² =0, df=1(I	P=0.96); I ² =0%				
Test for overall effect: Z=0.69(P=0.49)					
Test for subgroup differences: Not ap	plicable				
	Fa	vours Misoprostol 0.	.01 0.1 1 10 10	⁰⁰ Favours Misoprostol	plus oxytocin



Comparison 47. Carbetocin vs Injectable prostaglandins (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 48. Injectable prostaglandins vs Ergometrine (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.1 Vaginal birth	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	4	384	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.58]
3.1 Vaginal birth	4	384	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.58]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	399	Risk Ratio (IV, Random, 95% CI)	1.42 [0.64, 3.17]
6.1 Vaginal birth	3	399	Risk Ratio (IV, Random, 95% CI)	1.42 [0.64, 3.17]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	5	684	Risk Ratio (IV, Random, 95% CI)	0.78 [0.40, 1.51]
7.1 Vaginal birth	5	684	Risk Ratio (IV, Random, 95% CI)	0.78 [0.40, 1.51]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	8	1195	Mean Difference (IV, Random, 95% CI)	-16.08 [-57.17, 25.00]
8.1 Vaginal birth	8	1195	Mean Difference (IV, Random, 95% CI)	-16.08 [-57.17, 25.00]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	2	300	Mean Difference (IV, Random, 95% CI)	-1.28 [-6.58, 4.01]
9.1 Vaginal birth	2	300	Mean Difference (IV, Random, 95% CI)	-1.28 [-6.58, 4.01]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	5	684	Risk Ratio (IV, Random, 95% CI)	1.70 [0.98, 2.94]
11.1 Vaginal birth	5	684	Risk Ratio (IV, Random, 95% CI)	1.70 [0.98, 2.94]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	5	684	Risk Ratio (IV, Random, 95% CI)	2.70 [0.97, 7.55]
12.1 Vaginal birth	5	684	Risk Ratio (IV, Random, 95% CI)	2.70 [0.97, 7.55]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	80	Risk Ratio (IV, Random, 95% CI)	2.0 [0.39, 10.31]
13.1 Vaginal birth	1	80	Risk Ratio (IV, Random, 95% CI)	2.0 [0.39, 10.31]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	3	384	Risk Ratio (IV, Random, 95% CI)	1.70 [0.07, 40.44]
14.1 Vaginal birth	3	384	Risk Ratio (IV, Random, 95% CI)	1.70 [0.07, 40.44]

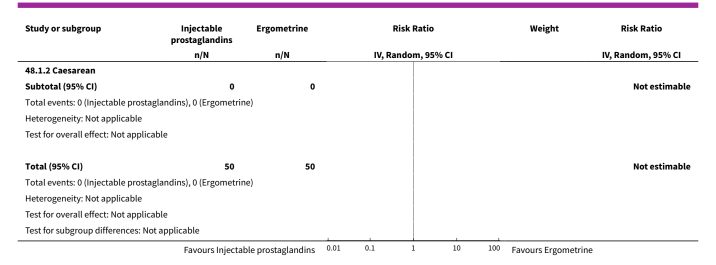


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	2	315	Risk Ratio (IV, Random, 95% CI)	0.16 [0.02, 1.37]
15.1 Vaginal birth	2	315	Risk Ratio (IV, Random, 95% CI)	0.16 [0.02, 1.37]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	434	Risk Ratio (IV, Random, 95% CI)	0.38 [0.16, 0.91]
16.1 Vaginal birth	3	434	Risk Ratio (IV, Random, 95% CI)	0.38 [0.16, 0.91]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	4	534	Risk Ratio (IV, Random, 95% CI)	4.52 [0.76, 26.80]
17.1 Vaginal birth	4	534	Risk Ratio (IV, Random, 95% CI)	4.52 [0.76, 26.80]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	5	664	Risk Ratio (IV, Random, 95% CI)	10.09 [2.75, 37.00]
18.1 Vaginal birth	5	664	Risk Ratio (IV, Random, 95% CI)	10.09 [2.75, 37.00]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

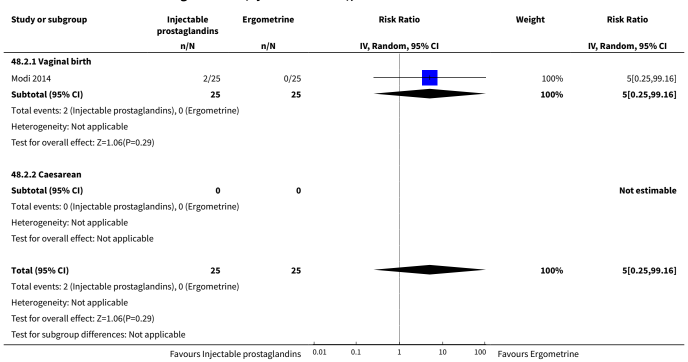
Analysis 48.1. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
48.1.1 Vaginal birth									
Supe 2016	0/50	0/50							Not estimable
Subtotal (95% CI)	50	50							Not estimable
Total events: 0 (Injectable prostagla	ındins), 0 (Ergometrin	e)							
Heterogeneity: Not applicable									
Test for overall effect: Not applicabl	e								
						1			
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	





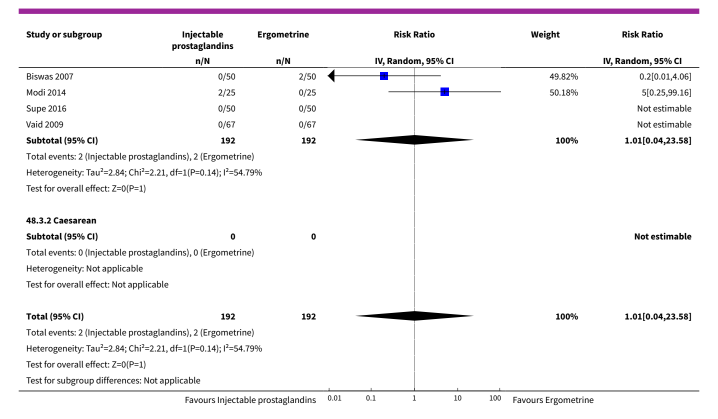
Analysis 48.2. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 2 PPH >= 1000 mL.



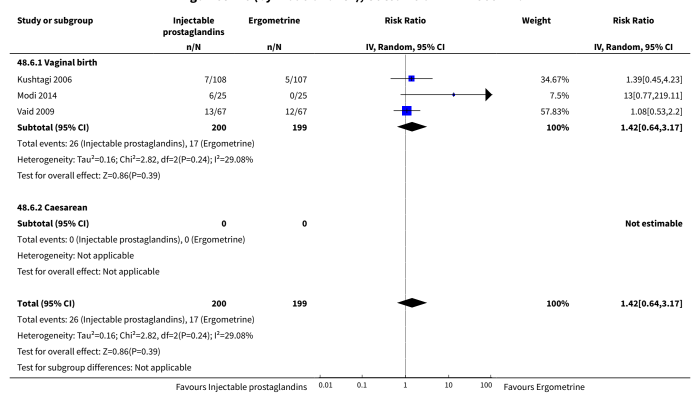
Analysis 48.3. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, Ra	ndom, 95	% CI			IV, Random, 95% CI
48.3.1 Vaginal birth									
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	



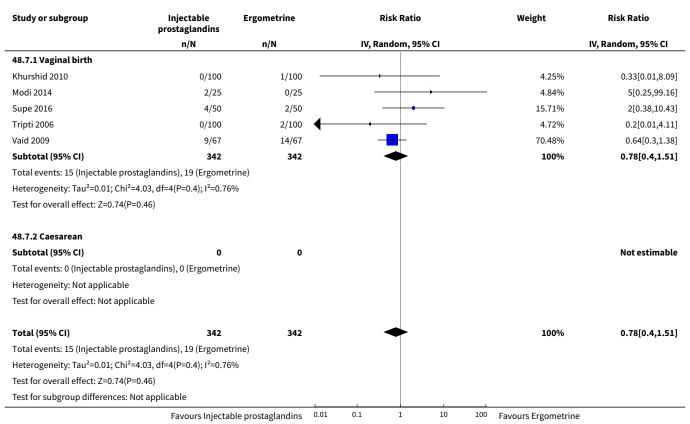


Analysis 48.6. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 6 PPH >= 500 mL.





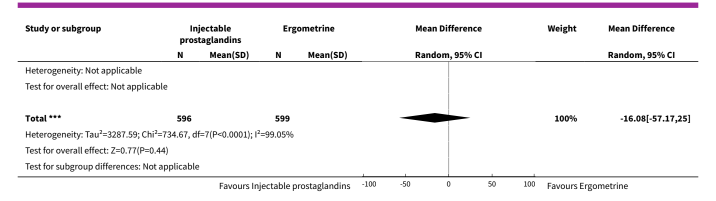
Analysis 48.7. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 7 Additional uterotonics.



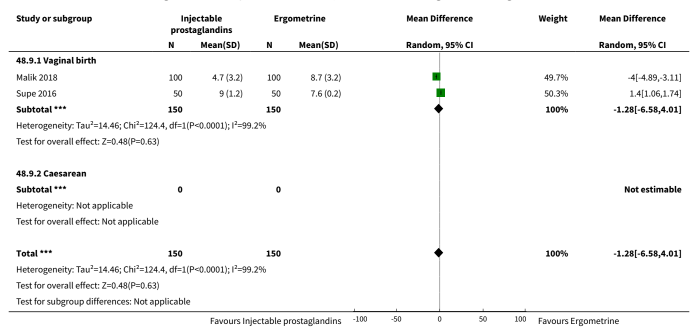
Analysis 48.8. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 8 Blood loss.

Study or subgroup		njectable Ergometrine Staglandins		ometrine	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
48.8.1 Vaginal birth							
Abdel-Aleem 1993	73	179 (59)	77	319 (52.3)	◀	13.04%	-140[-157.88,-122.12]
Khurshid 2010	100	63.6 (10.1)	100	83.6 (14.1)	+	13.35%	-20[-23.4,-16.6]
Kushtagi 2006	108	235.7 (99.3)	107	214.1 (110)	+	12.58%	21.6[-6.42,49.62]
Malik 2018	100	129 (27.3)	100	250 (35.2)	•	13.29%	-121[-129.73,-112.27]
Modi 2014	25	435 (147.6)	25	131 (72)		10.06%	304[239.63,368.37]
Reddy 2001	40	113 (127)	40	202 (84)	+	11.36%	-89[-136.19,-41.81]
Supe 2016	50	153.8 (43.5)	50	152.2 (49.3)		13.02%	1.6[-16.61,19.81]
Tripti 2006	100	74.9 (27.2)	100	93.6 (32.7)	+	13.29%	-18.74[-27.07,-10.41]
Subtotal ***	596		599			100%	-16.08[-57.17,25]
Heterogeneity: Tau ² =3287.59; Chi ²	=734.67, d	=7(P<0.0001); I ² =	99.05%				
Test for overall effect: Z=0.77(P=0.	44)						
48.8.2 Caesarean							
Subtotal ***	0		0				Not estimable
		Favours Inje	ectable p	rostaglandins	-100 -50 0 5	0 100 Favours Erg	ometrine





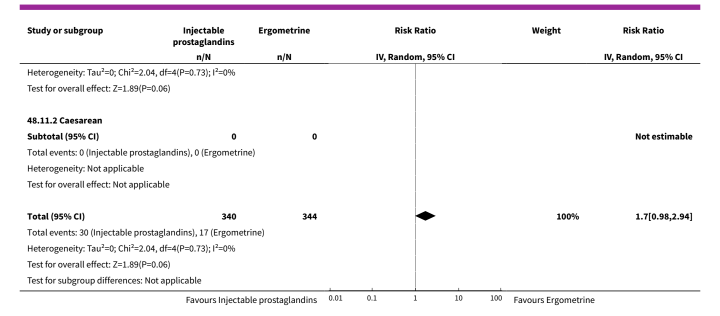
Analysis 48.9. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 9 Change in haemoglobin.



Analysis 48.11. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI
48.11.1 Vaginal birth								
Abdel-Aleem 1993	1/73	1/77			\rightarrow		3.97%	1.05[0.07,16.55]
Biswas 2007	3/50	0/50		_	+		3.49%	7[0.37,132.1]
Malik 2018	24/100	14/100			-		84.22%	1.71[0.94,3.12]
Supe 2016	0/50	1/50		+	 		2.98%	0.33[0.01,7.99]
Vaid 2009	2/67	1/67			+	_	5.33%	2[0.19,21.53]
Subtotal (95% CI)	340	344			•		100%	1.7[0.98,2.94]
Total events: 30 (Injectable p	prostaglandins), 17 (Ergomet	rine)				1		
	Favours Injectal	ole prostaglandins	0.01	0.1	1 10	100	Favours Ergometrine	



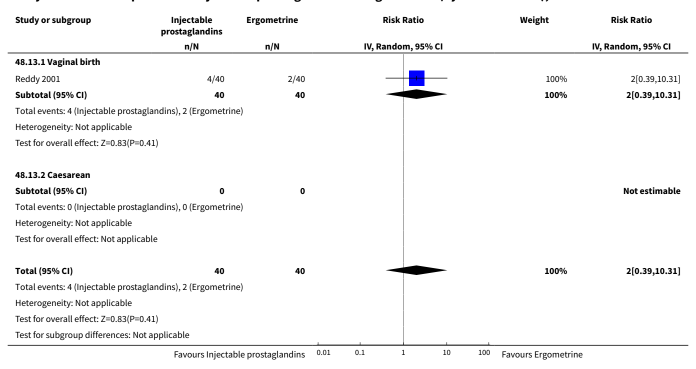


Analysis 48.12. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 12 Vomiting.

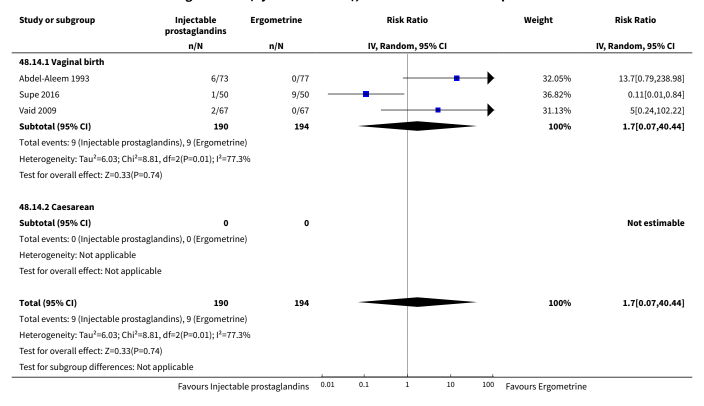
Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
48.12.1 Vaginal birth					
Abdel-Aleem 1993	12/73	1/77		17.95%	12.66[1.69,94.91]
Biswas 2007	1/50	2/50		14.21%	0.5[0.05,5.34]
Malik 2018	12/100	6/100		38.97%	2[0.78,5.12]
Supe 2016	8/50	1/50		17.62%	8[1.04,61.62]
Vaid 2009	1/67	1/67		11.25%	1[0.06,15.66]
Subtotal (95% CI)	340	344	~	100%	2.7[0.97,7.55]
Total events: 34 (Injectable prostag	landins), 11 (Ergometri	ne)			
Heterogeneity: Tau ² =0.48; Chi ² =6.1	5, df=4(P=0.19); I ² =34.9	9%			
Test for overall effect: Z=1.89(P=0.0	6)				
48.12.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Ergometrine	•)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	340	344	•	100%	2.7[0.97,7.55]
Total events: 34 (Injectable prostag	landins), 11 (Ergometri	ne)			
Heterogeneity: Tau ² =0.48; Chi ² =6.1	5, df=4(P=0.19); l ² =34.9	9%	ĺ		
Test for overall effect: Z=1.89(P=0.0	6)		ĺ		
Test for subgroup differences: Not a	applicable				
	Favours Injectab	e prostaglandins 0.01	0.1 1 10 10	DO Favours Ergometrin	e



Analysis 48.13. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 13 Headache.

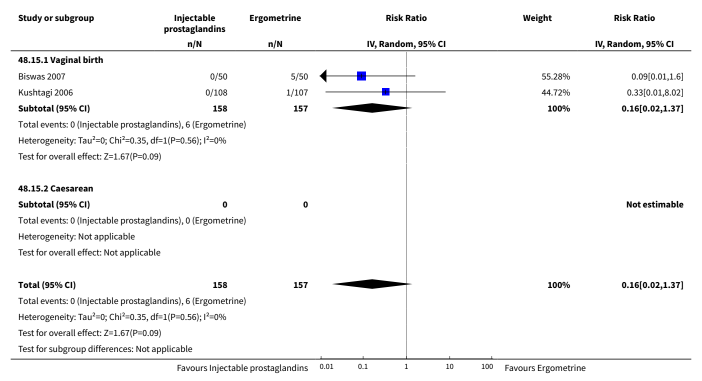


Analysis 48.14. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 14 Abdominal pain.





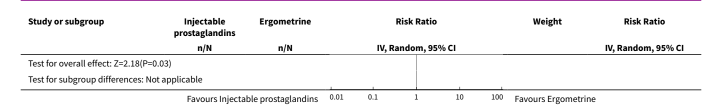
Analysis 48.15. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 15 Hypertension.



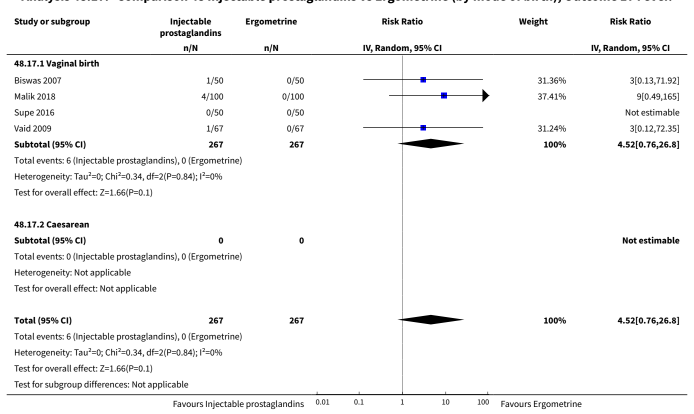
Analysis 48.16. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
48.16.1 Vaginal birth					
Malik 2018	6/100	14/100	- 1	90.95%	0.43[0.17,1.07]
Supe 2016	0/50	0/50			Not estimable
Vaid 2009	0/67	4/67		9.05%	0.11[0.01,2.02]
Subtotal (95% CI)	217	217	•	100%	0.38[0.16,0.91]
Total events: 6 (Injectable prostage	landins), 18 (Ergometri	ne)			
Heterogeneity: Tau ² =0; Chi ² =0.76,	df=1(P=0.38); I ² =0%				
Test for overall effect: Z=2.18(P=0.0	03)				
48.16.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostag	landins), 0 (Ergometrin	e)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	ole				
Total (95% CI)	217	217	•	100%	0.38[0.16,0.91]
Total events: 6 (Injectable prostag	landins), 18 (Ergometri	ne)	İ		
Heterogeneity: Tau ² =0; Chi ² =0.76,	df=1(P=0.38); I ² =0%				
	Favours Injectal	ole prostaglandins	0.01 0.1 1 10 1	00 Favours Ergometrine	





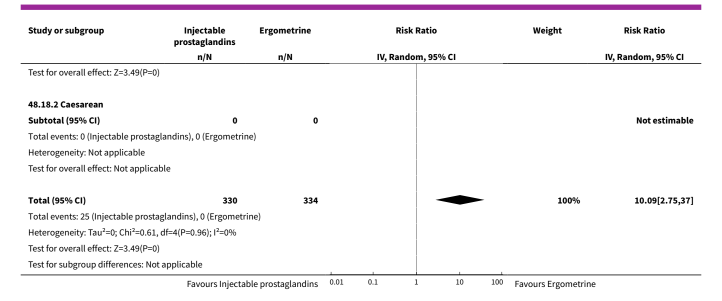
Analysis 48.17. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 17 Fever.



Analysis 48.18. Comparison 48 Injectable prostaglandins vs Ergometrine (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ratio		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	ľ	/, Random, 95% CI			IV, Random, 95% CI		
48.18.1 Vaginal birth									
Abdel-Aleem 1993	2/73	0/77		+		18.53%	5.27[0.26,107.96]		
Malik 2018	2/100	0/100		-		18.48%	5[0.24,102.85]		
Reddy 2001	7/40	0/40		-		21.1%	15[0.89,254.13]		
Supe 2016	7/50	0/50		+		21%	15[0.88,255.78]		
Vaid 2009	7/67	0/67		+		20.9%	15[0.87,257.48]		
Subtotal (95% CI)	330	334			-	100%	10.09[2.75,37]		
Total events: 25 (Injectable p	rostaglandins), 0 (Ergometri	ne)							
Heterogeneity: Tau ² =0; Chi ² =	0.61, df=4(P=0.96); I ² =0%								
	Favours Injectal	ole prostaglandins	0.01 0.1	1 10	100	Favours Ergometrine			





Comparison 49. Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	598	Risk Ratio (IV, Random, 95% CI)	0.80 [0.17, 3.78]
2.1 Vaginal birth	1	69	Risk Ratio (IV, Random, 95% CI)	0.36 [0.11, 1.23]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Both caesarean and vaginal birth	1	529	Risk Ratio (IV, Random, 95% CI)	1.77 [0.52, 5.97]
3 Blood transfusion	1	69	Risk Ratio (IV, Random, 95% CI)	0.65 [0.17, 2.53]
3.1 Vaginal birth	1	69	Risk Ratio (IV, Random, 95% CI)	0.65 [0.17, 2.53]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	598	Risk Ratio (IV, Random, 95% CI)	1.29 [0.92, 1.79]
6.1 Vaginal birth	1	69	Risk Ratio (IV, Random, 95% CI)	1.09 [0.66, 1.81]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 Both caesarean and vaginal birth	1	529	Risk Ratio (IV, Random, 95% CI)	1.45 [0.94, 2.24]
7 Additional uterotonics	1	112	Risk Ratio (IV, Random, 95% CI)	1.07 [0.16, 7.36]
7.1 Vaginal birth	1	112	Risk Ratio (IV, Random, 95% CI)	1.07 [0.16, 7.36]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	598	Mean Difference (IV, Random, 95% CI)	-26.18 [-104.63, 52.27]
8.1 Vaginal birth	1	69	Mean Difference (IV, Random, 95% CI)	-149.0 [-421.73, 123.73]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Both caesarean and vaginal birth	1	529	Mean Difference (IV, Random, 95% CI)	-15.10 [-97.01, 66.81]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	529	Risk Ratio (IV, Random, 95% CI)	5.06 [1.12, 22.86]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.3 Both caesarean and vaginal birth	1	529	Risk Ratio (IV, Random, 95% CI)	5.06 [1.12, 22.86]
12 Vomiting	1	529	Risk Ratio (IV, Random, 95% CI)	0.92 [0.40, 2.13]
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

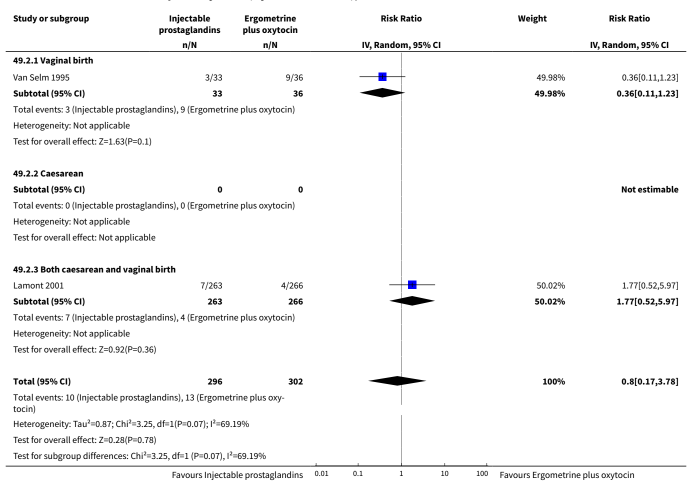


Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.3 Both caesarean and vaginal birth	1	529	Risk Ratio (IV, Random, 95% CI)	0.92 [0.40, 2.13]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	641	Risk Ratio (IV, Random, 95% CI)	23.70 [7.55, 74.45]
18.1 Vaginal birth	1	112	Risk Ratio (IV, Random, 95% CI)	17.19 [2.36, 125.22]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.3 Both caesarean and vaginal birth	1	529	Risk Ratio (IV, Random, 95% CI)	27.81 [6.86, 112.85]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

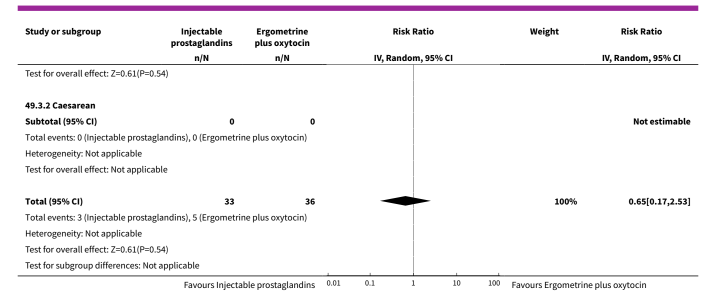
Analysis 49.2. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.



Analysis 49.3. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N	ı	V, Random, 95%	6 CI			IV, Random, 95% CI
49.3.1 Vaginal birth								
Van Selm 1995	3/33	5/36					100%	0.65[0.17,2.53]
Subtotal (95% CI)	33	36					100%	0.65[0.17,2.53]
Total events: 3 (Injectable pro	staglandins), 5 (Ergometrir	ne plus oxytocin)						
Heterogeneity: Not applicable	2							
	Favours Injectal	ble prostaglandins	0.01 0.1	1	10	100	Favours Ergometrine	e plus oxytocin



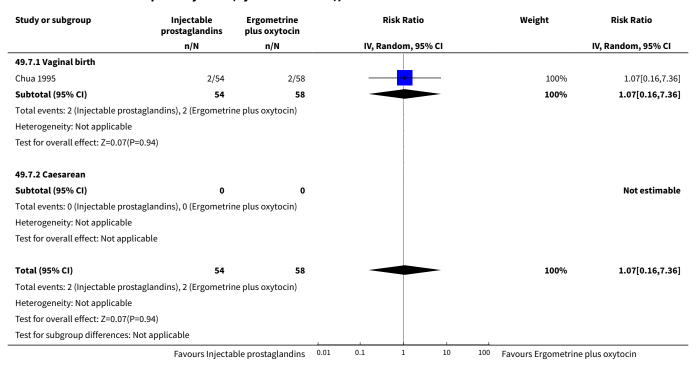


Analysis 49.6. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

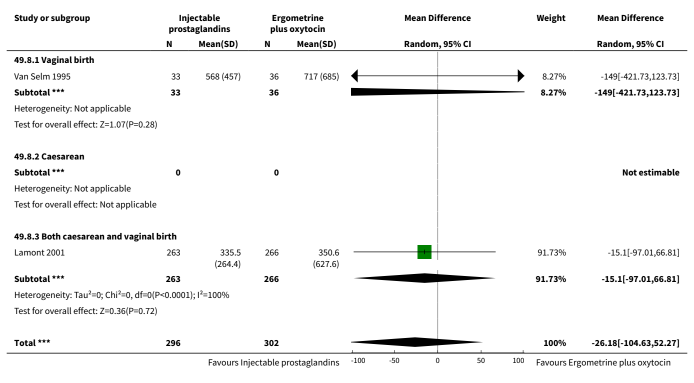
Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV,	Random, 95% CI		IV, Random, 95% CI
49.6.1 Vaginal birth						
Van Selm 1995	16/33	16/36		-	42.28%	1.09[0.66,1.81]
Subtotal (95% CI)	33	36		•	42.28%	1.09[0.66,1.81]
Total events: 16 (Injectable prostag tocin)	glandins), 16 (Ergome	trine plus oxy-				
Heterogeneity: Not applicable						
Test for overall effect: Z=0.34(P=0.7	(4)					
49.6.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Ergometrii	ne plus oxytocin)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicab	le					
49.6.3 Both caesarean and vagina	al birth					
Lamont 2001	43/263	30/266		-	57.72%	1.45[0.94,2.24]
Subtotal (95% CI)	263	266		•	57.72%	1.45[0.94,2.24]
Total events: 43 (Injectable prostag tocin)	glandins), 30 (Ergome	trine plus oxy-				
Heterogeneity: Not applicable						
Test for overall effect: Z=1.68(P=0.0	9)					
Total (95% CI)	296	302		•	100%	1.29[0.92,1.79]
Total events: 59 (Injectable prostag tocin)	(landins), 46 (Ergome	trine plus oxy-				, -
Heterogeneity: Tau ² =0; Chi ² =0.7, df	=1(P=0.4); I ² =0%					
Test for overall effect: Z=1.49(P=0.1	4)					
Test for subgroup differences: Chi ²	=0.7, df=1 (P=0.4), I ² =0	0%				
	Favours Injecta	ble prostaglandins	0.01 0.1	1 10	100 Favours Ergometri	ne plus oxytocin



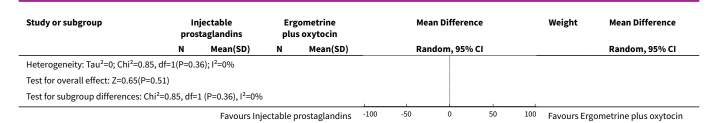
Analysis 49.7. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 7 Additional uterotonics.



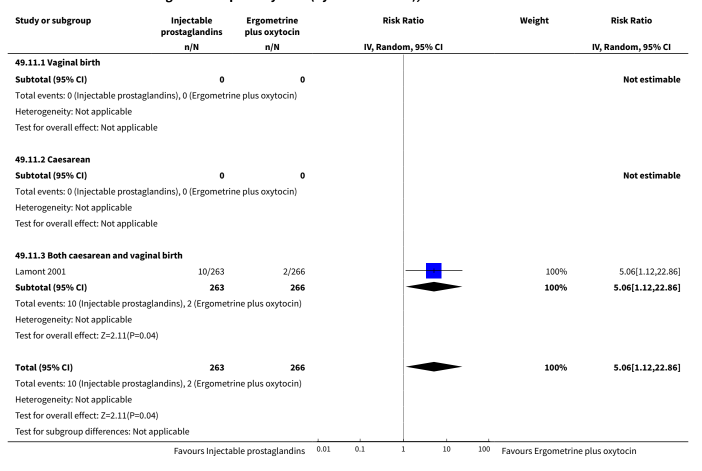
Analysis 49.8. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 8 Blood loss.







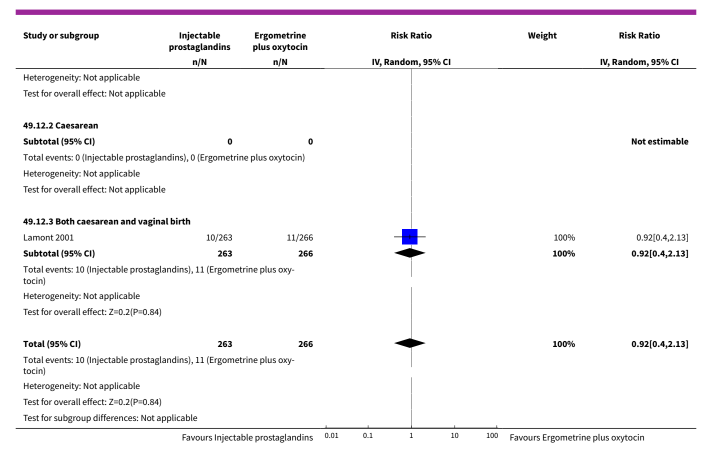
Analysis 49.11. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 11 Nausea.



Analysis 49.12. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Ri	isk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Ran	ndom, 95% (CI			IV, Random, 95% CI
49.12.1 Vaginal birth									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Injectable pro	staglandins), 0 (Ergometrin	e plus oxytocin)							
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Ergometrin	e plus oxytocin

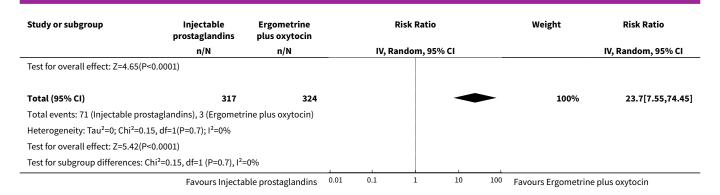




Analysis 49.18. Comparison 49 Injectable prostaglandins vs Ergometrine plus oxytocin (by mode of birth), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
49.18.1 Vaginal birth					
Chua 1995	16/54	1/58	-	33.21%	17.19[2.36,125.22]
Subtotal (95% CI)	54	58		33.21%	17.19[2.36,125.22]
Total events: 16 (Injectable prosta	glandins), 1 (Ergometr	ine plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=2.81(P=0.0	01)				
49.18.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostag	landins), 0 (Ergometrir	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applical	ble				
49.18.3 Both caesarean and vagi	nal birth				
Lamont 2001	55/263	2/266		66.79%	27.81[6.86,112.85]
Subtotal (95% CI)	263	266		66.79%	27.81[6.86,112.85]
Total events: 55 (Injectable prosta	glandins), 2 (Ergometr	ine plus oxytocin)			
Heterogeneity: Not applicable					
	Favours Injecta	ble prostaglandins 0	0.01 0.1 1 10 10	⁰ Favours Ergometrin	e plus oxytocin





Comparison 50. Misoprostol plus oxytocin vs Injectable prostaglandins (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title			Statistical method	Effect size	
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 51. Ergometrine vs Carbetocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 52. Carbetocin vs Ergometrine plus oxytocin (by mode of birth)

Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	4	1210	Risk Ratio (IV, Random, 95% CI)	0.34 [0.13, 0.86]
2.1 Vaginal birth	3	910	Risk Ratio (IV, Random, 95% CI)	0.69 [0.11, 4.38]
2.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	0.27 [0.09, 0.78]
3 Blood transfusion	4	1030	Risk Ratio (IV, Random, 95% CI)	1.42 [0.42, 4.82]
3.1 Vaginal birth	4	1030	Risk Ratio (IV, Random, 95% CI)	1.42 [0.42, 4.82]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	4	1030	Risk Ratio (IV, Random, 95% CI)	0.96 [0.44, 2.09]
6.1 Vaginal birth	4	1030	Risk Ratio (IV, Random, 95% CI)	0.96 [0.44, 2.09]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	6	1530	Risk Ratio (IV, Random, 95% CI)	0.54 [0.30, 1.00]
7.1 Vaginal birth	5	1230	Risk Ratio (IV, Random, 95% CI)	0.73 [0.44, 1.21]
7.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	0.19 [0.08, 0.49]



Outcome or sub- group title	No. of studies	No. of partici- pants	Statistical method	Effect size	
8 Blood loss	5	1330	Mean Difference (IV, Random, 95% CI)	-44.08 [-82.41, -5.75]	
8.1 Vaginal birth	4	1030	Mean Difference (IV, Random, 95% CI)	-48.84 [-94.82, -2.85]	
8.2 Caesarean	1	300	Mean Difference (IV, Random, 95% CI)	-24.0 [-68.42, 20.42]	
9 Change in haemo- globin	5	1160	Mean Difference (IV, Random, 95% CI)	-2.57 [-4.32, -0.82]	
9.1 Vaginal birth	4	860	Mean Difference (IV, Random, 95% CI)	-2.86 [-4.81, -0.90]	
9.2 Caesarean	1	300	Mean Difference (IV, Random, 95% CI)	-1.0 [-3.83, 1.83]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	6	1530	Risk Ratio (IV, Random, 95% CI)	0.29 [0.19, 0.45]	
11.1 Vaginal birth	5	1230	Risk Ratio (IV, Random, 95% CI)	0.27 [0.16, 0.43]	
11.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	0.45 [0.16, 1.28]	
12 Vomiting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.27 [0.16, 0.46]	
12.1 Vaginal birth	5	1230	Risk Ratio (IV, Random, 95% CI)	0.24 [0.13, 0.44]	
12.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	0.4 [0.13, 1.25]	
13 Headache	5	1330	Risk Ratio (IV, Random, 95% CI)	1.41 [0.45, 4.38]	
13.1 Vaginal birth	4	1030	Risk Ratio (IV, Random, 95% CI)	0.83 [0.46, 1.49]	
13.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	8.5 [2.00, 36.15]	
14 Abdominal pain	4	930	Risk Ratio (IV, Random, 95% CI)	0.57 [0.36, 0.91]	
14.1 Vaginal birth	4	930	Risk Ratio (IV, Random, 95% CI)	0.57 [0.36, 0.91]	
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15 Hypertension	3	660	Risk Ratio (IV, Random, 95% CI)	0.56 [0.22, 1.39]	
15.1 Vaginal birth	3	660	Risk Ratio (IV, Random, 95% CI)	0.56 [0.22, 1.39]	
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16 Shivering	5	1290	Risk Ratio (IV, Random, 95% CI)	0.45 [0.26, 0.80]	
16.1 Vaginal birth	4	990	Risk Ratio (IV, Random, 95% CI)	0.40 [0.22, 0.74]	
16.2 Caesarean	1	300	Risk Ratio (IV, Random, 95% CI)	1.0 [0.21, 4.88]	



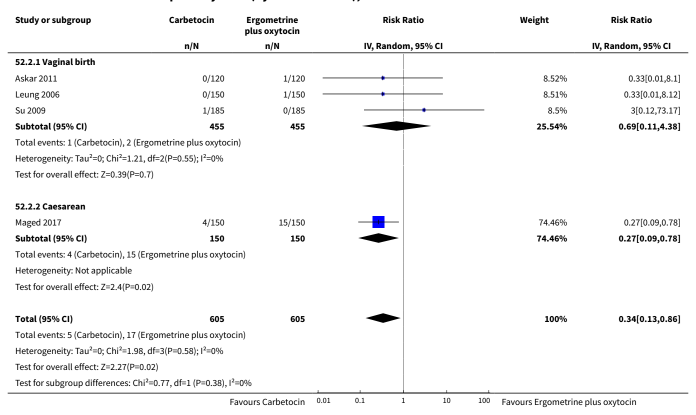
Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 52.1. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 1 Death.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk	Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI	
52.1.1 Vaginal birth							
Nirmala 2009	0/60	0/60				Not estimable	
Samimi 2013	0/100	0/100				Not estimable	
Subtotal (95% CI)	160	160				Not estimable	
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	٦)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
52.1.2 Caesarean							
Subtotal (95% CI)	0	0				Not estimable	
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	۱)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Total (95% CI)	160	160				Not estimable	
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	n)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Test for subgroup differences: Not app	olicable						
	Fa	avours Carbetocin	0.01 0.1	1 10 10	⁰ Favours Ergometr	ine plus oxytocin	



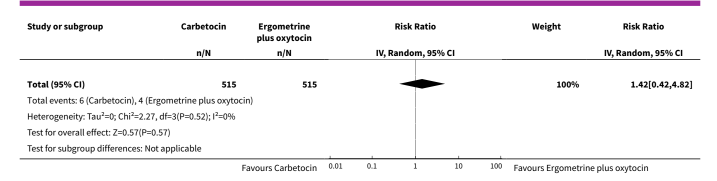
Analysis 52.2. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.



Analysis 52.3. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	R	isk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Ra	ndom, 95% CI		IV, Random, 95% CI
52.3.1 Vaginal birth						
Askar 2011	0/120	1/120	+		14.57%	0.33[0.01,8.1]
Leung 2006	5/150	2/150			56.23%	2.5[0.49,12.69]
Nirmala 2009	0/60	1/60	+		14.66%	0.33[0.01,8.02]
Su 2009	1/185	0/185		+	14.54%	3[0.12,73.17]
Subtotal (95% CI)	515	515			100%	1.42[0.42,4.82]
Total events: 6 (Carbetocin), 4 (Ergon	netrine plus oxytocin	1)				
Heterogeneity: Tau ² =0; Chi ² =2.27, df=	=3(P=0.52); I ² =0%					
Test for overall effect: Z=0.57(P=0.57)						
52.3.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocin	1)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
	Fa	vours Carbetocin	0.01 0.1	1 10	100 Favours Ergometrine	plus oxytocin





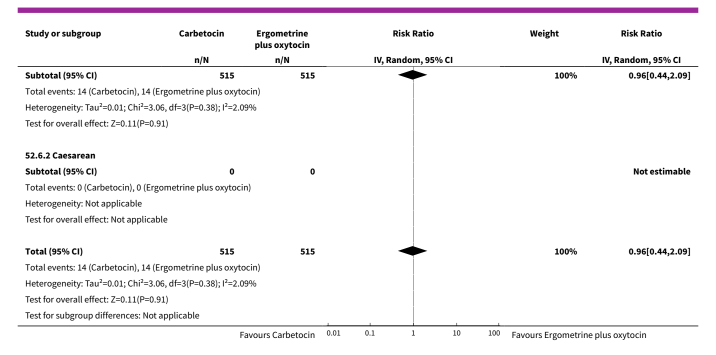
Analysis 52.4. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	or subgroup Carbetocin Ergometrine Risk Ratio plus oxytocin		Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
52.4.1 Vaginal birth						
Nirmala 2009	0/60	0/60				Not estimable
Samimi 2013	0/100	0/100				Not estimable
Subtotal (95% CI)	160	160				Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocin	1)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
52.4.2 Caesarean						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocin	1)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	160	160				Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocin	1)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not ap	plicable					
	Fa	vours Carbetocin	0.01 0.1	1 10 100	⁾ Favours Ergometri	ne plus oxytocin

Analysis 52.6. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin Ergometrine plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, Random, 95	% CI			IV, Random, 95% CI
52.6.1 Vaginal birth								
Askar 2011	2/120	3/120			_		19.18%	0.67[0.11,3.92]
Leung 2006	6/150	2/150		+-			23.88%	3[0.62,14.63]
Nirmala 2009	3/60	6/60					33.16%	0.5[0.13,1.91]
Su 2009	3/185	3/185			-	1	23.79%	1[0.2,4.89]
	Favours Carbetocin		0.01	0.1 1	10	100	Favours Ergometrine	plus oxytocin





Analysis 52.7. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin Ergometrine plus oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
52.7.1 Vaginal birth					
Askar 2011	18/120	21/120		21.97%	0.86[0.48,1.53]
Leung 2006	13/150	10/150	-	18.68%	1.3[0.59,2.87]
Nirmala 2009	3/60	9/60		12.61%	0.33[0.09,1.17]
Samimi 2013	1/100	11/100 —		6.76%	0.09[0.01,0.69]
Su 2009	25/185	31/185		23.31%	0.81[0.5,1.31]
Subtotal (95% CI)	615	615	•	83.32%	0.73[0.44,1.21]
Total events: 60 (Carbetocin), 82 (Erg	gometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0.15; Chi ² =7.78	, df=4(P=0.1); I ² =48.5	5%			
Test for overall effect: Z=1.24(P=0.22	2)				
52.7.2 Caesarean					
Maged 2017	5/150	26/150		16.68%	0.19[0.08,0.49]
Subtotal (95% CI)	150	150	•	16.68%	0.19[0.08,0.49]
Total events: 5 (Carbetocin), 26 (Erg	ometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
Test for overall effect: Z=3.47(P=0)					
Total (95% CI)	765	765	•	100%	0.54[0.3,1]
Total events: 65 (Carbetocin), 108 (E	rgometrine plus oxyt	ocin)			
Heterogeneity: Tau ² =0.35; Chi ² =15.7	6, df=5(P=0.01); l ² =68	.28%			
Test for overall effect: Z=1.97(P=0.05	i)				
Test for subgroup differences: Chi ² =	6.02, df=1 (P=0.01), I ²	=83.39%			
	F	avours Carbetocin 0.0	1 0.1 1 10	100 Favours Ergometrin	e plus oxytocin



Analysis 52.8. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Carbetocin		Ergometrine plus oxytocin		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
52.8.1 Vaginal birth							
Askar 2011	120	224.6 (110.6)	120	306.1 (95.7)		21.48%	-81.5[-107.66,-55.34]
Leung 2006	150	232 (122)	150	249 (175)		20.07%	-17[-51.14,17.14]
Nirmala 2009	60	244 (114)	60	343 (143)	—	17.71%	-99[-145.27,-52.73]
Su 2009	185	217.4 (99.2)	185	223.1 (76.3)		22.67%	-5.7[-23.73,12.33]
Subtotal ***	515		515			81.93%	-48.84[-94.82,-2.85]
Heterogeneity: Tau ² =1934.74; Chi ²	=30.65, df=	=3(P<0.0001); I ² =9	90.21%				
Test for overall effect: Z=2.08(P=0.0	04)						
52.8.2 Caesarean							
Maged 2017	150	578 (178)	150	602 (213)		18.07%	-24[-68.42,20.42]
Subtotal ***	150		150			18.07%	-24[-68.42,20.42]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.06(P=0.2	29)						
Total ***	665		665			100%	-44.08[-82.41,-5.75]
Heterogeneity: Tau ² =1602.31; Chi ²	=30.82, df=	=4(P<0.0001); I ² =8	87.02%				
Test for overall effect: Z=2.25(P=0.0	02)						
Test for subgroup differences: Chi ²	=0.58, df=	1 (P=0.45), I ² =0%					
			Favoi	urs Carbetocin	-100 -50 0 50	100 Favours Erg	ometrine plus oxytocin

Analysis 52.9. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup	Ca	Carbetocin		ometrine oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
52.9.1 Vaginal birth							
Askar 2011	120	8 (2)	120	11 (3)	-#-	24.41%	-3[-3.65,-2.35]
Leung 2006	150	14 (11)	150	15 (13)		15.67%	-1[-3.73,1.73]
Nirmala 2009	60	3 (2)	60	4 (2)	-	24.23%	-1[-1.72,-0.28]
Samimi 2013	100	4.1 (3.6)	100	10.4 (7.8)		20.47%	-6.3[-7.98,-4.62]
Subtotal ***	430		430		•	84.79%	-2.86[-4.81,-0.9]
Heterogeneity: Tau ² =3.37; Chi ² =39	.64, df=3(P	<0.0001); I ² =92.4	3%				
Test for overall effect: Z=2.86(P=0)							
52.9.2 Caesarean							
Maged 2017	150	11 (12)	150	12 (13)	-+-	15.21%	-1[-3.83,1.83]
Subtotal ***	150		150			15.21%	-1[-3.83,1.83]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.69(P=0.4	49)						
Total ***	580		580		•	100%	-2.57[-4.32,-0.82]
Heterogeneity: Tau ² =3.16; Chi ² =40	.53, df=4(P	<0.0001); I ² =90.1	3%				
Test for overall effect: Z=2.88(P=0)							
Test for subgroup differences: Chi ²	=1.12, df=1	(P=0.29), I ² =10.	57%				
			Favou	rs Carbetocin -	10 -5 0 5	10 Favours Erg	ometrine plus oxytocin



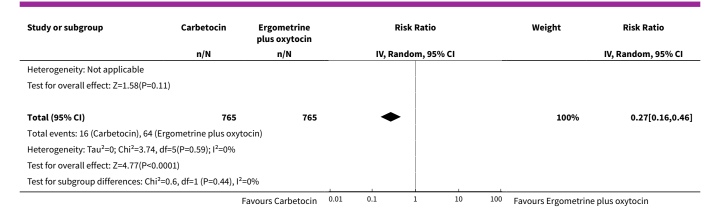
Analysis 52.11. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 11 Nausea.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
52.11.1 Vaginal birth					
Askar 2011	4/120	13/120		16.13%	0.31[0.1,0.92]
Leung 2006	2/150	11/150		8.66%	0.18[0.04,0.81]
Nirmala 2009	0/60	1/60	+	1.9%	0.33[0.01,8.02]
Samimi 2013	2/100	3/100		6.15%	0.67[0.11,3.9]
Su 2009	11/185	46/185		49.13%	0.24[0.13,0.45]
Subtotal (95% CI)	615	615	•	81.97%	0.27[0.16,0.43]
Total events: 19 (Carbetocin), 74	(Ergometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0; Chi ² =1.49	, df=4(P=0.83); I ² =0%				
Test for overall effect: Z=5.36(P<0	.0001)				
52.11.2 Caesarean					
Maged 2017	5/150	11/150		18.03%	0.45[0.16,1.28]
Subtotal (95% CI)	150	150		18.03%	0.45[0.16,1.28]
Total events: 5 (Carbetocin), 11 (E	Ergometrine plus oxytoci	n)			
Heterogeneity: Not applicable					
Test for overall effect: Z=1.5(P=0.3	13)				
Total (95% CI)	765	765	•	100%	0.29[0.19,0.45]
Total events: 24 (Carbetocin), 85	(Ergometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0; Chi ² =2.34	, df=5(P=0.8); I ² =0%				
Test for overall effect: Z=5.49(P<0	.0001)				
Test for subgroup differences: Ch	i ² =0.85, df=1 (P=0.36), I ² =	=0%			
<u> </u>	Fa	vours Carbetocin 0	.01 0.1 1 10	100 Favours Ergometrin	e plus oxytocin

Analysis 52.12. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
52.12.1 Vaginal birth					
Askar 2011	3/120	12/120		18.89%	0.25[0.07,0.86]
Leung 2006	1/150	10/150		6.95%	0.1[0.01,0.77]
Nirmala 2009	0/60	2/60		3.19%	0.2[0.01,4.08]
Samimi 2013	1/100	0/100		2.85%	3[0.12,72.77]
Su 2009	7/185	30/185		45.68%	0.23[0.11,0.52]
Subtotal (95% CI)	615	615	•	77.56%	0.24[0.13,0.44]
Total events: 12 (Carbetocin),	54 (Ergometrine plus oxyto	ocin)			
Heterogeneity: Tau ² =0; Chi ² =3	.14, df=4(P=0.53); I ² =0%				
Test for overall effect: Z=4.57(F	P<0.0001)				
52.12.2 Caesarean					
Maged 2017	4/150	10/150	-	22.44%	0.4[0.13,1.25]
Subtotal (95% CI)	150	150		22.44%	0.4[0.13,1.25]
Total events: 4 (Carbetocin), 1	0 (Ergometrine plus oxytoc	in)			
	F	avours Carbetocin	0.01 0.1 1 10	100 Favours Ergometrin	e plus oxytocin





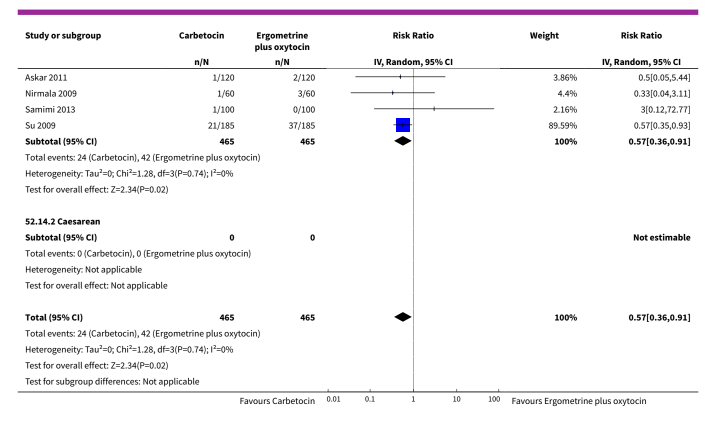
Analysis 52.13. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 13 Headache.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	•		Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
52.13.1 Vaginal birth					
Askar 2011	1/120	2/120	+	14.16%	0.5[0.05,5.44]
Leung 2006	1/150	2/150	+	14.14%	0.5[0.05,5.46]
Nirmala 2009	2/60	1/60		14.27%	2[0.19,21.47]
Su 2009	15/185	18/185		33.9%	0.83[0.43,1.6]
Subtotal (95% CI)	515	515	•	76.47%	0.83[0.46,1.49]
Total events: 19 (Carbetocin), 23 (I	Ergometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0; Chi ² =0.87,	df=3(P=0.83); I ² =0%				
Test for overall effect: Z=0.63(P=0.	53)				
52.13.2 Caesarean					
Maged 2017	17/150	2/150		23.53%	8.5[2,36.15]
Subtotal (95% CI)	150	150		23.53%	8.5[2,36.15]
Total events: 17 (Carbetocin), 2 (En	rgometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
Test for overall effect: Z=2.9(P=0)					
Total (95% CI)	665	665	•	100%	1.41[0.45,4.38]
Total events: 36 (Carbetocin), 25 (I	Ergometrine plus oxyto	cin)			
Heterogeneity: Tau ² =0.87; Chi ² =9.4	41, df=4(P=0.05); l ² =57.4	49%			
Test for overall effect: Z=0.6(P=0.5	5)				
			· ·		

Analysis 52.14. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 14 Abdominal pain.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio					Weight Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI		IV, Random, 95% CI
52.14.1 Vaginal birth								
		avours Carbetocin	0.01	0.1	1	10	100	Favours Ergometrine plus oxytocin



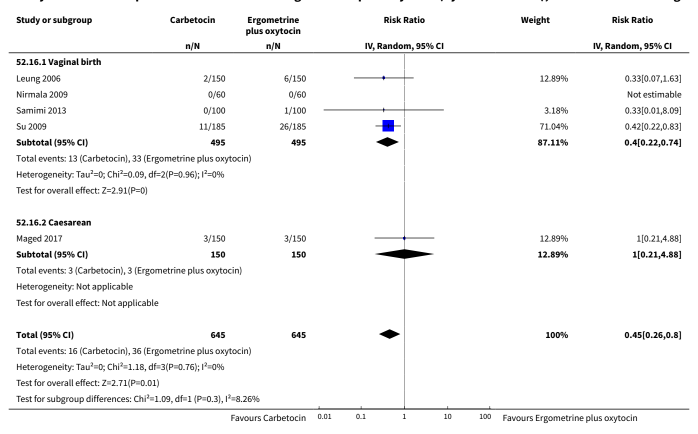


Analysis 52.15. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 15 Hypertension.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
52.15.1 Vaginal birth					
Askar 2011	2/120	4/120		25.6%	0.5[0.09,2.68]
Leung 2006	0/150	6/150	•	9.63%	0.08[0,1.35]
Nirmala 2009	7/60	9/60		64.77%	0.78[0.31,1.95]
Subtotal (95% CI)	330	330		100%	0.56[0.22,1.39]
Total events: 9 (Carbetocin), 19 (Ergo	metrine plus oxytoc	in)			
Heterogeneity: Tau ² =0.11; Chi ² =2.32,	df=2(P=0.31); I ² =13.	85%			
Test for overall effect: Z=1.26(P=0.21)					
52.15.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocii	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	330	330	•	100%	0.56[0.22,1.39]
Total events: 9 (Carbetocin), 19 (Ergo	metrine plus oxytoc	in)			
Heterogeneity: Tau ² =0.11; Chi ² =2.32,	df=2(P=0.31); I ² =13.	85%			
Test for overall effect: Z=1.26(P=0.21)					
Test for subgroup differences: Not ap	plicable		į		
	F	avours Carbetocin	0.01 0.1 1 10 1	00 Favours Ergometrine	e plus oxytocin



Analysis 52.16. Comparison 52 Carbetocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 16 Shivering.



Comparison 53. Misoprostol plus oxytocin vs Carbetocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	380	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	380	Risk Ratio (IV, Random, 95% CI)	4.00 [0.45, 35.46]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	4.00 [0.45, 35.46]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	1	380	Risk Ratio (IV, Random, 95% CI)	1.19 [0.74, 1.93]
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	1.19 [0.74, 1.93]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	1	380	Mean Difference (IV, Random, 95% CI)	-1.70 [-3.72, 0.32]
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	1	380	Mean Difference (IV, Random, 95% CI)	-1.70 [-3.72, 0.32]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	380	Risk Ratio (IV, Random, 95% CI)	1.21 [0.62, 2.39]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	1.21 [0.62, 2.39]
12 Vomiting	1	380	Risk Ratio (IV, Random, 95% CI)	1.33 [0.58, 3.09]



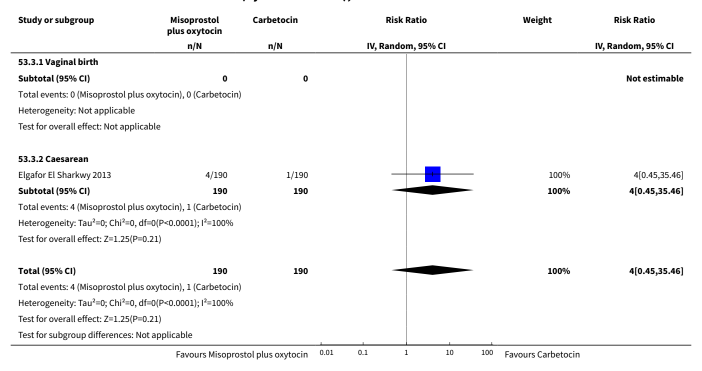
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	1.33 [0.58, 3.09]
13 Headache	1	380	Risk Ratio (IV, Random, 95% CI)	2.0 [0.37, 10.79]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	2.0 [0.37, 10.79]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	380	Risk Ratio (IV, Random, 95% CI)	7.83 [3.43, 17.88]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	7.83 [3.43, 17.88]
17 Fever	1	380	Risk Ratio (IV, Random, 95% CI)	8.5 [1.99, 36.28]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	1	380	Risk Ratio (IV, Random, 95% CI)	8.5 [1.99, 36.28]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 53.1. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 1 Death.

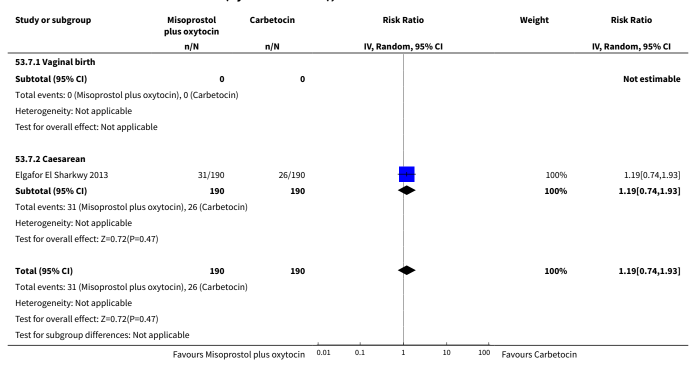
Study or subgroup	Misoprostol plus oxytocin	Carbetocin		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
53.1.1 Vaginal birth								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol plus oxyt	cocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
53.1.2 Caesarean								
Elgafor El Sharkwy 2013	0/190	0/190						Not estimable
Subtotal (95% CI)	190	190						Not estimable
Total events: 0 (Misoprostol plus oxyt	cocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total (95% CI)	190	190						Not estimable
Total events: 0 (Misoprostol plus oxyt	cocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Test for subgroup differences: Not ap	plicable							
	Favours Misopros	tol plus oxytocin	0.01	0.1	1 10	100 Fa	avours Carbetocin	

Analysis 53.3. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 3 Blood transfusion.





Analysis 53.7. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 7 Additional uterotonics.

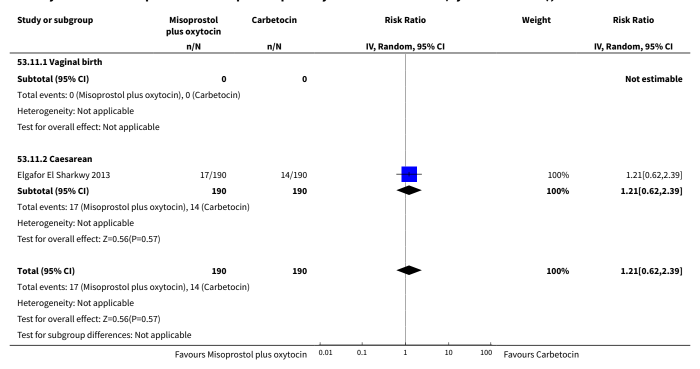


Analysis 53.9. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup		oprostol oxytocin	Ca	rbetocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
53.9.1 Vaginal birth							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
53.9.2 Caesarean							
Elgafor El Sharkwy 2013	190	13.4 (8.7)	190	15.1 (11.2)	+	100%	-1.7[-3.72,0.32]
Subtotal ***	190		190		•	100%	-1.7[-3.72,0.32]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.65(P=0.1)							
Total ***	190		190		•	100%	-1.7[-3.72,0.32]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.65(P=0.1)							
Test for subgroup differences: Not ap	plicable						
		Favours Mis	oprostol	plus oxytocin -100	-50 0 50	100 Favours Carl	betocin



Analysis 53.11. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 11 Nausea.



Analysis 53.12. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
53.12.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Carbetocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	le				
53.12.2 Caesarean					
Elgafor El Sharkwy 2013	12/190	9/190	-	100%	1.33[0.58,3.09]
Subtotal (95% CI)	190	190		100%	1.33[0.58,3.09]
Total events: 12 (Misoprostol plus o	xytocin), 9 (Carbetocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.67(P=0.5)				
Total (95% CI)	190	190	•	100%	1.33[0.58,3.09]
Total events: 12 (Misoprostol plus o	xytocin), 9 (Carbetocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.67(P=0.5)				
Test for subgroup differences: Not a	pplicable				
	Favours Misopros	tol plus oxytocin 0.0	01 0.1 1 10	100 Favours Carbetocin	



Analysis 53.13. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 13 Headache.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
53.13.1 Vaginal birth					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus oxy	tocin), 0 (Carbetocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
53.13.2 Caesarean					
Elgafor El Sharkwy 2013	4/190	2/190	- 	100%	2[0.37,10.79]
Subtotal (95% CI)	190	190		100%	2[0.37,10.79]
Total events: 4 (Misoprostol plus oxy	tocin), 2 (Carbetocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.81(P=0.42))				
Total (95% CI)	190	190		100%	2[0.37,10.79]
Total events: 4 (Misoprostol plus oxy	tocin), 2 (Carbetocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.81(P=0.42))				
Test for subgroup differences: Not ap	plicable				
	Favours Misoprost	col plus oxytocin 0.01	0.1 1 10 10	DO Favours Carbetocin	

Analysis 53.16. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rando	m, 95% CI			IV, Random, 95% CI
53.16.1 Vaginal birth								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol plus	oxytocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applic	cable							
53.16.2 Caesarean								
Elgafor El Sharkwy 2013	47/190	6/190			_		100%	7.83[3.43,17.88]
Subtotal (95% CI)	190	190			•		100%	7.83[3.43,17.88]
Total events: 47 (Misoprostol plu	us oxytocin), 6 (Carbetocin	1)						
Heterogeneity: Not applicable								
Test for overall effect: Z=4.89(P<	0.0001)							
Total (95% CI)	190	190			•		100%	7.83[3.43,17.88]
Total events: 47 (Misoprostol plu	us oxytocin), 6 (Carbetocin	1)						
Heterogeneity: Not applicable								
Test for overall effect: Z=4.89(P<	0.0001)							
Test for subgroup differences: No	ot applicable							
	Favours Misopros	stal plus axytacin	0.01	0.1	1 10	100	Favours Carbetocin	



Analysis 53.17. Comparison 53 Misoprostol plus oxytocin vs Carbetocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Randoı	n, 95% CI			IV, Random, 95% CI
53.17.1 Vaginal birth								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	e							
53.17.2 Caesarean								
Elgafor El Sharkwy 2013	17/190	2/190				-	100%	8.5[1.99,36.28]
Subtotal (95% CI)	190	190				-	100%	8.5[1.99,36.28]
Total events: 17 (Misoprostol plus o	xytocin), 2 (Carbetocir	n)						
Heterogeneity: Not applicable								
Test for overall effect: Z=2.89(P=0)								
Total (95% CI)	190	190			-	-	100%	8.5[1.99,36.28]
Total events: 17 (Misoprostol plus o	xytocin), 2 (Carbetocir	۱)						
Heterogeneity: Not applicable								
Test for overall effect: Z=2.89(P=0)								
Test for subgroup differences: Not a	pplicable							
	Favours Misopro	stol plus oxytocin	0.01 0.1	1 1	10	100	Favours Carbetocin	

Comparison 54. Ergometrine vs Ergometrine plus oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



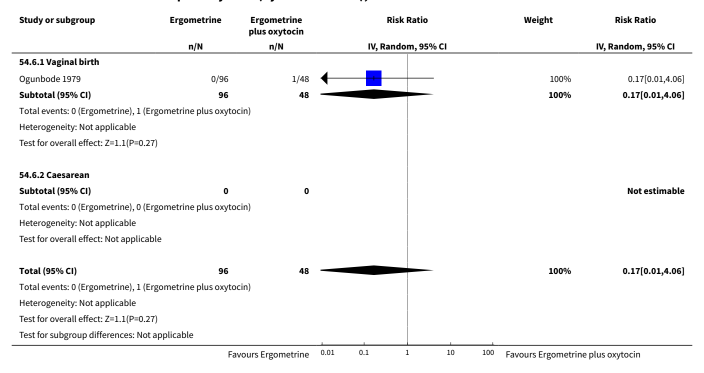
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	1	144	Risk Ratio (IV, Random, 95% CI)	0.17 [0.01, 4.06]	
6.1 Vaginal birth	1	144	Risk Ratio (IV, Random, 95% CI)	0.17 [0.01, 4.06]	
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	1	144	Mean Difference (IV, Random, 95% CI)	20.10 [5.76, 34.44]	
8.1 Vaginal birth	1	144	Mean Difference (IV, Random, 95% CI)	20.10 [5.76, 34.44]	
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 54.6. Comparison 54 Ergometrine vs Ergometrine plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.



Analysis 54.8. Comparison 54 Ergometrine vs Ergometrine plus oxytocin (by mode of birth), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	_	ometrine oxytocin	Mean	Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Rando	om, 95% CI		Random, 95% CI
54.8.1 Vaginal birth								
Ogunbode 1979	96	96 (54.2)	48	75.9 (33.2)		-	100%	20.1[5.76,34.44]
Subtotal ***	96		48			→	100%	20.1[5.76,34.44]
Heterogeneity: Not applicable								
Test for overall effect: Z=2.75(P=0.03	1)							
54.8.2 Caesarean								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
Test for overall effect: Not applicabl	e							
Total ***	96		48			•	100%	20.1[5.76,34.44]
Heterogeneity: Not applicable								
Test for overall effect: Z=2.75(P=0.03	1)							
Test for subgroup differences: Not a	pplicable	!						
			Favour	s Ergometrine -100	-50	0 50	100 Favours Erg	ometrine plus oxytocin



Comparison 55. Misoprostol plus oxytocin vs Ergometrine (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.1 Vaginal birth	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies No. of participants		Statistical method	Effect size
18.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 56. Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.41 [0.68, 2.94]
2.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	1.41 [0.68, 2.94]
2.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1605	Risk Ratio (IV, Random, 95% CI)	0.89 [0.36, 2.19]
3.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	0.89 [0.36, 2.19]
3.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.39 [0.65, 2.99]
6.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	1.39 [0.65, 2.99]
6.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	1605	Risk Ratio (IV, Random, 95% CI)	1.48 [0.82, 2.69]
7.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	1.48 [0.82, 2.69]
7.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	802	Mean Difference (IV, Random, 95% CI)	-16.0 [-40.24, 8.24]
8.1 Vaginal birth	1	802	Mean Difference (IV, Random, 95% CI)	-16.0 [-40.24, 8.24]
8.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglo- bin	2	1605	Mean Difference (IV, Random, 95% CI)	-0.50 [-1.72, 0.72]
9.1 Vaginal birth	2	1605	Mean Difference (IV, Random, 95% CI)	-0.50 [-1.72, 0.72]
9.2 Caesarean	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.04 [0.23, 4.77]
12.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	1.04 [0.23, 4.77]
12.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

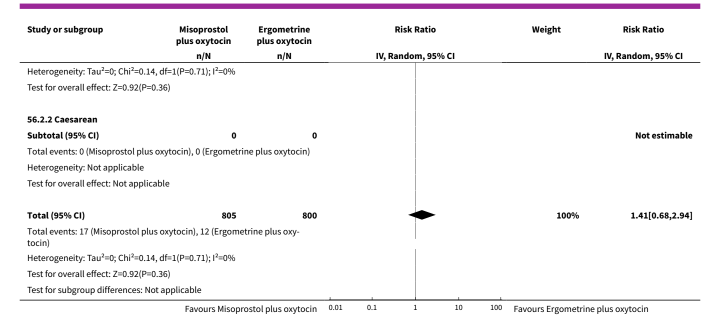


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Vaginal birth	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	1605	Risk Ratio (IV, Random, 95% CI)	2.95 [2.02, 4.29]
16.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	2.95 [2.02, 4.29]
16.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1605	Risk Ratio (IV, Random, 95% CI)	2.89 [1.51, 5.54]
17.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	2.89 [1.51, 5.54]
17.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	1605	Risk Ratio (IV, Random, 95% CI)	0.81 [0.46, 1.41]
18.1 Vaginal birth	2	1605	Risk Ratio (IV, Random, 95% CI)	0.81 [0.46, 1.41]
18.2 Caesarean	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Vaginal birth	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Caesarean	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 56.2. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
56.2.1 Vaginal birth									
Caliskan 2002	11/401	7/402			 			61.25%	1.58[0.62,4.02]
Caliskan 2003	6/404	5/398			-			38.75%	1.18[0.36,3.84]
Subtotal (95% CI)	805	800						100%	1.41[0.68,2.94]
Total events: 17 (Misoprostol tocin)	l plus oxytocin), 12 (Ergome	trine plus oxy-							
	Favours Misopro	ostol plus oxytocin	0.01	0.1	1	10	100	Favours Ergometrin	e plus oxytocin



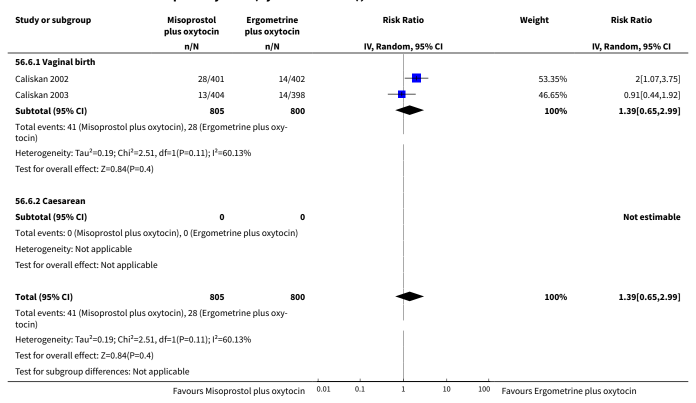


Analysis 56.3. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
56.3.1 Vaginal birth							
Caliskan 2002	4/401	4/402				42.22%	1[0.25,3.98]
Caliskan 2003	5/404	6/398				57.78%	0.82[0.25,2.67]
Subtotal (95% CI)	805	800		*		100%	0.89[0.36,2.19]
Total events: 9 (Misoprostol plus ox	ytocin), 10 (Ergometi	rine plus oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0.05, d	f=1(P=0.83); I ² =0%						
Test for overall effect: Z=0.25(P=0.8))						
56.3.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Ergometri	ne plus oxytocin)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicabl	le						
Total (95% CI)	805	800		*		100%	0.89[0.36,2.19]
Total events: 9 (Misoprostol plus ox	ytocin), 10 (Ergometi	rine plus oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0.05, d	f=1(P=0.83); I ² =0%						
Test for overall effect: Z=0.25(P=0.8))						
Test for subgroup differences: Not a	pplicable						
	Favours Misopr	ostol plus oxytocin	0.01	0.1 1 10	100 Fa	avours Ergometrine	plus oxytocin



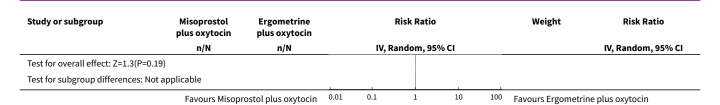
Analysis 56.6. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 6 PPH >= 500 mL.



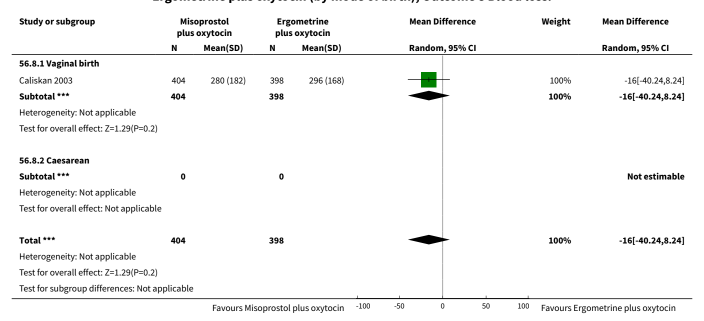
Analysis 56.7. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
56.7.1 Vaginal birth					
Caliskan 2002	17/401	9/402	 	55.55%	1.89[0.85,4.2]
Caliskan 2003	10/404	9/398	-	44.45%	1.09[0.45,2.67]
Subtotal (95% CI)	805	800	*	100%	1.48[0.82,2.69]
Total events: 27 (Misoprostol plus ox tocin)	kytocin), 18 (Ergomet	rine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.81, df	=1(P=0.37); I ² =0%				
Test for overall effect: Z=1.3(P=0.19)					
56.7.2 Caesarean					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus oxy	rtocin), 0 (Ergometrin	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	9				
Total (95% CI)	805	800	•	100%	1.48[0.82,2.69]
Total events: 27 (Misoprostol plus ox tocin)	kytocin), 18 (Ergomet	rine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.81, df	=1(P=0.37); I ² =0%				
	Favours Misopro	ostol plus oxytocin	0.01 0.1 1 10 100	Favours Ergometrin	e plus oxytocin





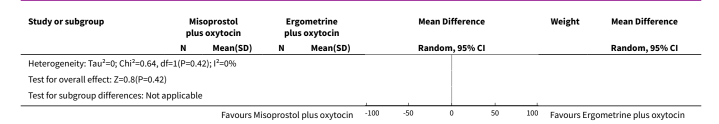
Analysis 56.8. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 8 Blood loss.



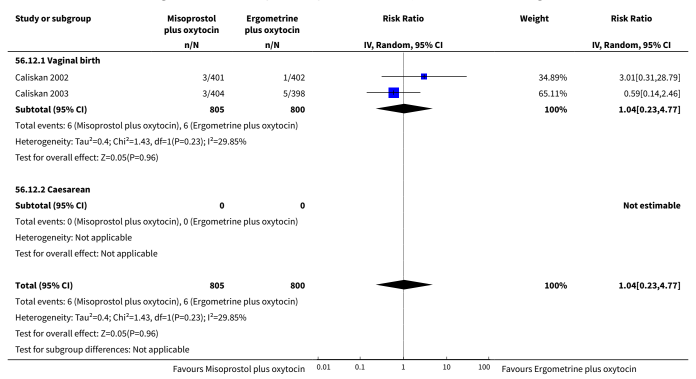
Analysis 56.9. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 9 Change in haemoglobin.

Study or subgroup		oprostol oxytocin	U	ometrine oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
56.9.1 Vaginal birth							
Caliskan 2002	401	15 (13)	402	15 (12)	•	50.02%	0[-1.73,1.73]
Caliskan 2003	404	14 (13)	398	15 (12)	•	49.98%	-1[-2.73,0.73]
Subtotal ***	805		800		•	100%	-0.5[-1.72,0.72]
Heterogeneity: Tau ² =0; Chi ² =0.	64, df=1(P=0.4	2); I ² =0%					
Test for overall effect: Z=0.8(P=	0.42)						
56.9.2 Caesarean							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not appl	icable						
Total ***	805		800			100%	-0.5[-1.72,0.72]
		Favours Mis	oprostol	plus oxytocin -100	-50 0 50	100 Favours Erg	ometrine plus oxytocin





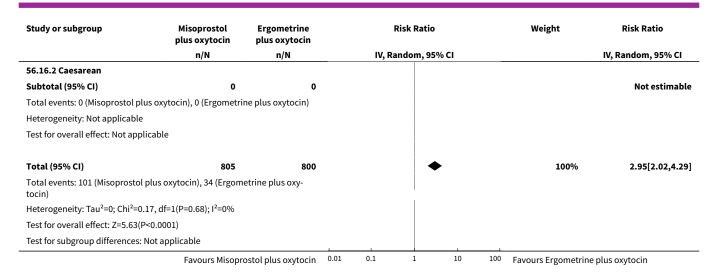
Analysis 56.12. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 12 Vomiting.



Analysis 56.16. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 16 Shivering.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin				Weight	Risk Ratio	
	n/N	n/N		IV, I	Random, 95% CI			IV, Random, 95% CI
56.16.1 Vaginal birth								
Caliskan 2002	52/401	19/402			-		55.1%	2.74[1.65,4.55]
Caliskan 2003	49/404	15/398			-		44.9%	3.22[1.84,5.64]
Subtotal (95% CI)	805	800			•		100%	2.95[2.02,4.29]
Total events: 101 (Misoprosto tocin)	l plus oxytocin), 34 (Ergom	etrine plus oxy-						
Heterogeneity: Tau ² =0; Chi ² =0	0.17, df=1(P=0.68); I ² =0%							
Test for overall effect: Z=5.63(P<0.0001)							
	Favours Misopro	ostol plus oxytocin	0.01	0.1	1 10	100	Favours Ergometrin	e plus oxytocin



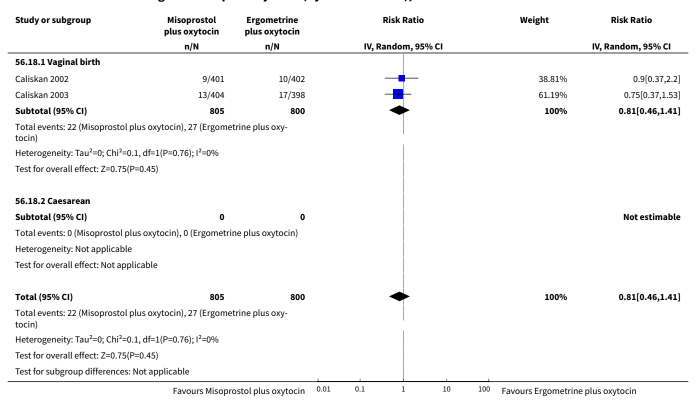


Analysis 56.17. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	6 CI		IV, Random, 95% CI
56.17.1 Vaginal birth							
Caliskan 2002	19/401	6/402		- 		51.12%	3.17[1.28,7.87]
Caliskan 2003	16/404	6/398		-		48.88%	2.63[1.04,6.64]
Subtotal (95% CI)	805	800		•	>	100%	2.89[1.51,5.54]
Total events: 35 (Misoprostol plus ox tocin)	ytocin), 12 (Ergome	trine plus oxy-					
Heterogeneity: Tau ² =0; Chi ² =0.08, df	=1(P=0.77); I ² =0%						
Test for overall effect: Z=3.21(P=0)							
56.17.2 Caesarean							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Misoprostol plus oxy	tocin), 0 (Ergometri	ne plus oxytocin)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Total (050), CI\	805	800			_	100%	2 90[1 51 5 54]
Total (95% CI)						100%	2.89[1.51,5.54]
Total events: 35 (Misoprostol plus ox tocin)	ytocin), 12 (Ergome	trine plus oxy-					
Heterogeneity: Tau ² =0; Chi ² =0.08, df=	=1(P=0.77); I ² =0%						
Test for overall effect: Z=3.21(P=0)							
Test for subgroup differences: Not ap	plicable						
	Favours Misopr	ostol plus oxytocin	0.01	0.1 1	10 100	Favours Ergometrine	e plus oxvtocin



Analysis 56.18. Comparison 56 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by mode of birth), Outcome 18 Diarrhoea.



Comparison 57. Oxytocin vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	3	1920	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	130	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	1	1569	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Both community and hospital setting	1	221	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	11	8782	Risk Ratio (IV, Random, 95% CI)	0.61 [0.51, 0.72]
2.1 Hospital setting	8	5306	Risk Ratio (IV, Random, 95% CI)	0.64 [0.52, 0.79]
2.2 Community setting	2	3255	Risk Ratio (IV, Random, 95% CI)	0.47 [0.22, 0.98]
2.3 Both community and hospital setting	1	221	Risk Ratio (IV, Random, 95% CI)	0.80 [0.34, 1.87]
3 Blood transfusion	8	6717	Risk Ratio (IV, Random, 95% CI)	0.75 [0.51, 1.12]
3.1 Hospital setting	6	4810	Risk Ratio (IV, Random, 95% CI)	0.71 [0.44, 1.14]



Outcome or subgroup title	group title No. of studies No. of partic pants		Statistical method	Effect size	
3.2 Community setting	1	1686	Risk Ratio (IV, Random, 95% CI)	0.82 [0.36, 1.88]	
3.3 Both community and hospital setting	1	221	Risk Ratio (IV, Random, 95% CI)	1.22 [0.21, 7.16]	
4 Severe maternal morbidity: intensive care admissions	1	1950	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.1 Hospital setting	1	1950	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	9	7021	Risk Ratio (IV, Random, 95% CI)	0.61 [0.52, 0.71]	
6.1 Hospital setting	6	3545	Risk Ratio (IV, Random, 95% CI)	0.54 [0.46, 0.63]	
6.2 Community setting	2	3255	Risk Ratio (IV, Random, 95% CI)	0.64 [0.43, 0.95]	
6.3 Both community and hospital setting	1	221	Risk Ratio (IV, Random, 95% CI)	0.83 [0.57, 1.22]	
7 Additional uterotonics	8	5047	Risk Ratio (IV, Random, 95% CI)	0.43 [0.32, 0.58]	
7.1 Hospital setting	6	3154	Risk Ratio (IV, Random, 95% CI)	0.38 [0.27, 0.53]	
7.2 Community setting	1	1672	Risk Ratio (IV, Random, 95% CI)	0.40 [0.31, 0.51]	
7.3 Both community and hospital setting	1	221	Risk Ratio (IV, Random, 95% CI)	0.99 [0.55, 1.78]	
8 Blood loss	8	3521	Mean Difference (IV, Random, 95% CI)	-118.52 [-141.40, -95.64]	
8.1 Hospital setting	7	3300	Mean Difference (IV, Random, 95% CI)	-122.08 [-145.37, -98.79]	
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8.3 Both community and hos- pital setting	1	221	Mean Difference (IV, Random, 95% CI)	-21.0 [-142.93, 100.93]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
9 Change in haemoglobin	5	2304	Mean Difference (IV, Random, 95% CI)	-2.68 [-4.47, -0.89]	
9.1 Hospital setting	5	2304	Mean Difference (IV, Random, 95% CI)	-2.68 [-4.47, -0.89]	
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.3 Both community and hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	1	1540	Risk Ratio (IV, Random, 95% CI)	1.00 [0.95, 1.05]	
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Community setting	1	1540	Risk Ratio (IV, Random, 95% CI)	1.00 [0.95, 1.05]	
10.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	3	1788	Risk Ratio (IV, Random, 95% CI)	0.82 [0.47, 1.42]	
11.1 Hospital setting	2	126	Risk Ratio (IV, Random, 95% CI)	0.27 [0.03, 2.10]	
11.2 Community setting	1	1662	Risk Ratio (IV, Random, 95% CI)	0.89 [0.51, 1.58]	
11.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
12 Vomiting	3	2150	Risk Ratio (IV, Random, 95% CI)	1.40 [0.44, 4.41]	
12.1 Hospital setting	2	490	Risk Ratio (IV, Random, 95% CI)	1.12 [0.07, 17.83]	
12.2 Community setting	1	1660	Risk Ratio (IV, Random, 95% CI)	1.46 [0.41, 5.17]	
12.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
13 Headache	2	1704	Risk Ratio (IV, Random, 95% CI)	1.56 [0.52, 4.74]	
13.1 Hospital setting	1	51	Risk Ratio (IV, Random, 95% CI)	6.74 [0.37, 124.21]	
13.2 Community setting	1	1653	Risk Ratio (IV, Random, 95% CI)	1.26 [0.63, 2.51]	
13.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14 Abdominal pain	1	1665	Risk Ratio (IV, Random, 95% CI)	0.89 [0.80, 1.00]	
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14.2 Community setting	1	1665	Risk Ratio (IV, Random, 95% CI)	0.89 [0.80, 1.00]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	1	416	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.3 Both community and hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal sense of well-being	1		Risk Ratio (IV, Random, 95% CI)	Totals not select- ed
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting - Women's perceptions of well- being at 3 months postpar- tum: Less energy than before birth	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.3 Community setting - Women's perceptions of well- being at 3 months postpar- tum: Experiencing (some) fa- tigue	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

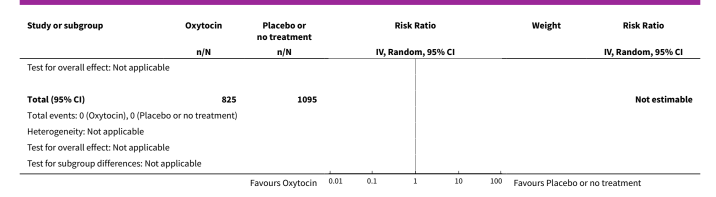


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
19.4 Both community and hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not select- ed
20.1 Hospital setting - Did management influence positively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Hospital setting - Did management influence negatively the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Hospital setting - Did management make no difference in the childbirth experiences of the mothers?	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.4 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.5 Both community and hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 57.1. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 1 Death.

Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI		IV, Random, 95% CI
57.1.1 Hospital setting						
Jerbi 2007	0/65	0/65				Not estimable
Subtotal (95% CI)	65	65				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
57.1.2 Community setting						
Stanton 2013	0/682	0/887				Not estimable
Subtotal (95% CI)	682	887				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
57.1.3 Both community and hospita	l setting					
de Groot 1996	0/78	0/143				Not estimable
Subtotal (95% CI)	78	143				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
		Favours Oxytocin	0.01	0.1 1 10	100 Favours Placebo	or no treatment



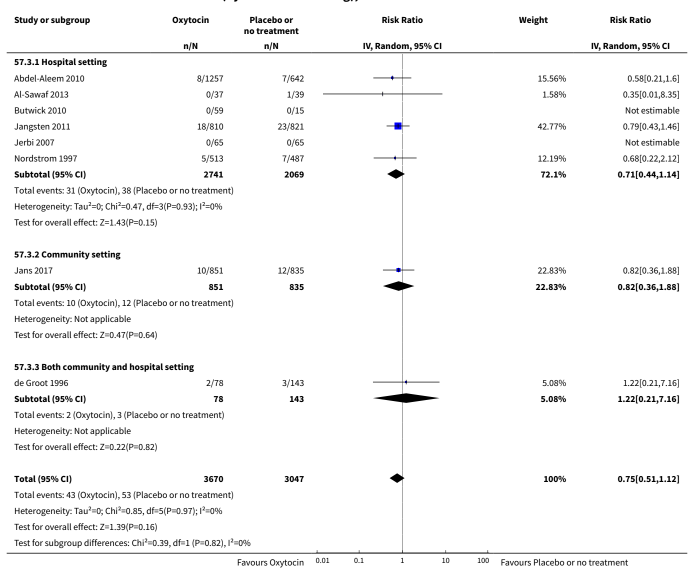


Analysis 57.2. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
57.2.1 Hospital setting					
Abdel-Aleem 2010	4/1291	4/659		1.5%	0.51[0.13,2.03
Al-Sawaf 2013	1/37	6/39	•	0.67%	0.18[0.02,1.39
Benchimol 2001	12/196	13/220		4.96%	1.04[0.48,2.22
Jangsten 2011	82/810	138/821	-	43.96%	0.6[0.47,0.78
Jerbi 2007	0/65	0/65			Not estimabl
Nordstrom 1997	32/513	43/487		14.79%	0.71[0.45,1.1
Poeschmann 1991	2/28	3/24		0.99%	0.57[0.1,3.14
Rosseland 2013	0/26	0/25			Not estimabl
Subtotal (95% CI)	2966	2340	◆	66.86%	0.64[0.52,0.79
Total events: 133 (Oxytocin), 20	7 (Placebo or no treatme	nt)			
Heterogeneity: Tau²=0; Chi²=3.5	7, df=5(P=0.61); I ² =0%				
Test for overall effect: Z=4.24(P<	:0.0001)				
57.2.2 Community setting					
Jans 2017	54/851	99/835		28.45%	0.54[0.39,0.74
Stanton 2013	1/682	8/887 -		0.67%	0.16[0.02,1.3
Subtotal (95% CI)	1533	1722	•	29.12%	0.47[0.22,0.98
Total events: 55 (Oxytocin), 107	(Placebo or no treatmen	t)			
Heterogeneity: Tau²=0.14; Chi²=	1.24, df=1(P=0.27); I ² =19.	09%			
Test for overall effect: Z=2.01(P=	:0.04)				
57.2.3 Both community and ho	ospital setting				
de Groot 1996	7/78	16/143		4.02%	0.8[0.34,1.8]
Subtotal (95% CI)	78	143	•	4.02%	0.8[0.34,1.87
Total events: 7 (Oxytocin), 16 (P	lacebo or no treatment)				
Heterogeneity: Tau²=0; Chi²=0, c	df=0(P<0.0001); I ² =100%				
Test for overall effect: Z=0.51(P=	:0.61)				
Total (95% CI)	4577	4205	•	100%	0.61[0.51,0.72
Total events: 195 (Oxytocin), 330	0 (Placebo or no treatme	nt)			·
Heterogeneity: Tau²=0; Chi²=6.3	7, df=8(P=0.61); I ² =0%				
Test for overall effect: Z=5.77(P<					
Test for subgroup differences: C	•	!=0%			



Analysis 57.3. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 3 Blood transfusion.



Analysis 57.4. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Oxytocin	n Placebo or no treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	CI			IV, Random, 95% CI
57.4.1 Hospital setting									
Abdel-Aleem 2010	0/1291	0/659							Not estimable
Subtotal (95% CI)	1291	659							Not estimable
Total events: 0 (Oxytocin), 0 (Placeb	o or no treatment)								
Heterogeneity: Not applicable									
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or	no treatment

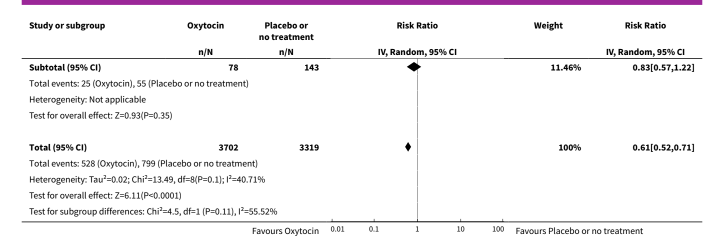


Study or subgroup	Oxytocin	Placebo or no treatment	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	om, 95% CI		IV, Random, 95% CI
Test for overall effect: Not applicable	!					
57.4.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	!					
57.4.3 Both community and hospit	al setting					
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	o or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	•					
Total (95% CI)	1291	659				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	o or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	!					
Test for subgroup differences: Not ap	plicable					
* *		Favours Oxytocin	0.01 0.1	1 10	100 Favours Placebo or	no tre

Analysis 57.6. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
57.6.1 Hospital setting					
Abdel-Aleem 2010	63/1291	65/659	-+ -	13.7%	0.49[0.35,0.69]
Al-Sawaf 2013	2/37	8/39		1.12%	0.26[0.06,1.16]
Benchimol 2001	29/196	60/220		10.87%	0.54[0.36,0.81]
Nordstrom 1997	104/513	175/487	-	21.6%	0.56[0.46,0.7]
Poeschmann 1991	7/28	10/24		3.58%	0.6[0.27,1.33]
Rosseland 2013	4/26	8/25		2.09%	0.48[0.17,1.4]
Subtotal (95% CI)	2091	1454	•	52.95%	0.54[0.46,0.63]
Total events: 209 (Oxytocin), 32	6 (Placebo or no treatmer	t)			
Heterogeneity: Tau ² =0; Chi ² =1. ⁴	45, df=5(P=0.92); I ² =0%				
Test for overall effect: Z=7.75(P-	<0.0001)				
57.6.2 Community setting					
Jans 2017	276/851	369/835	•	28.46%	0.73[0.65,0.83]
Stanton 2013	18/682	49/887		7.13%	0.48[0.28,0.81]
Subtotal (95% CI)	1533	1722	•	35.59%	0.64[0.43,0.95]
Total events: 294 (Oxytocin), 41	.8 (Placebo or no treatmer	t)			
Heterogeneity: Tau ² =0.05; Chi ² =	=2.38, df=1(P=0.12); I ² =58.0	03%			
Test for overall effect: Z=2.23(P	=0.03)				
F7 C 2 Dath as	ospital setting				
57.6.3 Both community and h					





Analysis 57.7. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
57.7.1 Hospital setting						
Abdel-Aleem 2010	41/1260	55/641		19.26%	0.38[0.26,0.56	
Al-Sawaf 2013	2/37	8/39		3.59%	0.26[0.06,1.16]	
Butwick 2010	8/59	7/15		8.83%	0.29[0.13,0.67	
Nordstrom 1997	40/513	67/487		19.97%	0.57[0.39,0.82	
Poeschmann 1991	0/28	2/24		0.97%	0.17[0.01,3.42	
Rosseland 2013	5/26	23/25		9.51%	0.21[0.09,0.46	
Subtotal (95% CI)	1923	1231	•	62.14%	0.38[0.27,0.53	
Total events: 96 (Oxytocin), 162 (Placebo or no treatment)					
Heterogeneity: Tau²=0.05; Chi²=7	7.13, df=5(P=0.21); I ² =29.8	5%				
Test for overall effect: Z=5.69(P<0	0.0001)					
57.7.2 Community setting						
Jans 2017	79/842	195/830	-	24.2%	0.4[0.31,0.51	
Subtotal (95% CI)	842	830	•	24.2%	0.4[0.31,0.51	
Total events: 79 (Oxytocin), 195 (Placebo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Z=7.4(P<0.	0001)					
57.7.3 Both community and ho	spital setting					
de Groot 1996	14/78	26/143	-	13.67%	0.99[0.55,1.78	
Subtotal (95% CI)	78	143	*	13.67%	0.99[0.55,1.78	
Total events: 14 (Oxytocin), 26 (P	lacebo or no treatment)					
Heterogeneity: Tau²=0; Chi²=0, di	f=0(P<0.0001); I ² =100%					
Test for overall effect: Z=0.04(P=0	0.97)					
Total (95% CI)	2843	2204	•	100%	0.43[0.32,0.58	
Total events: 189 (Oxytocin), 383	(Placebo or no treatment)				
Heterogeneity: Tau²=0.08; Chi²=1	.5.29, df=7(P=0.03); I ² =54.2	22%				
Test for overall effect: Z=5.53(P<0	0.0001)					
Test for subgroup differences: Ch	ni ² =8.58, df=1 (P=0.01), I ² =	76.7%				



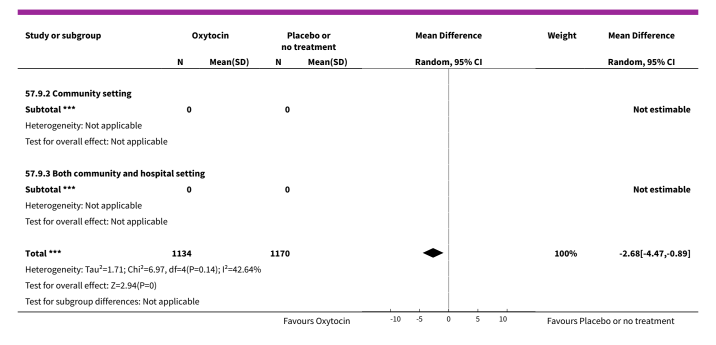
Analysis 57.8. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	0	xytocin		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
57.8.1 Hospital setting							
Al-Sawaf 2013	37	314.7 (94.6)	39	438.6 (130.2)		20.14%	-123.9[-174.88,-72.92]
Benchimol 2001	196	278 (254)	220	382 (269.5)		20.66%	-104[-154.32,-53.68]
Butwick 2010	59	750.8 (160.9)	15	800 (256)		2.83%	-49.22[-185.12,86.68]
Jangsten 2011	810	535 (414.5)	821	680 (486.7)		27.21%	-145[-188.85,-101.15]
Nordstrom 1997	513	409 (345)	487	527 (412)		23.46%	-118[-165.23,-70.77]
Poeschmann 1991	28	374 (279)	24	548 (376)	—	1.57%	-174[-356.51,8.51]
Rosseland 2013	26	841 (556)	25	853 (518)	+	0.6%	-12[-306.8,282.8]
Subtotal ***	1669		1631		•	96.48%	-122.08[-145.37,-98.79]
Heterogeneity: Tau ² =0; Chi ² =3.53	, df=6(P=0.7	4); I ² =0%					
Test for overall effect: Z=10.27(P<	0.0001)						
57.8.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applica	ible						
57.8.3 Both community and hos	pital settir	g					
de Groot 1996	78	499 (454)	143	520 (419)		3.52%	-21[-142.93,100.93]
Subtotal ***	78		143			3.52%	-21[-142.93,100.93]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.34(P=0	.74)						
			1774		•	100%	-118.52[-141.4,-95.64]
Total ***	1747		1117		•		
Total *** Heterogeneity: Tau ² =0; Chi ² =6.08		3); I²=0%	1//4				- , -
	, df=7(P=0.5	3); I ² =0%	1114				

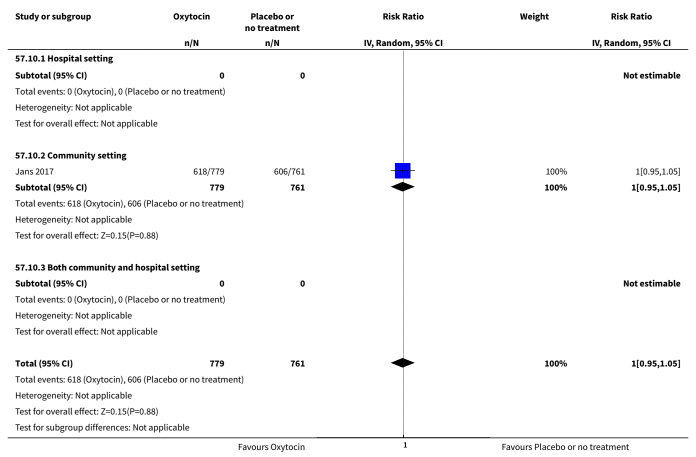
Analysis 57.9. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	O	kytocin		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
57.9.1 Hospital setting							
Al-Sawaf 2013	37	12 (9)	39	13 (6)		17.27%	-1[-4.46,2.46]
Benchimol 2001	196	6.7 (20.1)	220	10.2 (23.9)		13.08%	-3.5[-7.73,0.73]
Jangsten 2011	810	10.6 (13.4)	821	13.5 (13.2)	-	38.86%	-2.95[-4.24,-1.66]
Jerbi 2007	65	5.1 (12.3)	65	12 (13)		12.54%	-6.9[-11.25,-2.55]
Rosseland 2013	26	8.2 (6.7)	25	8.4 (5.3)		18.26%	-0.2[-3.51,3.11]
Subtotal ***	1134		1170		•	100%	-2.68[-4.47,-0.89]
Heterogeneity: Tau ² =1.71; Ch	ni ² =6.97, df=4(P=	0.14); I ² =42.64%					
Test for overall effect: Z=2.94	(P=0)						
			Fav	ours Oxytocin	-10 -5 0 5 10	Favours Pla	cebo or no treatment



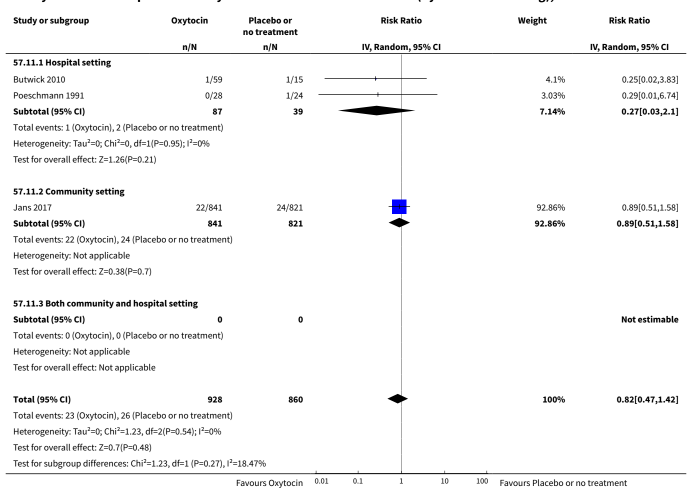


Analysis 57.10. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 10 Breastfeeding.





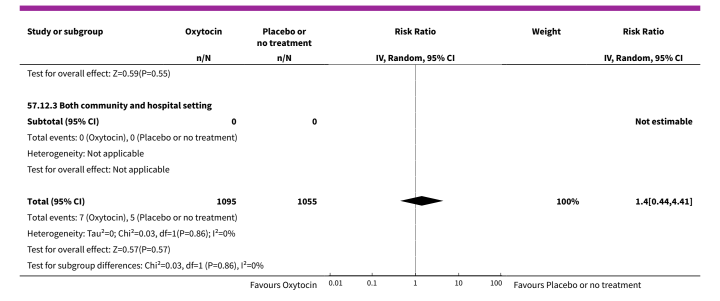
Analysis 57.11. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 11 Nausea.



Analysis 57.12. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Oxytocin	Placebo or no treatment		Ris	sk Ratio	Weight	Risk Ratio	
	n/N	n/N		IV, Ran	dom, 95% CI		IV, Random, 95% CI	
57.12.1 Hospital setting								
Benchimol 2001	1/196	1/220			•	17.23%	1.12[0.07,17.83]	
Butwick 2010	0/59	0/15					Not estimable	
Subtotal (95% CI)	255	235				17.23%	1.12[0.07,17.83]	
Total events: 1 (Oxytocin), 1 (Placeb	o or no treatment)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.08(P=0.93	3)							
57.12.2 Community setting								
Jans 2017	6/840	4/820		_	- - 	82.77%	1.46[0.41,5.17]	
Subtotal (95% CI)	840	820		-		82.77%	1.46[0.41,5.17]	
Total events: 6 (Oxytocin), 4 (Placeb	o or no treatment)							
Heterogeneity: Not applicable								
		Favours Oxytocin	0.01	0.1	1 10	100 Favours Placebo or r	no treatment	



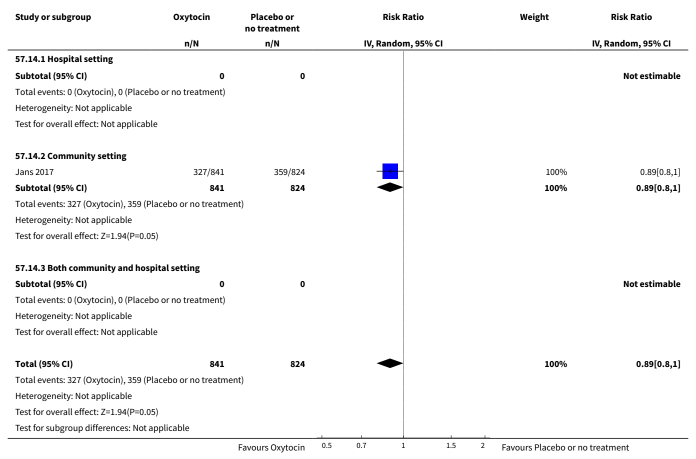


Analysis 57.13. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 13 Headache.

Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
3/26	0/25	+	13.04%	6.74[0.37,124.21]
26	25		13.04%	6.74[0.37,124.21]
bo or no treatment)				
2)				
18/836	14/817	-	86.96%	1.26[0.63,2.51]
836	817	•	86.96%	1.26[0.63,2.51]
cebo or no treatment)				
52)				
spital setting				
0	0			Not estimable
bo or no treatment)				
ble				
862	842		100%	1.56[0.52,4.74]
cebo or no treatment)				
1, df=1(P=0.27); l ² =17.26	6%			
43)				
=1.21, df=1 (P=0.27), I ² =	17.26%			
2	n/N 3/26 26 bb or no treatment) 2) 18/836 836 scebo or no treatment) 52) spital setting 0 bb or no treatment) ole 862 scebo or no treatment) 21, df=1(P=0.27); l ² =17.26 43)	18/836 14/817 25 26 25 25 26 25 26 25 26 25 26 27 26 27 27 27 27 27	no treatment n/N	no treatment n/N n/N 1V, Random, 95% CI 3/26 0/25 26 25 13.04% bb or no treatment) 2) 18/836 14/817 36.96% 836 817 86.96% scebo or no treatment) 52) spital setting 0 0 0 bb or no treatment) ple 862 842 ccebo or no treatment) 100% ccebo or no treatment) 21, df=1(P=0.27); l²=17.26% 43)



Analysis 57.14. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 14 Abdominal pain.



Analysis 57.16. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
57.16.1 Hospital setting					
Benchimol 2001	0/196	0/220			Not estimable
Subtotal (95% CI)	196	220			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
57.16.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
57.16.3 Both community and hospi	tal setting				
Subtotal (95% CI)	0	0			Not estimable
		Favours Oxytocin 0.0	01 0.1 1 10	100 Favours Placebo or	no treatment



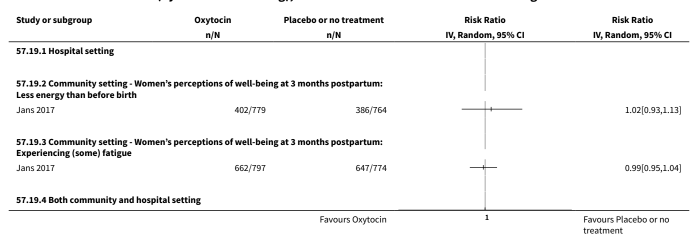
Study or subgroup	Oxytocin	Placebo or no treatment		Risk Ratio		Ratio V		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Total (95% CI)	196	220							Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Test for subgroup differences: Not app	licable								
		Favours Oxytocin	0.01	0.1	1	10	100	Favours Placebo or I	no treatment

Analysis 57.17. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 17 Fever.

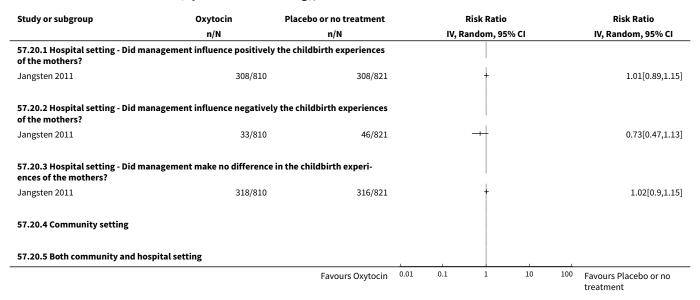
Study or subgroup	Oxytocin	Placebo or no treatment			Weight	Risk Ratio
	n/N	n/N	IV, Rando	n, 95% CI		IV, Random, 95% CI
57.17.1 Hospital setting						
Benchimol 2001	0/196	0/220				Not estimable
Subtotal (95% CI)	196	220				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
57.17.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
57.17.3 Both community and hospit	tal setting					
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	196	220				Not estimable
Total events: 0 (Oxytocin), 0 (Placebo	or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not ap	plicable					
		Favours Oxytocin	0.01 0.1	10 100	Favours Placebo or	no treatment



Analysis 57.19. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 19 Maternal sense of well-being.



Analysis 57.20. Comparison 57 Oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 20 Maternal satisfaction.



Comparison 58. Carbetocin vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.1 Hospital setting	1	50	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	50	Risk Ratio (IV, Random, 95% CI)	0.75 [0.30, 1.85]
6.1 Hospital setting	1	50	Risk Ratio (IV, Random, 95% CI)	0.75 [0.30, 1.85]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	169	Risk Ratio (IV, Random, 95% CI)	0.19 [0.12, 0.32]
7.1 Hospital setting	2	169	Risk Ratio (IV, Random, 95% CI)	0.19 [0.12, 0.32]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	50	Mean Difference (IV, Random, 95% CI)	-274.0 [-591.60, 43.60]
8.1 Hospital setting	1	50	Mean Difference (IV, Random, 95% CI)	-274.0 [-591.60, 43.60]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	1	50	Mean Difference (IV, Random, 95% CI)	-3.40 [-7.23, 0.43]
9.1 Hospital setting	1	50	Mean Difference (IV, Random, 95% CI)	-3.40 [-7.23, 0.43]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
13.1 Hospital setting	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

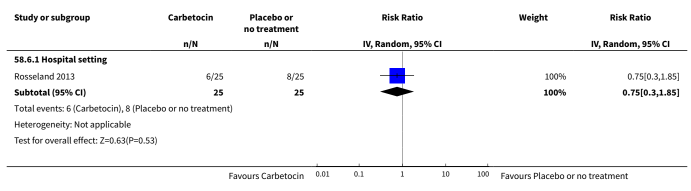


Outcome or subgroup title	No. of studies No. of participants		Statistical method	Effect size
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

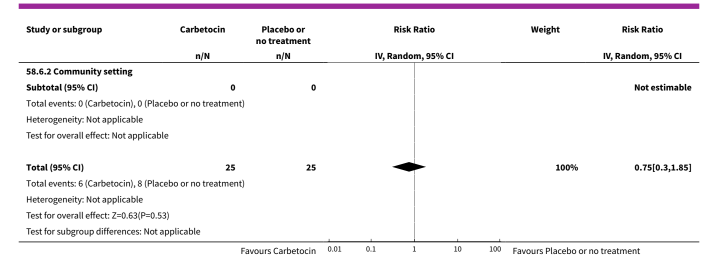
Analysis 58.2. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Placebo or no treatment	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
58.2.1 Hospital setting						
Rosseland 2013	0/25	0/25				Not estimable
Subtotal (95% CI)	25	25				Not estimable
Total events: 0 (Carbetocin), 0 (Placeb	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
58.2.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Placeb	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	25	25				Not estimable
Total events: 0 (Carbetocin), 0 (Placeb	oo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not app	plicable					
	Fa	vours Carbetocin	0.01 0.1	1 10 10	Favours Placebo o	r no treatment

Analysis 58.6. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.







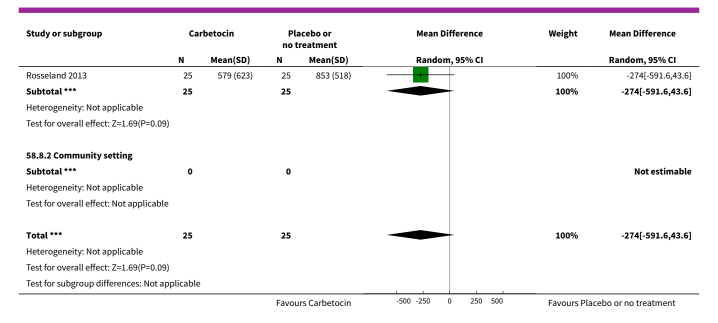
Analysis 58.7. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Carbetocin	Placebo or no treatment	Risk	Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI	
58.7.1 Hospital setting							
Barton 1996	8/62	41/57	-		58.55%	0.18[0.09,0.35]	
Rosseland 2013	5/25	23/25			41.45%	0.22[0.1,0.48]	
Subtotal (95% CI)	87	82	•		100%	0.19[0.12,0.32]	
Total events: 13 (Carbetocin), 64 (Pla	acebo or no treatmen	t)					
Heterogeneity: Tau ² =0; Chi ² =0.13, df	=1(P=0.72); I ² =0%						
Test for overall effect: Z=6.29(P<0.00	01)						
58.7.2 Community setting							
Subtotal (95% CI)	0	0				Not estimable	
Total events: 0 (Carbetocin), 0 (Place	ebo or no treatment)						
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	2						
Total (95% CI)	87	82	•		100%	0.19[0.12,0.32]	
Total events: 13 (Carbetocin), 64 (Pla	acebo or no treatmen	t)					
Heterogeneity: Tau ² =0; Chi ² =0.13, df	=1(P=0.72); I ² =0%						
Test for overall effect: Z=6.29(P<0.00	01)						
Test for subgroup differences: Not a	oplicable						
	Fa	avours Carbetocin	0.01 0.1	1 10	100 Favours Placebo or	no treatment	

Analysis 58.8. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Ca	arbetocin	Placebo or no treatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
58.8.1 Hospital setting							
			Favo	urs Carbetocin	-500 -250 0 250 500	Favours Pla	cebo or no treatment





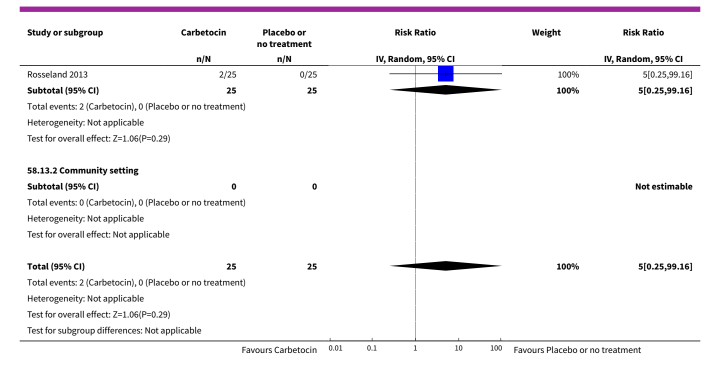
Analysis 58.9. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Ca	rbetocin		acebo or reatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI		Random, 95% CI
58.9.1 Hospital setting								
Rosseland 2013	25	5 (8.2)	25	8.4 (5.3)		+	100%	-3.4[-7.23,0.43]
Subtotal ***	25		25			•	100%	-3.4[-7.23,0.43]
Heterogeneity: Not applicable								
Test for overall effect: Z=1.74(P=0.08)								
58.9.2 Community setting								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total ***	25		25			•	100%	-3.4[-7.23,0.43]
Heterogeneity: Not applicable								
Test for overall effect: Z=1.74(P=0.08)								
Test for subgroup differences: Not ap	plicable							
			Favou	ırs Carbetocin	100 -50	0 50	100 Favours Pla	cebo or no treatment

Analysis 58.13. Comparison 58 Carbetocin vs Placebo or no treatment (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Carbetocin	Placebo or no treatment		Risk Ratio				Weight Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI		IV, Random, 95% CI
58.13.1 Hospital setting				1				
		Favours Carbetocin	0.01	0.1	1	10	100	Favours Placebo or no treatment





Comparison 59. Misoprostol vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3497	Risk Ratio (IV, Random, 95% CI)	1.00 [0.10, 9.59]
1.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	1.00 [0.10, 9.59]
2 PPH >= 1000 mL	8	5467	Risk Ratio (IV, Random, 95% CI)	0.73 [0.56, 0.95]
2.1 Hospital setting	5	2114	Risk Ratio (IV, Random, 95% CI)	0.85 [0.63, 1.15]
2.2 Community setting	3	3353	Risk Ratio (IV, Random, 95% CI)	0.59 [0.39, 0.88]
3 Blood transfusion	6	3134	Risk Ratio (IV, Random, 95% CI)	0.46 [0.15, 1.47]
3.1 Hospital setting	5	1514	Risk Ratio (IV, Random, 95% CI)	0.77 [0.19, 3.10]
3.2 Community setting	1	1620	Risk Ratio (IV, Random, 95% CI)	0.14 [0.02, 1.15]
4 Severe maternal morbidity: intensive care admissions	1	1620	Risk Ratio (IV, Random, 95% CI)	1.00 [0.14, 7.05]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	1	1620	Risk Ratio (IV, Random, 95% CI)	1.00 [0.14, 7.05]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	7	4047	Risk Ratio (IV, Random, 95% CI)	0.75 [0.59, 0.94]
6.1 Hospital setting	4	694	Risk Ratio (IV, Random, 95% CI)	0.60 [0.29, 1.25]
6.2 Community setting	3	3353	Risk Ratio (IV, Random, 95% CI)	0.73 [0.56, 0.96]
7 Additional uterotonics	8	3746	Risk Ratio (IV, Random, 95% CI)	0.67 [0.52, 0.87]
7.1 Hospital setting	7	2126	Risk Ratio (IV, Random, 95% CI)	0.68 [0.52, 0.88]
7.2 Community setting	1	1620	Risk Ratio (IV, Random, 95% CI)	0.50 [0.12, 1.98]
8 Blood loss	8	4146	Mean Difference (IV, Random, 95% CI)	-42.07 [-52.47, -31.68]
8.1 Hospital setting	5	794	Mean Difference (IV, Random, 95% CI)	-41.82 [-61.16, -22.49]
8.2 Community setting	3	3352	Mean Difference (IV, Random, 95% CI)	-43.79 [-58.09, -29.49]
9 Change in haemoglobin	5	2334	Mean Difference (IV, Random, 95% CI)	-2.53 [-3.53, -1.52]
9.1 Hospital setting	3	573	Mean Difference (IV, Random, 95% CI)	-2.05 [-4.30, 0.19]
9.2 Community setting	2	1761	Mean Difference (IV, Random, 95% CI)	-2.12 [-3.46, -0.77]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3997	Risk Ratio (IV, Random, 95% CI)	1.18 [0.78, 1.78]
11.1 Hospital setting	1	600	Risk Ratio (IV, Random, 95% CI)	5.00 [0.59, 42.54]
11.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	1.12 [0.74, 1.70]
12 Vomiting	6	4949	Risk Ratio (IV, Random, 95% CI)	1.41 [0.92, 2.16]
12.1 Hospital setting	3	1552	Risk Ratio (IV, Random, 95% CI)	2.79 [0.85, 9.15]
12.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	1.27 [0.80, 2.01]
13 Headache	1	1116	Risk Ratio (IV, Random, 95% CI)	0.94 [0.32, 2.77]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	1	1116	Risk Ratio (IV, Random, 95% CI)	0.94 [0.32, 2.77]

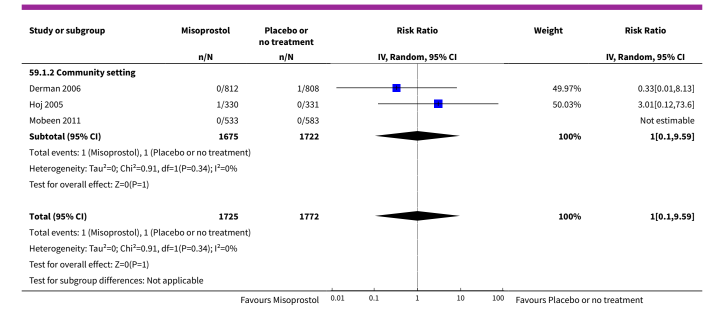


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14 Abdominal pain	4	1894	Risk Ratio (IV, Random, 95% CI)	0.98 [0.29, 3.37]
14.1 Hospital setting	4	1894	Risk Ratio (IV, Random, 95% CI)	0.98 [0.29, 3.37]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	5286	Risk Ratio (IV, Random, 95% CI)	2.94 [2.38, 3.64]
16.1 Hospital setting	6	1889	Risk Ratio (IV, Random, 95% CI)	3.55 [2.13, 5.90]
16.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	2.71 [2.33, 3.15]
17 Fever	5	4396	Risk Ratio (IV, Random, 95% CI)	4.09 [2.01, 8.32]
17.1 Hospital setting	2	999	Risk Ratio (IV, Random, 95% CI)	6.38 [4.01, 10.14]
17.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	2.87 [0.90, 9.18]
18 Diarrhoea	6	4791	Risk Ratio (IV, Random, 95% CI)	2.80 [1.21, 6.49]
18.1 Hospital setting	3	1394	Risk Ratio (IV, Random, 95% CI)	1.0 [0.06, 15.91]
18.2 Community setting	3	3397	Risk Ratio (IV, Random, 95% CI)	3.11 [1.28, 7.51]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 59.1. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95%	6 CI			IV, Random, 95% CI
59.1.1 Hospital setting									
Supe 2016	0/50	0/50							Not estimable
Subtotal (95% CI)	50	50							Not estimable
Total events: 0 (Misoprostol), 0 (Pla	cebo or no treatment)							
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	le								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or i	no treatment



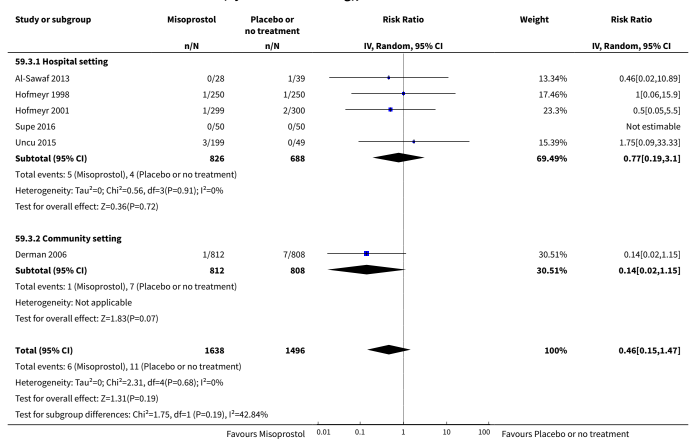


Analysis 59.2. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.

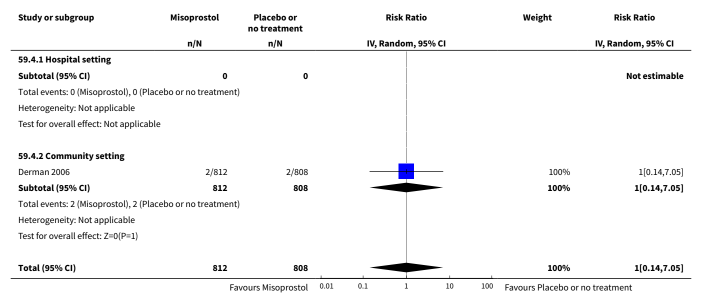
Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
59.2.1 Hospital setting					
Al-Sawaf 2013	2/28	6/39		2.78%	0.46[0.1,2.13]
Bamigboye 1998a	13/270	19/272	-+	11.94%	0.69[0.35,1.37]
Benchimol 2001	16/186	13/220	+-	11.37%	1.46[0.72,2.95]
Hofmeyr 1998	15/250	23/250	-+	13.82%	0.65[0.35,1.22]
Hofmeyr 2001	27/300	29/299	-	19.58%	0.93[0.56,1.53]
Subtotal (95% CI)	1034	1080	*	59.49%	0.85[0.63,1.15]
Total events: 73 (Misoprostol), 9	90 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0; Chi ² =4,	df=4(P=0.41); I ² =0.1%				
Test for overall effect: Z=1.03(P	=0.3)				
59.2.2 Community setting					
Derman 2006	2/812	10/808		2.82%	0.2[0.04,0.91]
Hoj 2005	37/330	56/331	-	27.59%	0.66[0.45,0.98]
Mobeen 2011	10/514	19/558		10.1%	0.57[0.27,1.22]
Subtotal (95% CI)	1656	1697	•	40.51%	0.59[0.39,0.88]
Total events: 49 (Misoprostol), 8	85 (Placebo or no treatme	nt)			
Heterogeneity: Tau ² =0.02; Chi ² =	=2.3, df=2(P=0.32); I ² =13.2	1%			
Test for overall effect: Z=2.58(P=	=0.01)				
Total (95% CI)	2690	2777	•	100%	0.73[0.56,0.95]
Total events: 122 (Misoprostol)	, 175 (Placebo or no treatr	nent)			
Heterogeneity: Tau ² =0.02; Chi ² =	=8.52, df=7(P=0.29); l ² =17.	89%			
Test for overall effect: Z=2.36(P	=0.02)				
Test for subgroup differences: 0	Chi ² =2.11, df=1 (P=0.15), I ²	=52.53%			
	Fa	vours Misoprostol 0.01	0.1 1 10	100 Favours Placebo or	no treatment



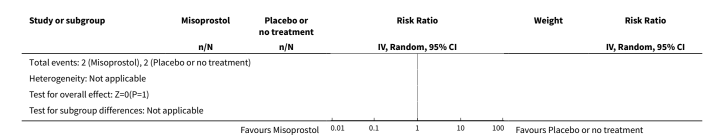
Analysis 59.3. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 3 Blood transfusion.



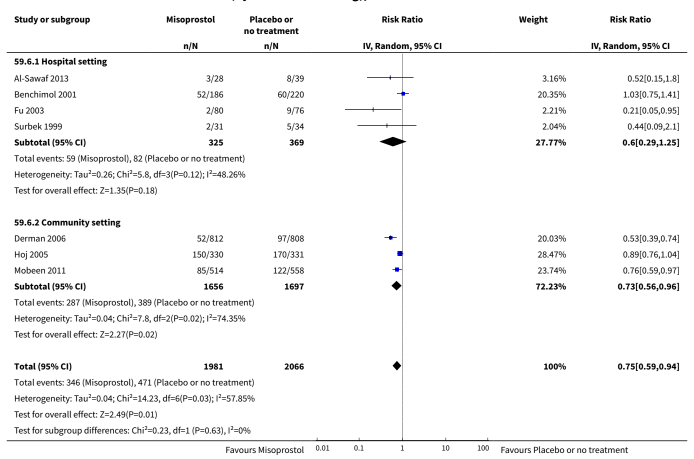
Analysis 59.4. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.







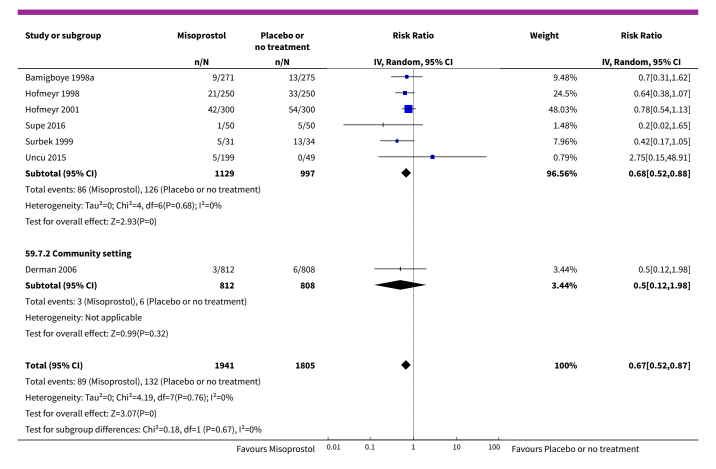
Analysis 59.6. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.



Analysis 59.7. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Placebo or no treatment		Risk Ratio			Weight	Risk Ratio
	n/N	n/N	IN.	/, Random, 9	5% CI			IV, Random, 95% CI
59.7.1 Hospital setting								
Al-Sawaf 2013	3/28	8/39	-	+	1		4.32%	0.52[0.15,1.8]
	Fav	Favours Misoprostol		1	10	100	Favours Placebo or n	o treatment

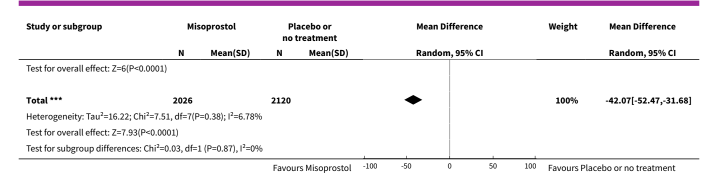




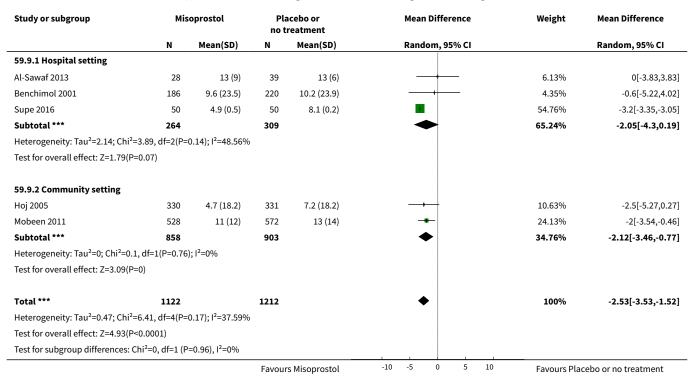
Analysis 59.8. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Mis	Misoprostol !		acebo or reatment		Mean Differe	nce	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95%	6 CI		Random, 95% CI
59.8.1 Hospital setting									
Al-Sawaf 2013	28	348 (112)	39	438.6 (130.2)	•			3.13%	-90.6[-148.83,-32.37]
Benchimol 2001	186	374 (238.4)	220	382 (269.5)	-	+		4.32%	-8[-57.42,41.42]
Fu 2003	76	212.3 (75)	80	242.9 (87)				15.21%	-30.64[-56.1,-5.18]
Supe 2016	50	124.4 (34.7)	50	167.4 (53)	-	_		29.19%	-43[-60.55,-25.45]
Surbek 1999	31	345 (108.6)	34	417 (151)	\leftarrow			2.64%	-72[-135.54,-8.46]
Subtotal ***	371		423		-	◆		54.48%	-41.82[-61.16,-22.49]
Heterogeneity: Tau ² =159.11; Chi ²	² =6.1, df=4(P	=0.19); I ² =34.439	6						
Test for overall effect: Z=4.24(P<0	0.0001)								
59.8.2 Community setting									
Derman 2006	811	214.3 (144.6)	808	262.3 (203.2)	_	-		30.22%	-48[-65.19,-30.81]
Hoj 2005	330	443 (338.3)	331	496 (380.6)	-	+		3.51%	-53[-107.89,1.89]
Mobeen 2011	514	337 (226)	558	366 (262)	-	+		11.79%	-29[-58.23,0.23]
Subtotal ***	1655		1697		•	•		45.52%	-43.79[-58.09,-29.49]
Heterogeneity: Tau ² =0; Chi ² =1.32	2, df=2(P=0.5	2); I ² =0%			1		1	1	
			Favou	rs Misoprostol	-100	50 0	50	100 Favours Pla	cebo or no treatment





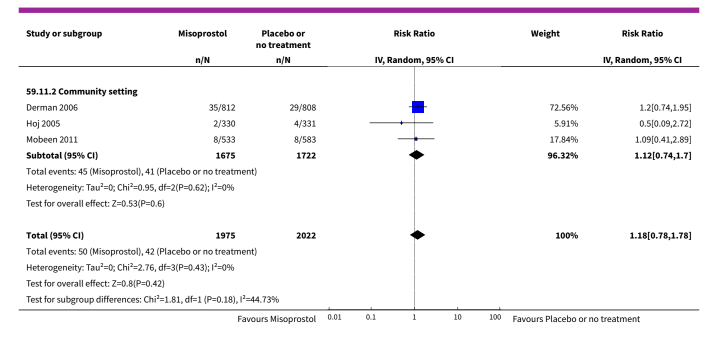
Analysis 59.9. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.



Analysis 59.11. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Misoprostol	no treatment			Risk Ratio			Weight	Risk Ratio	
	n/N			IV, Random, 95% CI					IV, Random, 95% CI	
59.11.1 Hospital setting										
Hofmeyr 2001	5/300	1/300			-	+	_	3.68%	5[0.59,42.54]	
Subtotal (95% CI)	300	300					_	3.68%	5[0.59,42.54]	
Total events: 5 (Misoprostol), 1	(Placebo or no treatment)								
Heterogeneity: Tau ² =0; Chi ² =0	, df=0(P<0.0001); I ² =100%									
Test for overall effect: Z=1.47(F	P=0.14)									
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or r	no treatment	



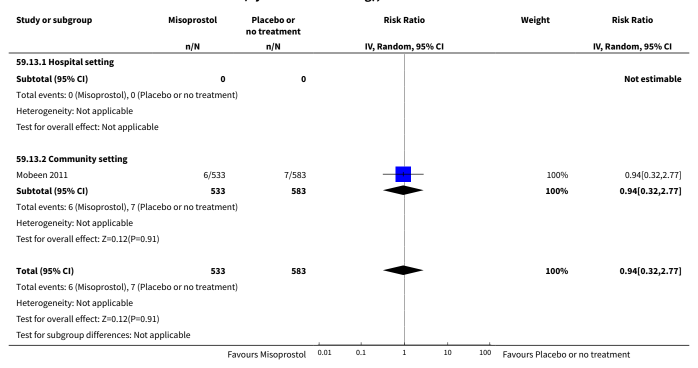


Analysis 59.12. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
59.12.1 Hospital setting					
Bamigboye 1998a	1/271	1/275		2.41%	1.01[0.06,16.14]
Benchimol 2001	7/186	1/220	+	4.24%	8.28[1.03,66.68]
Hofmeyr 2001	4/300	2/300		6.47%	2[0.37,10.84]
Subtotal (95% CI)	757	795		13.13%	2.79[0.85,9.15]
Total events: 12 (Misoprostol), 4 (Placebo or no treatmer	it)			
Heterogeneity: Tau ² =0; Chi ² =1.71,	, df=2(P=0.43); I ² =0%				
Test for overall effect: Z=1.7(P=0.0	09)				
59.12.2 Community setting					
Derman 2006	28/812	25/808	- 	65.64%	1.11[0.66,1.89]
Hoj 2005	10/330	4/331	+	13.98%	2.51[0.79,7.92]
Mobeen 2011	3/533	3/583		7.25%	1.09[0.22,5.4]
Subtotal (95% CI)	1675	1722	*	86.87%	1.27[0.8,2.01]
Total events: 41 (Misoprostol), 32	(Placebo or no treatme	ent)			
Heterogeneity: Tau ² =0; Chi ² =1.61,	, df=2(P=0.45); I ² =0%				
Test for overall effect: Z=1.01(P=0.	.31)				
Total (95% CI)	2432	2517	•	100%	1.41[0.92,2.16]
Total events: 53 (Misoprostol), 36	(Placebo or no treatme	ent)			
Heterogeneity: Tau ² =0; Chi ² =4.8, o	df=5(P=0.44); I ² =0%				
Test for overall effect: Z=1.56(P=0	.12)				
Test for subgroup differences: Chi	i ² =1.48, df=1 (P=0.22), I ²	=32.51%			
	Fa	vours Misoprostol 0.01	0.1 1 10	100 Favours Placebo or	no treatment



Analysis 59.13. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 13 Headache.



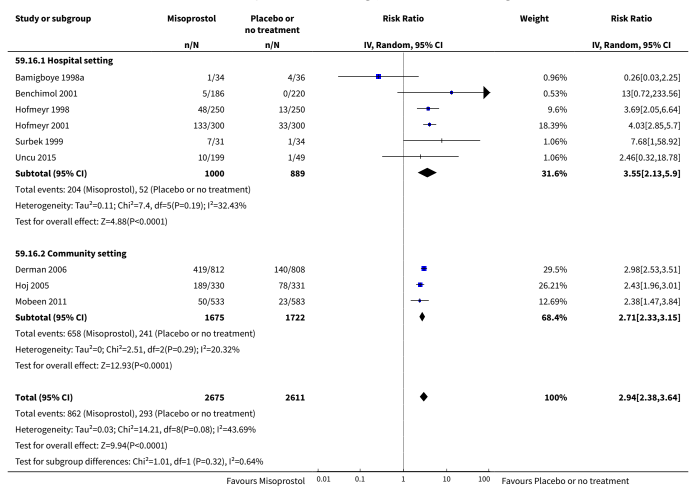
Analysis 59.14. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
59.14.1 Hospital setting						
Bamigboye 1998a	1/271	0/275	-	11.94%	3.04[0.12,74.4]	
Hofmeyr 1998	2/250	7/250		30.86%	0.29[0.06,1.36]	
Hofmeyr 2001	47/300	31/300		57.2%	1.52[0.99,2.32]	
Uncu 2015	0/199	0/49			Not estimable	
Subtotal (95% CI)	1020	874		100%	0.98[0.29,3.37]	
Total events: 50 (Misoprostol), 38 (P	lacebo or no treatme	nt)				
Heterogeneity: Tau ² =0.64; Chi ² =4.33	, df=2(P=0.11); l ² =53.8	31%				
Test for overall effect: Z=0.03(P=0.98	3)					
59.14.2 Community setting						
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Misoprostol), 0 (Plac	ebo or no treatment)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable	e					
Total (95% CI)	1020	874	•	100%	0.98[0.29,3.37]	
Total events: 50 (Misoprostol), 38 (P	lacebo or no treatme	nt)				
Heterogeneity: Tau ² =0.64; Chi ² =4.33	, df=2(P=0.11); I ² =53.8	31%				
Test for overall effect: Z=0.03(P=0.98	3)					
	Fa	vours Misoprostol 0.	01 0.1 1 10	100 Favours Placebo or I	no treatment	



Study or subgroup			lacebo or Risk Ratio treatment					Weight Risk Ratio
	n/N	n/N		IV, R	andom, 9	5% CI		IV, Random, 95% CI
Test for subgroup differences		_				_		
		Favours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or no treatment

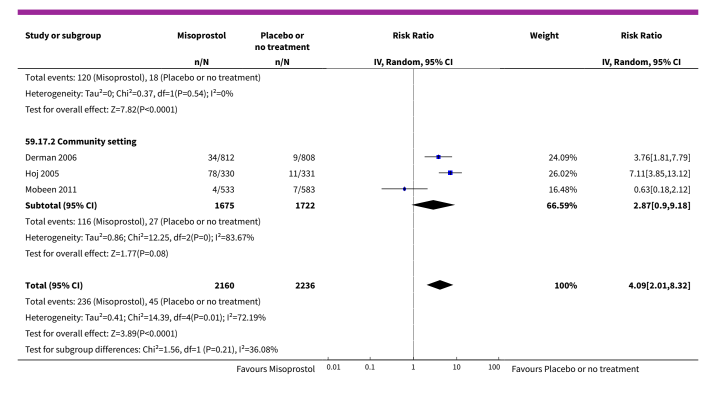
Analysis 59.16. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 16 Shivering.



Analysis 59.17. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol	Misoprostol Placebo or no treatment			Risk Rati	0	Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
59.17.1 Hospital setting									
Benchimol 2001	6/186	0/220			 	+	\rightarrow	5.14%	15.36[0.87,270.93]
Hofmeyr 2001	114/299	18/294			ĺ	-		28.27%	6.23[3.89,9.97]
Subtotal (95% CI)	485	514				•		33.41%	6.38[4.01,10.14]
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Placebo or i	no treatment





Analysis 59.18. Comparison 59 Misoprostol vs Placebo or no treatment (by healthcare setting), Outcome 18 Diarrhoea.

Misoprostol	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
0/271	0/275			Not estimable
1/300	1/300		9.24%	1[0.06,15.91]
0/199	0/49			Not estimable
770	624		9.24%	1[0.06,15.91]
Placebo or no treatment))			
able				
9/812	2/808		30.27%	4.48[0.97,20.66]
10/330	4/331	 	53.57%	2.51[0.79,7.92]
1/533	0/583	+	- 6.92%	3.28[0.13,80.36]
1675	1722	-	90.76%	3.11[1.28,7.51]
(Placebo or no treatmen	t)			
5, df=2(P=0.84); I ² =0%				
0.01)				
2445	2346	•	100%	2.8[1.21,6.49]
(Placebo or no treatmen	t)			
1, df=3(P=0.82); I ² =0%				
02)				
ni ² =0.58, df=1 (P=0.44), I ²	=0%			
	n/N 0/271 1/300 0/199 770 Placebo or no treatment) able 9/812 10/330 1/533 1675 (Placebo or no treatmen 6, df=2(P=0.84); l²=0% 0.01) 2445 (Placebo or no treatmen 8, df=3(P=0.82); l²=0% 0.02)	no treatment n/N n/N 0/271 0/275 1/300 1/300 0/199 0/49 770 624 Placebo or no treatment) able 9/812 2/808 10/330 4/331 1/533 0/583 1675 1722 (Placebo or no treatment) 6, df=2(P=0.84); l²=0% 0.01) 2445 2346 (Placebo or no treatment) 6, df=3(P=0.82); l²=0%	no treatment n/N n/N IV, Random, 95% CI 0/271 0/275 1/300 1/300 0/199 0/49 770 624 Placebo or no treatment) able 9/812 2/808 10/330 4/331 1/533 0/583 1675 1722 (Placebo or no treatment) 6, df=2(P=0.84); l²=0% 0.01) 2445 2346 (Placebo or no treatment) 8, df=3(P=0.82); l²=0% 002)	no treatment n/N n/N IV, Random, 95% CI 0/271 0/275 1/300 1/300 0/199 0/49 770 624 Placebo or no treatment) able 9/812 2/808 10/330 4/331 1/533 0/583 1675 1722 90.76% (Placebo or no treatment) 6, df=2(P=0.84); l²=0% 0.01) 2445 2346 (Placebo or no treatment) 0, df=3(P=0.82); l²=0% 02)



Comparison 60. Injectable prostaglandins vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.04, 3.24]
2.1 Hospital setting	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.04, 3.24]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal mor- bidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	46	Risk Ratio (IV, Random, 95% CI)	0.55 [0.22, 1.35]
6.1 Hospital setting	1	46	Risk Ratio (IV, Random, 95% CI)	0.55 [0.22, 1.35]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	146	Risk Ratio (IV, Random, 95% CI)	0.66 [0.21, 2.09]
7.1 Hospital setting	2	146	Risk Ratio (IV, Random, 95% CI)	0.66 [0.21, 2.09]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	146	Mean Difference (IV, Random, 95% CI)	-95.17 [-296.09, 105.75]
8.1 Hospital setting	2	146	Mean Difference (IV, Random, 95% CI)	-95.17 [-296.09, 105.75]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	1	100	Mean Difference (IV, Random, 95% CI)	0.90 [0.56, 1.24]
9.1 Hospital setting	1	100	Mean Difference (IV, Random, 95% CI)	0.90 [0.56, 1.24]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.02, 8.46]
11.1 Hospital setting	1	46	Risk Ratio (IV, Random, 95% CI)	0.36 [0.02, 8.46]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



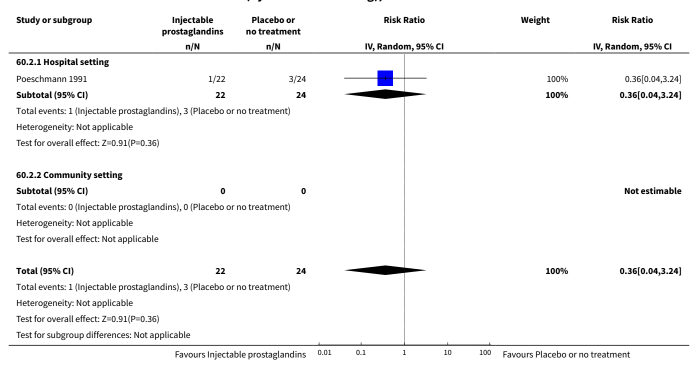
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 60.1. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		I	Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ra	ındom, 95% CI			IV, Random, 95% CI
60.1.1 Hospital setting								
Supe 2016	0/50	0/50						Not estimable
Subtotal (95% CI)	50	50						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
60.1.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	•							
Total (95% CI)	50	50						Not estimable
Total events: 0 (Injectable prostaglar	ndins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	<u> </u>							
Test for subgroup differences: Not ap	plicable							
	Favours Injecta	ble prostaglandins	0.01	0.1	1 10	100	Favours Placebo o	r no treatment



Analysis 60.2. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.

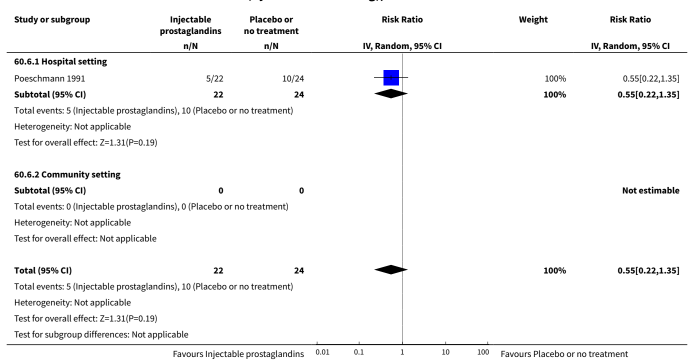


Analysis 60.3. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
60.3.1 Hospital setting					
Supe 2016	0/50	0/50			Not estimable
Subtotal (95% CI)	50	50			Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Placebo or	no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
60.3.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Placebo or	no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	50	50			Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Placebo or	no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Test for subgroup differences: Not ap	pplicable				
	Favours Injecta	ble prostaglandins 0.0.	1 0.1 1 10	100 Favours Placebo	or no treatment



Analysis 60.6. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.

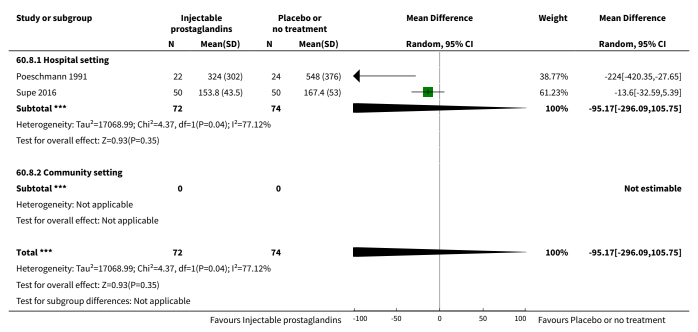


Analysis 60.7. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Placebo or no treatment		ı	Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95% CI			IV, Random, 95% CI
60.7.1 Hospital setting								
Poeschmann 1991	0/22	2/24					15.04%	0.22[0.01,4.29]
Supe 2016	4/50	5/50		_			84.96%	0.8[0.23,2.81]
Subtotal (95% CI)	72	74		-			100%	0.66[0.21,2.09]
Total events: 4 (Injectable prostagla	ndins), 7 (Placebo or	no treatment)						
Heterogeneity: Tau ² =0; Chi ² =0.62, df	=1(P=0.43); I ² =0%							
Test for overall effect: Z=0.71(P=0.48	3)							
60.7.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Placebo or	no treatment)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	e							
Total (95% CI)	72	74		-			100%	0.66[0.21,2.09]
Total events: 4 (Injectable prostagla	ndins), 7 (Placebo or	no treatment)						
Heterogeneity: Tau ² =0; Chi ² =0.62, df	f=1(P=0.43); I ² =0%							
Test for overall effect: Z=0.71(P=0.48	3)							
Test for subgroup differences: Not a	pplicable							
	Favours Injecta	ble prostaglandins	0.01	0.1	1 10	100	Favours Placebo or i	no treatment



Analysis 60.8. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

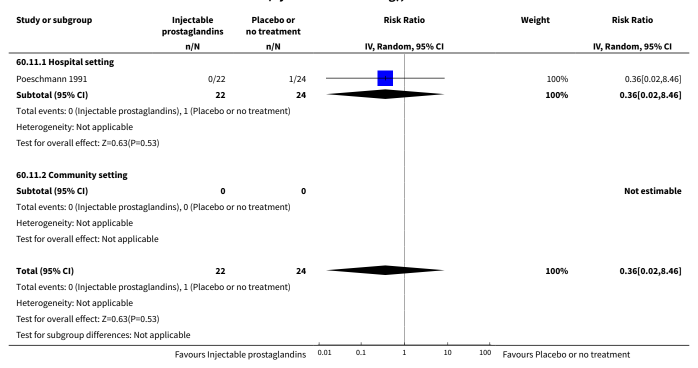


Analysis 60.9. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup		ectable taglandins		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
60.9.1 Hospital setting							
Supe 2016	50	9 (1.2)	50	8.1 (0.2)	-	100%	0.9[0.56,1.24]
Subtotal ***	50		50		◆	100%	0.9[0.56,1.24]
Heterogeneity: Not applicable							
Test for overall effect: Z=5.23(P<0.0	0001)						
60.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	le						
Total ***	50		50		•	100%	0.9[0.56,1.24]
Heterogeneity: Not applicable							
Test for overall effect: Z=5.23(P<0.0	0001)						
Test for subgroup differences: Not	applicable						
		Favours Inj	ectable p	rostaglandins -4	-2 0 2	4 Favours Pla	cebo or no treatment



Analysis 60.11. Comparison 60 Injectable prostaglandins vs Placebo or no treatment (by healthcare setting), Outcome 11 Nausea.



Comparison 61. Ergometrine vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.09 [0.01, 0.72]
2.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	0.09 [0.01, 0.72]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1529	Risk Ratio (IV, Random, 95% CI)	0.34 [0.04, 3.28]
3.1 Hospital setting	2	1529	Risk Ratio (IV, Random, 95% CI)	0.34 [0.04, 3.28]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbid- ity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	1429	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.42]
6.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	0.24 [0.14, 0.42]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	1529	Risk Ratio (IV, Random, 95% CI)	0.18 [0.09, 0.37]
7.1 Hospital setting	2	1529	Risk Ratio (IV, Random, 95% CI)	0.18 [0.09, 0.37]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	1529	Mean Difference (IV, Random, 95% CI)	-50.64 [-119.92, 18.65]
8.1 Hospital setting	2	1529	Mean Difference (IV, Random, 95% CI)	-50.64 [-119.92, 18.65]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	1	100	Mean Difference (IV, Random, 95% CI)	-0.5 [-0.58, -0.42]
9.1 Hospital setting	1	100	Mean Difference (IV, Random, 95% CI)	-0.5 [-0.58, -0.42]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	1429	Risk Ratio (IV, Random, 95% CI)	42.10 [2.55, 694.80]
11.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	42.10 [2.55, 694.80]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	1	1429	Risk Ratio (IV, Random, 95% CI)	25.67 [1.52, 432.78]
12.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	25.67 [1.52, 432.78]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [0.37, 138.91]



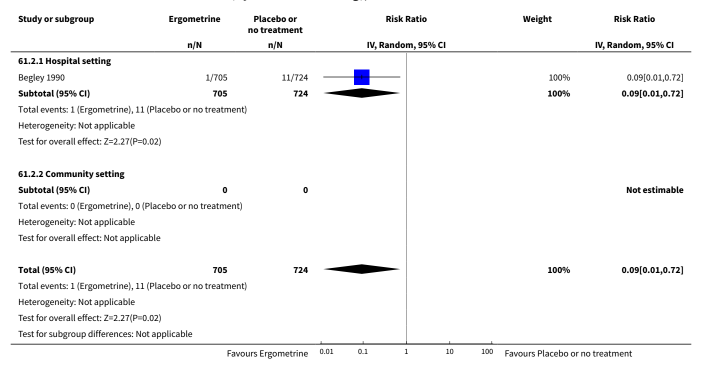
Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
13.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [0.37, 138.91]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	1	1429	Risk Ratio (IV, Random, 95% CI)	2.05 [1.04, 4.08]
14.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	2.05 [1.04, 4.08]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [2.83, 18.24]
15.1 Hospital setting	1	1429	Risk Ratio (IV, Random, 95% CI)	7.19 [2.83, 18.24]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Analysis 61.1. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 1 Death.

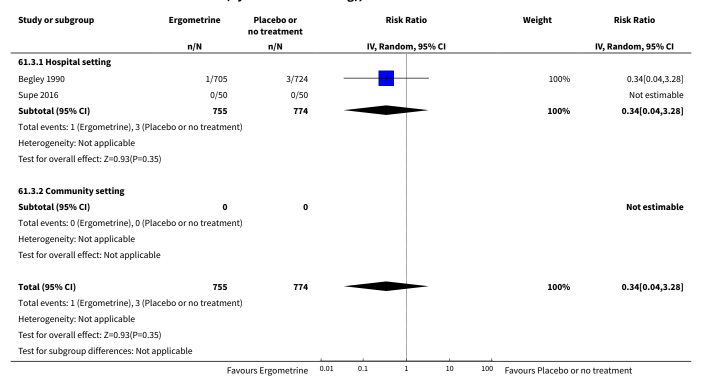
Study or subgroup	Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
61.1.1 Hospital setting					
Supe 2016	0/50	0/50			Not estimable
Subtotal (95% CI)	50	50			Not estimable
Total events: 0 (Ergometrine), 0 (Plac	ebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
61.1.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Plac	ebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	50	50			Not estimable
Total events: 0 (Ergometrine), 0 (Plac	ebo or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Test for subgroup differences: Not ap	plicable				
	Fav	ours Ergometrine 0.0	1 0.1 1 10	100 Favours Placebo o	r no treatment

Analysis 61.2. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.





Analysis 61.3. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 3 Blood transfusion.

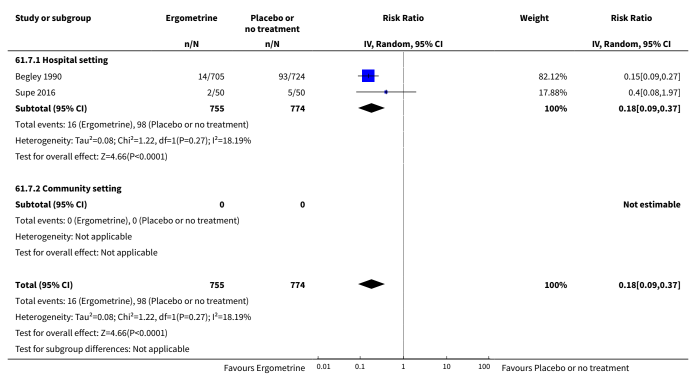


Analysis 61.6. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine	Placebo or no treatment		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IN	/, Random, 95% CI		IV, Random, 95% CI
61.6.1 Hospital setting						
Begley 1990	14/705	60/724	-	1	100%	0.24[0.14,0.42]
Subtotal (95% CI)	705	724	-	•	100%	0.24[0.14,0.42]
Total events: 14 (Ergometrine), 60 (Pla	acebo or no treatm	ent)				
Heterogeneity: Not applicable						
Test for overall effect: Z=4.89(P<0.000	1)					
61.6.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Ergometrine), 0 (Place	ebo or no treatmen	t)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	705	724	•	•	100%	0.24[0.14,0.42]
Total events: 14 (Ergometrine), 60 (Pla	acebo or no treatm	ent)				
Heterogeneity: Not applicable						
Test for overall effect: Z=4.89(P<0.000	1)					
Test for subgroup differences: Not app	plicable					
	Fa	vours Ergometrine	0.01 0.1	1 10	100 Favours Placebo or	no treatment



Analysis 61.7. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.

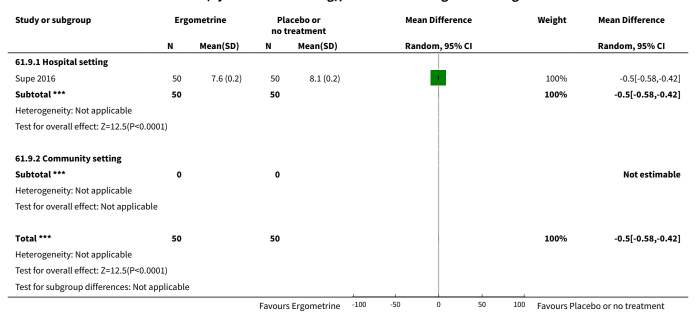


Analysis 61.8. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine		icebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
61.8.1 Hospital setting							
Begley 1990	705	148.9 (127.1)	724	234.8 (223.9)		50.13%	-85.9[-104.72,-67.08]
Supe 2016	50	152.2 (49.3)	50	167.4 (53)	-	49.87%	-15.2[-35.25,4.85]
Subtotal ***	755		774			100%	-50.64[-119.92,18.65]
Heterogeneity: Tau ² =2400.83; Chi ² =	=25.4, df=1	.(P<0.0001); I ² =96	6.06%				
Test for overall effect: Z=1.43(P=0.1	L5)						
61.8.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	ole						
Total ***	755		774			100%	-50.64[-119.92,18.65]
Heterogeneity: Tau ² =2400.83; Chi ² =	=25.4, df=1	.(P<0.0001); I ² =96	5.06%				
Test for overall effect: Z=1.43(P=0.1	L5)						
Test for subgroup differences: Not	applicable						
			Favours	Ergometrine	-100 -50 0 50	100 Favours Pla	acebo or no treatment



Analysis 61.9. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.

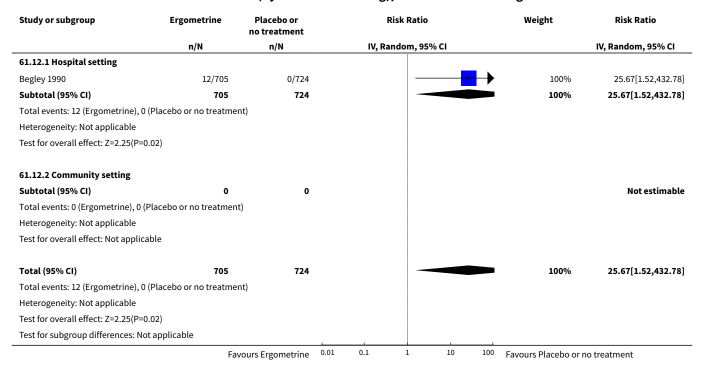


Analysis 61.11. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 11 Nausea.

Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
20/705	0/724		100%	42.1[2.55,694.8]
705	724		100%	42.1[2.55,694.8]
Placebo or no treatme	nt)			
01)				
0	0			Not estimable
lacebo or no treatmen	t)	İ		
ole				
705	724		100%	42.1[2.55,694.8]
Placebo or no treatme	nt)	İ		
		İ		
01)		į		
applicable		į		
	n/N 20/705 705 Placebo or no treatme 01) 0 lacebo or no treatmen ole 705 Placebo or no treatme	no treatment n/N n/N 20/705 0/724 705 724 Placebo or no treatment) 0 0 dacebo or no treatment) ole 705 724 Placebo or no treatment)	no treatment n/N n/N iV, Random, 95% CI 20/705 0/724 705 724 Placebo or no treatment) 0 0 lacebo or no treatment) ole 705 724 Placebo or no treatment)	no treatment n/N n/N iV, Random, 95% CI 20/705 0/724 705 724 Placebo or no treatment) 0 0 lacebo or no treatment) ple 705 724 Placebo or no treatment) 100%



Analysis 61.12. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 12 Vomiting.

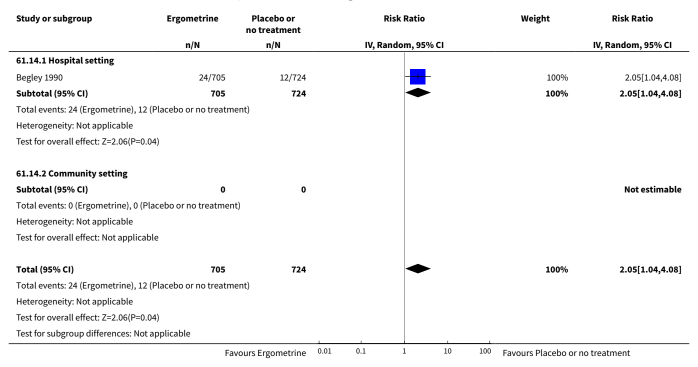


Analysis 61.13. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
61.13.1 Hospital setting					
Begley 1990	3/705	0/724		100%	7.19[0.37,138.91]
Subtotal (95% CI)	705	724		100%	7.19[0.37,138.91]
Total events: 3 (Ergometrine), 0 (Plac	ebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.31(P=0.19))				
61.13.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Plac	ebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	705	724		100%	7.19[0.37,138.91]
Total events: 3 (Ergometrine), 0 (Plac	ebo or no treatment)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.31(P=0.19))				
Test for subgroup differences: Not ap	plicable				
	Favo	ours Ergometrine 0.0	1 0.1 1 10 1	100 Favours Placebo or	no treatment



Analysis 61.14. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 14 Abdominal pain.



Analysis 61.15. Comparison 61 Ergometrine vs Placebo or no treatment (by healthcare setting), Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
61.15.1 Hospital setting					
Begley 1990	35/705	5/724		100%	7.19[2.83,18.24]
Subtotal (95% CI)	705	724		100%	7.19[2.83,18.24]
Total events: 35 (Ergometrine), 5 (Pla	cebo or no treatme	nt)			
Heterogeneity: Not applicable					
Test for overall effect: Z=4.15(P<0.000	1)				
61.15.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Place	ebo or no treatmen	t)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	705	724	•	100%	7.19[2.83,18.24]
Total events: 35 (Ergometrine), 5 (Pla	cebo or no treatme	nt)			
Heterogeneity: Not applicable					
Test for overall effect: Z=4.15(P<0.000	1)				
Test for subgroup differences: Not ap	plicable				
	Fa	vours Ergometrine 0.	01 0.1 1 10 1	100 Favours Placebo or	no treatment



Comparison 62. Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.44 [0.18, 1.05]
2.1 Hospital setting	2	3207	Risk Ratio (IV, Random, 95% CI)	0.44 [0.18, 1.05]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	3	3400	Risk Ratio (IV, Random, 95% CI)	0.34 [0.18, 0.66]
3.1 Hospital setting	3	3400	Risk Ratio (IV, Random, 95% CI)	0.34 [0.18, 0.66]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	3207	Risk Ratio (IV, Random, 95% CI)	0.37 [0.30, 0.46]
6.1 Hospital setting	2	3207	Risk Ratio (IV, Random, 95% CI)	0.37 [0.30, 0.46]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	3	3400	Risk Ratio (IV, Random, 95% CI)	0.19 [0.15, 0.24]
7.1 Hospital setting	3	3400	Risk Ratio (IV, Random, 95% CI)	0.19 [0.15, 0.24]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	1705	Mean Difference (IV, Random, 95% CI)	-35.02 [-101.63, 31.59]
8.1 Hospital setting	2	1705	Mean Difference (IV, Random, 95% CI)	-35.02 [-101.63, 31.59]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	1454	Mean Difference (IV, Random, 95% CI)	-3.57 [-6.50, -0.63]
9.1 Hospital setting	2	1454	Mean Difference (IV, Random, 95% CI)	-3.57 [-6.50, -0.63]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	2	3207	Risk Ratio (IV, Random, 95% CI)	1.03 [0.99, 1.07]
10.1 Hospital setting	2	3207	Risk Ratio (IV, Random, 95% CI)	1.03 [0.99, 1.07]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	1512	Risk Ratio (IV, Random, 95% CI)	1.95 [1.38, 2.76]
11.1 Hospital setting	1	1512	Risk Ratio (IV, Random, 95% CI)	1.95 [1.38, 2.76]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	3207	Risk Ratio (IV, Random, 95% CI)	2.15 [1.46, 3.18]
12.1 Hospital setting	2	3207	Risk Ratio (IV, Random, 95% CI)	2.15 [1.46, 3.18]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	2	3207	Risk Ratio (IV, Random, 95% CI)	1.65 [0.78, 3.48]
13.1 Hospital setting	2	3207	Risk Ratio (IV, Random, 95% CI)	1.65 [0.78, 3.48]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

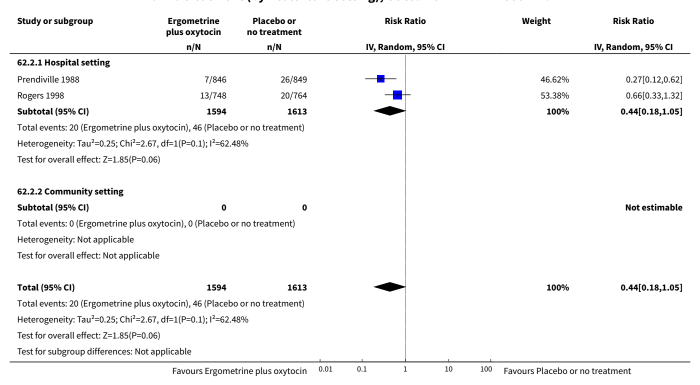


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting - General health at 6 weeks postpartum (Worse than prepregnancy)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Hospital setting - General health at 6 weeks postpartum (Exhausted since birth)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.3 Hospital setting - General health at 6 weeks postpartum (Exhausted at 6 weeks)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.4 Hospital setting - General health at 6 weeks postpartum (Blues)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.5 Hospital setting - General health at 6 weeks postpartum (Depressed)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.6 Hospital setting - General health at 6 weeks postpartum (Help for depression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.7 Hospital setting - General health at 6 weeks postpartum (Admission to hospital for de- pression)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.8 Hospital setting - General health at 6 weeks postpartum (No health problems reported)	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.9 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting - Satisfied with third-stage management	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

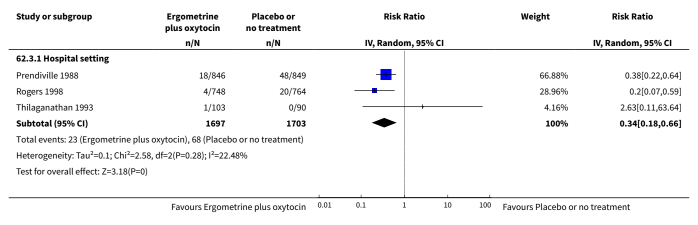


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
20.2 Hospital setting - Felt in control during third stage	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

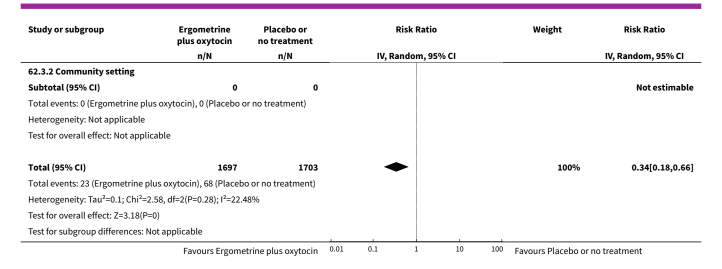
Analysis 62.2. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 2 PPH >= 1000 mL.



Analysis 62.3. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 3 Blood transfusion.





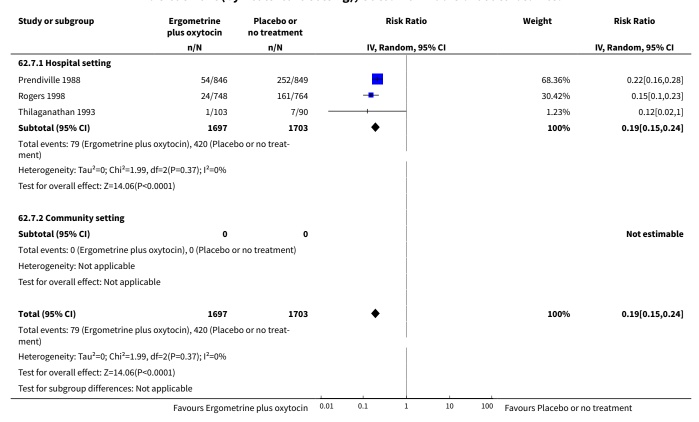


Analysis 62.6. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
62.6.1 Hospital setting					
Prendiville 1988	50/846	152/849		50.67%	0.33[0.24,0.45]
Rogers 1998	51/748	126/764	-	49.33%	0.41[0.3,0.56]
Subtotal (95% CI)	1594	1613	•	100%	0.37[0.3,0.46]
Total events: 101 (Ergometrine plus ment)	oxytocin), 278 (Place	ebo or no treat-			
Heterogeneity: Tau ² =0; Chi ² =1.03, d	f=1(P=0.31); I ² =3%				
Test for overall effect: Z=8.86(P<0.0	001)				
62.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus o	xytocin), 0 (Placebo c	or no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	1594	1613	◆	100%	0.37[0.3,0.46]
Total events: 101 (Ergometrine plus ment)	oxytocin), 278 (Place	ebo or no treat-			
Heterogeneity: Tau ² =0; Chi ² =1.03, d	f=1(P=0.31); I ² =3%				
Test for overall effect: Z=8.86(P<0.0	001)				
Test for subgroup differences: Not a	pplicable				
	Favours Ergome	trine plus oxytocin 0	0.01 0.1 1 10	100 Favours Placebo or	no treatment



Analysis 62.7. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 7 Additional uterotonics.



Analysis 62.8. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup		ometrine oxytocin		acebo or reatment	Me	ean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Ra	andom, 95% CI		Random, 95% CI
62.8.1 Hospital setting								
Rogers 1998	748	268.5 (246.1)	764	336.5 (243.2)			51.5%	-68[-92.67,-43.33]
Thilaganathan 1993	103	200 (74.1)	90	200 (148.3)	_		48.5%	0[-33.81,33.81]
Subtotal ***	851		854	-			100%	-35.02[-101.63,31.59]
Heterogeneity: Tau ² =2083.99; Chi ²	=10.14, df=	1(P=0); I ² =90.14 ⁰	%					
Test for overall effect: Z=1.03(P=0.3	3)							
62.8.2 Community setting								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
Test for overall effect: Not applicab	ole							
Total ***	851		854	-			100%	-35.02[-101.63,31.59]
Heterogeneity: Tau ² =2083.99; Chi ²	=10.14, df=	1(P=0); I ² =90.14	%					
Test for overall effect: Z=1.03(P=0.3	3)							
Test for subgroup differences: Not	applicable							
		Favours Ergo	ometrine	plus oxytocin -1	00 -50	0 50	100 Favours Pla	acebo or no treatment



Analysis 62.9. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 9 Change in haemoglobin.

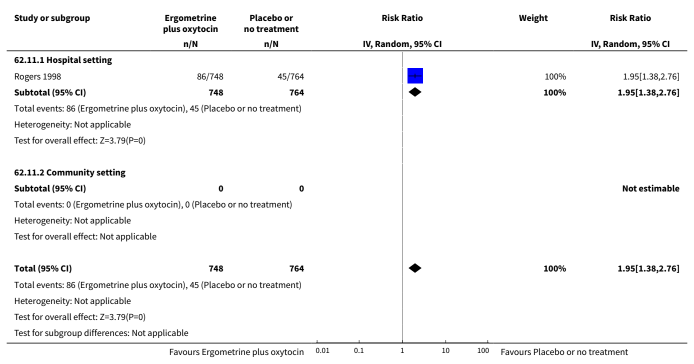
Study or subgroup		ometrine oxytocin		acebo or reatment	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
62.9.1 Hospital setting							
Prendiville 1988	634	1 (21)	627	6 (13)	•	52.23%	-5[-6.93,-3.07]
Thilaganathan 1993	103	5 (8.1)	90	7 (8.1)	•	47.77%	-2[-4.29,0.29]
Subtotal ***	737		717		♦	100%	-3.57[-6.5,-0.63]
Heterogeneity: Tau ² =3.33; Chi ² =3.	86, df=1(P=	0.05); I ² =74.1%					
Test for overall effect: Z=2.38(P=0.	.02)						
62.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applica	ble						
Total ***	737		717		•	100%	-3.57[-6.5,-0.63]
Heterogeneity: Tau ² =3.33; Chi ² =3.	86, df=1(P=	0.05); I ² =74.1%					
Test for overall effect: Z=2.38(P=0.	.02)						
Test for subgroup differences: Not	applicable						
		Favours Fro	metrine	plus oxytocin -100	-50 0 50	100 Favours Pla	cebo or no treatment

Analysis 62.10. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 10 Breastfeeding.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV	, Random, 95% CI			IV, Random, 95% CI
62.10.1 Hospital setting							
Prendiville 1988	637/846	632/849		•		57.43%	1.01[0.96,1.07]
Rogers 1998	546/748	531/764		•		42.57%	1.05[0.99,1.12]
Subtotal (95% CI)	1594	1613				100%	1.03[0.99,1.07]
Total events: 1183 (Ergometrine plus ment)	oxytocin), 1163 (Pla	acebo or no treat-		İ			
Heterogeneity: Tau ² =0; Chi ² =0.76, df	=1(P=0.38); I ² =0%						
Test for overall effect: Z=1.29(P=0.2)							
62.10.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Ergometrine plus ox	ytocin), 0 (Placebo o	r no treatment)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	<u> </u>						
Total (95% CI)	1594	1613				100%	1.03[0.99,1.07]
Total events: 1183 (Ergometrine plus ment)	oxytocin), 1163 (Pla	acebo or no treat-					
Heterogeneity: Tau ² =0; Chi ² =0.76, df	=1(P=0.38); I ² =0%						
Test for overall effect: Z=1.29(P=0.2)							
Test for subgroup differences: Not ap	plicable						
	Favours Ergome	trine plus oxytocin	0.01 0.1	1 10	100	Favours Placebo or r	no treatment



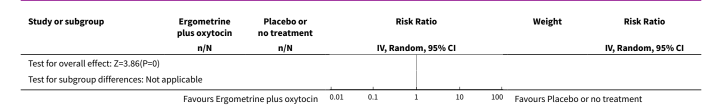
Analysis 62.11. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 11 Nausea.



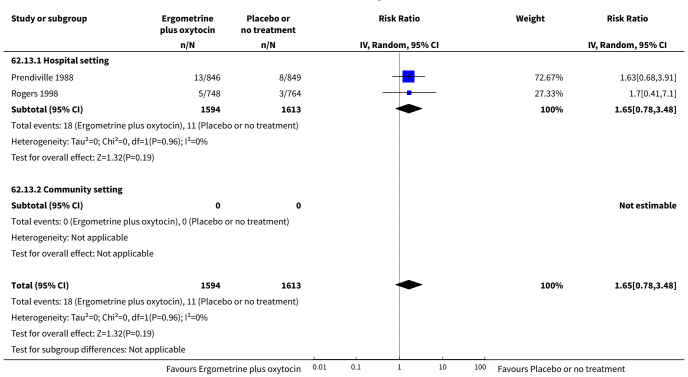
Analysis 62.12. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	ogroup Ergometrine Placebo or Risk Ratio plus oxytocin no treatment		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
62.12.1 Hospital setting					
Prendiville 1988	102/846	55/849	-	64.91%	1.86[1.36,2.55]
Rogers 1998	47/748	17/764	-	35.09%	2.82[1.64,4.87]
Subtotal (95% CI)	1594	1613	•	100%	2.15[1.46,3.18]
Total events: 149 (Ergometrine plument)	us oxytocin), 72 (Placel	oo or no treat-			
Heterogeneity: Tau ² =0.04; Chi ² =1.	69, df=1(P=0.19); I ² =40	.7%			
Test for overall effect: Z=3.86(P=0))				
62.12.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus	oxytocin), 0 (Placebo o	r no treatment)			
Heterogeneity: Not applicable					
Test for overall effect: Not applica	ble				
Total (95% CI)	1594	1613	•	100%	2.15[1.46,3.18]
Total events: 149 (Ergometrine plument)	us oxytocin), 72 (Placel	oo or no treat-			
Heterogeneity: Tau ² =0.04; Chi ² =1.	69, df=1(P=0.19); I ² =40.	.7%			
	Favours Ergome	trine plus oxytocin 0.01	0.1 1 10	100 Favours Placebo or	no treatment

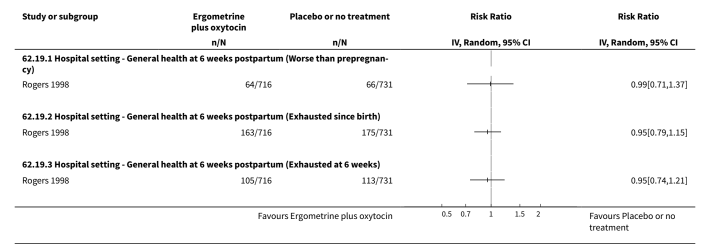




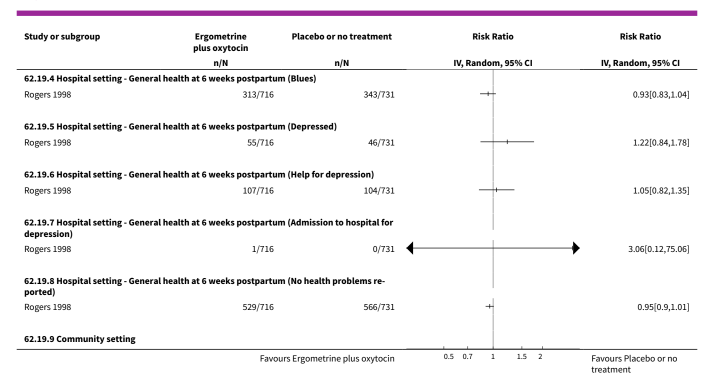
Analysis 62.13. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 13 Headache.



Analysis 62.19. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 19 Maternal well-being.







Analysis 62.20. Comparison 62 Ergometrine plus oxytocin vs Placebo or no treatment (by healthcare setting), Outcome 20 Maternal satisfaction.

Study or subgroup	Ergometrine plus oxytocin	Placebo or no treatment Risk Ratio		Risk Ratio	
	n/N	n/N	IV, Random, 95% CI	IV, Random, 95% CI	
62.20.1 Hospital setting - Sati	sfied with third-stage manageme	ent			
Rogers 1998	721/745	718/762	-	1.03[1,1.05]	
62.20.2 Hospital setting - Felt	in control during third stage				
Rogers 1998	621/745	667/762		0.95[0.91,0.99]	
62.20.3 Community setting					
	Favor	urs Ergometrine plus oxytocin	1	Favours Placebo or no treatment	

Comparison 63. Misoprostol plus oxytocin vs Placebo or no treatment (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of participants	Statistical method	Effect size
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of participants	Statistical method	Effect size
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 64. Misoprostol vs Oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	24	28520	Risk Ratio (IV, Random, 95% CI)	0.62 [0.14, 2.74]
1.1 Hospital setting	23	28127	Risk Ratio (IV, Random, 95% CI)	0.62 [0.14, 2.74]
1.2 Community setting	1	393	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	38	34261	Risk Ratio (IV, Random, 95% CI)	1.26 [1.11, 1.43]
2.1 Hospital setting	38	34261	Risk Ratio (IV, Random, 95% CI)	1.26 [1.11, 1.43]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	42	35467	Risk Ratio (IV, Random, 95% CI)	0.81 [0.65, 1.00]
3.1 Hospital setting	42	35467	Risk Ratio (IV, Random, 95% CI)	0.81 [0.65, 1.00]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	10	21698	Risk Ratio (IV, Random, 95% CI)	1.16 [0.55, 2.43]
4.1 Hospital setting	10	21698	Risk Ratio (IV, Random, 95% CI)	1.16 [0.55, 2.43]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	44	36920	Risk Ratio (IV, Random, 95% CI)	1.08 [0.94, 1.24]
6.1 Hospital setting	44	36920	Risk Ratio (IV, Random, 95% CI)	1.08 [0.94, 1.24]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	47	35981	Risk Ratio (IV, Random, 95% CI)	1.01 [0.85, 1.20]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size	
7.1 Hospital setting	47	35981	Risk Ratio (IV, Random, 95% CI)	1.01 [0.85, 1.20]	
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	43	35239	Mean Difference (IV, Random, 95% CI)	-8.90 [-23.45, 5.65]	
8.1 Hospital setting	43	35239	239 Mean Difference (IV, Random, 95% CI)		
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglobin	31	12028	Mean Difference (IV, Random, 95% CI)	-0.14 [-0.74, 0.47]	
9.1 Hospital setting	30	11724	Mean Difference (IV, Random, 95% CI)	-0.15 [-0.77, 0.46]	
9.2 Community setting	1	304	Mean Difference (IV, Random, 95% CI)	0.76 [-3.21, 4.73]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	33	29732	Risk Ratio (IV, Random, 95% CI)	1.22 [0.93, 1.60]	
11.1 Hospital setting	32	29339	Risk Ratio (IV, Random, 95% CI)	1.22 [0.93, 1.61]	
11.2 Community setting	1	393	Risk Ratio (IV, Random, 95% CI)	0.84 [0.14, 4.96]	
12 Vomiting	41	32687	Risk Ratio (IV, Random, 95% CI)	1.51 [1.19, 1.91]	
12.1 Hospital setting	40	32294	Risk Ratio (IV, Random, 95% CI)	1.48 [1.16, 1.89]	
12.2 Community setting	1	393	Risk Ratio (IV, Random, 95% CI)	2.81 [0.14, 58.05]	
13 Headache	10	4079	Risk Ratio (IV, Random, 95% CI)	0.88 [0.54, 1.42]	
13.1 Hospital setting	10	4079	Risk Ratio (IV, Random, 95% CI)	0.88 [0.54, 1.42]	
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
14 Abdominal pain	8	3382	Risk Ratio (IV, Random, 95% CI)	0.91 [0.79, 1.06]	
14.1 Hospital setting	8	3382	Risk Ratio (IV, Random, 95% CI)	0.91 [0.79, 1.06]	
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
15 Hypertension	3	1028	Risk Ratio (IV, Random, 95% CI)	3.64 [0.60, 22.27]	
15.1 Hospital setting	3	1028	Risk Ratio (IV, Random, 95% CI)	3.64 [0.60, 22.27]	
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
16 Shivering	49	34865	Risk Ratio (IV, Random, 95% CI)	4.02 [3.23, 4.99]	

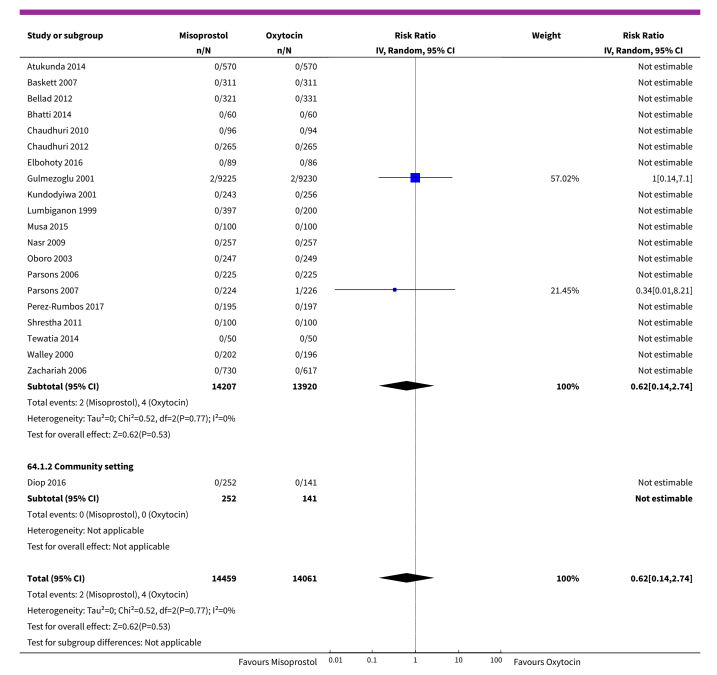


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
16.1 Hospital setting	48	34534	Risk Ratio (IV, Random, 95% CI)	3.82 [3.09, 4.74]
16.2 Community setting	1	331	Risk Ratio (IV, Random, 95% CI)	16.13 [7.81, 33.31]
17 Fever	41	33008	Risk Ratio (IV, Random, 95% CI)	3.75 [2.73, 5.15]
17.1 Hospital setting	40	32615	Risk Ratio (IV, Random, 95% CI)	3.80 [2.76, 5.25]
17.2 Community setting	1	393	Risk Ratio (IV, Random, 95% CI)	2.24 [0.48, 10.40]
18 Diarrhoea	26	30733	Risk Ratio (IV, Random, 95% CI)	2.13 [1.55, 2.93]
18.1 Hospital setting	25	30340	Risk Ratio (IV, Random, 95% CI)	2.17 [1.57, 2.99]
18.2 Community setting	1	393	Risk Ratio (IV, Random, 95% CI)	0.56 [0.04, 8.88]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting - Complaints about or prob- lems with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.3 Community setting - Satisfied or very satisfied with drug	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.4 Community setting - Would take drug again af- ter subsequent deliveries	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.5 Community setting - Would recommend drug to a friend	1		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 64.1. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol	Oxytocin			Risk Ratio)		Weight	Risk Ratio
	n/N	n/N		IV, Ra	andom, 9	5% CI			IV, Random, 95% CI
64.1.1 Hospital setting									
Afolabi 2010	0/100	0/100							Not estimable
Alwani 2014	0/100	1/100						21.54%	0.33[0.01,8.09]
Amin 2014	0/100	0/100							Not estimable
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Oxytocin	

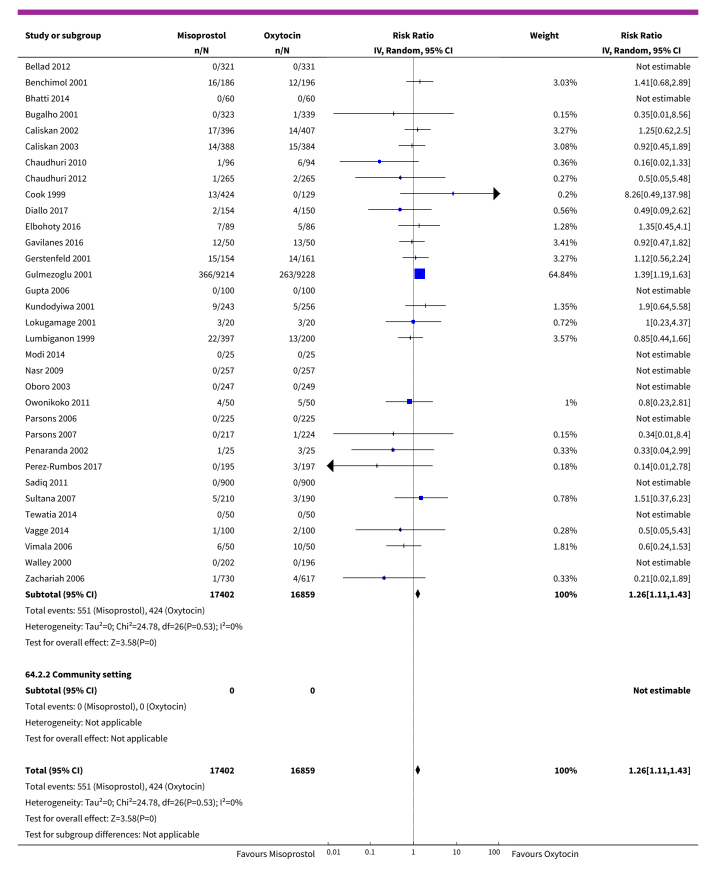




Analysis 64.2. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Misoprostol	Risk Ratio					Weight	Risk Ratio	
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
64.2.1 Hospital setting									
Acharya 2001	1/30	1/30			-			0.21%	1[0.07,15.26]
Afolabi 2010	0/100	0/100							Not estimable
Al-Sawaf 2013	2/28	1/37		-	+			0.29%	2.64[0.25,27.7]
Atukunda 2014	18/570	14/570			+			3.32%	1.29[0.65,2.56]
Baskett 2007	14/311	7/311			+-	. ,		1.97%	2[0.82,4.89]
	Fav	ours Misoprostol	0.01	0.1	1	10	100	Favours Oxytocin	



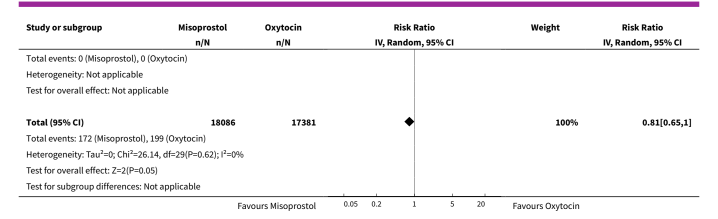




Analysis 64.3. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.3.1 Hospital setting					
Acharya 2001	1/30	1/30		0.6%	1[0.07,15.26
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	0/28	0/37			Not estimable
Alwani 2014	2/100	5/100		1.71%	0.4[0.08,2.01]
Atukunda 2014	7/570	16/570		5.75%	0.44[0.18,1.06
Baskett 2007	0/311	0/311			Not estimable
Bellad 2012	1/321	1/331		0.58%	1.03[0.06,16.42]
Bhatti 2014	1/60	1/60		0.59%	1[0.06,15.62]
Bugalho 2001	2/323	1/339		- 0.78%	2.1[0.19,23.04]
Caliskan 2002	12/396	13/407		7.48%	0.95[0.44,2.05]
Caliskan 2003	14/388	13/384		8.11%	1.07[0.51,2.24]
Chaudhuri 2010	0/96	3/94	<u> </u>	0.51%	0.14[0.01,2.67]
Chaudhuri 2012	5/265	3/265		2.21%	1.67[0.4,6.9]
Cook 1999	5/424	2/129		1.68%	0.76[0.15,3.87]
Diallo 2017	5/154	7/150		3.52%	0.7[0.23,2.14
Dutta 2016	5/200	4/200		2.64%	1.25[0.34,4.59
Elbohoty 2016	1/89	1/86		0.59%	0.97[0.06,15.21]
Fazel 2013	0/50	0/50		0.5570	Not estimable
Gerstenfeld 2001	2/159	0/166		0.49%	5.22[0.25,107.86]
				•	
Gulmezoglu 2001	72/9221	97/9226	•	48.43%	0.74[0.55,1.01
Gupta 2006	0/100	0/100			Not estimable
Karkanis 2002	0/110	0/113		0.700/	Not estimable
Kundodyiwa 2001	2/243	1/256	_	0.78%	2.11[0.19,23.09
Lokugamage 2001	1/20	0/20		0.45%	3[0.13,69.52]
Lumbiganon 1999	0/397	0/200			Not estimable
Modi 2014	0/25	0/25			Not estimable
Nankaly 2016	1/122	5/63		0.99%	0.1[0.01,0.87]
Nasr 2009	8/257	4/257	+	3.16%	2[0.61,6.56]
Oboro 2003	0/247	0/249			Not estimable
Owonikoko 2011	1/50	0/50	•	0.44%	3[0.13,71.92]
Parsons 2006	1/222	2/221	-	0.78%	0.5[0.05,5.45]
Parsons 2007	1/217	5/221	•	0.98%	0.2[0.02,1.73]
Perez-Rumbos 2017	2/195	3/197	+	1.41%	0.67[0.11,3.99]
Rajaei 2014	1/200	4/200 —		0.94%	0.25[0.03,2.22]
Sadiq 2011	0/900	0/884			Not estimable
Shady 2017	13/120	0/120		0.56%	27[1.62,449.1
Singh 2009	0/150	0/75			Not estimable
Sultana 2007	4/210	3/190		2.03%	1.21[0.27,5.32]
Tewatia 2014	0/50	0/50			Not estimable
Vagge 2014	1/100	1/100		0.59%	1[0.06,15.77]
Walley 2000	0/136	1/138	-	0.44%	0.34[0.01,8.23
Zachariah 2006	1/730	2/617		0.78%	0.42[0.04,4.65
Subtotal (95% CI)	18086	17381	•	100%	0.81[0.65,1
Total events: 172 (Misoprostol), 1			Ť		
Heterogeneity: Tau ² =0; Chi ² =26.1	· · ·				
Test for overall effect: Z=2(P=0.05					
64.3.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable





Analysis 64.4. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.

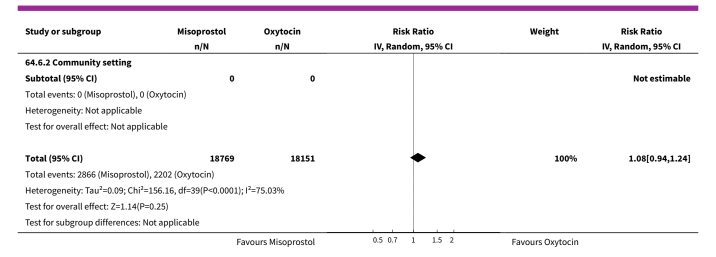
Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.4.1 Hospital setting					
Afolabi 2010	0/100	0/100			Not estimable
Amin 2014	0/100	0/100			Not estimable
Atukunda 2014	11/570	8/570	— 	67.92%	1.38[0.56,3.39]
Chaudhuri 2010	0/96	0/94			Not estimable
Gulmezoglu 2001	4/9224	5/9231		32.08%	0.8[0.22,2.98]
Kundodyiwa 2001	0/243	0/256			Not estimable
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	0/257	0/257			Not estimable
Shrestha 2011	0/100	0/100			Not estimable
Tewatia 2014	0/50	0/50			Not estimable
Subtotal (95% CI)	10840	10858	*	100%	1.16[0.55,2.43]
Total events: 15 (Misoprostol), 13 (O	xytocin)				
Heterogeneity: Tau²=0; Chi²=0.44, di	f=1(P=0.51); I ² =0%				
Test for overall effect: Z=0.38(P=0.7)					
64.4.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Oxy	tocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	10840	10858	•	100%	1.16[0.55,2.43]
Total events: 15 (Misoprostol), 13 (O	xytocin)				
Heterogeneity: Tau²=0; Chi²=0.44, di	f=1(P=0.51); I ² =0%				
Test for overall effect: Z=0.38(P=0.7)					
Test for subgroup differences: Not a	nnlicable				



Analysis 64.6. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.6.1 Hospital setting	0/400	0/400			
Afolabi 2010	0/100	0/100			Not estimable
Al-Sawaf 2013	3/28	2/37		0.58%	1.98[0.35,11.08
Amin 2014	4/100	3/100		0.76%	1.33[0.31,5.81
Asmat 2017	123/839	120/839		4.94%	1.02[0.81,1.29
Atukunda 2014	89/570	57/570		4.44%	1.56[1.14,2.13
Bellad 2012	10/321	30/331		2.32%	0.34[0.17,0.69
Benchimol 2001	52/186	29/196		3.83%	1.89[1.26,2.84
Bhatti 2014	2/60	5/60	•	0.66%	0.4[0.08,1.98
Bugalho 2001	10/323	15/339		2.01%	0.7[0.32,1.53
Caliskan 2002	39/396	33/407	+	3.61%	1.21[0.78,1.89
Caliskan 2003	35/388	28/384	- +	3.41%	1.24[0.77,1.99
Chaudhuri 2010	38/96	51/94		4.46%	0.73[0.54,0.99
Chaudhuri 2012	16/265	15/265		2.38%	1.07[0.54,2.11
Cook 1999	63/424	1/129	—	0.45%	19.17[2.68,136.84
Diallo 2017	10/154	14/150		2.02%	0.7[0.32,1.52
Dutta 2016	6/200	4/200		1%	1.5[0.43,5.23
Elbohoty 2016	49/89	34/86	<u> </u>	4.38%	1.39[1.01,1.92
Gavilanes 2016	33/50	26/50		4.31%	1.27[0.91,1.77
Gerstenfeld 2001	70/154	61/161	 • • • • • • • • • • • • • • • • • • •	4.76%	1.2[0.92,1.56
Gulmezoglu 2001	1793/9213	1248/9227	+	5.67%	1.44[1.35,1.54
Gupta 2006	1/100	0/100		0.18%	3[0.12,72.77
Kundodyiwa 2001	37/243	34/256	'	3.68%	1.15[0.74,1.76
_okugamage 2001	17/20	17/20		4.77%	1[0.77,1.3
Lumbiganon 1999	96/397	52/200		4.57%	0.93[0.69,1.24
Modi 2014	0/25	0/25		4.51 /0	Not estimabl
Mukta 2013	8/100	6/100		1.38%	1.33[0.48,3.7
Musa 2015	15/100	14/100		2.42%	1.07[0.55,2.1
Nasr 2009	17/257	12/257		2.24%	1.42[0.69,2.91
Oboro 2003	3/247	1/249		0.35%	3.02[0.32,28.88
Owonikoko 2011	34/50	27/50		4.4%	1.26[0.92,1.73
Parsons 2006	0/225	5/225		0.22%	0.09[0.01,1.63
Parsons 2007	3/217	6/224	-	0.86%	0.52[0.13,2.04
Penaranda 2002	7/25	8/25		1.81%	0.88[0.37,2.05
Perez-Rumbos 2017	1/195	6/197		0.4%	0.17[0.02,1.39
Sadiq 2011	80/900	161/900		4.82%	0.5[0.39,0.64
Shady 2017	20/120	2/120		0.8%	10[2.39,41.84
Shrestha 2011	4/100	6/100	+	1.02%	0.67[0.19,2.29
Singh 2009	0/150	0/75			Not estimabl
Sultana 2007	8/210	5/190		1.23%	1.45[0.48,4.35
Tewatia 2014	0/50	0/50			Not estimabl
/agge 2014	4/100	3/100	-	0.76%	1.33[0.31,5.81
/imala 2006	47/50	46/50	-	5.55%	1.02[0.92,1.14
Walley 2000	0/202	2/196		0.2%	0.19[0.01,4.02
Zachariah 2006	19/730	13/617		2.33%	1.24[0.62,2.48
Subtotal (95% CI)	18769	18151	•	100%	1.08[0.94,1.24
Fotal events: 2866 (Misoprostol)					
Heterogeneity: Tau ² =0.09; Chi ² =	· · · · · · · · · · · · · · · · · · ·	²=75.03%			
Test for overall effect: Z=1.14(P=					
	/				

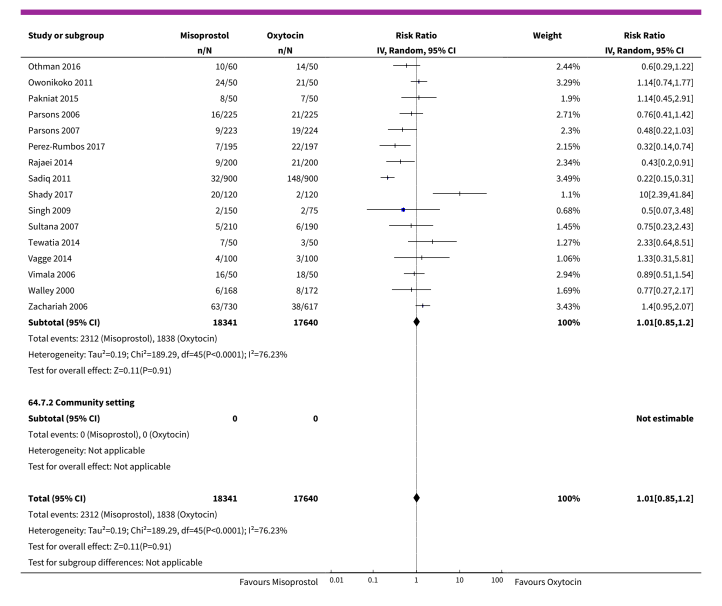




Analysis 64.7. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.7.1 Hospital setting					
Acharya 2001	2/30	3/30		0.83%	0.67[0.12,3.71]
Afolabi 2010	3/100	4/100		1.06%	0.75[0.17,3.27]
Al-Sawaf 2013	3/28	2/37		0.83%	1.98[0.35,11.08]
Alwani 2014	4/100	9/100		1.5%	0.44[0.14,1.4]
Atukunda 2014	47/570	31/570	 +-	3.28%	1.52[0.98,2.35]
Baskett 2007	159/311	126/311	+	3.97%	1.26[1.06,1.5]
Bellad 2012	1/321	8/331 -	•	0.61%	0.13[0.02,1.02]
Bhatti 2014	1/60	3/60		0.54%	0.33[0.04,3.11]
Bugalho 2001	7/323	7/339		1.69%	1.05[0.37,2.96]
Caliskan 2002	33/396	26/407	+-	3.11%	1.3[0.8,2.14]
Caliskan 2003	23/388	26/384	 -	2.96%	0.88[0.51,1.51]
Chaudhuri 2010	11/96	14/94		2.39%	0.77[0.37,1.61]
Chaudhuri 2012	20/265	23/265	-	2.86%	0.87[0.49,1.54]
Cook 1999	95/424	6/129		2.22%	4.82[2.16,10.73]
Diallo 2017	7/154	6/150		1.63%	1.14[0.39,3.3]
Eftekhari 2009	7/50	16/50		2.23%	0.44[0.2,0.97]
Elbohoty 2016	20/89	11/86	 	2.57%	1.76[0.9,3.45]
Gavilanes 2016	10/50	12/50		2.37%	0.83[0.4,1.75]
Gerstenfeld 2001	36/159	18/166		3.02%	2.09[1.24,3.52]
Gulmezoglu 2001	1398/9225	1002/9228	+	4.1%	1.4[1.29,1.51]
Gupta 2006	5/100	1/100	-	0.58%	5[0.59,42.04]
Karkanis 2002	28/110	20/113	+-	3.06%	1.44[0.86,2.4]
Kundodyiwa 2001	13/243	7/256	+	1.97%	1.96[0.79,4.82]
Lokugamage 2001	6/20	1/20	+	0.64%	6[0.79,45.42]
Lumbiganon 1999	41/397	28/200	-+	3.25%	0.74[0.47,1.16]
Modi 2014	0/25	0/25			Not estimable
Mukta 2013	22/100	16/100	+	2.84%	1.38[0.77,2.46]
Musa 2015	20/100	19/100	+	2.9%	1.05[0.6,1.85]
Nankaly 2016	15/122	9/63	- 	2.3%	0.86[0.4,1.86]
Nasr 2009	6/257	4/257		1.33%	1.5[0.43,5.25]
Oboro 2003	31/247	27/249	+	3.14%	1.16[0.71,1.88]

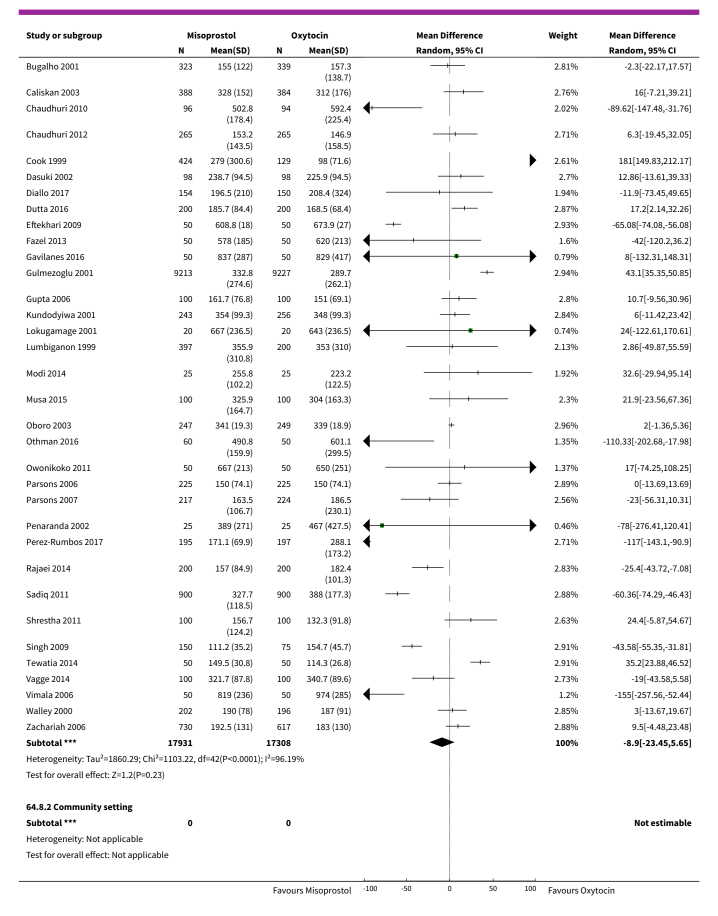




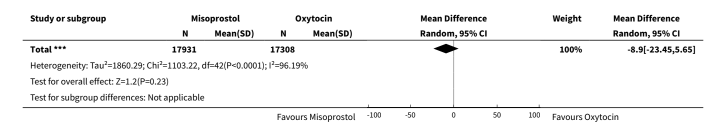
Analysis 64.8. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Mis	oprostol	0	xytocin		Me	ean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Ra	ndom, 95% CI				Random, 95% CI
64.8.1 Hospital setting											
Acharya 2001	30	545 (192.8)	30	533 (296.2)	\leftarrow				\rightarrow	0.91%	12[-114.47,138.47]
Afolabi 2010	100	153.2 (58)	100	155.6 (58)			-			2.86%	-2.4[-18.47,13.67]
Al-Sawaf 2013	28	348 (112)	37	314.7 (94.6)			+		_	2.16%	33.3[-18.18,84.78]
Amin 2014	100	300 (262.8)	100	250 (262.8)				-	\rightarrow	1.7%	50[-22.83,122.83]
Asmat 2017	839	322 (199.9)	839	337 (211.4)		_	+			2.81%	-15[-34.69,4.69]
Atukunda 2014	570	341.5 (206.2)	570	304.2 (190.8)						2.76%	37.3[14.24,60.36]
Bellad 2012	321	192 (124)	331	366 (135.9)	•					2.81%	-174[-193.96,-154.04]
Benchimol 2001	186	374 (238.4)	196	278 (254)				-	-	2.21%	96[46.63,145.37]
Bhatti 2014	60	200 (125)	60	360 (136)	4					2.27%	-160[-206.74,-113.26]
			Favou	rs Misoprostol	-100	-50	0	50	100	Favours Oxy	rtocin .





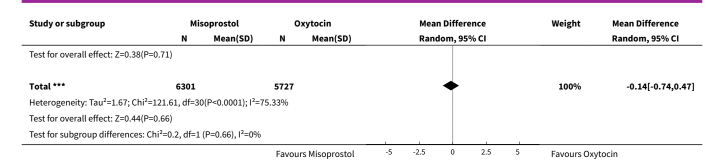




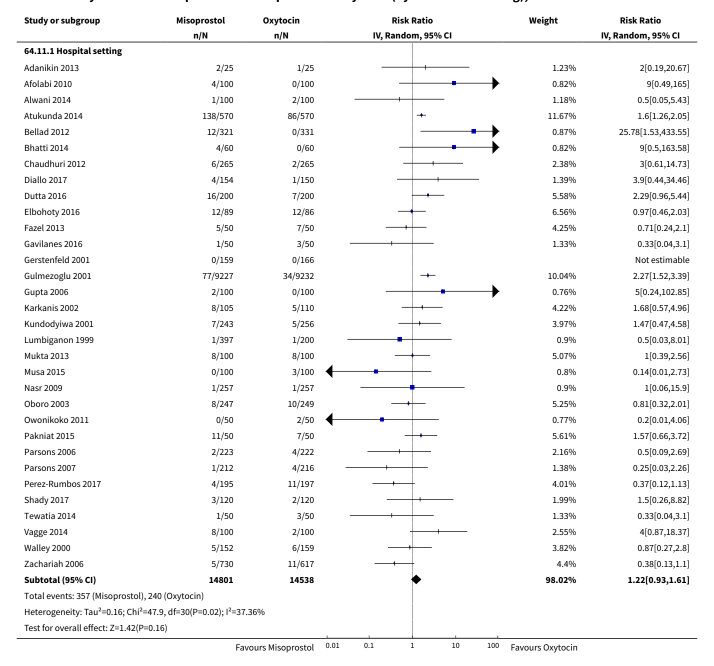
Analysis 64.9. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol	0:	kytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
64.9.1 Hospital setting							
Acharya 2001	30	12 (8.4)	30	11 (7)		1.7%	1[-2.9,4.9]
Afolabi 2010	100	3 (10.2)	100	4 (10.2)		2.55%	-1[-3.83,1.83]
Al-Sawaf 2013	28	13 (9)	37	12 (9)		1.42%	1[-3.42,5.42]
Alwani 2014	100	12 (13.1)	100	11.2 (9)		2.28%	0.8[-2.32,3.92]
Atukunda 2014	570	9.4 (14.3)	569	9.6 (16.7)		3.8%	-0.2[-2.01,1.61]
Benchimol 2001	186	9.6 (23.5)	196	6.7 (20.1)		1.43%	2.9[-1.5,7.3]
Caliskan 2002	396	15 (12)	407	14 (14)	+	3.81%	1[-0.8,2.8]
Caliskan 2003	388	14 (11)	384	14 (14)		3.84%	0[-1.78,1.78]
Chaudhuri 2010	96	9.3 (5.7)	94	11.4 (9.9)		3.14%	-2.1[-4.4,0.2]
Chaudhuri 2012	265	6.1 (6.2)	265	6.3 (4.8)		5.04%	-0.2[-1.14,0.74]
Cook 1999	424	6.9 (17.5)	129	14.7 (12.1)		2.71%	-7.8[-10.47,-5.13]
Diallo 2017	154	3.8 (9)	150	2.9 (13)		2.88%	0.9[-1.62,3.42]
Dutta 2016	200	9 (3.2)	200	8.4 (2.8)	+	5.44%	0.59[0,1.18]
Eftekhari 2009	50	7.4 (5.4)	50	6.6 (5.6)		3.32%	0.8[-1.36,2.96]
Gupta 2006	100	2.5 (2.2)	100	2.3 (2.2)	+	5.42%	0.23[-0.38,0.84]
Karkanis 2002	110	16 (10.7)	113	14 (10.8)	-	2.55%	2[-0.83,4.83]
Lokugamage 2001	19	14.3 (10.1)	19	13.2 (10.1)		0.77%	1.1[-5.35,7.55]
Musa 2015	100	8.8 (5.8)	100	8.1 (6)		4.05%	0.7[-0.94,2.34]
Oboro 2003	247	7.1 (9.4)	249	6.8 (7.8)		4.22%	0.3[-1.22,1.82]
Pakniat 2015	50	11.4 (2.9)	50	13.8 (1.3)		5.12%	-2.4[-3.28,-1.52]
Parsons 2006	225	10.7 (11.4)	225	10 (10.4)	- +	3.51%	0.7[-1.32,2.72]
Parsons 2007	220	11.9 (13)	220	11.6 (11)		3.2%	0.3[-1.95,2.55]
Perez-Rumbos 2017	195	24 (19)	197	25 (17)		1.92%	-1[-4.57,2.57]
Rajaei 2014	200	7 (9)	200	8 (11)		3.57%	-1[-2.97,0.97]
Sadiq 2011	530	5.5 (6.9)	418	7.1 (6.1)	→	5.18%	-1.6[-2.43,-0.77]
Shrestha 2011	100	10 (21)	100	9 (19)	-	0.99%	1[-4.55,6.55]
Tewatia 2014	50	5 (2)	50	3 (2)	-	5.24%	2[1.22,2.78]
Vimala 2006	50	4 (16)	50	6 (18)		0.72%	-2[-8.68,4.68]
Walley 2000	200	6 (9.3)	192	5.5 (7.3)		4.02%	0.5[-1.15,2.15]
Zachariah 2006	730	13.8 (10.7)	617	14.9 (13.6)		4.5%	-1.1[-2.42,0.22]
Subtotal ***	6113		5611		•	98.34%	-0.15[-0.77,0.46]
Heterogeneity: Tau ² =1.69; Chi ² =1	21.46, df=29	(P<0.0001); I ² =7	6.12%				
Test for overall effect: Z=0.49(P=0	.63)						
64.9.2 Community setting							
Diop 2016	188	3.5 (16.1)	116	2.7 (17.8)		1.66%	0.76[-3.21,4.73]
Subtotal ***	188		116			1.66%	0.76[-3.21,4.73]
Heterogeneity: Not applicable							

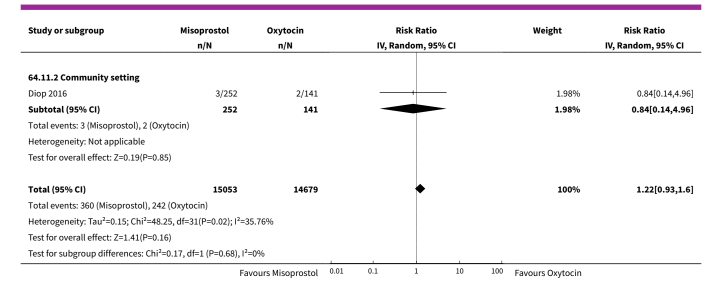




Analysis 64.11. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 11 Nausea.



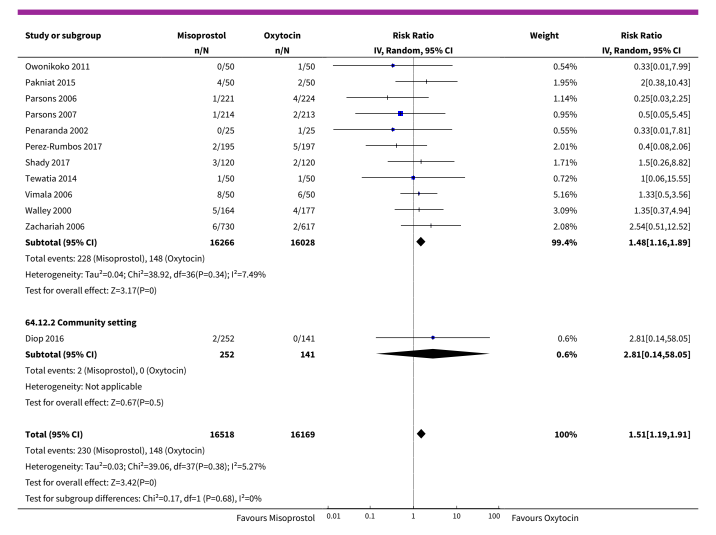




Analysis 64.12. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.12.1 Hospital setting					
Acharya 2001	2/30	3/30		1.81%	0.67[0.12,3.71]
Adanikin 2013	2/25	2/25		1.52%	1[0.15,6.55]
Afolabi 2010	0/100	0/100			Not estimable
Alwani 2014	1/100	1/100		0.72%	1[0.06,15.77]
Amin 2014	12/100	2/100		2.44%	6[1.38,26.12]
Atukunda 2014	35/569	19/570		13.68%	1.85[1.07,3.19]
Bellad 2012	5/321	0/331	+	0.65%	11.34[0.63,204.28]
Benchimol 2001	7/186	1/196	 	1.24%	7.38[0.92,59.38]
Bhatti 2014	3/60	1/60	- 	1.09%	3[0.32,28.03]
Bugalho 2001	2/323	1/337		0.95%	2.09[0.19,22.9]
Caliskan 2002	2/396	2/407		1.41%	1.03[0.15,7.26]
Caliskan 2003	4/388	3/384		2.38%	1.32[0.3,5.86]
Chaudhuri 2010	2/96	3/94		1.72%	0.65[0.11,3.82]
Chaudhuri 2012	5/265	2/265	- 	2%	2.5[0.49,12.77]
Diallo 2017	2/154	1/150		0.95%	1.95[0.18,21.26]
Dutta 2016	4/200	3/200		2.4%	1.33[0.3,5.88]
Elbohoty 2016	9/89	7/86		5.57%	1.24[0.48,3.19]
Fazel 2013	2/50	3/50		1.76%	0.67[0.12,3.82]
Gavilanes 2016	1/50	1/50		0.72%	1[0.06,15.55]
Gerstenfeld 2001	0/159	0/166			Not estimable
Gulmezoglu 2001	66/9227	25/9232	-	17.44%	2.64[1.67,4.18]
Karkanis 2002	6/105	4/110	- - 	3.38%	1.57[0.46,5.41]
Kundodyiwa 2001	2/243	2/256		1.41%	1.05[0.15,7.42]
Lumbiganon 1999	0/397	1/200		0.54%	0.17[0.01,4.11]
Mukta 2013	4/100	2/100	- +	1.9%	2[0.37,10.67]
Musa 2015	0/100	0/100			Not estimable
Nasr 2009	6/257	5/257	- 	3.73%	1.2[0.37,3.88]
Oboro 2003	12/247	9/249	-	6.73%	1.34[0.58,3.13]
Othman 2016	1/60	15/50	— —	1.36%	0.06[0.01,0.41]

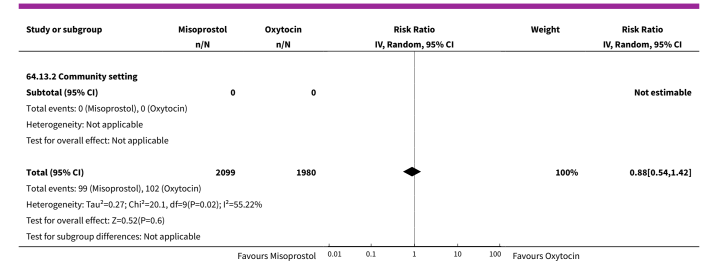




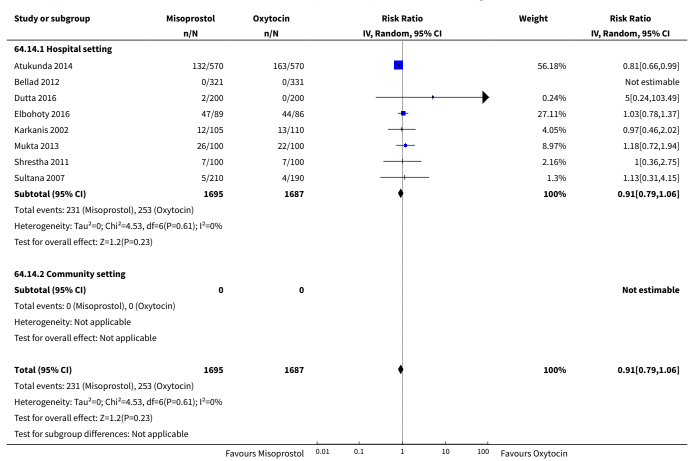
Analysis 64.13. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
64.13.1 Hospital setting						
Atukunda 2014	10/570	11/570		13.25%	0.91[0.39,2.12]	
Dutta 2016	5/200	3/200	- •	7.61%	1.67[0.4,6.88]	
Elbohoty 2016	24/89	24/86	+	18.39%	0.97[0.6,1.56]	
Gavilanes 2016	0/50	1/50	+	2.08%	0.33[0.01,7.99]	
Karkanis 2002	9/105	4/110	+-	9.87%	2.36[0.75,7.42]	
Othman 2016	3/60	20/50		9.81%	0.13[0.04,0.4]	
Owonikoko 2011	1/50	3/50		3.85%	0.33[0.04,3.1]	
Perez-Rumbos 2017	39/195	26/197	-	18.77%	1.52[0.96,2.39]	
Vimala 2006	6/50	8/50		11.6%	0.75[0.28,2]	
Zachariah 2006	2/730	2/617		4.76%	0.85[0.12,5.98]	
Subtotal (95% CI)	2099	1980	*	100%	0.88[0.54,1.42]	
Total events: 99 (Misoprostol), 102	(Oxytocin)					
Heterogeneity: Tau ² =0.27; Chi ² =20.	.1, df=9(P=0.02); I ² =55.2	2%				
Test for overall effect: Z=0.52(P=0.6	5)					
	Fav	ours Misoprostol 0	0.01 0.1 1 10 1	00 Favours Oxytocin		



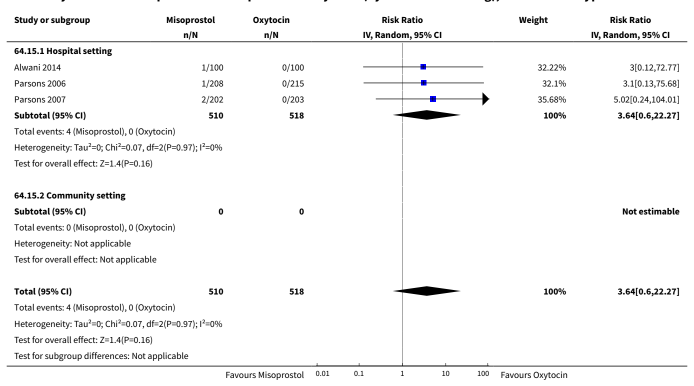


Analysis 64.14. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 14 Abdominal pain.





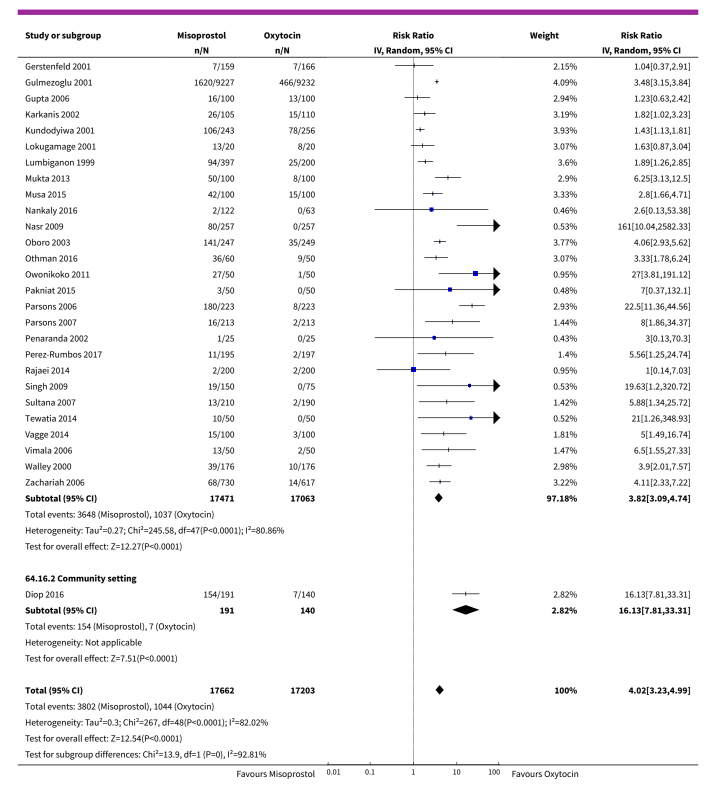
Analysis 64.15. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 15 Hypertension.



Analysis 64.16. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.16.1 Hospital setting					
Acharya 2001	2/30	2/30		1%	1[0.15,6.64]
Adanikin 2013	1/25	2/25		0.72%	0.5[0.05,5.17]
Afolabi 2010	4/100	2/100		1.2%	2[0.37,10.67]
Alwani 2014	7/100	1/100	-	0.86%	7[0.88,55.86]
Amin 2014	25/100	4/100		2.16%	6.25[2.26,17.31]
Atukunda 2014	321/570	168/570	+	4.04%	1.91[1.65,2.21]
Baskett 2007	21/311	0/311		0.53%	43[2.62,706.74]
Bellad 2012	173/321	14/331		3.33%	12.74[7.56,21.49]
Benchimol 2001	5/186	0/196		0.5%	11.59[0.65,208.12]
Bhatti 2014	32/60	3/60		1.95%	10.67[3.45,32.96]
Bugalho 2001	123/323	51/337	+	3.84%	2.52[1.89,3.36]
Caliskan 2002	47/396	16/407		3.26%	3.02[1.74,5.23]
Caliskan 2003	44/388	19/384		3.34%	2.29[1.36,3.85]
Chaudhuri 2010	8/96	1/94		0.88%	7.83[1,61.42]
Chaudhuri 2012	51/265	2/265		1.52%	25.5[6.27,103.67]
Dasuki 2002	13/98	2/98		1.44%	6.5[1.51,28.05]
Diallo 2017	11/154	3/150		1.73%	3.57[1.02,12.55]
Dutta 2016	41/200	4/200		2.18%	10.25[3.74,28.08]
Elbohoty 2016	28/89	16/86		3.29%	1.69[0.99,2.9]
Fazel 2013	8/50	1/50		0.89%	8[1.04,61.62]
Gavilanes 2016	33/50	1/50		0.95%	33[4.69,232.05]
	Fav	ours Misoprostol 0.	.01 0.1 1 10 10	DO Favours Oxytocin	



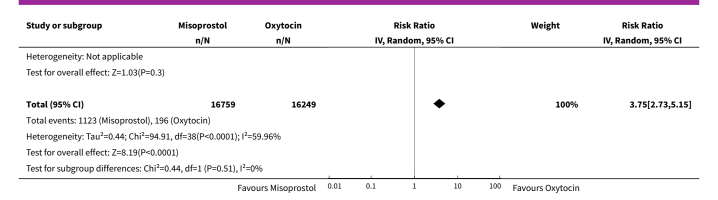




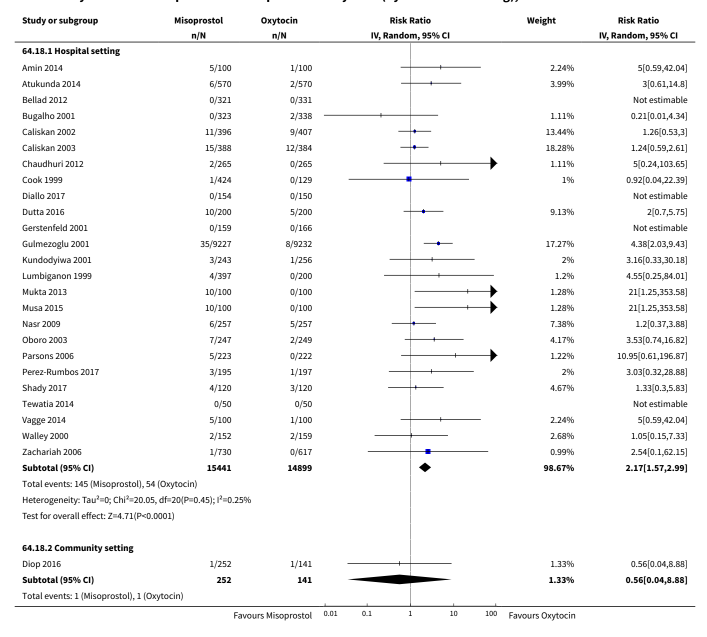
Analysis 64.17. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol	Oxytocin	Risk Ratio	Weight	Risk Ratio
CA 17 1 Hassital auttima	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
64.17.1 Hospital setting	4/25	2/25		2.240/	2[0.4.0.05
Adanikin 2013	4/25	2/25		2.34%	2[0.4,9.95
Afolabi 2010	0/100	0/100	.	2 220/	Not estimable
Alwani 2014	4/100	2/100		2.22%	2[0.37,10.67
Amin 2014	15/100	3/100		3.17%	5[1.49,16.74
Atukunda 2014	53/570	12/570	-	4.82%	4.42[2.39,8.18
Baskett 2007	39/311	1/311		1.78%	39[5.39,282.1
Bellad 2012	4/321	0/331	-	0.98%	9.28[0.5,171.66
Benchimol 2001	6/186	0/196	+ + +	1.01%	13.7[0.78,241.41
Bhatti 2014	2/60	0/60	-	0.93%	5[0.25,102
Caliskan 2002	16/396	6/407		3.91%	2.74[1.08,6.93
Caliskan 2003	17/388	5/384		3.75%	3.36[1.25,9.03
Chaudhuri 2010	2/96	4/94		2.23%	0.49[0.09,2.61
Chaudhuri 2012	6/265	0/265	+ +	1.01%	13[0.74,229.61
Diallo 2017	4/154	1/150	+	1.55%	3.9[0.44,34.46
Dutta 2016	17/200	1/200		1.75%	17[2.28,126.53
Elbohoty 2016	31/89	12/86		4.88%	2.5[1.37,4.53
Gulmezoglu 2001	559/9198	78/9205	+	5.71%	7.17[5.67,9.07
Gupta 2006	2/100	0/100	-	0.92%	5[0.24,102.85
Karkanis 2002	20/107	12/112	 	4.68%	1.74[0.9,3.39
Kundodyiwa 2001	18/243	1/256		1.75%	18.96[2.55,140.96
Lokugamage 2001	0/20	0/20			Not estimable
Lumbiganon 1999	19/394	6/199	+-	3.99%	1.6[0.65,3.94
Mukta 2013	6/100	2/100	+	2.39%	3[0.62,14.51
Musa 2015	16/100	4/100		3.55%	4[1.39,11.55
Nankaly 2016	2/122	2/63		1.84%	0.52[0.07,3.58
Nasr 2009	48/257	2/257			24[5.9,97.7
Oboro 2003	3/247	1/249		1.47%	3.02[0.32,28.88
Othman 2016	1/60	4/50		1.57%	0.21[0.02,1.8
Pakniat 2015	2/50	1/50		1.37%	2[0.19,21.36
Parsons 2006	24/211	0/218	<u> </u>	1.05%	50.62[3.1,827.06
Parsons 2007	8/200	4/209		3.23%	2.09[0.64,6.83
Perez-Rumbos 2017	4/195	7/197		3.16%	0.58[0.17,1.94
Rajaei 2014	29/200	4/200		3.64%	7.25[2.6,20.24
Shrestha 2011	16/100	4/100		3.55%	4[1.39,11.55
Singh 2009	25/150	0/75		1.06%	25.67[1.58,415.93
Tewatia 2014	13/50	0/50		1.05%	27[1.65,442.17
Vagge 2014	12/100	2/100	<u> </u>	2.59%	6[1.38,26.12
Vimala 2006	8/50	2/50	<u> </u>	2.54%	4[0.89,17.91
Walley 2000	12/162	5/152	<u> </u>	3.66%	
Zachariah 2006			<u> </u>	3.67%	2.25[0.81,6.24
	48/730	4/617	•		10.14[3.68,27.97
Subtotal (95% CI)	16507	16108	_	97.53%	3.8[2.76,5.25
Total events: 1115 (Misoprostol),		00.000/			
Heterogeneity: Tau²=0.45; Chi²=9 Test for overall effect: Z=8.13(P<0		=6U.66%			
	,				
64.17.2 Community setting	0/252	2/1/1		2.470/	2.24[0.40.10.4
Diop 2016	8/252	2/141		2.47%	2.24[0.48,10.4
Subtotal (95% CI)	252 Oxytocin)	141		2.47%	2.24[0.48,10.4

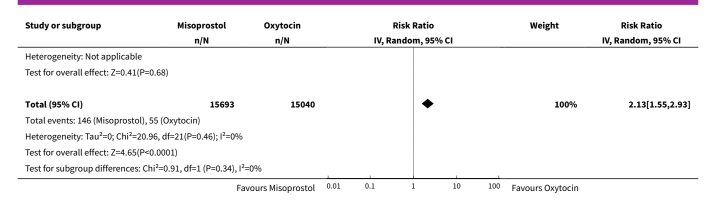




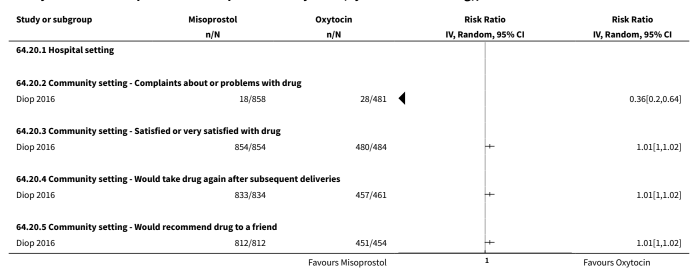
Analysis 64.18. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 18 Diarrhoea.







Analysis 64.20. Comparison 64 Misoprostol vs Oxytocin (by healthcare setting), Outcome 20 Maternal satisfaction.



Comparison 65. Injectable prostaglandins vs Oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	100	Risk Ratio (IV, Random, 95% CI)	1.43 [0.20, 10.31]
2.1 Hospital setting	2	100	Risk Ratio (IV, Random, 95% CI)	1.43 [0.20, 10.31]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	250	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.65]



Outcome or subgroup ti- tle	ome or subgroup ti- No. of studies No. of partici- pants		Statistical method	Effect size	
3.1 Hospital setting	2	250	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.65]	
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5 Severe maternal morbid- ity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
6 PPH >= 500 mL	4	500	Risk Ratio (IV, Random, 95% CI)	0.84 [0.26, 2.71]	
6.1 Hospital setting	4	500	Risk Ratio (IV, Random, 95% CI)	0.84 [0.26, 2.71]	
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
7 Additional uterotonics	4	500	Risk Ratio (IV, Random, 95% CI)	0.29 [0.09, 0.94]	
7.1 Hospital setting	4	500	Risk Ratio (IV, Random, 95% CI)	0.29 [0.09, 0.94]	
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
8 Blood loss	4	500	Mean Difference (IV, Random, 95% CI)	-15.83 [-152.28, 120.62]	
8.1 Hospital setting	4	500	Mean Difference (IV, Random, 95% CI)	-15.83 [-152.28, 120.62]	
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]	
11 Nausea	3	450	Risk Ratio (IV, Random, 95% CI)	1.17 [0.40, 3.41]	
11.1 Hospital setting	3	450	Risk Ratio (IV, Random, 95% CI)	1.17 [0.40, 3.41]	



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	400	Risk Ratio (IV, Random, 95% CI)	2.48 [0.57, 10.73]
12.1 Hospital setting	2	400	Risk Ratio (IV, Random, 95% CI)	2.48 [0.57, 10.73]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	200	Risk Ratio (IV, Random, 95% CI)	0.20 [0.01, 4.11]
13.1 Hospital setting	1	200	Risk Ratio (IV, Random, 95% CI)	0.20 [0.01, 4.11]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	400	Risk Ratio (IV, Random, 95% CI)	0.91 [0.11, 7.73]
16.1 Hospital setting	2	400	Risk Ratio (IV, Random, 95% CI)	0.91 [0.11, 7.73]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	200	Risk Ratio (IV, Random, 95% CI)	2.0 [0.18, 21.71]
17.1 Hospital setting	1	200	Risk Ratio (IV, Random, 95% CI)	2.0 [0.18, 21.71]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	400	Risk Ratio (IV, Random, 95% CI)	10.38 [1.96, 54.98]
18.1 Hospital setting	2	400	Risk Ratio (IV, Random, 95% CI)	10.38 [1.96, 54.98]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

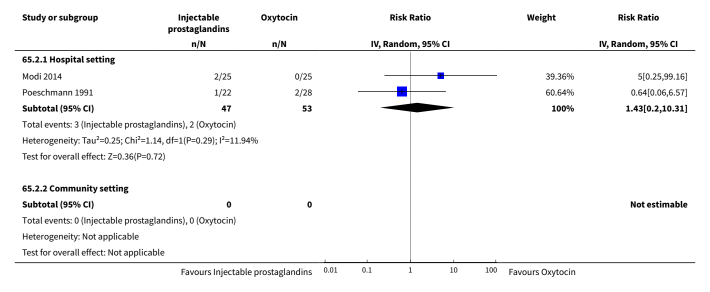


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

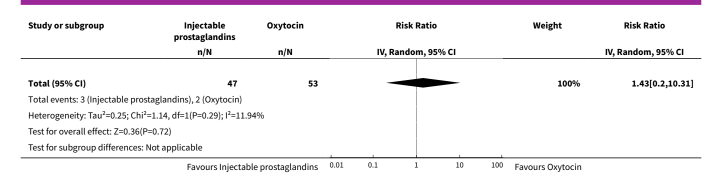
Analysis 65.1. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI				IV, Random, 95% CI
65.1.1 Hospital setting								
Kumar 2016	0/100	0/100						Not estimable
Subtotal (95% CI)	100	100						Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Oxytocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	e							
65.1.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Oxytocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	e							
Total (95% CI)	100	100						Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Oxytocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	e							
Test for subgroup differences: Not a	pplicable							
	Favours Injectab	e prostaglandins	0.01	0.1	1 10	100 Fa	vours Oxytocin	

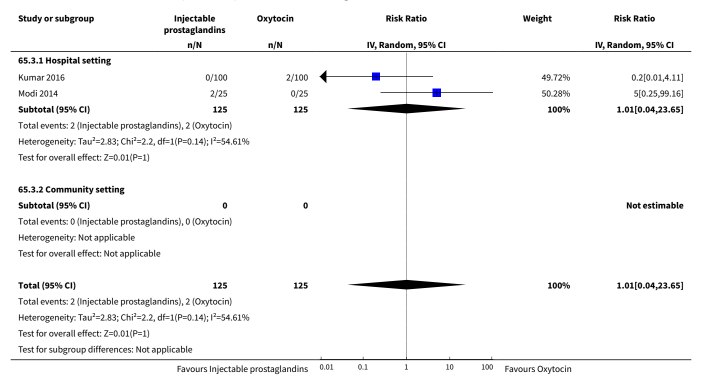
Analysis 65.2. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.







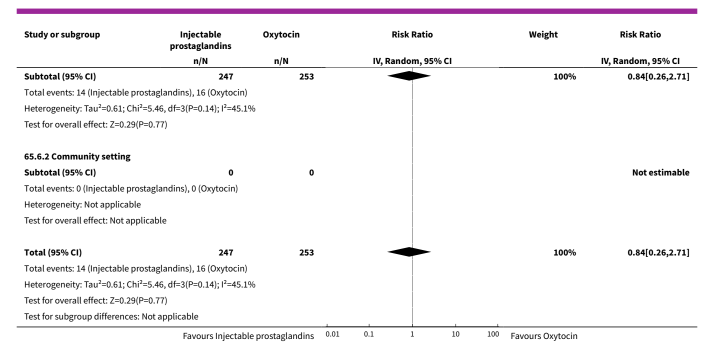
Analysis 65.3. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.



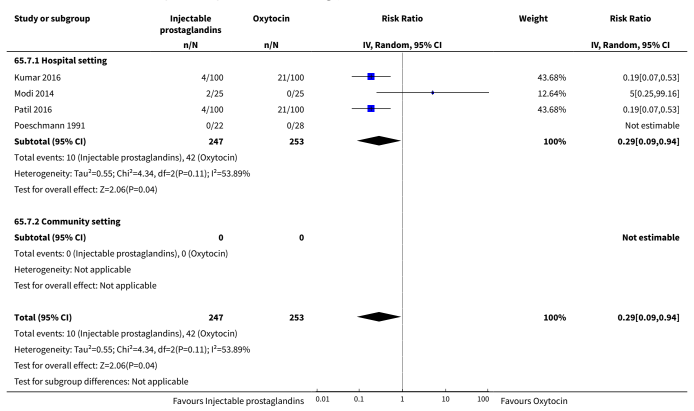
Analysis 65.6. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable Oxytocin prostaglandins				Risk Ratio		Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
65.6.1 Hospital setting									
Kumar 2016	3/100	7/100			•			33.6%	0.43[0.11,1.61]
Modi 2014	6/25	0/25			+		\rightarrow	13.32%	13[0.77,219.11]
Patil 2016	0/100	2/100	\leftarrow	+		_		11.97%	0.2[0.01,4.11]
Poeschmann 1991	5/22	7/28			-	1		41.11%	0.91[0.33,2.48]
	Favours Injectab	le prostaglandins	0.01	0.1	1	10	100	Favours Oxvtocin	



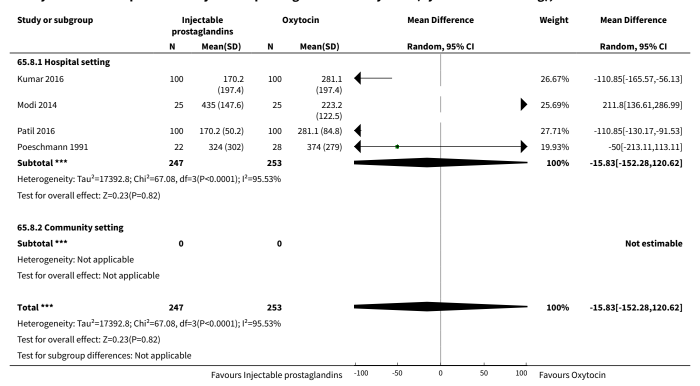


Analysis 65.7. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.





Analysis 65.8. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.

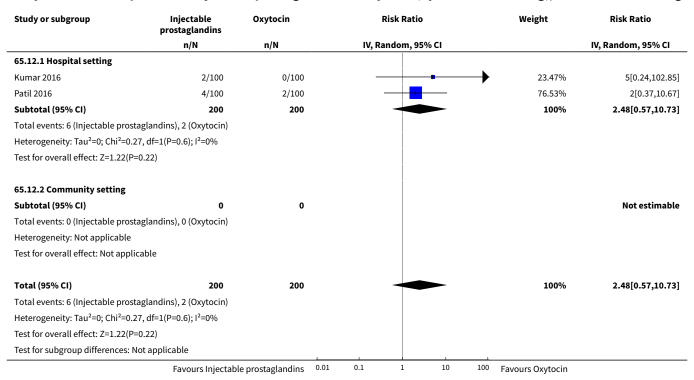


Analysis 65.11. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
65.11.1 Hospital setting					
Kumar 2016	2/100	2/100		30.5%	1[0.14,6.96]
Patil 2016	5/100	4/100	- 1	69.5%	1.25[0.35,4.52]
Poeschmann 1991	0/22	0/28			Not estimable
Subtotal (95% CI)	222	228		100%	1.17[0.4,3.41]
Total events: 7 (Injectable prostagla	indins), 6 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.04, d	f=1(P=0.85); I ² =0%				
Test for overall effect: Z=0.28(P=0.78	3)				
65.11.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	indins), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicabl	e				
Total (95% CI)	222	228	*	100%	1.17[0.4,3.41]
Total events: 7 (Injectable prostagla	indins), 6 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.04, d	f=1(P=0.85); I ² =0%				
Test for overall effect: Z=0.28(P=0.78	3)				
Test for subgroup differences: Not a	pplicable				
	Favours Injectab	e prostaglandins 0.01	0.1 1 10 1	LOO Favours Oxytocin	



Analysis 65.12. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.

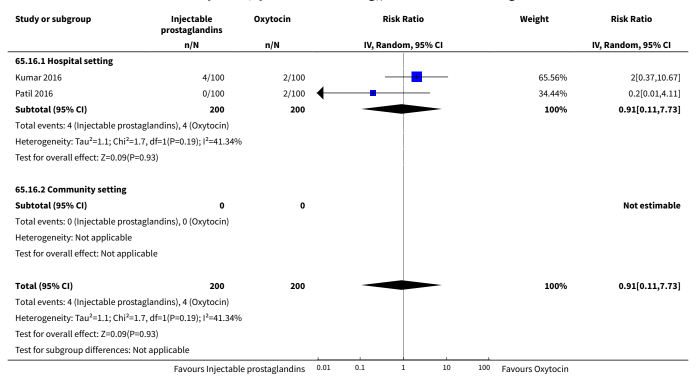


Analysis 65.13. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Injectable prostaglandins	Oxytocin		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV	, Random, 95% CI		IV, Random, 95% CI
65.13.1 Hospital setting						
Patil 2016	0/100	2/100			100%	0.2[0.01,4.11]
Subtotal (95% CI)	100	100			100%	0.2[0.01,4.11]
Total events: 0 (Injectable prostagla	andins), 2 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0, df=0	0(P<0.0001); I ² =100%					
Test for overall effect: Z=1.04(P=0.3)					
65.13.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Oxytocin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicab	le					
Total (95% CI)	100	100			100%	0.2[0.01,4.11]
Total events: 0 (Injectable prostagla	andins), 2 (Oxytocin)					
Heterogeneity: Tau ² =0; Chi ² =0, df=0	0(P<0.0001); I ² =100%			į		
Test for overall effect: Z=1.04(P=0.3)			į		
Test for subgroup differences: Not a	applicable			İ		
	Favours Injectabl	e prostaglandins	0.01 0.1	1 10	100 Favours Oxytocin	



Analysis 65.16. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 16 Shivering.

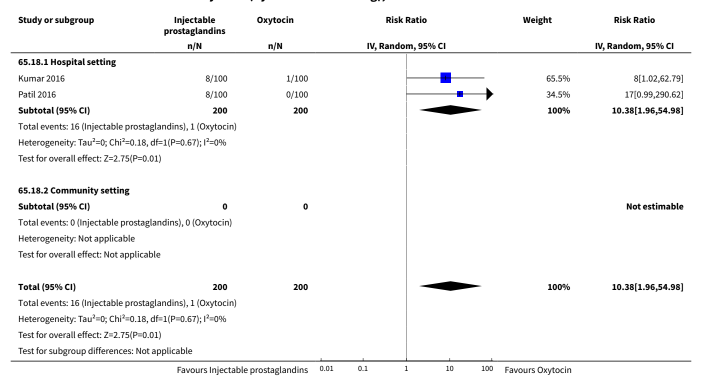


Analysis 65.17. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	subgroup Injectable Oxytocin Risk Ratio prostaglandins		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
65.17.1 Hospital setting					
Kumar 2016	2/100	1/100	- •	100%	2[0.18,21.71]
Subtotal (95% CI)	100	100		100%	2[0.18,21.71]
Total events: 2 (Injectable prostagla	andins), 1 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.57(P=0.5	7)				
65.17.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	100	100		100%	2[0.18,21.71]
Total events: 2 (Injectable prostagla	andins), 1 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.57(P=0.5	7)				
Test for subgroup differences: Not a	applicable				
	Favours Injectabl	e prostaglandins 0.0	1 0.1 1 10 1	00 Favours Oxytocin	



Analysis 65.18. Comparison 65 Injectable prostaglandins vs Oxytocin (by healthcare setting), Outcome 18 Diarrhoea.



Comparison 66. Carbetocin vs Oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	5	30327	Risk Ratio (IV, Random, 95% CI)	2.00 [0.37, 10.92]
1.1 Hospital setting	5	30327	Risk Ratio (IV, Random, 95% CI)	2.00 [0.37, 10.92]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	10	30633	Risk Ratio (IV, Random, 95% CI)	0.73 [0.45, 1.19]
2.1 Hospital setting	10	30633	Risk Ratio (IV, Random, 95% CI)	0.73 [0.45, 1.19]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	16	32115	Risk Ratio (IV, Random, 95% CI)	0.68 [0.38, 1.22]
3.1 Hospital setting	16	32115	Risk Ratio (IV, Random, 95% CI)	0.68 [0.38, 1.22]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	2	29847	Risk Ratio (IV, Random, 95% CI)	1.16 [0.67, 2.02]
4.1 Hospital setting	2	29847	Risk Ratio (IV, Random, 95% CI)	1.16 [0.67, 2.02]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal mor- bidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	30633	Risk Ratio (IV, Random, 95% CI)	0.75 [0.58, 0.98]
6.1 Hospital setting	11	30633	Risk Ratio (IV, Random, 95% CI)	0.75 [0.58, 0.98]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	22	32699	Risk Ratio (IV, Random, 95% CI)	0.48 [0.34, 0.68]
7.1 Hospital setting	22	32699	Risk Ratio (IV, Random, 95% CI)	0.48 [0.34, 0.68]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	16	2115	Mean Difference (IV, Random, 95% CI)	-92.73 [-154.97, -30.49]
8.1 Hospital setting	16	2115	Mean Difference (IV, Random, 95% CI)	-92.73 [-154.97, -30.49]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	12	2096	Mean Difference (IV, Random, 95% CI)	-1.66 [-3.81, 0.50]
9.1 Hospital setting	12	2096	Mean Difference (IV, Random, 95% CI)	-1.66 [-3.81, 0.50]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	2	190	Risk Ratio (IV, Random, 95% CI)	0.94 [0.86, 1.03]
10.1 Hospital setting	2	190	Risk Ratio (IV, Random, 95% CI)	0.94 [0.86, 1.03]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	12	2288	Risk Ratio (IV, Random, 95% CI)	1.11 [0.78, 1.56]
11.1 Hospital setting	12	2288	Risk Ratio (IV, Random, 95% CI)	1.11 [0.78, 1.56]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	13	31833	Risk Ratio (IV, Random, 95% CI)	0.90 [0.53, 1.50]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
12.1 Hospital setting	13	31833	Risk Ratio (IV, Random, 95% CI)	0.90 [0.53, 1.50]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	15	2620	Risk Ratio (IV, Random, 95% CI)	0.84 [0.63, 1.12]
13.1 Hospital setting	15	2620	Risk Ratio (IV, Random, 95% CI)	0.84 [0.63, 1.12]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	10	31293	Risk Ratio (IV, Random, 95% CI)	1.18 [0.97, 1.44]
14.1 Hospital setting	10	31293	Risk Ratio (IV, Random, 95% CI)	1.18 [0.97, 1.44]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	9	1998	Risk Ratio (IV, Random, 95% CI)	0.78 [0.49, 1.23]
16.1 Hospital setting	9	1998	Risk Ratio (IV, Random, 95% CI)	0.78 [0.49, 1.23]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	4	466	Risk Ratio (IV, Random, 95% CI)	1.58 [0.27, 9.35]
17.1 Hospital setting	4	466	Risk Ratio (IV, Random, 95% CI)	1.58 [0.27, 9.35]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



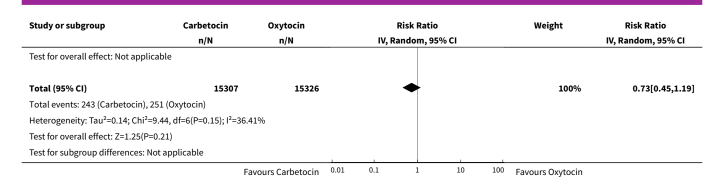
Analysis 66.1. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup C	arbetocin	Oxytocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
66.1.1 Hospital setting						
Attilakos 2010	0/188	0/189				Not estimable
Boucher 1998	0/29	0/28				Not estimable
El Behery 2015	0/90	0/90				Not estimable
Elbohoty 2016	0/88	0/86				Not estimable
Widmer 2018	4/14771	2/14768		1	100%	2[0.37,10.92]
Subtotal (95% CI)	15166	15161			100%	2[0.37,10.92]
Total events: 4 (Carbetocin), 2 (Oxytocin)						
Heterogeneity: Not applicable						
Test for overall effect: Z=0.8(P=0.42)						
66.1.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (Oxytocin)						
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Total (95% CI)	15166	15161			100%	2[0.37,10.92]
Total events: 4 (Carbetocin), 2 (Oxytocin)						
Heterogeneity: Not applicable						
Test for overall effect: Z=0.8(P=0.42)						
Test for subgroup differences: Not application	able					
	Fa	vours Carbetocin	0.01 0.1	1 10	100 Favours Oxytocin	

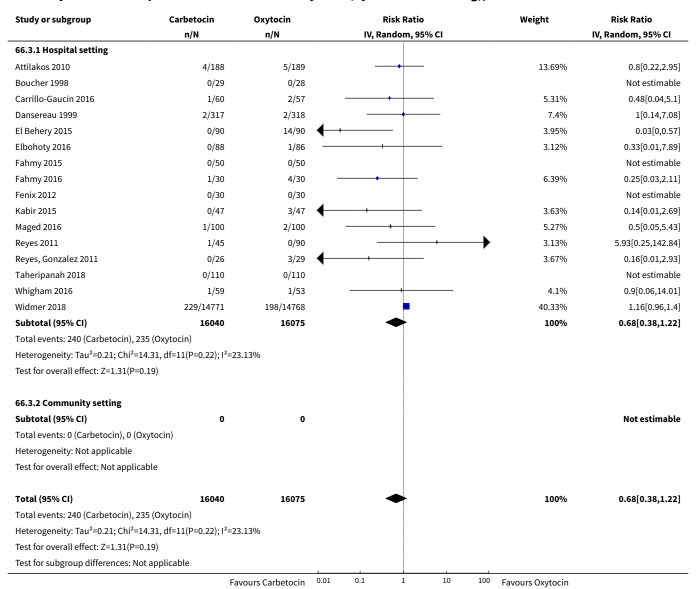
Analysis 66.2. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
66.2.1 Hospital setting					
Attilakos 2010	9/188	9/189		17.68%	1.01[0.41,2.48]
Boucher 1998	0/29	0/28			Not estimable
El Behery 2015	2/90	12/90		8.81%	0.17[0.04,0.72]
Elbohoty 2016	3/88	5/86		9.51%	0.59[0.14,2.38]
Fenix 2012	0/30	0/30			Not estimable
Kabir 2015	0/47	4/47		2.65%	0.11[0.01,2.01]
Maged 2016	0/100	1/100		2.21%	0.33[0.01,8.09]
Rosseland 2013	0/25	0/26			Not estimable
Whigham 2016	7/59	8/53		16.7%	0.79[0.31,2.02]
Widmer 2018	222/14651	212/14677	+	42.44%	1.05[0.87,1.26]
Subtotal (95% CI)	15307	15326	•	100%	0.73[0.45,1.19]
Total events: 243 (Carbetocin), 251	(Oxytocin)				
Heterogeneity: Tau ² =0.14; Chi ² =9.4	14, df=6(P=0.15); I ² =36.4	1%			
Test for overall effect: Z=1.25(P=0.2	21)				
66.2.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Oxy	ytocin)				
Heterogeneity: Not applicable					
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	



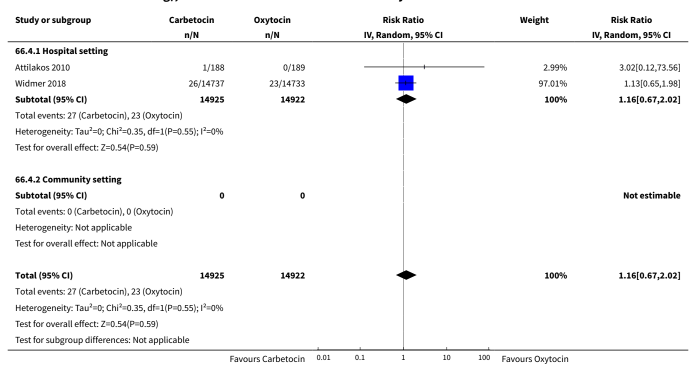


Analysis 66.3. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.





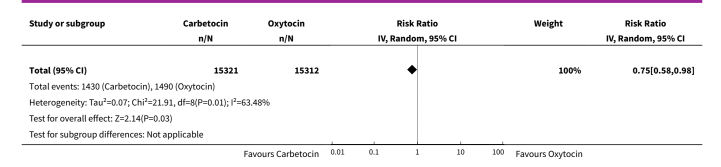
Analysis 66.4. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.



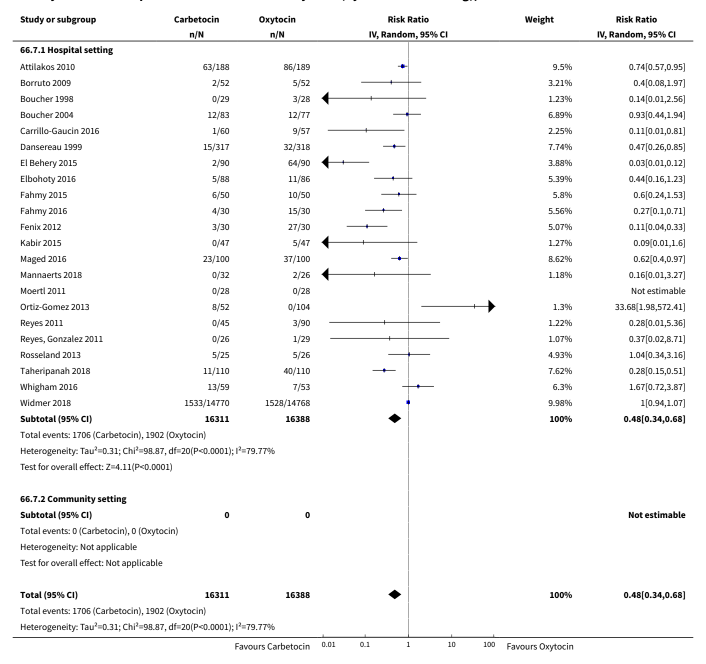
Analysis 66.6. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
66.6.1 Hospital setting					
Borruto 2009	12/52	23/52		11.45%	0.52[0.29,0.93]
Boucher 1998	0/29	0/28			Not estimable
Boucher 2004	10/64	11/67		7.81%	0.95[0.43,2.09]
El Behery 2015	8/90	19/90		7.98%	0.42[0.19,0.91]
Elbohoty 2016	21/88	34/86		14.71%	0.6[0.38,0.95]
Fenix 2012	0/30	0/30			Not estimable
Kabir 2015	0/47	3/47	*	0.77%	0.14[0.01,2.69]
Maged 2016	4/100	16/100		4.93%	0.25[0.09,0.72]
Rosseland 2013	6/25	4/26		4.37%	1.56[0.5,4.88]
Whigham 2016	42/59	37/53	+	21.78%	1.02[0.8,1.3]
Widmer 2018	1327/14737	1343/14733	•	26.19%	0.99[0.92,1.06]
Subtotal (95% CI)	15321	15312	•	100%	0.75[0.58,0.98]
Total events: 1430 (Carbetocin), 1	490 (Oxytocin)				
Heterogeneity: Tau ² =0.07; Chi ² =2.	1.91, df=8(P=0.01); I ² =63.	48%			
Test for overall effect: Z=2.14(P=0	.03)				
66.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Ox	kytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applica	ble				
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	





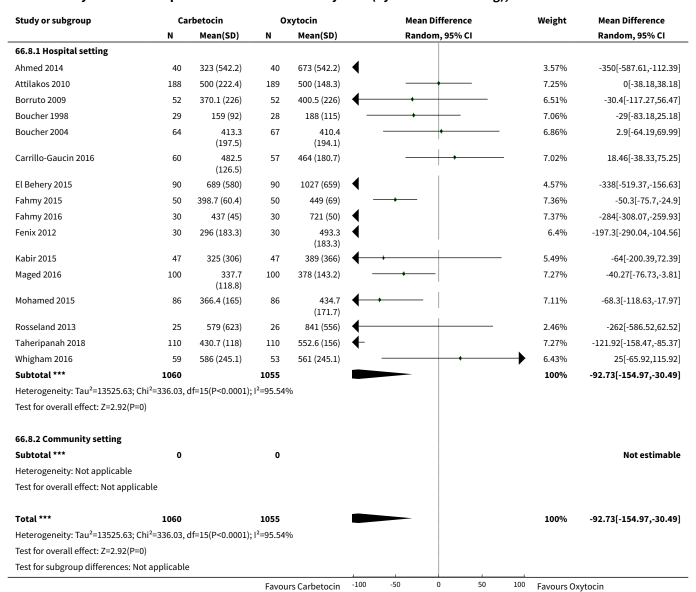
Analysis 66.7. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.





Study or subgroup Carbetocin		Oxytocin			Risk Ratio			Weight	Risk Ratio IV, Random, 95% CI
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Test for overall effect: Z=4.11	(P<0.0001)								
Test for subgroup differences	: Not applicable					1			
		Favours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 66.8. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.





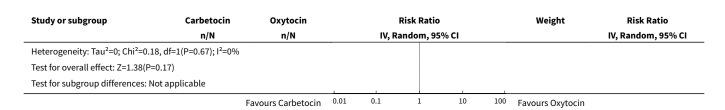
Analysis 66.9. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Caı	rbetocin	0:	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
66.9.1 Hospital setting							
Attilakos 2010	183	16 (2.2)	185	16 (2.2)	+	10.49%	0[-0.45,0.45]
Boucher 2004	82	12.8 (10.8)	73	15.9 (11.6)	+	8.2%	-3.1[-6.64,0.44]
Dansereau 1999	317	7.5 (10)	318	8.3 (10)	•	9.99%	-0.8[-2.36,0.76]
El Behery 2015	90	17.4 (8.7)	90	9.4 (6.7)		9.44%	8[5.73,10.27]
Fenix 2012	30	6 (4.6)	30	11 (4.6)	•	9.37%	-5[-7.35,-2.65]
Maged 2016	100	5.5 (3.5)	100	9.6 (6.2)	+	10.09%	-4.1[-5.5,-2.7]
Mannaerts 2018	32	14.5 (11)	26	15 (9)	+	6.58%	-0.5[-5.65,4.65]
Moertl 2011	28	11 (9.9)	28	11.4 (7.6)	+	7.1%	-0.4[-5.02,4.22]
Reyes, Gonzalez 2011	26	12.4 (8.7)	29	14.1 (11.2)	+	6.46%	-1.7[-6.97,3.57]
Rosseland 2013	25	5 (8.2)	26	8.2 (6.7)	+	7.61%	-3.2[-7.32,0.92]
Taheripanah 2018	110	11 (10.1)	110	20.5 (12)	+	8.82%	-9.5[-12.43,-6.57]
Whigham 2016	30	22 (11.5)	28	21 (11.5)	+	5.86%	1[-4.94,6.94]
Subtotal ***	1053		1043		•	100%	-1.66[-3.81,0.5]
Heterogeneity: Tau ² =11.48; Chi ² =1	38.97, df=1	1(P<0.0001); I ² =9	92.08%				
Test for overall effect: Z=1.51(P=0.1	L3)						
66.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	ole						
Total ***	1053		1043		•	100%	-1.66[-3.81,0.5]
Heterogeneity: Tau ² =11.48; Chi ² =1	38.97, df=1	1(P<0.0001); I ² =9	92.08%				
Test for overall effect: Z=1.51(P=0.1	L3)						
Test for subgroup differences: Not	applicable						
			Favou	rs Carbetocin -100	-50 0 50	100 Favours Oxy	/tocin

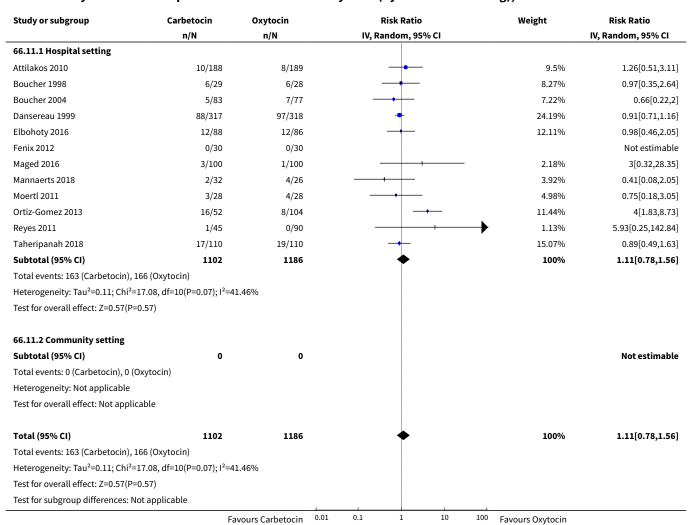
Analysis 66.10. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 10 Breastfeeding.

Study or subgroup	Carbetocin	Oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95%	CI		IV, Random, 95% CI
66.10.1 Hospital setting							
Reyes 2011	41/45	88/90		•		83.79%	0.93[0.85,1.03]
Reyes, Gonzalez 2011	22/26	25/29		+		16.21%	0.98[0.79,1.22]
Subtotal (95% CI)	71	119		•		100%	0.94[0.86,1.03]
Total events: 63 (Carbetocin), 113 (Oxy	tocin)						
Heterogeneity: Tau ² =0; Chi ² =0.18, df=1	(P=0.67); I ² =0%						
Test for overall effect: Z=1.38(P=0.17)							
66.10.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Carbetocin), 0 (Oxytoci	in)						
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Total (95% CI)	71	119		•		100%	0.94[0.86,1.03]
Total events: 63 (Carbetocin), 113 (Oxy	tocin)						
	Fa	vours Carbetocin	0.01 0.3	1 1	10 100	Favours Oxytocin	





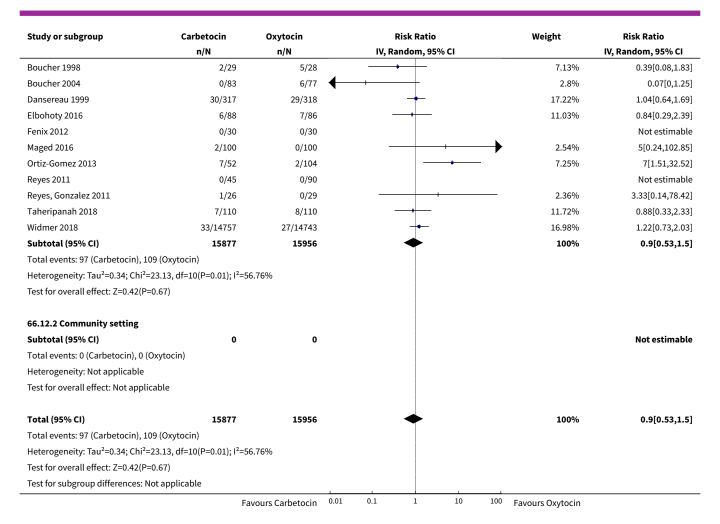
Analysis 66.11. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 11 Nausea.



Analysis 66.12. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Oxytocin		Risk Ratio IV, Random, 95% CI			Weight	Risk Ratio	
	n/N	n/N						IV, Random, 95% CI	
66.12.1 Hospital setting									
Attilakos 2010	5/188	5/189		-				9.48%	1.01[0.3,3.42]
Borruto 2009	4/52	20/52			-	1		11.49%	0.2[0.07,0.54]
	Fa	vours Carbetocin	0.01	0.1	1	10	100	Favours Oxytocin	

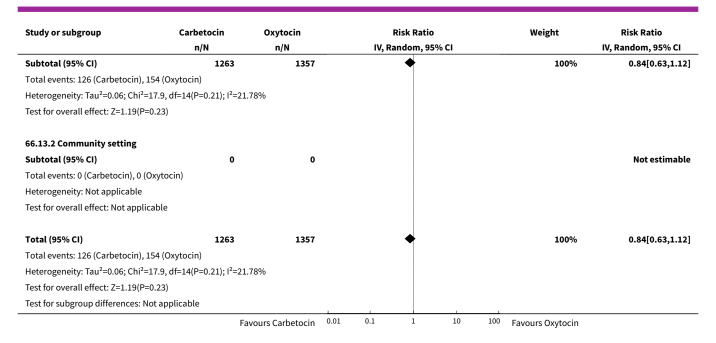




Analysis 66.13. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
66.13.1 Hospital setting					
Attilakos 2010	0/188	1/189		0.82%	0.34[0.01,8.17]
Borruto 2009	7/52	15/52		9.62%	0.47[0.21,1.05]
Boucher 1998	1/29	0/28		0.84%	2.9[0.12,68.33]
Boucher 2004	6/83	11/77		7.6%	0.51[0.2,1.3]
Dansereau 1999	46/317	43/318	-	22.49%	1.07[0.73,1.58]
El Behery 2015	23/90	30/90	-+ 	19.37%	0.77[0.48,1.21]
Elbohoty 2016	20/88	24/86	-+ 	17.26%	0.81[0.49,1.36]
Fenix 2012	1/30	1/30		1.12%	1[0.07,15.26]
Maged 2016	5/100	2/100	- +	3%	2.5[0.5,12.59]
Moertl 2011	2/28	2/28		2.25%	1[0.15,6.61]
Ortiz-Gomez 2013	6/52	8/104		6.89%	1.5[0.55,4.1]
Reyes 2011	2/45	1/90		1.46%	4[0.37,42.95]
Reyes, Gonzalez 2011	3/26	0/29		0.98%	7.78[0.42,143.81]
Rosseland 2013	2/25	3/26		2.73%	0.69[0.13,3.81]
Taheripanah 2018	2/110	13/110		3.59%	0.15[0.04,0.67]
	Fa	vours Carbetocin	0.01 0.1 1 10	100 Favours Oxytocin	



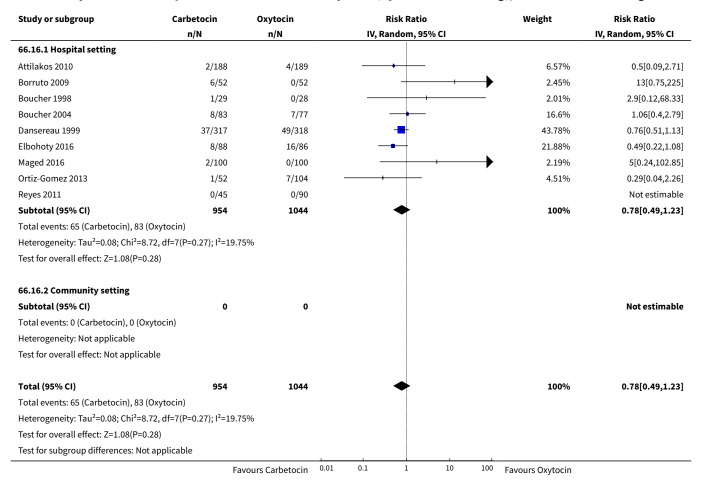


Analysis 66.14. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 14 Abdominal pain.

Study or subgroup	Carbetocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
66.14.1 Hospital setting					
Attilakos 2010	1/188	1/189		0.51%	1.01[0.06,15.95]
Borruto 2009	21/52	20/52		12.59%	1.05[0.65,1.69]
Boucher 1998	0/29	1/28 —	•	0.39%	0.32[0.01,7.59]
Boucher 2004	5/83	0/77		0.47%	10.21[0.57,181.71]
Dansereau 1999	131/317	127/318	+	32.1%	1.03[0.86,1.25]
Elbohoty 2016	72/88	44/86	-	28.24%	1.6[1.27,2.01]
Fenix 2012	7/30	8/30		4.62%	0.88[0.36,2.11]
Kabir 2015	4/47	5/47		2.4%	0.8[0.23,2.8]
Reyes 2011	0/45	1/90		0.39%	0.66[0.03,15.87]
Widmer 2018	63/14754	56/14743	+	18.28%	1.12[0.78,1.61]
Subtotal (95% CI)	15633	15660	♦	100%	1.18[0.97,1.44]
Total events: 304 (Carbetocin), 263 (Carbetocin)	Oxytocin)				
Heterogeneity: Tau ² =0.02; Chi ² =12.7,	df=9(P=0.18); I ² =29.1	1%			
Test for overall effect: Z=1.62(P=0.11))				
66.14.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Oxyto	ocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	15633	15660	•	100%	1.18[0.97,1.44]
Total events: 304 (Carbetocin), 263 (Carbetocin)	Oxytocin)				
Heterogeneity: Tau ² =0.02; Chi ² =12.7,	df=9(P=0.18); I ² =29.1	1%			
Test for overall effect: Z=1.62(P=0.11))				
Test for subgroup differences: Not ap	nlicable				



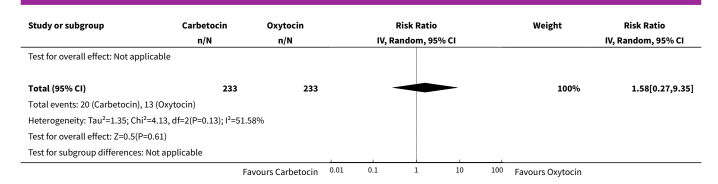
Analysis 66.16. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 16 Shivering.



Analysis 66.17. Comparison 66 Carbetocin vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Carbetocin	Oxytocin		1	Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ra	ndom, 95% CI			IV, Random, 95% CI
66.17.1 Hospital setting								
Boucher 1998	0/29	0/28						Not estimable
El Behery 2015	8/90	0/90			-		23.91%	17[1,290.19]
Elbohoty 2016	12/88	12/86			-		55.22%	0.98[0.46,2.05]
Reyes, Gonzalez 2011	0/26	1/29					20.87%	0.37[0.02,8.71]
Subtotal (95% CI)	233	233		_			100%	1.58[0.27,9.35]
Total events: 20 (Carbetocin), 13 (Oxyto	ocin)							
Heterogeneity: Tau ² =1.35; Chi ² =4.13, d	f=2(P=0.13); I ² =51.5	8%						
Test for overall effect: Z=0.5(P=0.61)								
66.17.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Carbetocin), 0 (Oxytoci	n)							
Heterogeneity: Not applicable								
	Fa	vours Carbetocin	0.01	0.1	1 10	100	Favours Oxytocin	





Comparison 67. Ergometrine vs Oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Death	1	1293	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	1293	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	6	2591	Risk Ratio (IV, Random, 95% CI)	1.30 [0.52, 3.27]
2.1 Hospital setting	6	2591	Risk Ratio (IV, Random, 95% CI)	1.30 [0.52, 3.27]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	5	1893	Risk Ratio (IV, Random, 95% CI)	1.44 [0.20, 10.23]
3.1 Hospital setting	5	1893	Risk Ratio (IV, Random, 95% CI)	1.44 [0.20, 10.23]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	10	3221	Risk Ratio (IV, Random, 95% CI)	1.31 [0.86, 1.99]
6.1 Hospital setting	10	3221	Risk Ratio (IV, Random, 95% CI)	1.31 [0.86, 1.99]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7 Additional uterotonics	6	2493	Risk Ratio (IV, Random, 95% CI)	1.46 [0.61, 3.48]
7.1 Hospital setting	6	2493	Risk Ratio (IV, Random, 95% CI)	1.46 [0.61, 3.48]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	10	3221	Mean Difference (IV, Random, 95% CI)	8.09 [-17.83, 34.00]
8.1 Hospital setting	10	3221	Mean Difference (IV, Random, 95% CI)	8.09 [-17.83, 34.00]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	1893	Mean Difference (IV, Random, 95% CI)	0.42 [-0.30, 1.13]
9.1 Hospital setting	2	1893	Mean Difference (IV, Random, 95% CI)	0.42 [-0.30, 1.13]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	6	2529	Risk Ratio (IV, Random, 95% CI)	4.56 [1.13, 18.44]
11.1 Hospital setting	6	2529	Risk Ratio (IV, Random, 95% CI)	4.56 [1.13, 18.44]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	5	2343	Risk Ratio (IV, Random, 95% CI)	3.83 [1.10, 13.28]
12.1 Hospital setting	5	2343	Risk Ratio (IV, Random, 95% CI)	3.83 [1.10, 13.28]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	4	2293	Risk Ratio (IV, Random, 95% CI)	5.63 [0.93, 33.96]
13.1 Hospital setting	4	2293	Risk Ratio (IV, Random, 95% CI)	5.63 [0.93, 33.96]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	1410	Risk Ratio (IV, Random, 95% CI)	13.39 [2.01, 89.44]
15.1 Hospital setting	3	1410	Risk Ratio (IV, Random, 95% CI)	13.39 [2.01, 89.44]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

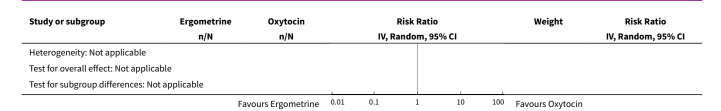


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
16 Shivering	3	1493	Risk Ratio (IV, Random, 95% CI)	1.73 [0.93, 3.25]
16.1 Hospital setting	3	1493	Risk Ratio (IV, Random, 95% CI)	1.73 [0.93, 3.25]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1443	Risk Ratio (IV, Random, 95% CI)	2.97 [0.97, 9.05]
17.1 Hospital setting	2	1443	Risk Ratio (IV, Random, 95% CI)	2.97 [0.97, 9.05]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	1393	Risk Ratio (IV, Random, 95% CI)	3.74 [0.42, 33.53]
18.1 Hospital setting	2	1393	Risk Ratio (IV, Random, 95% CI)	3.74 [0.42, 33.53]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

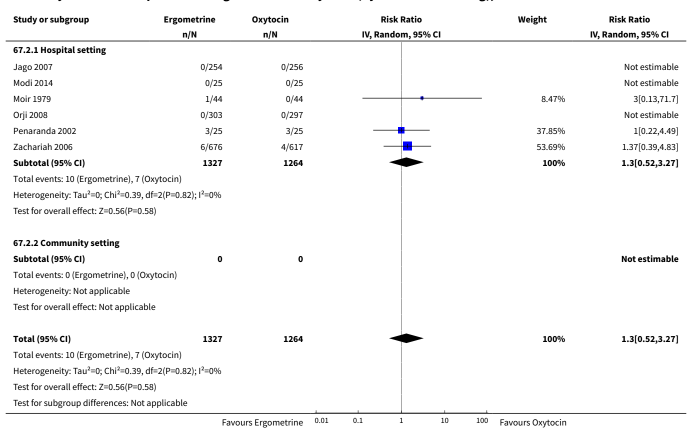
Analysis 67.1. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Ergometrine	Oxytocin			Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ra	andom, 95% CI			IV, Random, 95% CI
67.1.1 Hospital setting								
Zachariah 2006	0/676	0/617						Not estimable
Subtotal (95% CI)	676	617						Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
67.1.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable								
Total (95% CI)	676	617						Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)			1				
	Fav	ours Ergometrine	0.01	0.1	1 10	100	Favours Oxytocin	





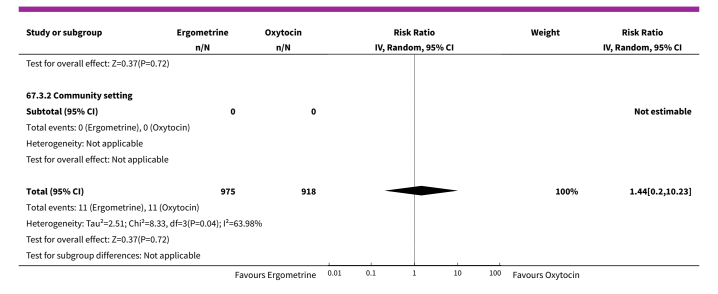
Analysis 67.2. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.



Analysis 67.3. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Ergometrine	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N n/N IV, Random, 95% CI				IV, Random, 95% CI				
67.3.1 Hospital setting									
Dhananjaya 2014	4/50	0/50			+	-	\rightarrow	21.32%	9[0.5,162.89]
Ezeama 2014	1/149	9/151		-				27.74%	0.11[0.01,0.88]
Modi 2014	0/25	0/25							Not estimable
Singh 2009	3/75	0/75		-	_	-	\rightarrow	20.98%	7[0.37,133.22]
Zachariah 2006	3/676	2/617		_	-			29.96%	1.37[0.23,8.17]
Subtotal (95% CI)	975	918			-			100%	1.44[0.2,10.23]
Total events: 11 (Ergometrine),	11 (Oxytocin)								
Heterogeneity: Tau ² =2.51; Chi ²	=8.33, df=3(P=0.04); I ² =63.9	8%							
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	



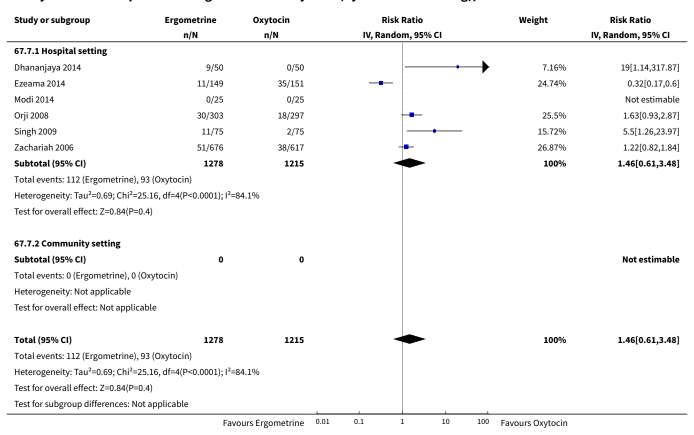


Analysis 67.6. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
67.6.1 Hospital setting					
Dhananjaya 2014	6/50	1/50	+	3.72%	6[0.75,48.05]
Ezeama 2014	3/149	12/151		9.07%	0.25[0.07,0.88]
Jago 2007	0/254	0/256			Not estimable
Modi 2014	0/25	0/25			Not estimable
Moir 1979	2/44	2/44		4.32%	1[0.15,6.79]
Moodie 1976	14/40	10/40	- -	20.72%	1.4[0.71,2.77]
Orji 2008	18/303	12/297	+	19.75%	1.47[0.72,3]
Penaranda 2002	12/25	8/25	+-	20.09%	1.5[0.74,3.03]
Singh 2009	2/75	0/75	- 	1.84%	5[0.24,102.42]
Zachariah 2006	20/676	13/617	- -	20.49%	1.4[0.7,2.8]
Subtotal (95% CI)	1641	1580	•	100%	1.31[0.86,1.99]
Total events: 77 (Ergometrine), 58 (O)xytocin)				
Heterogeneity: Tau ² =0.1; Chi ² =9.89, o	df=7(P=0.19); I ² =29.22	%			
Test for overall effect: Z=1.26(P=0.21)				
67.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Oxy	rtocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	2				
Total (95% CI)	1641	1580	•	100%	1.31[0.86,1.99]
Total events: 77 (Ergometrine), 58 (O)xytocin)				
Heterogeneity: Tau ² =0.1; Chi ² =9.89, o	df=7(P=0.19); I ² =29.22	%			
Test for overall effect: Z=1.26(P=0.21)				
Test for subgroup differences: Not ap	pplicable				
	Fave	ours Ergometrine 0.01	0.1 1 10	100 Favours Oxytocin	



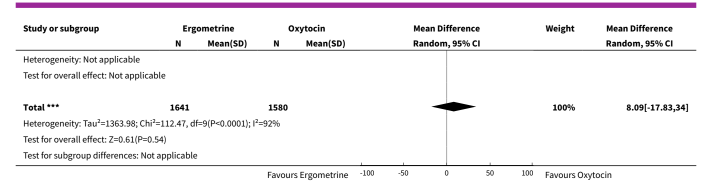
Analysis 67.7. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.



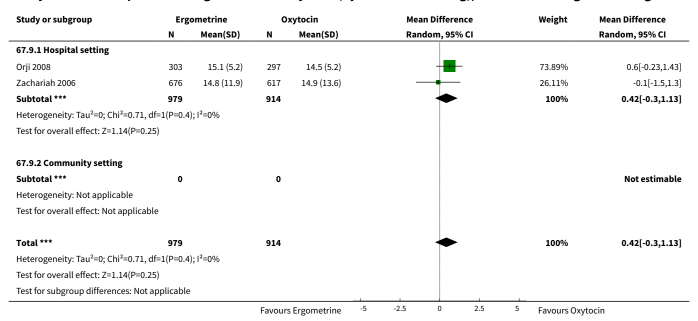
Analysis 67.8. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	0	xytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
67.8.1 Hospital setting							
Dhananjaya 2014	50	345 (109.5)	50	219 (86.3)	→	9.97%	126[87.35,164.65]
Ezeama 2014	149	287.1 (84.4)	151	301.8 (109.2)		11.73%	-14.7[-36.77,7.37]
Jago 2007	254	150.2 (63.6)	256	171.9 (81.6)		12.43%	-21.7[-34.39,-9.01]
Modi 2014	25	131 (72)	25	223.2 (122.5)	←	8.05%	-92.2[-147.92,-36.48]
Moir 1979	44	201 (50)	44	208 (58)		11.68%	-7[-29.63,15.63]
Moodie 1976	40	369 (118)	40	391 (129)		8.21%	-22[-76.18,32.18]
Orji 2008	303	246.6 (95.4)	297	245.7 (77.6)		12.36%	0.92[-12.98,14.82]
Penaranda 2002	25	546.8 (338.5)	25	467 (427.5)	←	1.32%	79.8[-133.95,293.55]
Singh 2009	75	223.5 (73.7)	75	154.7 (45.7)		11.94%	68.75[49.12,88.38]
Zachariah 2006	676	188 (138)	617	183 (130)	+-	12.31%	5[-9.61,19.61]
Subtotal ***	1641		1580			100%	8.09[-17.83,34]
Heterogeneity: Tau ² =1363.98; Ch	ii ² =112.47, d	f=9(P<0.0001); I ²	=92%				
Test for overall effect: Z=0.61(P=0	0.54)						
67.8.2 Community setting							
Subtotal ***	0		0				Not estimable
			Favour	s Ergometrine	-100 -50 0 50 100	Favours Ox	ytocin





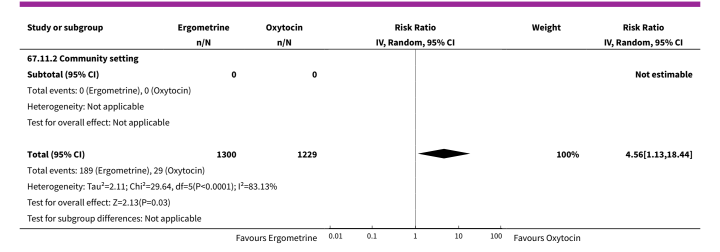
Analysis 67.9. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.



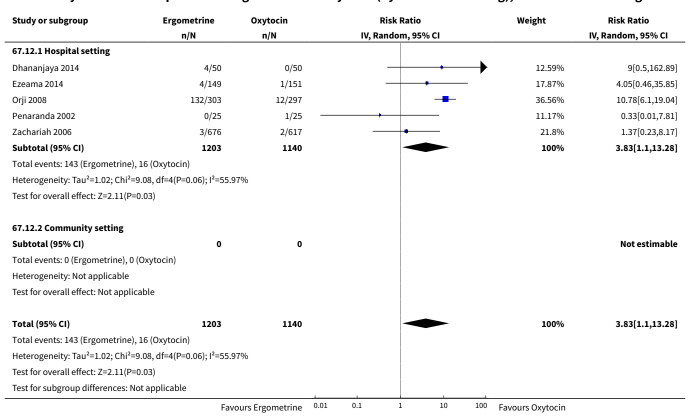
Analysis 67.11. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	ogroup Ergometrine Oxytocin Risk Ratio			Weight	Risk Ratio			
	n/N	n/N	IV, Random, 95% CI				IV, Random, 95% CI	
67.11.1 Hospital setting								
Dhananjaya 2014	1/50	0/50			+		10.73%	3[0.13,71.92]
Ezeama 2014	9/149	3/151		-	-		20.01%	3.04[0.84,11.01]
Moir 1979	6/44	0/44		_	+	\rightarrow	12.05%	13[0.75,223.98]
Moodie 1976	35/78	0/70				→	12.37%	63.81[3.99,1021.17]
Orji 2008	132/303	15/297			-		23.34%	8.63[5.18,14.36]
Zachariah 2006	6/676	11/617			<u></u>		21.5%	0.5[0.19,1.34]
Subtotal (95% CI)	1300	1229					100%	4.56[1.13,18.44]
Total events: 189 (Ergometrin	e), 29 (Oxytocin)							
Heterogeneity: Tau ² =2.11; Chi	² =29.64, df=5(P<0.0001); I ² =8	33.13%						
Test for overall effect: Z=2.13(P=0.03)							
	Fav	ours Ergometrine	0.01 (0.1	1 10	100	Favours Oxytocin	





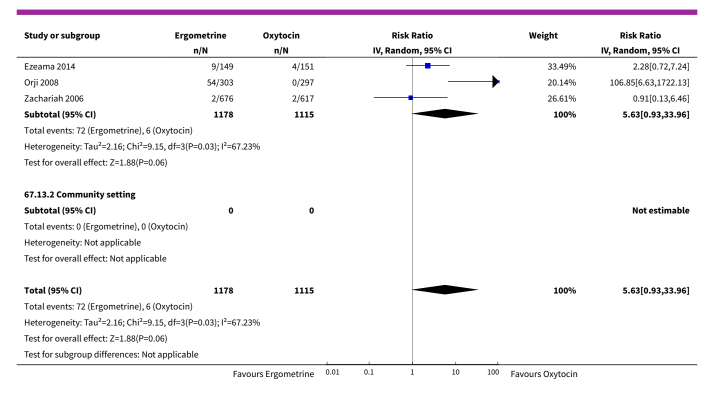
Analysis 67.12. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.



Analysis 67.13. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
67.13.1 Hospital setting									
Dhananjaya 2014	7/50	0/50					<u> </u>	19.75%	15[0.88,255.78]
	Favo	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	





Analysis 67.15. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
67.15.1 Hospital setting					
Ezeama 2014	7/149	0/151	+	23.98%	15.2[0.88,263.78]
Jago 2007	134/254	29/256		51.35%	4.66[3.24,6.69]
Orji 2008	54/303	0/297		24.66%	106.85[6.63,1722.13]
Subtotal (95% CI)	706	704		100%	13.39[2.01,89.44]
Total events: 195 (Ergometrine), 29 (Ox	ytocin)				
Heterogeneity: Tau²=1.79; Chi²=5.39, di	f=2(P=0.07); I ² =62.9	1%			
Test for overall effect: Z=2.68(P=0.01)					
67.15.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	cin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	706	704		100%	13.39[2.01,89.44]
Total events: 195 (Ergometrine), 29 (Ox	ytocin)				
Heterogeneity: Tau ² =1.79; Chi ² =5.39, di	f=2(P=0.07); I ² =62.9	1%			
Test for overall effect: Z=2.68(P=0.01)					
Test for subgroup differences: Not appl	icable				
	Favo	ours Ergometrine 0	0.01 0.1 1 10 100	Favours Oxytocin	



Analysis 67.16. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Ergometrine	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
67.16.1 Hospital setting					
Penaranda 2002	1/25	0/25		3.96%	3[0.13,70.3]
Singh 2009	0/75	0/75			Not estimable
Zachariah 2006	26/676	14/617	-	96.04%	1.7[0.89,3.22]
Subtotal (95% CI)	776	717	•	100%	1.73[0.93,3.25]
Total events: 27 (Ergometrine), 14 (Ox	ytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.12, df=	1(P=0.73); I ² =0%				
Test for overall effect: Z=1.72(P=0.09)					
67.16.2 Community setting					
Subtotal (95% CI)	0	0	ĺ		Not estimable
Total events: 0 (Ergometrine), 0 (Oxyto	ocin)		ĺ		
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	776	717	•	100%	1.73[0.93,3.25]
Total events: 27 (Ergometrine), 14 (Ox	ytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.12, df=	1(P=0.73); I ² =0%				
Test for overall effect: Z=1.72(P=0.09)					
Test for subgroup differences: Not app	olicable				
	Fav	ours Ergometrine 0.0	1 0.1 1 10 1	00 Favours Oxytocin	

Analysis 67.17. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup E	rgometrine	Oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
67.17.1 Hospital setting									
Singh 2009	0/75	0/75							Not estimable
Zachariah 2006	13/676	4/617			-	_		100%	2.97[0.97,9.05]
Subtotal (95% CI)	751	692				-		100%	2.97[0.97,9.05]
Total events: 13 (Ergometrine), 4 (Oxytoo	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.91(P=0.06)									
67.17.2 Community setting									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Ergometrine), 0 (Oxytoci	n)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Total (95% CI)	751	692				-		100%	2.97[0.97,9.05]
Total events: 13 (Ergometrine), 4 (Oxytoo	cin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.91(P=0.06)									
Test for subgroup differences: Not applic	able								
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Oxytocin	



Analysis 67.18. Comparison 67 Ergometrine vs Oxytocin (by healthcare setting), Outcome 18 Diarrhoea.

			Weight	
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
1/50	0/50		47.71%	3[0.13,71.92]
2/676	0/617		52.29%	4.56[0.22,94.88]
726	667		100%	3.74[0.42,33.53]
xytocin)				
df=1(P=0.85); I ² =0%				
24)				
0	0			Not estimable
xytocin)				
ole				
726	667		100%	3.74[0.42,33.53]
xytocin)				
df=1(P=0.85); I ² =0%				
24)				
applicable				
2	2/676 726 xytocin) df=1(P=0.85); l²=0% 24) 0 xytocin) ole 726 xytocin) df=1(P=0.85); l²=0% 24) applicable	2/676 0/617 726 667 xytocin) df=1(P=0.85); l²=0% 24) 0 0 xytocin) ole 726 667 xytocin) df=1(P=0.85); l²=0% 24)	2/676 0/617 726 667 xytocin) df=1(P=0.85); l²=0% 24) 0 0 xytocin) ole 726 667 xytocin) df=1(P=0.85); l²=0% 24) applicable	2/676 0/617 726 667 xytocin) df=1(P=0.85); l²=0% 24) 0 0 xytocin) ole 726 667 100% xytocin) df=1(P=0.85); l²=0% 24) applicable

Comparison 68. Ergometine plus oxytocin vs Oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	2	1314	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	2	1314	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	10	10990	Risk Ratio (IV, Random, 95% CI)	0.73 [0.57, 0.93]
2.1 Hospital setting	10	10990	Risk Ratio (IV, Random, 95% CI)	0.73 [0.57, 0.93]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	11	10985	Risk Ratio (IV, Random, 95% CI)	0.88 [0.53, 1.44]
3.1 Hospital setting	11	10985	Risk Ratio (IV, Random, 95% CI)	0.88 [0.53, 1.44]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	1	991	Risk Ratio (IV, Random, 95% CI)	2.99 [0.12, 73.32]
4.1 Hospital setting	1	991	Risk Ratio (IV, Random, 95% CI)	2.99 [0.12, 73.32]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	11	11090	Risk Ratio (IV, Random, 95% CI)	0.72 [0.57, 0.91]
6.1 Hospital setting	11	11090	Risk Ratio (IV, Random, 95% CI)	0.72 [0.57, 0.91]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	10	8968	Risk Ratio (IV, Random, 95% CI)	0.79 [0.59, 1.07]
7.1 Hospital setting	10	8968	Risk Ratio (IV, Random, 95% CI)	0.79 [0.59, 1.07]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	10	4248	Mean Difference (IV, Random, 95% CI)	-10.31 [-40.32, 19.70]
8.1 Hospital setting	10	4248	Mean Difference (IV, Random, 95% CI)	-10.31 [-40.32, 19.70]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	6	4324	Mean Difference (IV, Random, 95% CI)	-2.23 [-5.24, 0.77]
9.1 Hospital setting	6	4324	Mean Difference (IV, Random, 95% CI)	-2.23 [-5.24, 0.77]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	1	3483	Risk Ratio (IV, Random, 95% CI)	0.99 [0.96, 1.01]
10.1 Hospital setting	1	3483	Risk Ratio (IV, Random, 95% CI)	0.99 [0.96, 1.01]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	7	6931	Risk Ratio (IV, Random, 95% CI)	1.72 [0.84, 3.53]
11.1 Hospital setting	7	6931	Risk Ratio (IV, Random, 95% CI)	1.72 [0.84, 3.53]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	9	10227	Risk Ratio (IV, Random, 95% CI)	3.05 [1.76, 5.29]
12.1 Hospital setting	9	10227	Risk Ratio (IV, Random, 95% CI)	3.05 [1.76, 5.29]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	5	5105	Risk Ratio (IV, Random, 95% CI)	1.26 [0.79, 1.99]
13.1 Hospital setting	5	5105	Risk Ratio (IV, Random, 95% CI)	1.26 [0.79, 1.99]

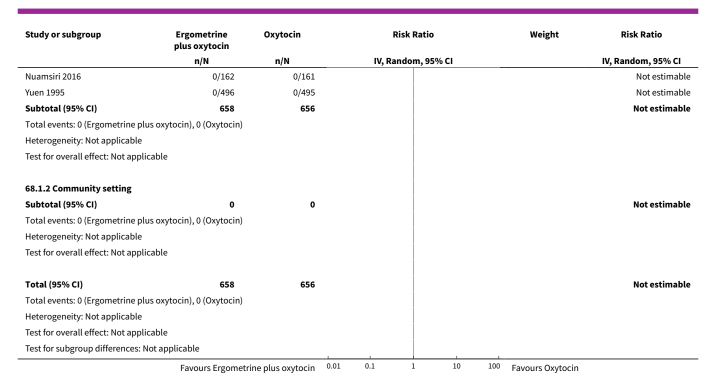


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	1362	Risk Ratio (IV, Random, 95% CI)	2.00 [0.29, 13.97]
15.1 Hospital setting	3	1362	Risk Ratio (IV, Random, 95% CI)	2.00 [0.29, 13.97]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	1591	Risk Ratio (IV, Random, 95% CI)	0.96 [0.60, 1.53]
16.1 Hospital setting	2	1591	Risk Ratio (IV, Random, 95% CI)	0.96 [0.60, 1.53]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1591	Risk Ratio (IV, Random, 95% CI)	1.08 [0.48, 2.43]
17.1 Hospital setting	2	1591	Risk Ratio (IV, Random, 95% CI)	1.08 [0.48, 2.43]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	3	2030	Risk Ratio (IV, Random, 95% CI)	1.26 [0.72, 2.22]
18.1 Hospital setting	3	2030	Risk Ratio (IV, Random, 95% CI)	1.26 [0.72, 2.22]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 68.1. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
68.1.1 Hospital setting				1					
	Favours Ergomet	rine plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	

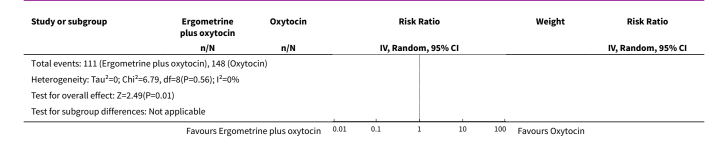




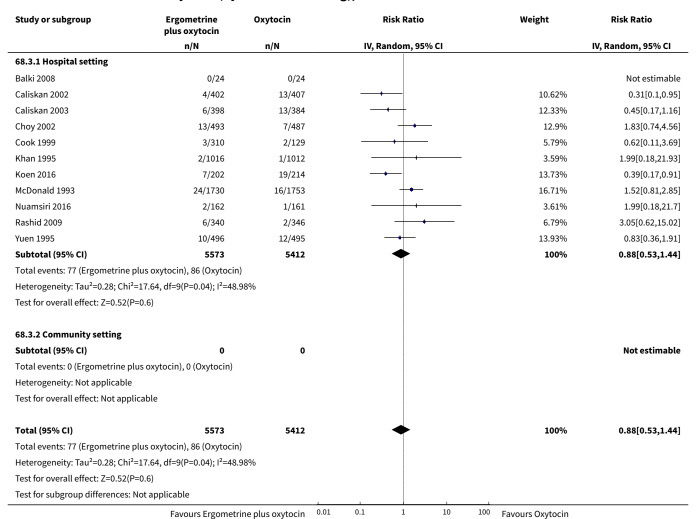
Analysis 68.2. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.2.1 Hospital setting					
Caliskan 2002	7/402	14/407		7.6%	0.51[0.21,1.24]
Caliskan 2003	5/398	15/384		6.09%	0.32[0.12,0.88]
Choy 2002	3/500	6/491		3.21%	0.49[0.12,1.95]
Cook 1999	7/310	0/129		0.75%	6.27[0.36,108.98]
Khan 1995	9/1016	11/1012		7.96%	0.81[0.34,1.96]
McDonald 1993	68/1730	83/1753	=	62.17%	0.83[0.61,1.14]
Mitchell 1993	0/228	1/230 —	•	0.6%	0.34[0.01,8.21]
Nuamsiri 2016	0/162	0/161			Not estimable
Rashid 2009	6/340	8/346		5.57%	0.76[0.27,2.18]
Yuen 1995	6/496	10/495		6.06%	0.6[0.22,1.63]
Subtotal (95% CI)	5582	5408	◆	100%	0.73[0.57,0.93]
Total events: 111 (Ergometrine	plus oxytocin), 148 (Oxyto	cin)			
Heterogeneity: Tau ² =0; Chi ² =6.7	'9, df=8(P=0.56); I ² =0%				
Test for overall effect: Z=2.49(P=	=0.01)				
68.2.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plu	us oxytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applie	cable				
Total (95% CI)	5582	5408	♦	100%	0.73[0.57,0.93]



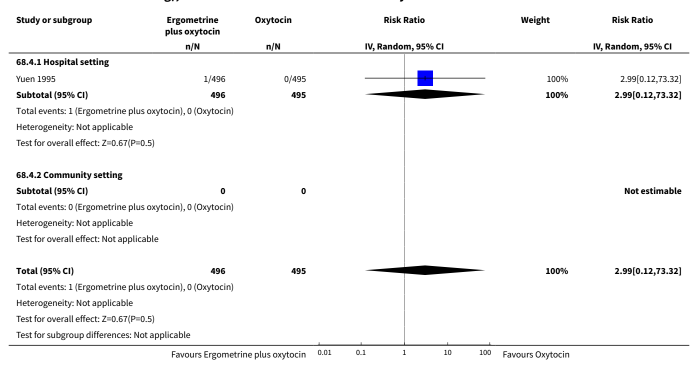


Analysis 68.3. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.





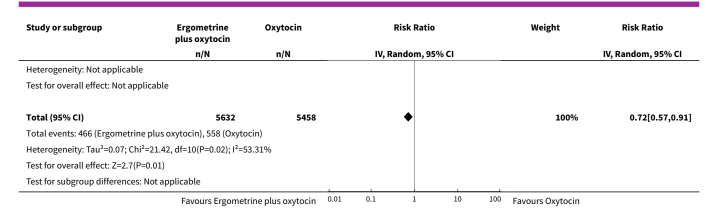
Analysis 68.4. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.



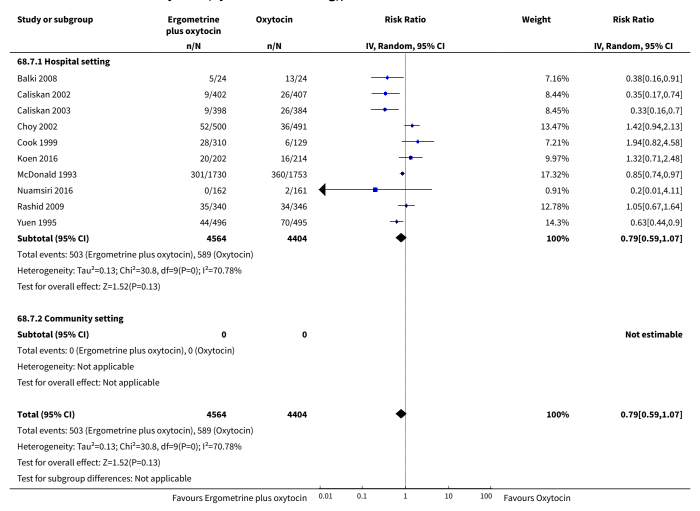
Analysis 68.6. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.6.1 Hospital setting					
Caliskan 2002	14/402	33/407		9.06%	0.43[0.23,0.79]
Caliskan 2003	14/398	28/384		8.78%	0.48[0.26,0.9]
Choy 2002	23/500	26/491		10.24%	0.87[0.5,1.5]
Cook 1999	23/310	1/129		1.35%	9.57[1.31,70.13]
Khan 1995	36/1016	41/1012	-+	12.66%	0.87[0.56,1.36]
McDonald 1993	286/1730	316/1753	+	20.46%	0.92[0.79,1.06]
Mitchell 1993	6/228	17/230		5.23%	0.36[0.14,0.89]
Nuamsiri 2016	0/162	1/161		0.55%	0.33[0.01,8.07]
Rashid 2009	8/340	9/346		4.99%	0.9[0.35,2.32]
Un Nisa 2012	20/50	26/50	-+ 	12.85%	0.77[0.5,1.18]
Yuen 1995	36/496	60/495		13.82%	0.6[0.4,0.89]
Subtotal (95% CI)	5632	5458	♦	100%	0.72[0.57,0.91]
Total events: 466 (Ergometrine	e plus oxytocin), 558 (Oxyto	cin)			
Heterogeneity: Tau ² =0.07; Chi	² =21.42, df=10(P=0.02); I ² =5.	3.31%			
Test for overall effect: Z=2.7(P=	=0.01)				
68.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine p	olus oxytocin), 0 (Oxytocin)				
	Favours Ergomet	rine plus oxytocin	0.01 0.1 1 10	100 Favours Oxytocin	



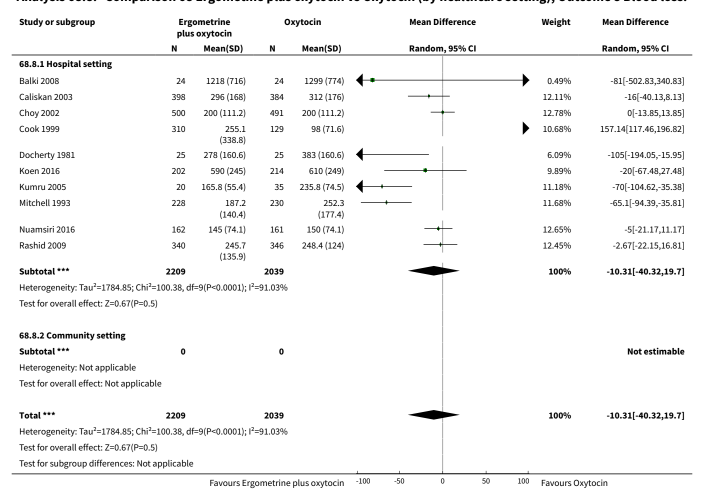


Analysis 68.7. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.





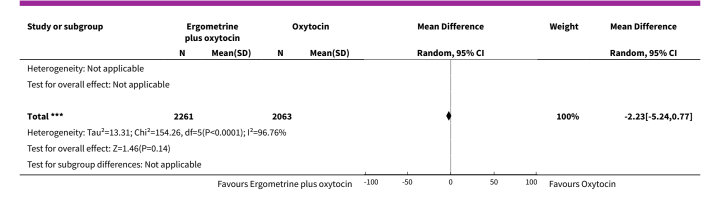
Analysis 68.8. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.



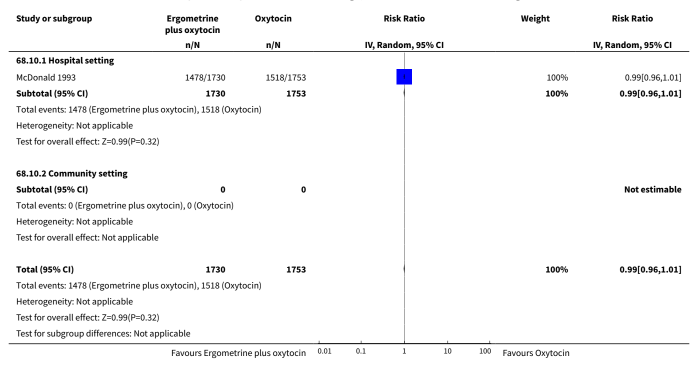
Analysis 68.9. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Ū	ometrine oxytocin	0:	xytocin	Mean Difference	e Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% C	I	Random, 95% CI
68.9.1 Hospital setting							
Caliskan 2002	402	15 (12)	407	14 (14)	•	16.59%	1[-0.8,2.8]
Caliskan 2003	398	15 (12)	384	14 (14)	•	16.55%	1[-0.83,2.83]
Choy 2002	493	10 (11.1)	487	8 (11.9)	•	16.95%	2[0.56,3.44]
Cook 1999	310	0.5 (14.1)	129	14.7 (12.1)	+	15.56%	-14.2[-16.81,-11.59]
Nuamsiri 2016	162	8 (7.4)	161	9 (7.4)	•	16.77%	-1[-2.62,0.62]
Yuen 1995	496	12 (2.8)	495	15 (2.8)	•	17.59%	-3[-3.35,-2.65]
Subtotal ***	2261		2063		•	100%	-2.23[-5.24,0.77]
Heterogeneity: Tau ² =13.31; Ch	i ² =154.26, df=5	(P<0.0001); I ² =96	6.76%				
Test for overall effect: Z=1.46(F	P=0.14)						
68.9.2 Community setting							
Subtotal ***	0		0	1			Not estimable
		Favours Erge	ometrine	plus oxytocin -100) -50 0	50 100 Favours Ox	ytocin





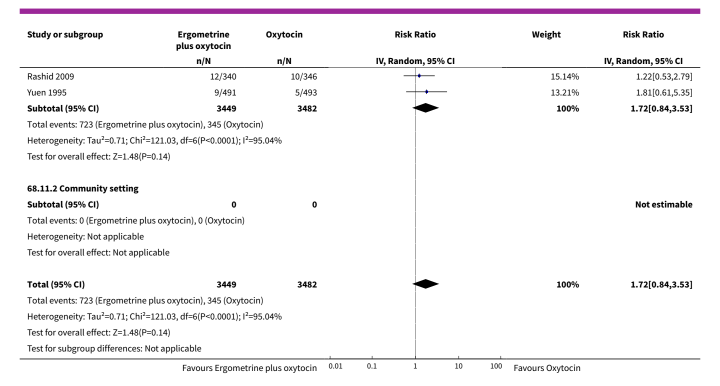
Analysis 68.10. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 10 Breastfeeding.



Analysis 68.11. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI				IV, Random, 95% CI
68.11.1 Hospital setting								
Balki 2008	10/24	2/24		-			10.95%	5[1.22,20.46]
Choy 2002	198/500	186/491		+			18.75%	1.05[0.89,1.22]
Koen 2016	26/202	23/214		+	-		17.17%	1.2[0.71,2.03]
McDonald 1993	467/1730	117/1753			+		18.67%	4.04[3.34,4.9]
Nuamsiri 2016	1/162	2/161		- +			6.11%	0.5[0.05,5.43]
	Favours Ergometr	rine plus oxytocin	0.01	0.1 1	10	100	Favours Oxytocin	

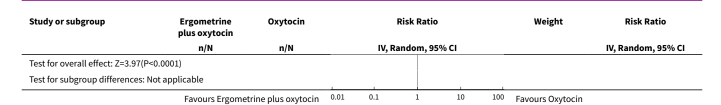




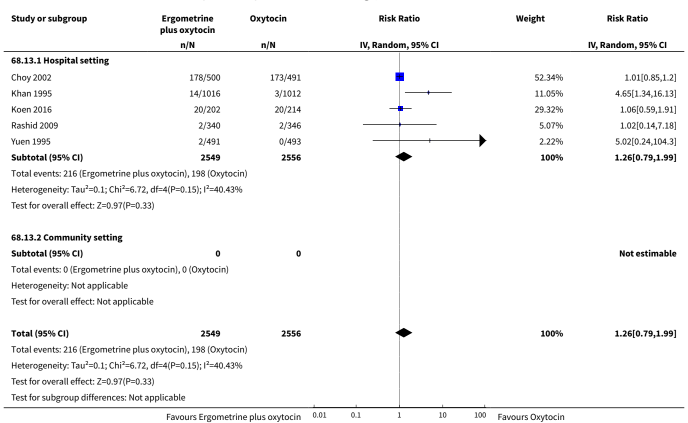
Analysis 68.12. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.12.1 Hospital setting					
Balki 2008	6/24	1/24	+	5.78%	6[0.78,46.14]
Caliskan 2002	1/402	2/407		4.44%	0.51[0.05,5.56]
Caliskan 2003	5/398	3/384		9.75%	1.61[0.39,6.68]
Choy 2002	8/500	2/491	+	8.74%	3.93[0.84,18.4]
Khan 1995	14/1016	4/1012		13.13%	3.49[1.15,10.55]
Koen 2016	11/202	5/214	 • -	14.01%	2.33[0.82,6.59]
McDonald 1993	358/1730	59/1753	-	26.25%	6.15[4.71,8.03]
Rashid 2009	4/340	1/346	-	5.17%	4.07[0.46,36.23]
Yuen 1995	7/491	5/493	- •	12.72%	1.41[0.45,4.4]
Subtotal (95% CI)	5103	5124	•	100%	3.05[1.76,5.29]
Total events: 414 (Ergometrine p	lus oxytocin), 82 (Oxytoci	in)			
Heterogeneity: Tau ² =0.28; Chi ² =1	15.63, df=8(P=0.05); l ² =48.	.81%			
Test for overall effect: Z=3.97(P<0	0.0001)				
68.12.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus	s oxytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applica	able				
Total (95% CI)	5103	5124	•	100%	3.05[1.76,5.29]
Total events: 414 (Ergometrine p	lus oxytocin), 82 (Oxytoci	in)			
Heterogeneity: Tau ² =0.28; Chi ² =1	L5.63, df=8(P=0.05); I ² =48.	.81%			





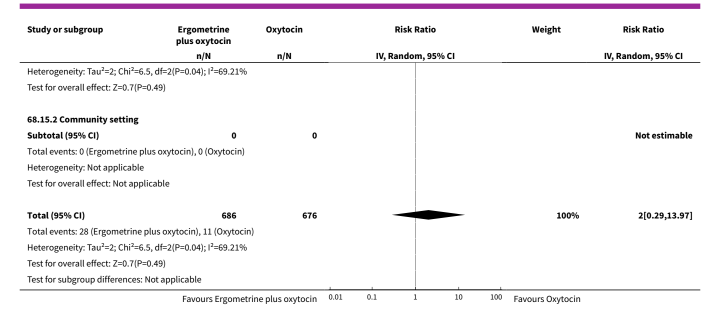
Analysis 68.13. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 13 Headache.



Analysis 68.15. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 15 Hypertension.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, Ra	ndom, 95	5% CI			IV, Random, 95% CI
68.15.1 Hospital setting									
Balki 2008	1/24	4/24	-	-				31.09%	0.25[0.03,2.08]
Choy 2002	17/500	7/491			-	—		44.78%	2.38[1,5.7]
Nuamsiri 2016	10/162	0/161					—	24.12%	20.87[1.23,353.21]
Subtotal (95% CI)	686	676		-	-			100%	2[0.29,13.97]
Total events: 28 (Ergometrine p	lus oxytocin), 11 (Oxytocin)			İ				
	Favours Ergometr	rine plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	





Analysis 68.16. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.16.1 Hospital setting					
Caliskan 2002	19/402	16/407	-	50.89%	1.2[0.63,2.3]
Caliskan 2003	15/398	19/384	_	49.11%	0.76[0.39,1.48]
Subtotal (95% CI)	800	791	*	100%	0.96[0.6,1.53]
Total events: 34 (Ergometrine plus or	xytocin), 35 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.93, df	=1(P=0.34); I ² =0%				
Test for overall effect: Z=0.17(P=0.87))				
68.16.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus oxy	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	!				
Total (95% CI)	800	791	•	100%	0.96[0.6,1.53]
Total events: 34 (Ergometrine plus of	xytocin), 35 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.93, df	=1(P=0.34); I ² =0%				
Test for overall effect: Z=0.17(P=0.87))				
Test for subgroup differences: Not ap	plicable				
	Favours Ergometrir	ne plus oxytocin 0.01	0.1 1 10 1	00 Favours Oxytocin	



Analysis 68.17. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.17.1 Hospital setting					
Caliskan 2002	6/402	6/407		52.4%	1.01[0.33,3.11]
Caliskan 2003	6/398	5/384		47.6%	1.16[0.36,3.76]
Subtotal (95% CI)	800	791	*	100%	1.08[0.48,2.43]
Total events: 12 (Ergometrine p	olus oxytocin), 11 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.0	03, df=1(P=0.87); I ² =0%				
Test for overall effect: Z=0.18(P	=0.85)				
68.17.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine pl	us oxytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not appl	icable				
Total (95% CI)	800	791	•	100%	1.08[0.48,2.43]
Total events: 12 (Ergometrine p	olus oxytocin), 11 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.0	03, df=1(P=0.87); I ² =0%				
Test for overall effect: Z=0.18(P	=0.85)				
Test for subgroup differences: N	Not applicable				
	Favours Ergometrii	ne plus oxytocin 0.01	. 0.1 1 10 10	00 Favours Oxytocin	

Analysis 68.18. Comparison 68 Ergometine plus oxytocin vs Oxytocin (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
68.18.1 Hospital setting					
Caliskan 2002	10/402	9/407		39.93%	1.12[0.46,2.74]
Caliskan 2003	17/398	12/384	-	60.07%	1.37[0.66,2.82]
Cook 1999	0/310	0/129			Not estimable
Subtotal (95% CI)	1110	920	•	100%	1.26[0.72,2.22]
Total events: 27 (Ergometrine plus o	xytocin), 21 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.11, df	=1(P=0.74); I ² =0%				
Test for overall effect: Z=0.82(P=0.41)				
68.18.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Ergometrine plus ox	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	2				
Total (95% CI)	1110	920	•	100%	1.26[0.72,2.22]
Total events: 27 (Ergometrine plus o	xytocin), 21 (Oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.11, df	=1(P=0.74); I ² =0%				
Test for overall effect: Z=0.82(P=0.41)				
Test for subgroup differences: Not ap	pplicable				
	Favours Ergometri	ne plus oxytocin 0.01	0.1 1 10 1	00 Favours Oxytocin	



Comparison 69. Misoprostol plus oxytocin vs Oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	9	4737	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	9	4737	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	17	8514	Risk Ratio (IV, Random, 95% CI)	0.87 [0.69, 1.09]
2.1 Hospital setting	17	8514	Risk Ratio (IV, Random, 95% CI)	0.87 [0.69, 1.09]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	19	8742	Risk Ratio (IV, Random, 95% CI)	0.50 [0.37, 0.67]
3.1 Hospital setting	19	8742	Risk Ratio (IV, Random, 95% CI)	0.50 [0.37, 0.67]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	3	1886	Risk Ratio (IV, Random, 95% CI)	0.50 [0.05, 5.47]
4.1 Hospital setting	3	1886	Risk Ratio (IV, Random, 95% CI)	0.50 [0.05, 5.47]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal mor- bidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	14	8148	Risk Ratio (IV, Random, 95% CI)	0.71 [0.59, 0.85]
6.1 Hospital setting	14	8148	Risk Ratio (IV, Random, 95% CI)	0.71 [0.59, 0.85]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	18	8391	Risk Ratio (IV, Random, 95% CI)	0.54 [0.44, 0.67]
7.1 Hospital setting	18	8391	Risk Ratio (IV, Random, 95% CI)	0.54 [0.44, 0.67]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	17	8690	Mean Difference (IV, Random, 95% CI)	-87.26 [-157.83, -16.69]
8.1 Hospital setting	17	8690	Mean Difference (IV, Random, 95% CI)	-87.26 [-157.83, -16.69]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	15	7929	Mean Difference (IV, Random, 95% CI)	-2.59 [-3.70, -1.48]
9.1 Hospital setting	15	7929	Mean Difference (IV, Random, 95% CI)	-2.59 [-3.70, -1.48]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	7	3798	Risk Ratio (IV, Random, 95% CI)	2.21 [1.19, 4.10]
11.1 Hospital setting	7	3798	Risk Ratio (IV, Random, 95% CI)	2.21 [1.19, 4.10]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	11	6718	Risk Ratio (IV, Random, 95% CI)	2.24 [1.52, 3.31]
12.1 Hospital setting	11	6718	Risk Ratio (IV, Random, 95% CI)	2.24 [1.52, 3.31]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	2	303	Risk Ratio (IV, Random, 95% CI)	1.26 [0.26, 6.23]
13.1 Hospital setting	2	303	Risk Ratio (IV, Random, 95% CI)	1.26 [0.26, 6.23]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	1	366	Risk Ratio (IV, Random, 95% CI)	1.93 [1.01, 3.67]
14.1 Hospital setting	1	366	Risk Ratio (IV, Random, 95% CI)	1.93 [1.01, 3.67]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	19	9458	Risk Ratio (IV, Random, 95% CI)	3.38 [2.50, 4.57]
16.1 Hospital setting	19	9458	Risk Ratio (IV, Random, 95% CI)	3.38 [2.50, 4.57]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	17	8607	Risk Ratio (IV, Random, 95% CI)	2.99 [2.00, 4.45]
17.1 Hospital setting	17	8607	Risk Ratio (IV, Random, 95% CI)	2.99 [2.00, 4.45]

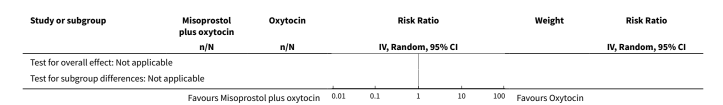


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	7	5649	Risk Ratio (IV, Random, 95% CI)	2.08 [0.99, 4.38]
18.1 Hospital setting	7	5649	Risk Ratio (IV, Random, 95% CI)	2.08 [0.99, 4.38]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

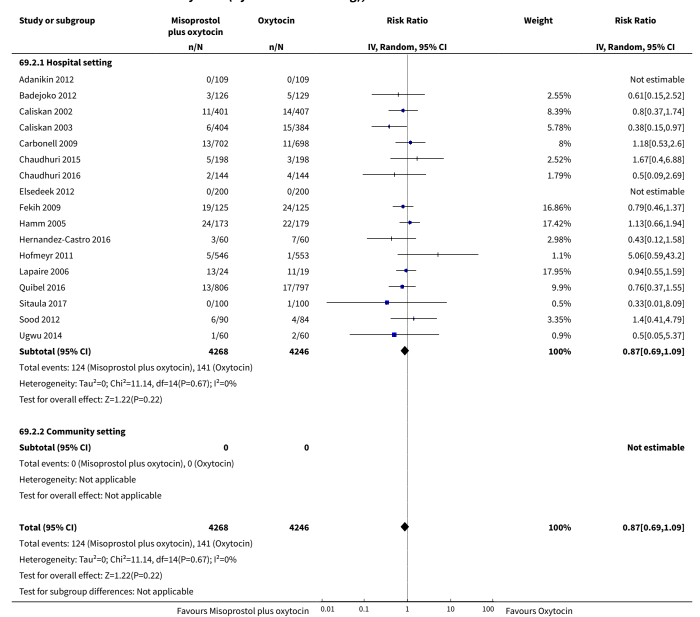
Analysis 69.1. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
69.1.1 Hospital setting		,		,	
Badejoko 2012	0/126	0/129			Not estimable
Bhullar 2004	0/377	0/379			Not estimable
Carbonell 2009	0/702	0/698			Not estimable
Chaudhuri 2015	0/198	0/198			Not estimable
Chaudhuri 2016	0/144	0/144			Not estimable
El Tahan 2012	0/179	0/187			Not estimable
Hofmeyr 2011	0/546	0/557			Not estimable
Lapaire 2006	0/28	0/25			Not estimable
Ugwu 2014	0/60	0/60			Not estimable
Subtotal (95% CI)	2360	2377			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
69.1.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	2360	2377			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
	Favours Misopros	tol plus oxytocin 0.01	0.1 1 10 1	LOO Favours Oxytocin	



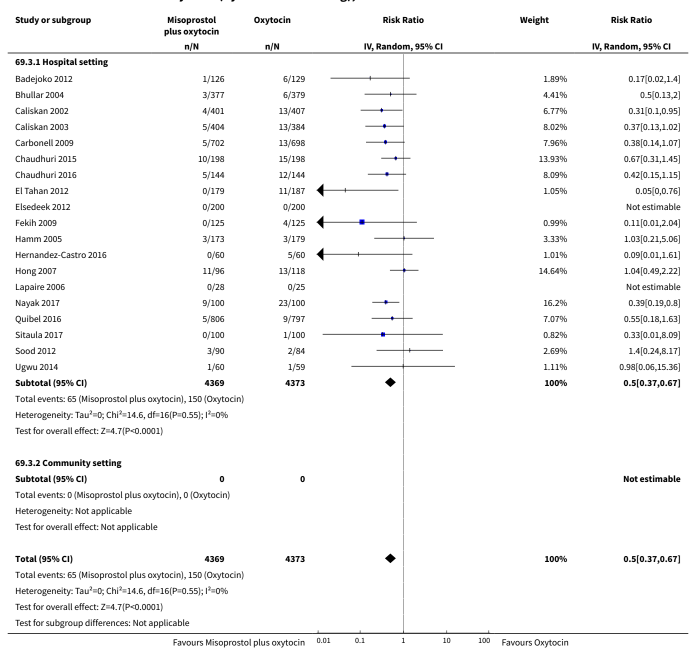


Analysis 69.2. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.





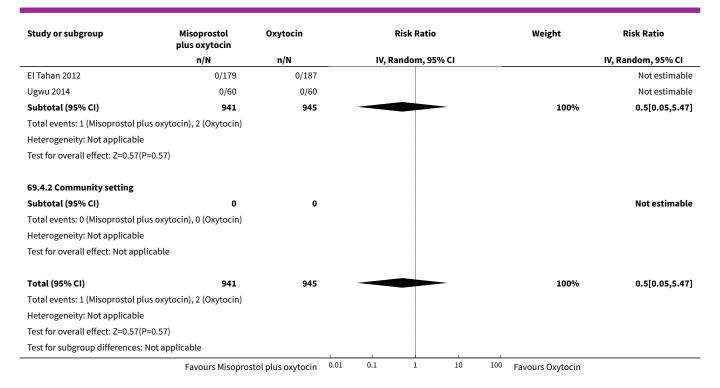
Analysis 69.3. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 3 Blood transfusion.



Analysis 69.4. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin			Risk Ratio	•		Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	5% CI			IV, Random, 95% CI
69.4.1 Hospital setting									
Carbonell 2009	1/702	2/698				_ ,		100%	0.5[0.05,5.47]
	Favours Misopros	stol plus oxytocin	0.01	0.1	1	10	100	Favours Oxytocin	

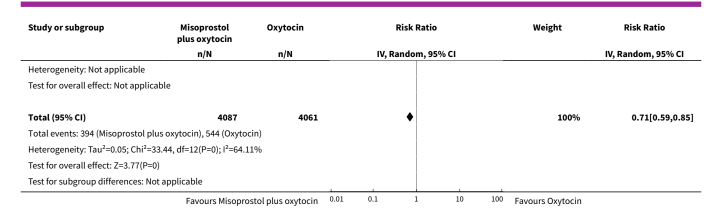




Analysis 69.6. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
69.6.1 Hospital setting					
Adanikin 2012	0/109	0/109			Not estimable
Badejoko 2012	28/126	27/129	+	7.6%	1.06[0.66,1.7]
Bhullar 2004	13/377	20/379	-+ 	4.82%	0.65[0.33,1.29]
Caliskan 2002	28/401	33/407		7.34%	0.86[0.53,1.4]
Caliskan 2003	13/404	28/384		5.24%	0.44[0.23,0.84]
Carbonell 2009	28/702	50/698		7.9%	0.56[0.35,0.87]
Chaudhuri 2015	79/198	132/198	+	13.09%	0.6[0.49,0.73]
Chaudhuri 2016	7/144	19/144		3.6%	0.37[0.16,0.85]
Hofmeyr 2011	22/546	35/553	 	6.79%	0.64[0.38,1.07]
Lapaire 2006	18/24	15/19	+	10.27%	0.95[0.68,1.32]
Nayak 2017	2/100	9/100		1.32%	0.22[0.05,1]
Quibel 2016	68/806	66/797	+	10.34%	1.02[0.74,1.41]
Sood 2012	73/90	77/84	+	14.53%	0.88[0.79,1]
Ugwu 2014	15/60	33/60		7.18%	0.45[0.28,0.75]
Subtotal (95% CI)	4087	4061	◆	100%	0.71[0.59,0.85]
Total events: 394 (Misoprostol p	olus oxytocin), 544 (Oxytoc	in)			
Heterogeneity: Tau ² =0.05; Chi ² =	=33.44, df=12(P=0); I ² =64.1	1%			
Test for overall effect: Z=3.77(P	=0)				
69.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plu	s oxytocin), 0 (Oxytocin)				

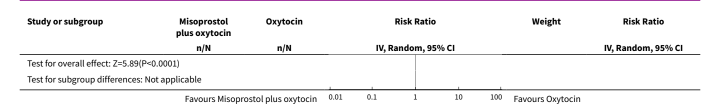




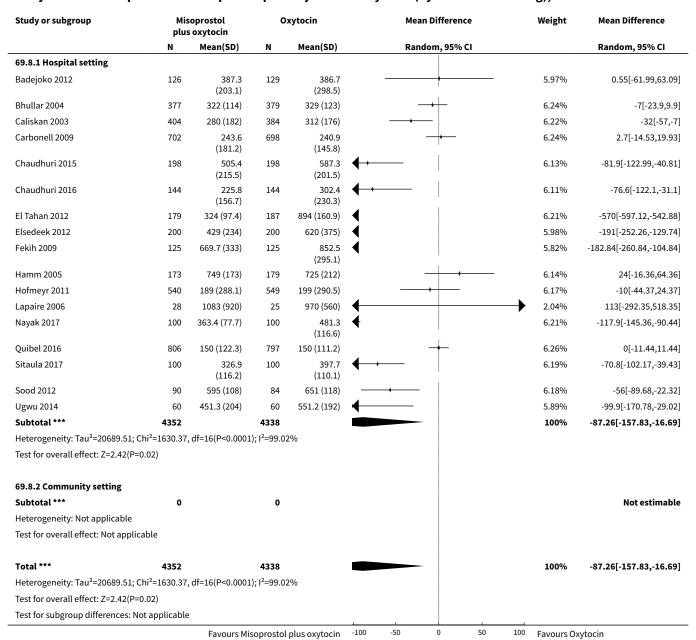
Analysis 69.7. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio IV, Random, 95% CI	
	n/N	n/N	IV, Random, 95% CI			
69.7.1 Hospital setting						
Badejoko 2012	6/126	5/129	- 1	2.48%	1.23[0.38,3.92]	
Bhullar 2004	10/377	13/379		4.2%	0.77[0.34,1.74	
Caliskan 2002	17/401	26/407	-+	6.08%	0.66[0.37,1.2]	
Caliskan 2003	10/404	26/384		4.94%	0.37[0.18,0.75]	
Carbonell 2009	33/702	54/698		8.23%	0.61[0.4,0.92]	
Chaudhuri 2015	18/198	45/198		7.06%	0.4[0.24,0.67]	
Chaudhuri 2016	12/144	22/144		5.39%	0.55[0.28,1.06]	
El Tahan 2012	12/179	52/187		6.1%	0.24[0.13,0.44]	
Elsedeek 2012	14/200	36/200		6.19%	0.39[0.22,0.7]	
Hamm 2005	45/173	76/179		9.88%	0.61[0.45,0.83]	
Hernandez-Castro 2016	6/60	24/60		4.15%	0.25[0.11,0.57]	
Hong 2007	28/96	31/118	-	8.04%	1.11[0.72,1.71]	
Lapaire 2006	0/28	0/25			Not estimable	
Nayak 2017	4/100	7/100		2.36%	0.57[0.17,1.89]	
Pakniat 2015	7/50	7/50		3.26%	1[0.38,2.64]	
Quibel 2016	19/806	25/797	-+	6.16%	0.75[0.42,1.35]	
Sood 2012	20/90	36/84		7.71%	0.52[0.33,0.82]	
Ugwu 2014	16/58	40/60		7.78%	0.41[0.26,0.65]	
Subtotal (95% CI)	4192	4199	♦	100%	0.54[0.44,0.67]	
Total events: 277 (Misoprostol p	us oxytocin), 525 (Oxytoc	in)				
Heterogeneity: Tau ² =0.09; Chi ² =3	32.83, df=16(P=0.01); l ² =5	1.26%				
Test for overall effect: Z=5.89(P<	0.0001)					
69.7.2 Community setting						
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Misoprostol plus	oxytocin), 0 (Oxytocin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applic	able					
Total (95% CI)	4192	4199	•	100%	0.54[0.44,0.67]	
Total events: 277 (Misoprostol p	us oxytocin), 525 (Oxytoc	in)				
Heterogeneity: Tau ² =0.09; Chi ² =3	32.83, df=16(P=0.01); l ² =5	1.26%				



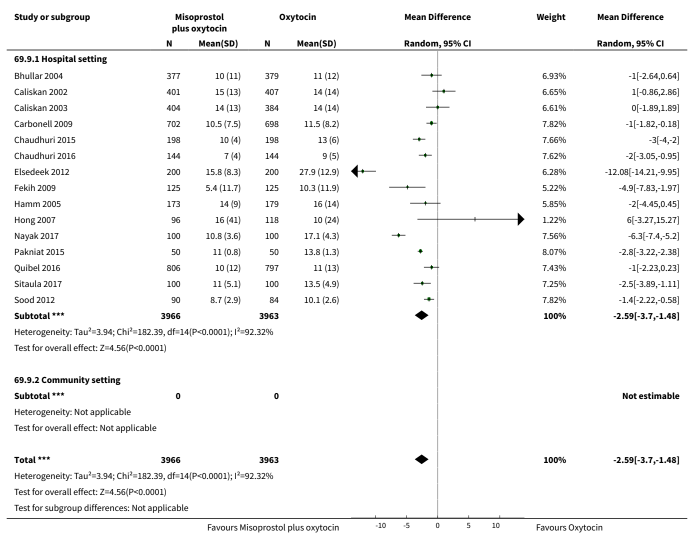


Analysis 69.8. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 8 Blood loss.





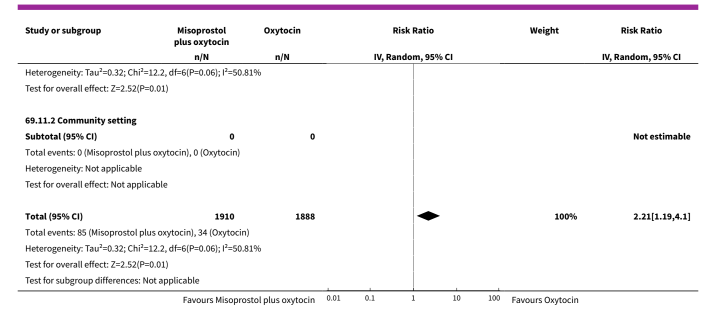
Analysis 69.9. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.



Analysis 69.11. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Misoprostol plus oxytocin	•		Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
69.11.1 Hospital setting						
Adanikin 2012	7/109	2/109	+	10.46%	3.5[0.74,16.47]	
Carbonell 2009	14/702	2/698		11.11%	6.96[1.59,30.51]	
Fekih 2009	26/125	6/125	_ 	19.32%	4.33[1.85,10.16]	
Lapaire 2006	0/28	1/25	+	3.4%	0.3[0.01,7.02]	
Pakniat 2015	6/50	7/50		16.71%	0.86[0.31,2.37]	
Quibel 2016	22/806	8/797		20.15%	2.72[1.22,6.07]	
Sood 2012	10/90	8/84	-	18.85%	1.17[0.48,2.82]	
Subtotal (95% CI)	1910	1888	•	100%	2.21[1.19,4.1]	
Total events: 85 (Misoprostol	plus oxytocin), 34 (Oxytocin)					
	Favours Misopros	stol plus oxytocin	0.01 0.1 1 10	100 Favours Oxytocin		



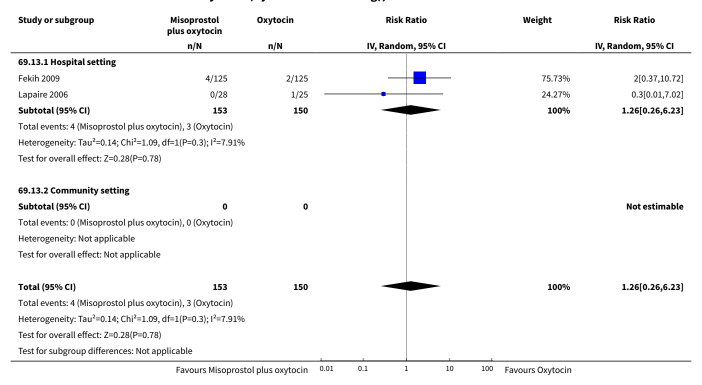


Analysis 69.12. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
69.12.1 Hospital setting						
Adanikin 2012	8/109	3/109	+	7.26%	2.67[0.73,9.79]	
Badejoko 2012	29/126	7/129		14.9%	4.24[1.93,9.33]	
Bhullar 2004	5/377	2/379	+	4.94%	2.51[0.49,12.87]	
Caliskan 2002	3/401	2/407		4.23%	1.52[0.26,9.06]	
Caliskan 2003	3/404	3/384		5.16%	0.95[0.19,4.68]	
Carbonell 2009	28/702	4/698		10.22%	6.96[2.45,19.74]	
El Tahan 2012	16/179	12/187	 • -	16.57%	1.39[0.68,2.86]	
Fekih 2009	15/125	10/125	 • -	15.54%	1.5[0.7,3.21]	
Pakniat 2015	1/50	2/50		2.52%	0.5[0.05,5.34]	
Quibel 2016	18/806	6/797		12.22%	2.97[1.18,7.43]	
Sood 2012	5/90	3/84		6.43%	1.56[0.38,6.31]	
Subtotal (95% CI)	3369	3349	•	100%	2.24[1.52,3.31]	
Total events: 131 (Misoprostol pl	lus oxytocin), 54 (Oxytocir	1)				
Heterogeneity: Tau ² =0.1; Chi ² =13	3.34, df=10(P=0.21); I ² =25.	06%				
Test for overall effect: Z=4.06(P<	0.0001)					
69.12.2 Community setting						
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Misoprostol plus	oxytocin), 0 (Oxytocin)					
Heterogeneity: Not applicable						
Test for overall effect: Not applic	able					
Total (95% CI)	3369	3349	•	100%	2.24[1.52,3.31]	
Total events: 131 (Misoprostol pl	lus oxytocin), 54 (Oxytocir	1)				
Heterogeneity: Tau ² =0.1; Chi ² =13	3.34, df=10(P=0.21); I ² =25.	06%				
Test for overall effect: Z=4.06(P<	0.0001)					
Test for subgroup differences: No	ot applicable					
	Favours Misopros	stol plus oxytocin 0.01	0.1 1 10	L00 Favours Oxytocin		



Analysis 69.13. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 13 Headache.



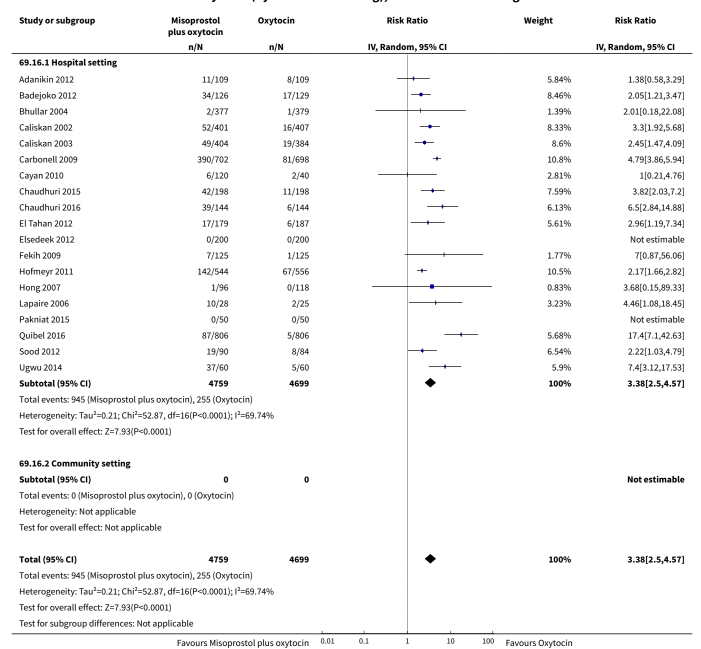
Analysis 69.14. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
69.14.1 Hospital setting					
El Tahan 2012	24/179	13/187	-	100%	1.93[1.01,3.67]
Subtotal (95% CI)	179	187	•	100%	1.93[1.01,3.67]
Total events: 24 (Misoprostol plus o	xytocin), 13 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2(P=0.05)					
69.14.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	179	187	•	100%	1.93[1.01,3.67]
Total events: 24 (Misoprostol plus o	xytocin), 13 (Oxytocin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=2(P=0.05)					
	Favours Misoprost	ol plus oxytocin 0.01	0.1 1 10 1	100 Favours Oxytocin	



Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
Test for subgroup differences: Not applicable			_						
Favours Misoprostol plus oxytocin			0.01	0.1	1	10	100	Favours Oxytocin	

Analysis 69.16. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 16 Shivering.





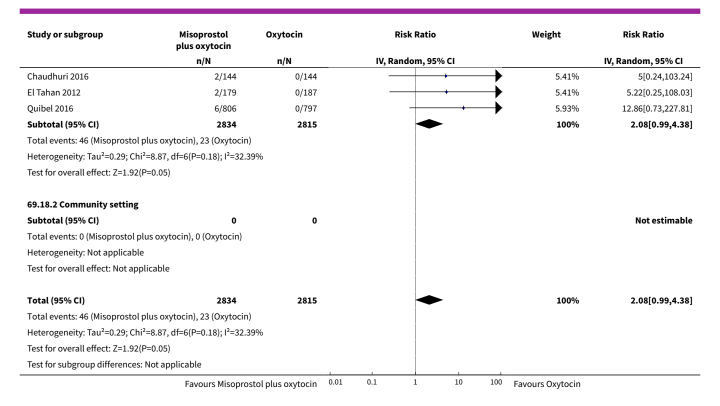
Analysis 69.17. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
69.17.1 Hospital setting						
Adanikin 2012	8/109	6/109		5.95%	1.33[0.48,3.71]	
Badejoko 2012	28/126	7/129		7.07%	4.1[1.86,9.03]	
Caliskan 2002	19/401	6/407		6.5%	3.21[1.3,7.96]	
Caliskan 2003	16/404	5/384		6.09%	3.04[1.13,8.22]	
Carbonell 2009	272/702	23/698		8.86%	11.76[7.78,17.76]	
Cayan 2010	6/120	1/40		2.66%	2[0.25,16.11]	
Chaudhuri 2015	13/198	4/198		5.6%	3.25[1.08,9.79]	
Chaudhuri 2016	8/144	2/144	 	4.01%	4[0.86,18.51]	
El Tahan 2012	16/179	8/187	 • •	6.91%	2.09[0.92,4.76]	
Elsedeek 2012	11/200	13/200		7.13%	0.85[0.39,1.84]	
Fekih 2009	9/125	2/125	<u> </u>	4.07%	4.5[0.99,20.41]	
Hofmeyr 2011	61/522	28/536		8.78%	2.24[1.45,3.44]	
Hong 2007	10/96	5/118	 	5.88%	2.46[0.87,6.95]	
Pakniat 2015	2/50	1/50		2.2%	2[0.19,21.36]	
Quibel 2016	245/806	50/806		9.31%	4.9[3.67,6.54]	
Sood 2012	10/90	6/84		6.21%	1.56[0.59,4.09]	
Ugwu 2014	9/60	1/60		- 2.76%	9[1.18,68.85]	
Subtotal (95% CI)	4332	4275	•	100%	2.99[2,4.45]	
Total events: 743 (Misoprosto	ol plus oxytocin), 168 (Oxytoc	in)				
Heterogeneity: Tau ² =0.42; Ch	i ² =63.65, df=16(P<0.0001); I ² =	74.86%				
Test for overall effect: Z=5.37((P<0.0001)					
69.17.2 Community setting						
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Misoprostol p	olus oxytocin), 0 (Oxytocin)					
Heterogeneity: Not applicable	e					
Test for overall effect: Not app						
Total (95% CI)	4332	4275	•	100%	2.99[2,4.45]	
Total events: 743 (Misoprosto	ol plus oxytocin), 168 (Oxytoc	in)				
Heterogeneity: Tau ² =0.42; Ch	i ² =63.65, df=16(P<0.0001); I ² =	74.86%				
Test for overall effect: Z=5.37((P<0.0001)					
Test for subgroup differences	: Not applicable					

Analysis 69.18. Comparison 69 Misoprostol plus oxytocin vs Oxytocin (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Misoprostol plus oxytocin	Oxytocin		Risk Ratio	Risk Ratio		
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
69.18.1 Hospital setting							
Caliskan 2002	9/401	9/407				28.56%	1.01[0.41,2.53]
Caliskan 2003	13/404	12/384		-		32.54%	1.03[0.48,2.23]
Carbonell 2009	11/702	2/698			_	16.5%	5.47[1.22,24.58]
Chaudhuri 2015	3/198	0/198			—	5.65%	7[0.36,134.64]
	Favours Misopros	stol plus oxytocin	0.01 0.1	. 1 10	100	Favours Oxytocin	





Comparison 70. Injectable prostaglandins vs Misoprostol (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	170	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.1 Hospital setting	2	170	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	4	403	Risk Ratio (IV, Random, 95% CI)	0.88 [0.14, 5.39]
3.1 Hospital setting	4	403	Risk Ratio (IV, Random, 95% CI)	0.88 [0.14, 5.39]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	303	Risk Ratio (IV, Random, 95% CI)	1.60 [0.57, 4.52]
6.1 Hospital setting	3	303	Risk Ratio (IV, Random, 95% CI)	1.60 [0.57, 4.52]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	4	403	Risk Ratio (IV, Random, 95% CI)	1.02 [0.28, 3.69]
7.1 Hospital setting	4	403	Risk Ratio (IV, Random, 95% CI)	1.02 [0.28, 3.69]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	3	270	Mean Difference (IV, Random, 95% CI)	52.99 [-39.74, 145.71]
8.1 Hospital setting	3	270	Mean Difference (IV, Random, 95% CI)	52.99 [-39.74, 145.71]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	220	Mean Difference (IV, Random, 95% CI)	2.28 [-1.35, 5.90]
9.1 Hospital setting	2	220	Mean Difference (IV, Random, 95% CI)	2.28 [-1.35, 5.90]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	2	233	Risk Ratio (IV, Random, 95% CI)	0.23 [0.06, 0.92]
11.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	0.23 [0.06, 0.92]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	233	Risk Ratio (IV, Random, 95% CI)	1.31 [0.01, 163.15]
12.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	1.31 [0.01, 163.15]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

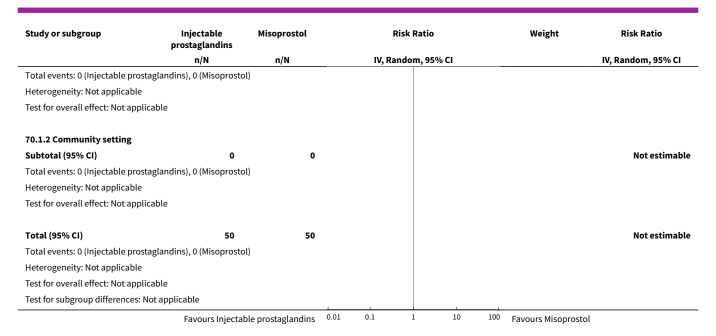


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	0.99 [0.20, 4.80]
14.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	0.99 [0.20, 4.80]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	353	Risk Ratio (IV, Random, 95% CI)	0.04 [0.01, 0.21]
16.1 Hospital setting	3	353	Risk Ratio (IV, Random, 95% CI)	0.04 [0.01, 0.21]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	233	Risk Ratio (IV, Random, 95% CI)	0.08 [0.01, 0.39]
17.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	0.08 [0.01, 0.39]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	233	Risk Ratio (IV, Random, 95% CI)	9.03 [1.70, 48.00]
18.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	9.03 [1.70, 48.00]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

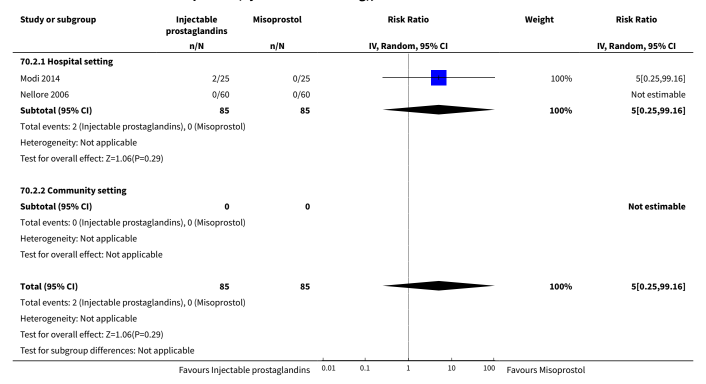
Analysis 70.1. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 1 Death.

Study or subgroup	Injectable prostaglandins	Misoprostol		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
70.1.1 Hospital setting									
Supe 2016	0/50	0/50							Not estimable
Subtotal (95% CI)	50	50							Not estimable
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	



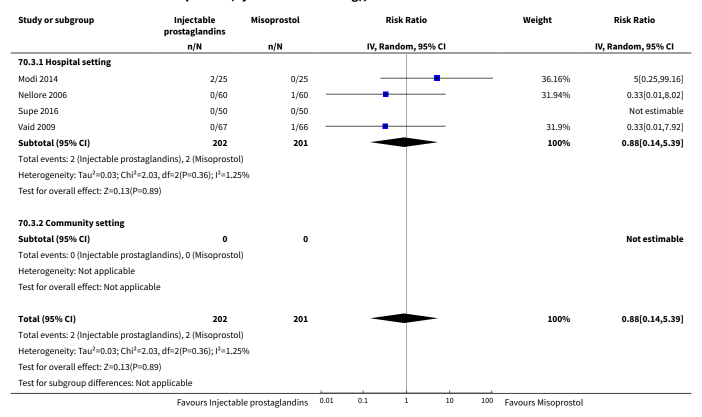


Analysis 70.2. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 2 PPH >= 1000 mL.





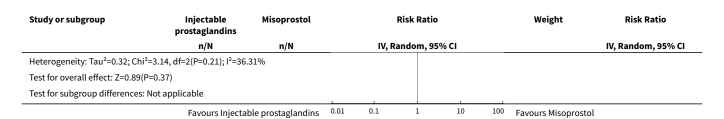
Analysis 70.3. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 3 Blood transfusion.



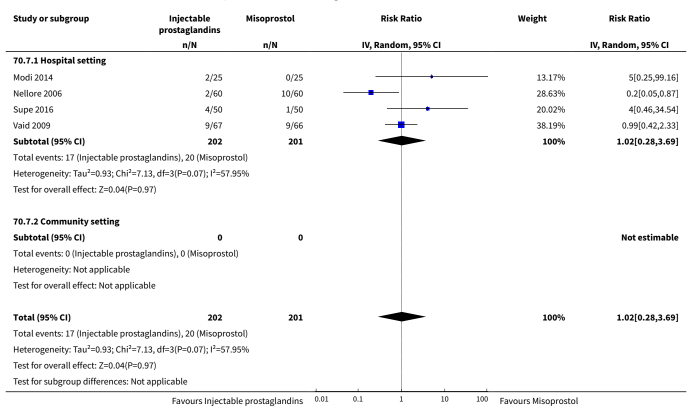
Analysis 70.6. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Misoprostol	R	isk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Ra	ndom, 95% CI			IV, Random, 95% CI
70.6.1 Hospital setting							
Modi 2014	6/25	0/25		+	\rightarrow	11.65%	13[0.77,219.11]
Nellore 2006	3/60	4/60		-		31.99%	0.75[0.18,3.21]
Vaid 2009	13/67	8/66		+		56.36%	1.6[0.71,3.61]
Subtotal (95% CI)	152	151		*		100%	1.6[0.57,4.52]
Total events: 22 (Injectable pr	rostaglandins), 12 (Misoprost	ol)					
Heterogeneity: Tau ² =0.32; Chi	i ² =3.14, df=2(P=0.21); l ² =36.31	L%					
Test for overall effect: Z=0.89((P=0.37)						
70.6.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Injectable pro	ostaglandins), 0 (Misoprostol)						
Heterogeneity: Not applicable	e						
Test for overall effect: Not app	olicable						
Total (95% CI)	152	151		•		100%	1.6[0.57,4.52]
Total events: 22 (Injectable pr	rostaglandins), 12 (Misoprosto	ol)					
	Favours Injectabl	e prostaglandins	0.01 0.1	1 10	100	Favours Misoprostol	





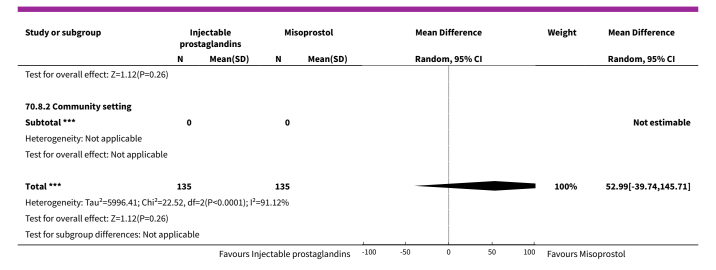
Analysis 70.7. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 7 Additional uterotonics.



Analysis 70.8. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Injectable Misoprostol Mear prostaglandins		Difference		Weight	Mean Difference				
	N	Mean(SD)	N	Mean(SD)		Rande	om, 95% CI			Random, 95% CI
70.8.1 Hospital setting										
Modi 2014	25	435 (147.6)	25	255.8 (102.2)				•	30.72%	179.2[108.84,249.56]
Nellore 2006	60	205 (175)	60	245 (158)		-			32.33%	-40[-99.66,19.66]
Supe 2016	50	153.8 (43.5)	50	124.4 (34.7)			_		36.95%	29.4[13.98,44.82]
Subtotal ***	135		135						100%	52.99[-39.74,145.71]
Heterogeneity: Tau ² =5996.41; C	:hi²=22.52, df=	2(P<0.0001); I ² =9	91.12%							
		Favours Inj	ectable p	rostaglandins	-100	-50	0 50	100	Favours Mis	soprostol





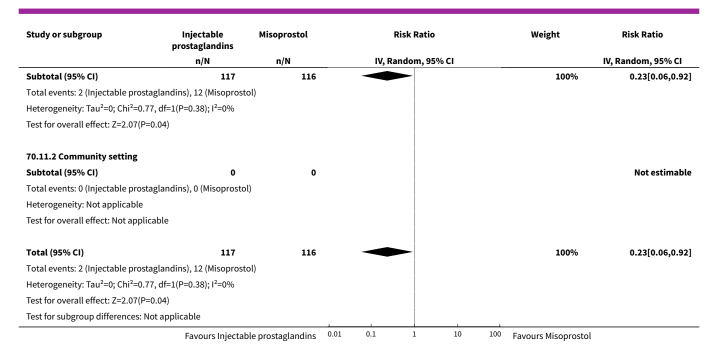
Analysis 70.9. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup		jectable Misopro taglandins		oprostol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
70.9.1 Hospital setting							
Nellore 2006	60	6.2 (2.8)	60	5.8 (2.3)	•	49.32%	0.4[-0.52,1.32]
Supe 2016	50	9 (1.2)	50	4.9 (0.5)		50.68%	4.1[3.74,4.46]
Subtotal ***	110		110		•	100%	2.28[-1.35,5.9]
Heterogeneity: Tau ² =6.72; Chi ² =54.	19, df=1(P	<0.0001); I ² =98.1	5%				
Test for overall effect: Z=1.23(P=0.2	22)						
70.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	le						
Total ***	110		110		•	100%	2.28[-1.35,5.9]
Heterogeneity: Tau ² =6.72; Chi ² =54.	19, df=1(P	<0.0001); I ² =98.1	5%				
Test for overall effect: Z=1.23(P=0.2	.2)						
Test for subgroup differences: Not	applicable						
		Favours Inj	ectable p	rostaglandins -100	-50 0	50 100 Favours Mis	oprostol

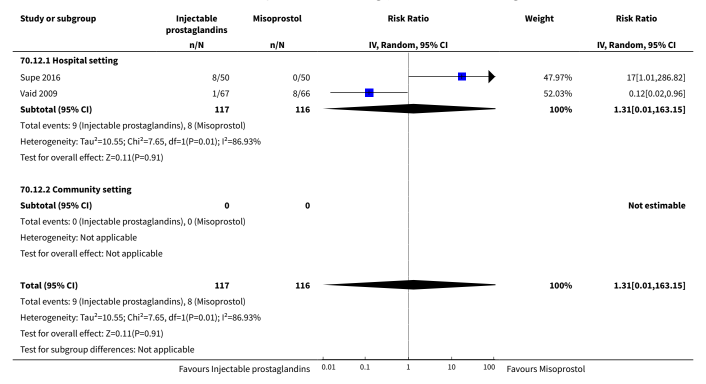
Analysis 70.11. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		IV, Ra	andom, 95	% CI			IV, Random, 95% CI
70.11.1 Hospital setting									
Supe 2016	0/50	6/50	-	-	 			23.14%	0.08[0,1.33]
Vaid 2009	2/67	6/66			- 			76.86%	0.33[0.07,1.57]
	Favours Injectal	ole prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	



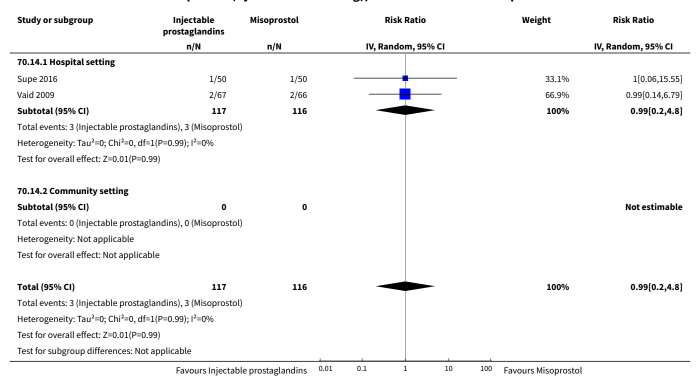


Analysis 70.12. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 12 Vomiting.





Analysis 70.14. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 14 Abdominal pain.



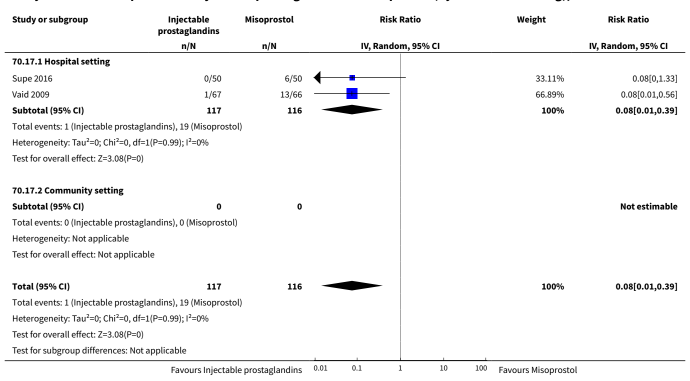
Analysis 70.16. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Injectable prostaglandins	Misoprostol	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI	
70.16.1 Hospital setting						
Nellore 2006	0/60	5/60	•	32.08%	0.09[0.01,1.61]	
Supe 2016	0/50	10/50	-	33.53%	0.05[0,0.79]	
Vaid 2009	0/67	29/66	—	34.39%	0.02[0,0.27]	
Subtotal (95% CI)	177	176		100%	0.04[0.01,0.21]	
Total events: 0 (Injectable prostage	landins), 44 (Misoprostol)				
Heterogeneity: Tau ² =0; Chi ² =0.71,	df=2(P=0.7); I ² =0%					
Test for overall effect: Z=3.85(P=0)						
70.16.2 Community setting						
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Injectable prostag	landins), 0 (Misoprostol)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicab	ole					
Total (95% CI)	177	176	-	100%	0.04[0.01,0.21]	
Total events: 0 (Injectable prostag	landins), 44 (Misoprostol)				
Heterogeneity: Tau ² =0; Chi ² =0.71,	df=2(P=0.7); I ² =0%					
Test for overall effect: Z=3.85(P=0)						
	Favours Injectable	prostaglandins	0.01 0.1 1 10	100 Favours Misoprostol		



Study or subgroup	Injectable prostaglandins	Misoprostol		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
Test for subgroup difference	s: Not applicable		_			1	_		
	Favours Injectab	ole prostaglandins	0.01	0.1	1	10	100	Favours Misoprostol	

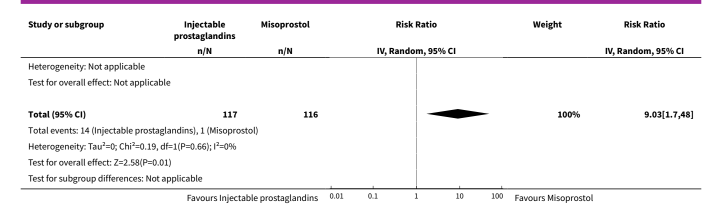
Analysis 70.17. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 17 Fever.



Analysis 70.18. Comparison 70 Injectable prostaglandins vs Misoprostol (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Misoprostol	soprostol Risk Ratio				Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95% CI			IV, Random, 95% CI
70.18.1 Hospital setting								
Supe 2016	7/50	0/50			-	-	34.7%	15[0.88,255.78]
Vaid 2009	7/67	1/66			-		65.3%	6.9[0.87,54.51]
Subtotal (95% CI)	117	116					100%	9.03[1.7,48]
Total events: 14 (Injectable pr	rostaglandins), 1 (Misoprost	ol)						
Heterogeneity: Tau ² =0; Chi ² =0	0.19, df=1(P=0.66); I ² =0%							
Test for overall effect: Z=2.58((P=0.01)							
70.18.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable pro	ostaglandins), 0 (Misoprosto	1)						
	Favours Injectab	le prostaglandins	0.01	0.1	1	100	Favours Misoprostol	





Comparison 71. Misoprostol vs Carbetocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	177	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [0.62, 8.64]
2.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [0.62, 8.64]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	177	Risk Ratio (IV, Random, 95% CI)	2.97 [0.12, 71.85]
3.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	2.97 [0.12, 71.85]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbid- ity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [1.52, 3.50]
6.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	2.31 [1.52, 3.50]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	1	177	Risk Ratio (IV, Random, 95% CI)	3.96 [1.55, 10.07]
7.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	3.96 [1.55, 10.07]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	177	Risk Ratio (IV, Random, 95% CI)	0.99 [0.47, 2.08]
11.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	0.99 [0.47, 2.08]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	1	177	Risk Ratio (IV, Random, 95% CI)	1.48 [0.55, 3.99]
12.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	1.48 [0.55, 3.99]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	177	Risk Ratio (IV, Random, 95% CI)	1.19 [0.71, 1.99]
13.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	1.19 [0.71, 1.99]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	1	177	Risk Ratio (IV, Random, 95% CI)	0.65 [0.52, 0.80]
14.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	0.65 [0.52, 0.80]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

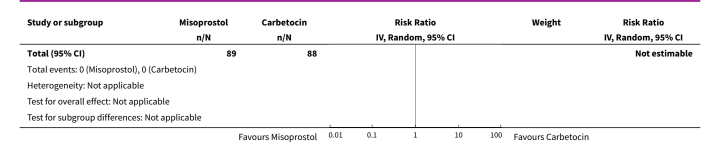


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	177	Risk Ratio (IV, Random, 95% CI)	3.46 [1.67, 7.17]
16.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	3.46 [1.67, 7.17]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	177	Risk Ratio (IV, Random, 95% CI)	2.55 [1.41, 4.64]
17.1 Hospital setting	1	177	Risk Ratio (IV, Random, 95% CI)	2.55 [1.41, 4.64]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

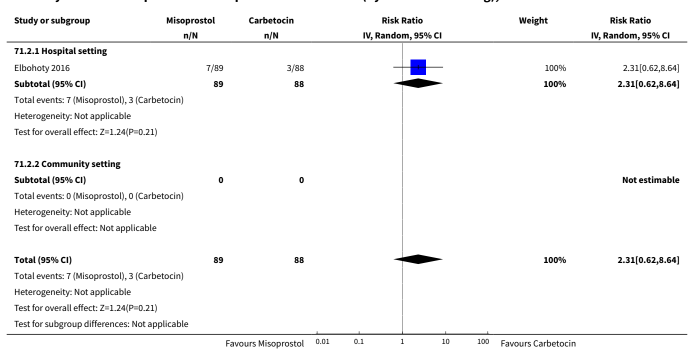
Analysis 71.1. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol	Carbetocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI					IV, Random, 95% CI
71.1.1 Hospital setting									
Elbohoty 2016	0/89	0/88							Not estimable
Subtotal (95% CI)	89	88							Not estimable
Total events: 0 (Misoprostol), 0 (Carbet	ocin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
71.1.2 Community setting									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Misoprostol), 0 (Carbet	ocin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	





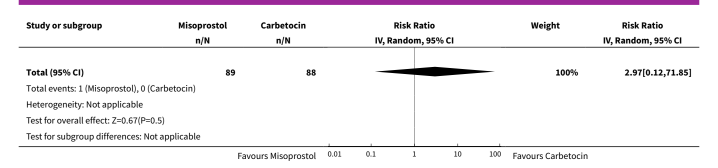
Analysis 71.2. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.



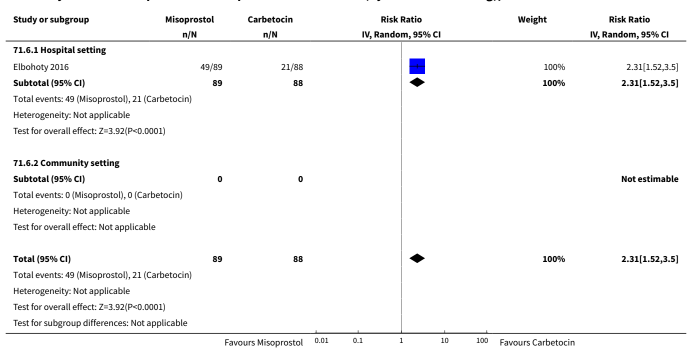
Analysis 71.3. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Carbetocin		Risk I	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Randoi	m, 95% CI			IV, Random, 95% CI
71.3.1 Hospital setting								
Elbohoty 2016	1/89	0/88			-		100%	2.97[0.12,71.85]
Subtotal (95% CI)	89	88				_	100%	2.97[0.12,71.85]
Total events: 1 (Misoprostol), 0 (Carbeto	ocin)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.67(P=0.5)								
71.3.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	ocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable					1			
	Fav	vours Misoprostol	0.01 0.	1 1	. 10	100	Favours Carbetocin	





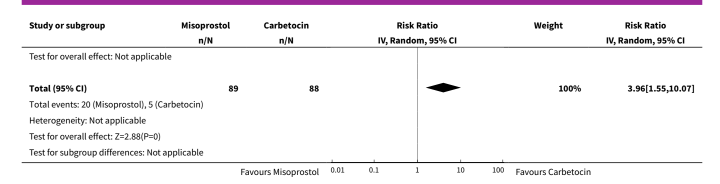
Analysis 71.6. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 6 PPH >= 500 mL.



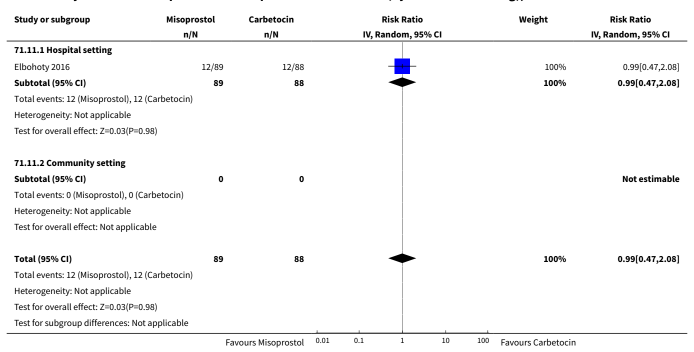
Analysis 71.7. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Carbetocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
71.7.1 Hospital setting									
Elbohoty 2016	20/89	5/88			-	1		100%	3.96[1.55,10.07]
Subtotal (95% CI)	89	88						100%	3.96[1.55,10.07]
Total events: 20 (Misoprostol), 5 (Carbe	tocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=2.88(P=0)									
71.7.2 Community setting									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	ocin)								
Heterogeneity: Not applicable									
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	





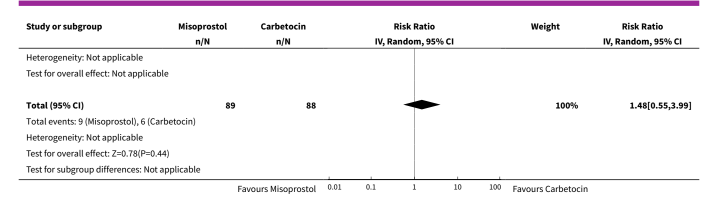
Analysis 71.11. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 11 Nausea.



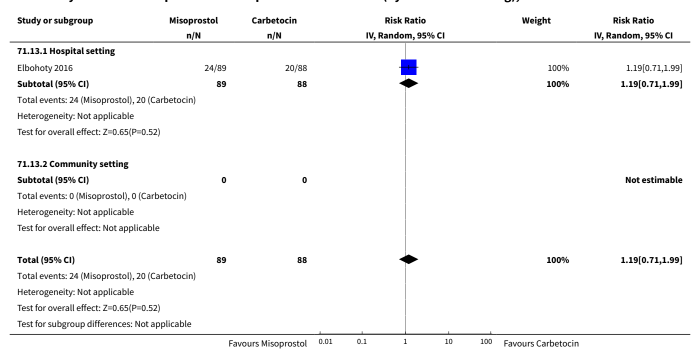
Analysis 71.12. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	dy or subgroup Misoprostol Ca				Risk Ratio			Weight	Risk Ratio
	n/N	n/N n/N		IV, I	Random, 95	% CI			IV, Random, 95% CI
71.12.1 Hospital setting									
Elbohoty 2016	9/89	6/88			-	_		100%	1.48[0.55,3.99]
Subtotal (95% CI)	89	88				-		100%	1.48[0.55,3.99]
Total events: 9 (Misoprostol), 6 (Carbeto	ocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.78(P=0.44)									
71.12.2 Community setting									
Subtotal (95% CI)	0	0							Not estimable
Total events: 0 (Misoprostol), 0 (Carbeto	ocin)								
	Fa	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	





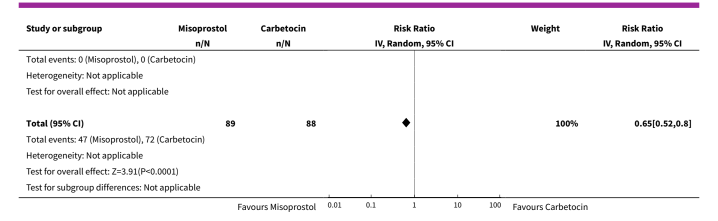
Analysis 71.13. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 13 Headache.



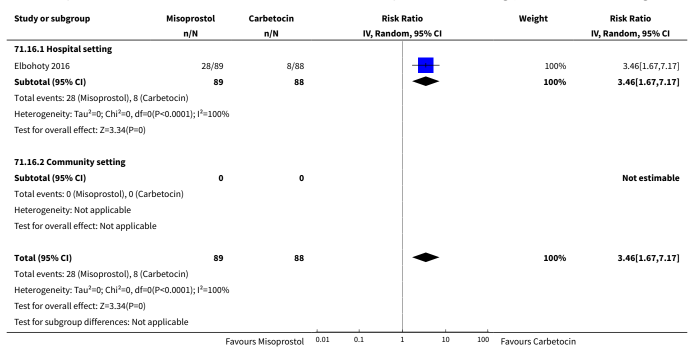
Analysis 71.14. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Misoprostol Carbetocin n/N n/N		Risk Rat	io	Weight	Risk Ratio IV, Random, 95% CI
	n/N			IV, Random,	95% CI		
71.14.1 Hospital setting							
Elbohoty 2016	47/89	72/88		+		100%	0.65[0.52,0.8]
Subtotal (95% CI)	89	88		•		100%	0.65[0.52,0.8]
Total events: 47 (Misoprostol), 7	'2 (Carbetocin)						
Heterogeneity: Not applicable							
Test for overall effect: Z=3.91(P<	<0.0001)						
71.14.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
	Fav	vours Misoprostol	0.01	0.1 1	10 100	Favours Carbetocin	





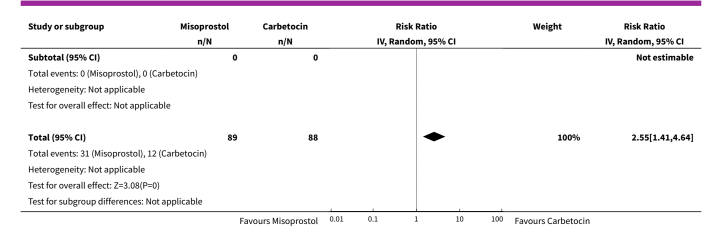
Analysis 71.16. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 16 Shivering.



Analysis 71.17. Comparison 71 Misoprostol vs Carbetocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol	Carbetocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 959	% CI			IV, Random, 95% CI
71.17.1 Hospital setting									
Elbohoty 2016	31/89	12/88			-	-		100%	2.55[1.41,4.64]
Subtotal (95% CI)	89	88				-		100%	2.55[1.41,4.64]
Total events: 31 (Misoprostol), 12 (Carbetocin)								
Heterogeneity: Not applicable									
Test for overall effect: Z=3.08(P=0)									
71.17.2 Community setting									
	Far	vours Misoprostol	0.01	0.1	1	10	100	Favours Carbetocin	





Comparison 72. Ergometrine vs Misoprostol (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	7	5254	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	6	4054	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	11	5562	Risk Ratio (IV, Random, 95% CI)	1.25 [0.45, 3.48]
2.1 Hospital setting	10	4362	Risk Ratio (IV, Random, 95% CI)	1.42 [0.47, 4.34]
2.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.17]
3 Blood transfusion	13	5308	Risk Ratio (IV, Random, 95% CI)	1.71 [0.65, 4.50]
3.1 Hospital setting	12	4108	Risk Ratio (IV, Random, 95% CI)	2.02 [0.73, 5.57]
3.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.17]
4 Severe maternal morbidity: intensive care admissions	1	864	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.16]
4.1 Hospital setting	1	864	Risk Ratio (IV, Random, 95% CI)	0.33 [0.01, 8.16]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	16	6459	Risk Ratio (IV, Random, 95% CI)	1.18 [0.79, 1.75]



Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
6.1 Hospital setting	15	5259	Risk Ratio (IV, Random, 95% CI)	1.17 [0.77, 1.79]
6.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	1.25 [0.34, 4.63]
7 Additional uterotonics	16	6565	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.60]
7.1 Hospital setting	15	5365	Risk Ratio (IV, Random, 95% CI)	1.06 [0.68, 1.65]
7.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.75 [0.17, 3.34]
8 Blood loss	15	6337	Mean Difference (IV, Random, 95% CI)	11.66 [-9.85, 33.17]
8.1 Hospital setting	14	5137	Mean Difference (IV, Random, 95% CI)	6.55 [-11.42, 24.51]
8.2 Community setting	1	1200	Mean Difference (IV, Random, 95% CI)	71.30 [60.86, 81.74]
9 Change in haemoglobin	8	4438	Mean Difference (IV, Random, 95% CI)	0.91 [-0.43, 2.26]
9.1 Hospital setting	8	4438	Mean Difference (IV, Random, 95% CI)	0.91 [-0.43, 2.26]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	13	5202	Risk Ratio (IV, Random, 95% CI)	1.53 [1.15, 2.04]
11.1 Hospital setting	12	4002	Risk Ratio (IV, Random, 95% CI)	1.43 [1.02, 2.00]
11.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	1.98 [1.49, 2.64]
12 Vomiting	14	6136	Risk Ratio (IV, Random, 95% CI)	1.22 [0.81, 1.83]
12.1 Hospital setting	13	4936	Risk Ratio (IV, Random, 95% CI)	1.26 [0.77, 2.05]
12.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	1.13 [0.70, 1.83]
13 Headache	8	3369	Risk Ratio (IV, Random, 95% CI)	1.70 [0.70, 4.12]
13.1 Hospital setting	8	3369	Risk Ratio (IV, Random, 95% CI)	1.70 [0.70, 4.12]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	2	233	Risk Ratio (IV, Random, 95% CI)	1.58 [0.04, 65.80]
14.1 Hospital setting	2	233	Risk Ratio (IV, Random, 95% CI)	1.58 [0.04, 65.80]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	2	300	Risk Ratio (IV, Random, 95% CI)	7.55 [0.94, 60.53]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
15.1 Hospital setting	2	300	Risk Ratio (IV, Random, 95% CI)	7.55 [0.94, 60.53]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	17	6860	Risk Ratio (IV, Random, 95% CI)	0.34 [0.26, 0.44]
16.1 Hospital setting	16	5660	Risk Ratio (IV, Random, 95% CI)	0.31 [0.22, 0.44]
16.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.39 [0.33, 0.47]
17 Fever	14	6330	Risk Ratio (IV, Random, 95% CI)	0.24 [0.17, 0.33]
17.1 Hospital setting	13	5130	Risk Ratio (IV, Random, 95% CI)	0.20 [0.15, 0.28]
17.2 Community setting	1	1200	Risk Ratio (IV, Random, 95% CI)	0.45 [0.29, 0.70]
18 Diarrhoea	8	4165	Risk Ratio (IV, Random, 95% CI)	1.04 [0.46, 2.34]
18.1 Hospital setting	8	4165	Risk Ratio (IV, Random, 95% CI)	1.04 [0.46, 2.34]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

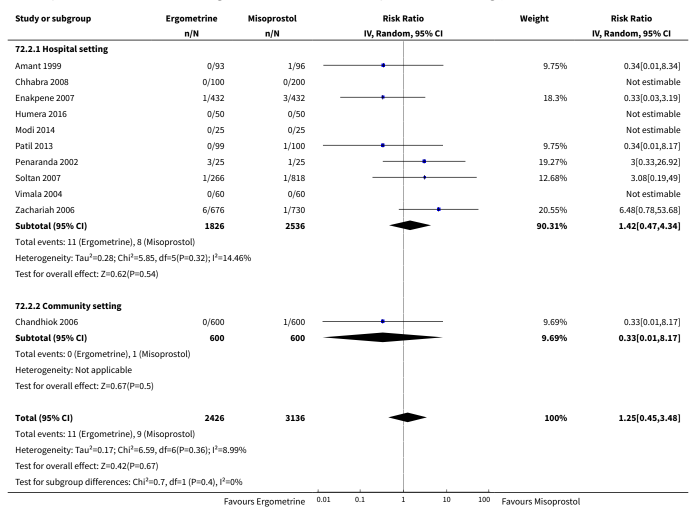
Analysis 72.1. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 1 Death.

Study or subgroup	Ergometrine	Misoprostol		Risk R	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Randon	n, 95% CI			IV, Random, 95% CI
72.1.1 Hospital setting								
Chhabra 2008	0/100	0/200						Not estimable
Enakpene 2007	0/432	0/432						Not estimable
Fawzy 2012	0/100	0/200						Not estimable
Soltan 2007	0/266	0/818						Not estimable
Supe 2016	0/50	0/50						Not estimable
Zachariah 2006	0/676	0/730						Not estimable
Subtotal (95% CI)	1624	2430						Not estimable
Total events: 0 (Ergometrine), 0 (Mis	oprostol)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
	Fav	ours Ergometrine	0.01	0.1 1	10	100	Favours Misoprostol	



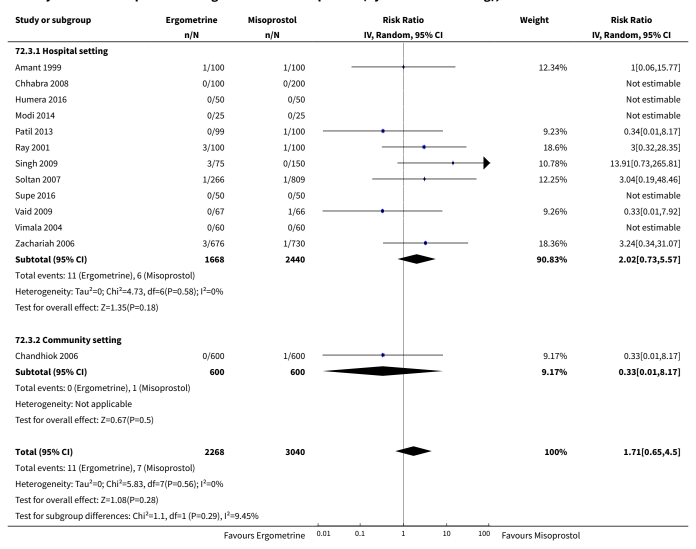
Study or subgroup E	rgometrine	Misoprostol			Risk Ratio			Weight	Risk Ratio
	n/N	n/N	n/N		IV, Random, 95% CI				IV, Random, 95% CI
72.1.2 Community setting									
Chandhiok 2006	0/600	0/600							Not estimable
Subtotal (95% CI)	600	600							Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Total (95% CI)	2224	3030							Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
Test for subgroup differences: Not applic	cable						1		
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	

Analysis 72.2. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 2 PPH >= 1000 mL.





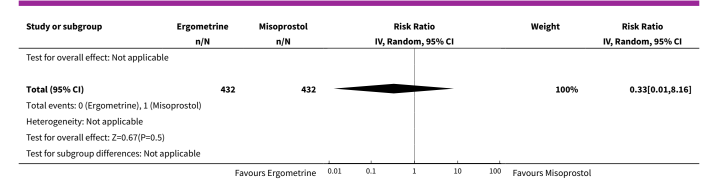
Analysis 72.3. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 3 Blood transfusion.



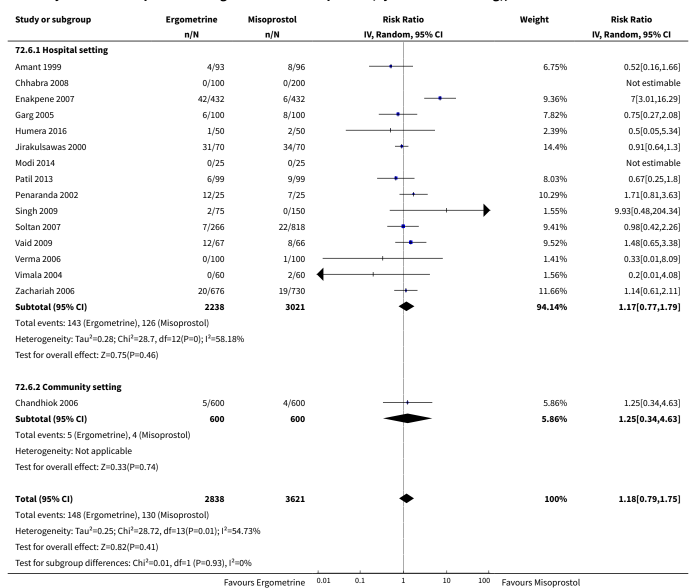
Analysis 72.4. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup E	rgometrine	Misoprostol		Risk	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rand	om, 95% CI			IV, Random, 95% CI
72.4.1 Hospital setting								
Enakpene 2007	0/432	1/432	-	-			100%	0.33[0.01,8.16]
Subtotal (95% CI)	432	432					100%	0.33[0.01,8.16]
Total events: 0 (Ergometrine), 1 (Misopro	ostol)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.67(P=0.5)								
72.4.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Ergometrine), 0 (Misopro	ostol)							
Heterogeneity: Not applicable			1	1				
	Fav	ours Ergometrine	0.01	0.1	1 10	100	Favours Misoprostol	



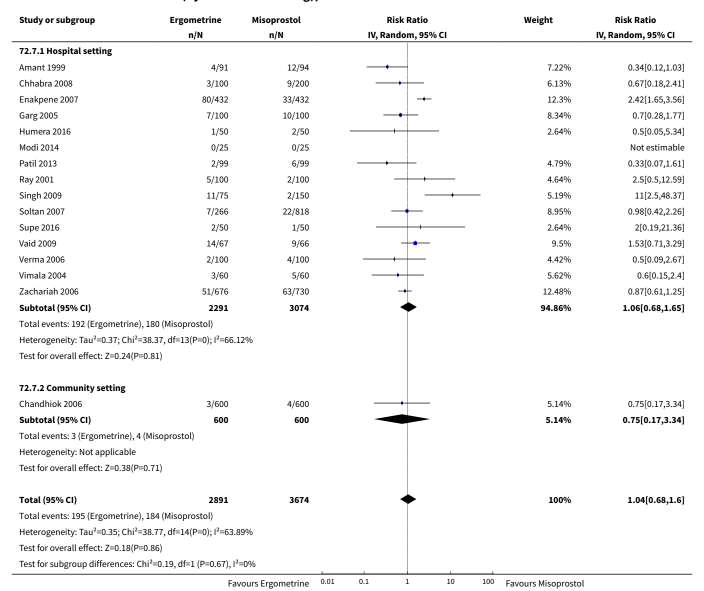


Analysis 72.6. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 6 PPH >= 500 mL.





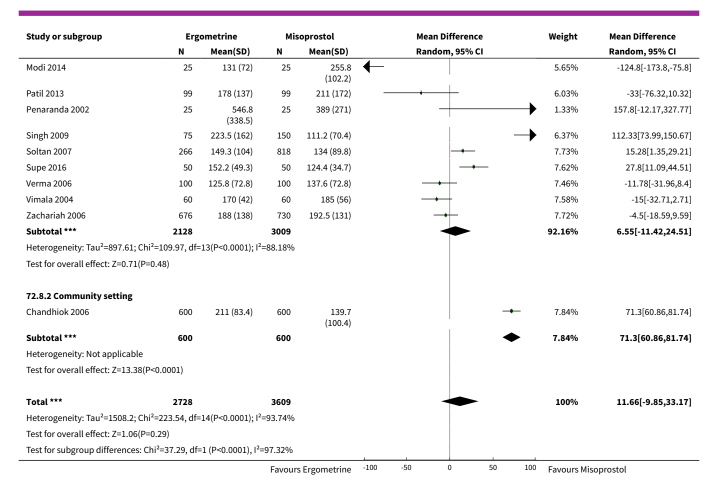
Analysis 72.7. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 7 Additional uterotonics.



Analysis 72.8. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	Mis	oprostol		М	ean Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Random, 95% CI				Random, 95% CI	
72.8.1 Hospital setting		,									
Chhabra 2008	100	150 (52)	200	150 (50)			+			7.78%	0[-12.32,12.32]
Enakpene 2007	432	246 (175.5)	432	191.6 (134.5)					-	7.43%	54.4[33.55,75.25]
Fawzy 2012	100	275.8 (165.5)	200	233.5 (132.9)				-	_	6.44%	42.22[4.92,79.52]
Humera 2016	50	172.8 (79.7)	50	195.1 (94.3)			+			6.64%	-22.3[-56.5,11.9]
Jirakulsawas 2000	70	484.7 (120.1)	70	490.5 (109.8)		_	*	_		6.38%	-5.79[-43.91,32.33]
			Favour	s Ergometrine	-100	-50	0	50	100	Favours Mis	oprostol

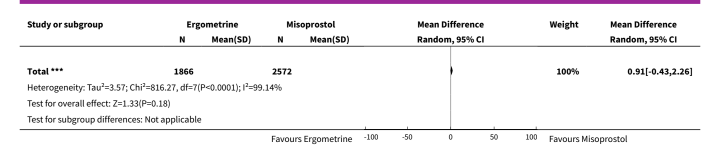




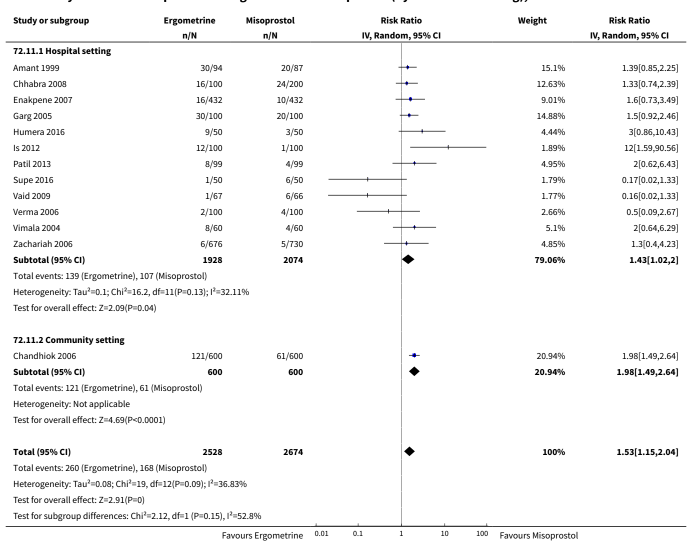
Analysis 72.9. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Ergo	ometrine	Mis	oprostol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
72.9.1 Hospital setting							
Chhabra 2008	100	8 (6.3)	200	7.8 (2)	+	11.87%	0.2[-1.07,1.47]
Enakpene 2007	432	4.1 (5.1)	432	1.4 (2)	•	13%	2.72[2.2,3.24]
Gore 2017	182	9.4 (0.7)	182	9.7 (0.8)	•	13.24%	-0.3[-0.45,-0.15]
Soltan 2007	266	4.6 (4)	818	3.6 (2.9)	•	13%	1[0.48,1.52]
Supe 2016	50	7.6 (0.2)	50	4.9 (0.5)	•	13.24%	2.7[2.55,2.85]
Verma 2006	100	2.5 (3)	100	3.1 (3)	•	12.62%	-0.6[-1.43,0.23]
Vimala 2004	60	8 (6.3)	60	7.6 (2)		11.01%	0.4[-1.27,2.07]
Zachariah 2006	676	14.8 (11.9)	730	13.8 (10.7)	•	12.02%	1[-0.19,2.19]
Subtotal ***	1866		2572			100%	0.91[-0.43,2.26]
Heterogeneity: Tau ² =3.57; Chi ² =81	16.27, df=7(F	P<0.0001); I ² =99	.14%				
Test for overall effect: Z=1.33(P=0.	.18)						
72.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applica	ble						
			Favour	s Ergometrine -1	00 -50 0 50	100 Favours Mis	oprostol



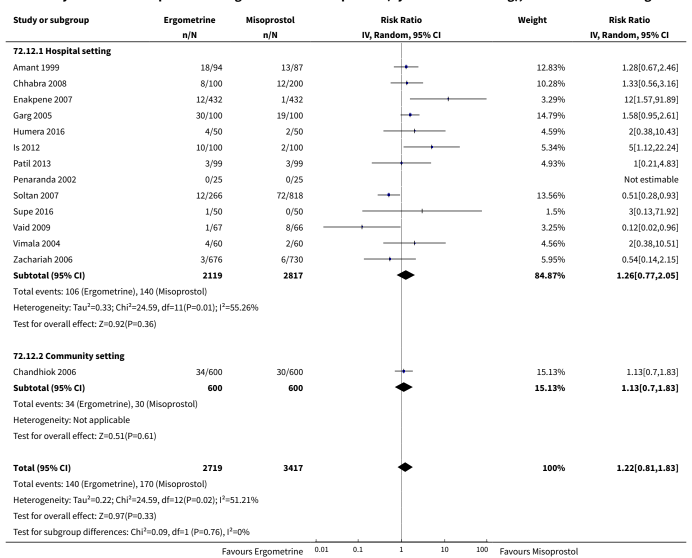


Analysis 72.11. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 11 Nausea.





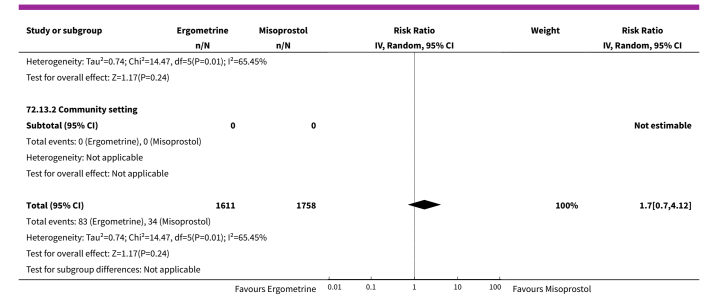
Analysis 72.12. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 12 Vomiting.



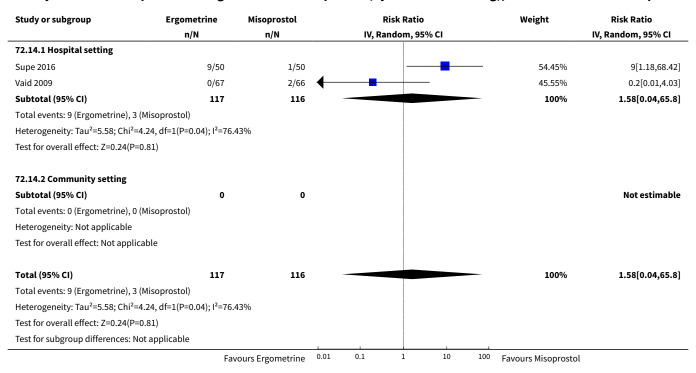
Analysis 72.13. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
72.13.1 Hospital setting					
Amant 1999	12/94	10/87	-	22.73%	1.11[0.51,2.44]
Chhabra 2008	8/100	14/200	-	22.23%	1.14[0.5,2.63]
Enakpene 2007	54/432	1/432		11.67%	54[7.5,388.59]
Garg 2005	3/100	4/100		15.71%	0.75[0.17,3.27]
Humera 2016	0/50	0/50			Not estimable
Patil 2013	0/99	0/99			Not estimable
Vimala 2004	4/60	3/60		15.87%	1.33[0.31,5.7]
Zachariah 2006	2/676	2/730		11.78%	1.08[0.15,7.64]
Subtotal (95% CI)	1611	1758	-	100%	1.7[0.7,4.12]
Total events: 83 (Ergometrine),	34 (Misoprostol)				
	Fav	ours Ergometrine	0.01 0.1 1 10	100 Favours Misoprostol	





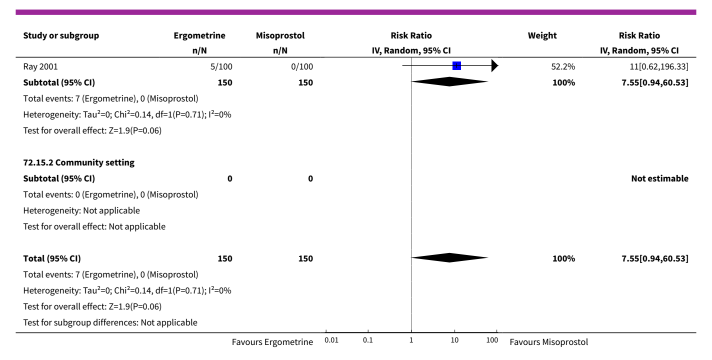
Analysis 72.14. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 14 Abdominal pain.



Analysis 72.15. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 15 Hypertension.

Study or subgroup	Ergometrine	Misoprostol		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, R	andom, 9	5% CI			IV, Random, 95% CI
72.15.1 Hospital setting									
Humera 2016	2/50	0/50				•	<u> </u>	47.8%	5[0.25,101.58]
	Fav	ours Ergometrine	0.01	0.1	1	10	100	Favours Misoprostol	_

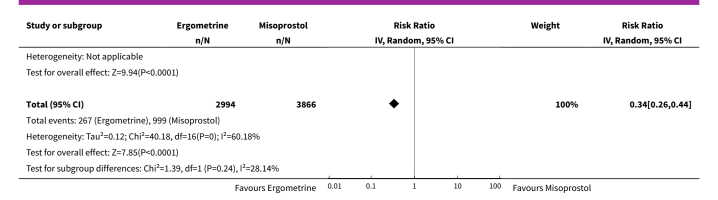




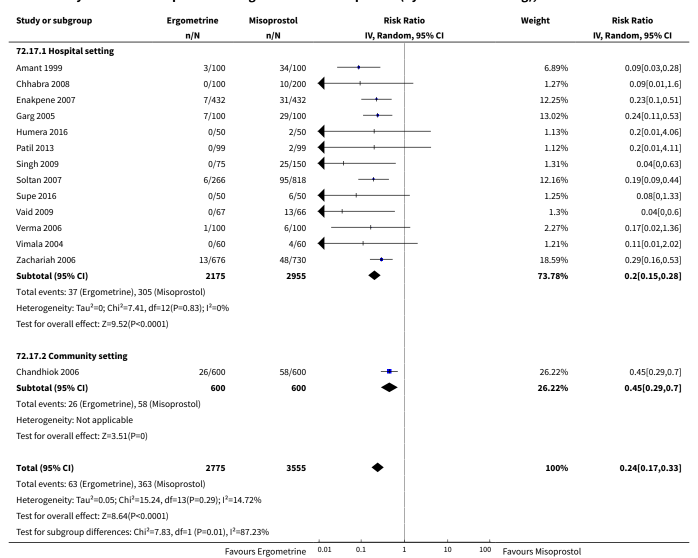
Analysis 72.16. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Ergometrine	Misoprostol	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
72.16.1 Hospital setting					
Amant 1999	38/94	66/86	+	13.5%	0.53[0.4,0.69]
Chhabra 2008	0/100	14/200	•	0.88%	0.07[0,1.14]
Enakpene 2007	21/432	23/432	-	9.15%	0.91[0.51,1.62]
Fawzy 2012	8/100	50/200		7.57%	0.32[0.16,0.65]
Garg 2005	10/100	31/100		8.14%	0.32[0.17,0.62]
Humera 2016	4/50	19/50		4.96%	0.21[0.08,0.57]
Is 2012	2/100	1/100		1.19%	2[0.18,21.71]
Patil 2013	2/99	36/99		3.03%	0.06[0.01,0.22]
Penaranda 2002	1/25	1/25		0.94%	1[0.07,15.12]
Singh 2009	0/75	19/150	—	0.89%	0.05[0,0.83]
Soltan 2007	32/266	309/818		12.59%	0.32[0.23,0.45]
Supe 2016	0/50	10/50		0.88%	0.05[0,0.79]
Vaid 2009	4/67	29/66		5.07%	0.14[0.05,0.37]
Verma 2006	4/100	18/100		4.68%	0.22[0.08,0.63]
Vimala 2004	0/60	13/60	—	0.88%	0.04[0,0.61]
Zachariah 2006	26/676	68/730		11.07%	0.41[0.27,0.64]
Subtotal (95% CI)	2394	3266	◆	85.42%	0.31[0.22,0.44]
Total events: 152 (Ergometrine)	, 707 (Misoprostol)				
Heterogeneity: Tau ² =0.21; Chi ² =	=40.18, df=15(P=0); l ² =62.6	57%			
Test for overall effect: Z=6.63(P-	<0.0001)				
72.16.2 Community setting					
Chandhiok 2006	115/600	292/600	+	14.58%	0.39[0.33,0.47]
Subtotal (95% CI)	600	600	•	14.58%	0.39[0.33,0.47]
Total events: 115 (Ergometrine)	, 292 (Misoprostol)				





Analysis 72.17. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 17 Fever.





Analysis 72.18. Comparison 72 Ergometrine vs Misoprostol (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Ergometrine	Misoprostol		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95% CI			IV, Random, 95% CI
72.18.1 Hospital setting							
Amant 1999	0/94	1/86		-		6.44%	0.31[0.01,7.39]
Enakpene 2007	0/432	0/432					Not estimable
Garg 2005	3/100	3/100				26.34%	1[0.21,4.84]
Patil 2013	0/99	0/99					Not estimable
Soltan 2007	4/266	10/818				49.38%	1.23[0.39,3.89]
Supe 2016	0/50	0/50					Not estimable
Vaid 2009	0/67	1/66	-	-+		6.46%	0.33[0.01,7.92]
Zachariah 2006	2/676	1/730				11.38%	2.16[0.2,23.76]
Subtotal (95% CI)	1784	2381		*		100%	1.04[0.46,2.34]
Total events: 9 (Ergometrine), 16 (Mis	soprostol)						
Heterogeneity: Tau ² =0; Chi ² =1.51, df	=4(P=0.82); I ² =0%			İ			
Test for overall effect: Z=0.1(P=0.92)							
72.18.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Ergometrine), 0 (Misc	oprostol)						
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	2						
Total (95% CI)	1784	2381		•		100%	1.04[0.46,2.34]
Total events: 9 (Ergometrine), 16 (Mis	soprostol)						
Heterogeneity: Tau ² =0; Chi ² =1.51, df	=4(P=0.82); I ² =0%						
Test for overall effect: Z=0.1(P=0.92)							
Test for subgroup differences: Not ap	plicable						
	Fav	ours Ergometrine	0.01 0.1	1	10 100	Favours Misoprostol	

Comparison 73. Misoprostol vs Ergometrine plus oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	4	3258	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	4	3258	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	9	6028	Risk Ratio (IV, Random, 95% CI)	1.61 [1.08, 2.40]
2.1 Hospital setting	9	6028	Risk Ratio (IV, Random, 95% CI)	1.61 [1.08, 2.40]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	8	6335	Risk Ratio (IV, Random, 95% CI)	1.41 [0.91, 2.21]
3.1 Hospital setting	8	6335	Risk Ratio (IV, Random, 95% CI)	1.41 [0.91, 2.21]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal mor- bidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	10	6492	Risk Ratio (IV, Random, 95% CI)	1.75 [1.35, 2.26]
6.1 Hospital setting	10	6492	Risk Ratio (IV, Random, 95% CI)	1.75 [1.35, 2.26]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	9	6395	Risk Ratio (IV, Random, 95% CI)	1.87 [1.49, 2.36]
7.1 Hospital setting	9	6395	Risk Ratio (IV, Random, 95% CI)	1.87 [1.49, 2.36]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	8	5634	Mean Difference (IV, Random, 95% CI)	22.86 [4.50, 41.22]
8.1 Hospital setting	8	5634	Mean Difference (IV, Random, 95% CI)	22.86 [4.50, 41.22]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	9	5588	Mean Difference (IV, Random, 95% CI)	1.09 [-0.49, 2.67]
9.1 Hospital setting	9	5588	Mean Difference (IV, Random, 95% CI)	1.09 [-0.49, 2.67]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	4	3399	Risk Ratio (IV, Random, 95% CI)	0.71 [0.61, 0.84]
11.1 Hospital setting	4	3399	Risk Ratio (IV, Random, 95% CI)	0.71 [0.61, 0.84]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	6	4983	Risk Ratio (IV, Random, 95% CI)	0.72 [0.50, 1.04]
12.1 Hospital setting	6	4983	Risk Ratio (IV, Random, 95% CI)	0.72 [0.50, 1.04]



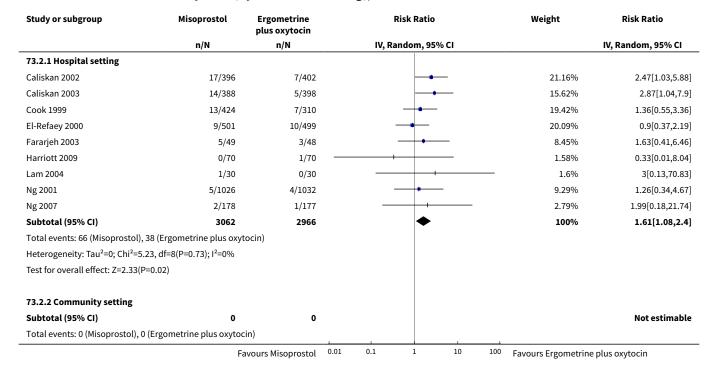
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	3	3259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.47, 1.76]
13.1 Hospital setting	3	3259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.47, 1.76]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	1	846	Risk Ratio (IV, Random, 95% CI)	0.90 [0.79, 1.02]
14.1 Hospital setting	1	846	Risk Ratio (IV, Random, 95% CI)	0.90 [0.79, 1.02]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	2553	Risk Ratio (IV, Random, 95% CI)	0.50 [0.18, 1.41]
15.1 Hospital setting	3	2553	Risk Ratio (IV, Random, 95% CI)	0.50 [0.18, 1.41]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	6	4980	Risk Ratio (IV, Random, 95% CI)	2.71 [1.95, 3.76]
16.1 Hospital setting	6	4980	Risk Ratio (IV, Random, 95% CI)	2.71 [1.95, 3.76]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	7	5045	Risk Ratio (IV, Random, 95% CI)	5.33 [2.87, 9.88]
17.1 Hospital setting	7	5045	Risk Ratio (IV, Random, 95% CI)	5.33 [2.87, 9.88]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	6	3659	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.57]
18.1 Hospital setting	6	3659	Risk Ratio (IV, Random, 95% CI)	1.04 [0.68, 1.57]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	1		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
20.1 Hospital setting	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]



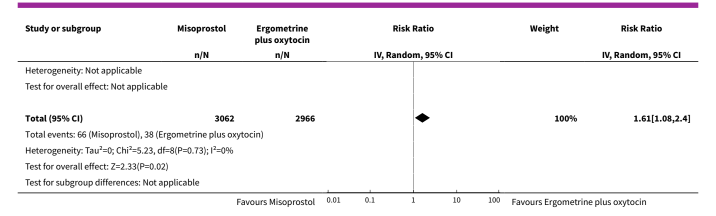
Analysis 73.1. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
73.1.1 Hospital setting					
El-Refaey 2000	0/501	0/499			Not estimable
Harriott 2009	0/70	0/70			Not estimable
Lam 2004	0/30	0/30			Not estimable
Ng 2001	0/1026	0/1032			Not estimable
Subtotal (95% CI)	1627	1631			Not estimable
Total events: 0 (Misoprostol), 0 (Ergor	metrine plus oxytoci	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
73.1.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Ergor	metrine plus oxytoci	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	1627	1631			Not estimable
Total events: 0 (Misoprostol), 0 (Ergor	metrine plus oxytoci	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Test for subgroup differences: Not ap	plicable				
	Fa	ours Misoprostol	0.01 0.1 1 10	100 Favours Ergometr	ine plus oxytocin

Analysis 73.2. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.





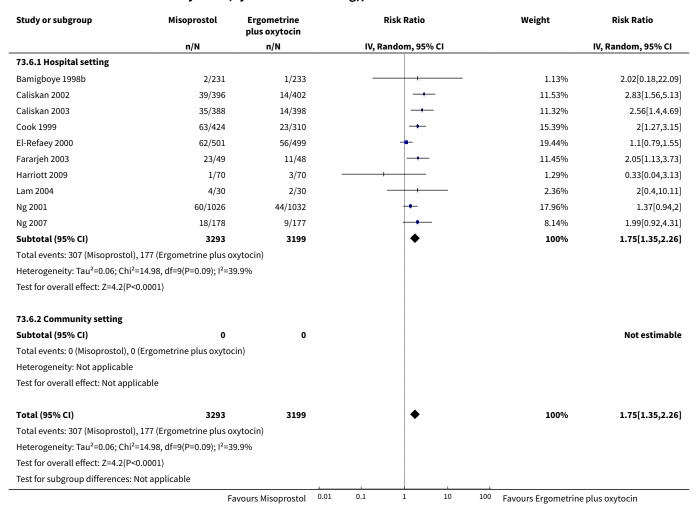


Analysis 73.3. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
73.3.1 Hospital setting					
Bamigboye 1998b	0/231	0/233			Not estimable
Caliskan 2002	12/396	4/402		13.34%	3.05[0.99,9.36]
Caliskan 2003	14/388	6/398		17.71%	2.39[0.93,6.16]
Cook 1999	5/424	3/310		8.79%	1.22[0.29,5.06]
El-Refaey 2000	9/501	11/499		20.13%	0.81[0.34,1.95]
Harriott 2009	0/70	0/70			Not estimable
Ng 2001	15/1026	16/1032		27.83%	0.94[0.47,1.9]
Ng 2007	8/178	4/177		12.21%	1.99[0.61,6.49]
Subtotal (95% CI)	3214	3121	•	100%	1.41[0.91,2.21]
Total events: 63 (Misoprostol), 4	14 (Ergometrine plus oxyt	ocin)			
Heterogeneity: Tau ² =0.06; Chi ² =	-6.15, df=5(P=0.29); l ² =18.	66%			
Test for overall effect: Z=1.53(P=	=0.13)				
73.3.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0	(Ergometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
	cable				
Test for overall effect: Not appli					
Total (95% CI)	3214	3121	•	100%	1.41[0.91,2.21]
			•	100%	1.41[0.91,2.21]
Total (95% CI)	14 (Ergometrine plus oxyt	ocin)	•	100%	1.41[0.91,2.21]
Total (95% CI) Total events: 63 (Misoprostol), 4	14 (Ergometrine plus oxyt =6.15, df=5(P=0.29); I ² =18.	ocin)	•	100%	1.41[0.91,2.21]



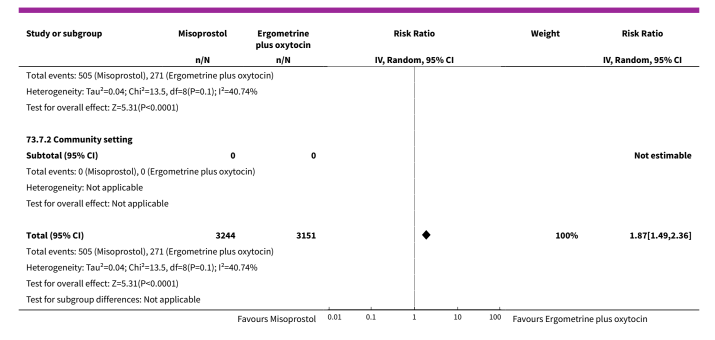
Analysis 73.6. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.



Analysis 73.7. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk	Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI			IV, Random, 95% CI
73.7.1 Hospital setting							
Bamigboye 1998b	4/231	1/233				1.09%	4.03[0.45,35.83]
Caliskan 2002	33/396	9/402		-		7.89%	3.72[1.8,7.68]
Caliskan 2003	23/388	9/398				7.35%	2.62[1.23,5.59]
Cook 1999	95/424	28/310		-		17.13%	2.48[1.67,3.68]
El-Refaey 2000	68/501	50/499		-		19.5%	1.35[0.96,1.91]
Harriott 2009	6/70	6/70				4.05%	1[0.34,2.95]
Lam 2004	3/30	0/30		+	\longrightarrow	0.62%	7[0.38,129.93]
Ng 2001	232/1026	144/1032		•		27.76%	1.62[1.34,1.96]
Ng 2007	41/178	24/177				14.6%	1.7[1.07,2.69]
Subtotal (95% CI)	3244	3151		•		100%	1.87[1.49,2.36]
	F:	avours Misoprostol	0.01 0.1	1 10	100 F;	avours Ergometrine	e nlus axvtacin





Analysis 73.8. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

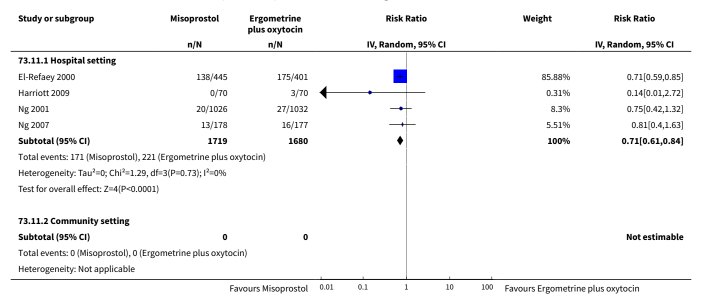
Study or subgroup	Mis	oprostol	_	ometrine oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
73.8.1 Hospital setting							
Bamigboye 1998b	231	187 (92)	233	183 (68)	-+-	18.03%	4[-10.73,18.73]
Caliskan 2003	388	328 (152)	398	296 (168)		15.65%	32[9.61,54.39]
Cook 1999	424	279 (300.6)	310	255.1 (338.8)	-	8.66%	23.86[-23.48,71.2]
El-Refaey 2000	501	256 (137)	499	251 (136.8)		17.37%	5[-11.97,21.97]
Fararjeh 2003	49	588 (360)	48	387.1 (273.4)		1.89%	200.87[73.82,327.92]
Harriott 2009	70	180.1 (120)	70	197 (177)		8.1%	-16.9[-67,33.2]
Ng 2001	1026	296 (160)	1032	254 (157)		18.32%	42[28.3,55.7]
Ng 2007	178	289 (178)	177	255 (149)	+	11.96%	34[-0.14,68.14]
Subtotal ***	2867		2767		•	100%	22.86[4.5,41.22]
Heterogeneity: Tau ² =430.06; Ch	i ² =28.94, df=7	'(P=0); I ² =75.81%	6				
Test for overall effect: Z=2.44(P=	=0.01)						
73.8.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applie	cable						
Total ***	2867		2767		•	100%	22.86[4.5,41.22]
Heterogeneity: Tau ² =430.06; Ch	i ² =28.94, df=7	'(P=0); I ² =75.81%	6				
Test for overall effect: Z=2.44(P=	=0.01)						
Test for subgroup differences: N	lot applicable	!					
			Favou	rs Misoprostol -100	-50 0 50	100 Favours Erg	ometrine plus oxytocin



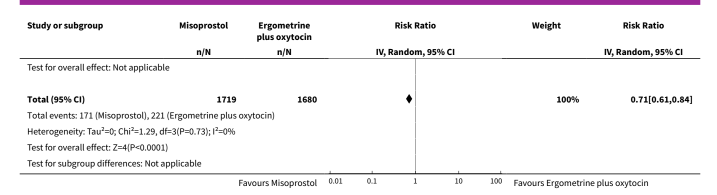
Analysis 73.9. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Mis	oprostol	·	ometrine oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
73.9.1 Hospital setting							
Bamigboye 1998b	83	2.3 (16)	83	2.8 (19)	+	5.57%	-0.5[-5.84,4.84]
Caliskan 2002	396	15 (12)	402	15 (12)	†	13.07%	0[-1.67,1.67]
Caliskan 2003	388	14 (11)	398	15 (12)	+	13.19%	-1[-2.61,0.61]
Cook 1999	424	6.9 (17.5)	310	0.5 (14.1)	+	11.58%	6.4[4.11,8.69]
El-Refaey 2000	218	10 (12)	236	10 (16)	+	10.84%	0[-2.59,2.59]
Fararjeh 2003	49	17.5 (7.3)	48	12.8 (7.3)	+	10.08%	4.7[1.79,7.61]
Harriott 2009	70	5 (7.7)	70	5.6 (7.2)	+	11.13%	-0.6[-3.07,1.87]
Ng 2001	1026	13.4 (12.7)	1032	13.4 (12.7)	<u> </u>	14.23%	0[-1.1,1.1]
Ng 2007	178	17 (14)	177	16 (13)	+	10.31%	1[-1.81,3.81]
Subtotal ***	2832		2756		•	100%	1.09[-0.49,2.67]
Heterogeneity: Tau ² =4.26; Chi ² =	39.15, df=8(P	<0.0001); I ² =79.5	57%				
Test for overall effect: Z=1.35(P=	=0.18)						
73.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not appli	cable						
Total ***	2832		2756		•	100%	1.09[-0.49,2.67]
Heterogeneity: Tau ² =4.26; Chi ² =	39.15, df=8(P	<0.0001); I ² =79.5	57%				
Test for overall effect: Z=1.35(P=	=0.18)						
Test for subgroup differences: N	lot applicable						

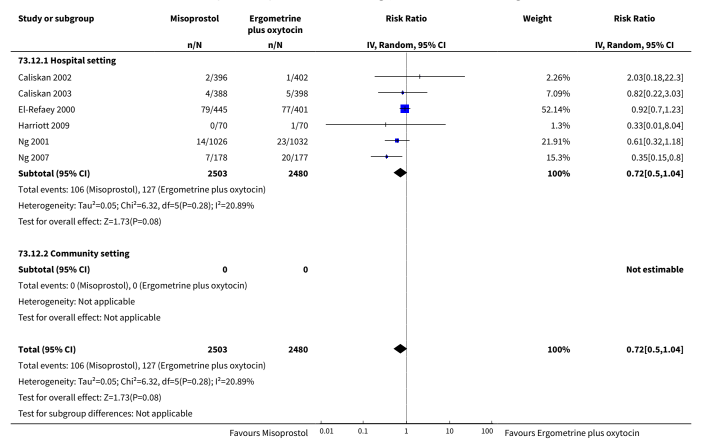
Analysis 73.11. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 11 Nausea.





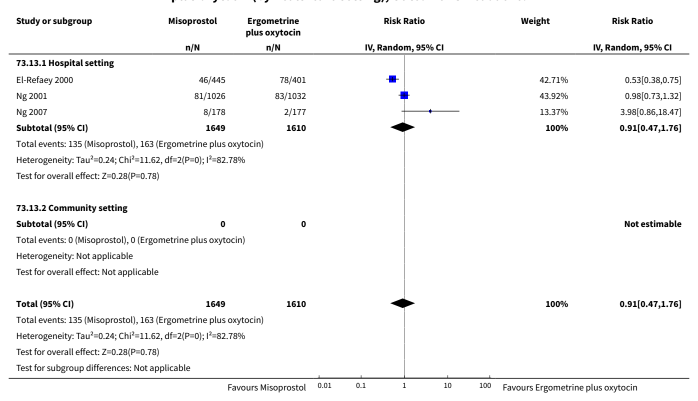


Analysis 73.12. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 12 Vomiting.





Analysis 73.13. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 13 Headache.

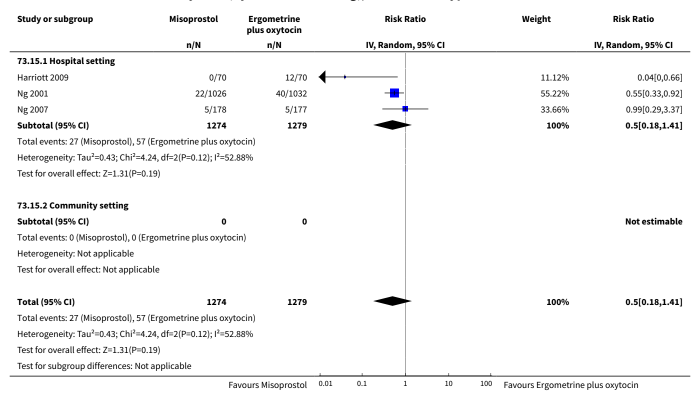


Analysis 73.14. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 14 Abdominal pain.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
73.14.1 Hospital setting					
El-Refaey 2000	217/445	218/401	+	100%	0.9[0.79,1.02]
Subtotal (95% CI)	445	401	•	100%	0.9[0.79,1.02]
Total events: 217 (Misoprostol), 218	(Ergometrine plus ox	sytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=1.63(P=0.1)					
73.14.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol), 0 (Ergo	ometrine plus oxytoc	in)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	445	401	•	100%	0.9[0.79,1.02]
Total events: 217 (Misoprostol), 218	(Ergometrine plus ox	ytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=1.63(P=0.1)					
Test for subgroup differences: Not a	pplicable				
	Fa	vours Misoprostol 0.0	0.1 1 10	100 Favours Ergometrin	e plus oxytocin



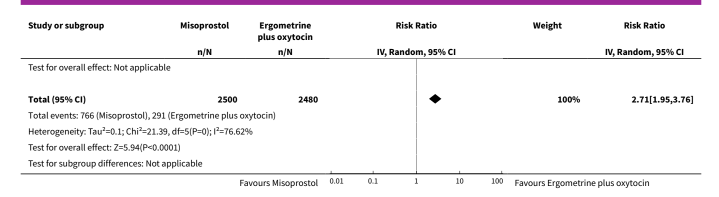
Analysis 73.15. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 15 Hypertension.



Analysis 73.16. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Misoprostol	Ergometrine plus oxytocin		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Ran	dom, 95% CI			IV, Random, 95% CI
73.16.1 Hospital setting								
Caliskan 2002	47/396	19/402			-		17.06%	2.51[1.5,4.2]
Caliskan 2003	44/388	15/398					15.6%	3.01[1.7,5.32]
El-Refaey 2000	319/445	147/401					27.81%	1.96[1.7,2.25]
Harriott 2009	11/67	6/70			+-		8.67%	1.92[0.75,4.89]
Ng 2001	310/1026	102/1032			-		26.28%	3.06[2.49,3.76]
Ng 2007	35/178	2/177					4.58%	17.4[4.25,71.25]
Subtotal (95% CI)	2500	2480			•		100%	2.71[1.95,3.76]
Total events: 766 (Misoprostol	l), 291 (Ergometrine plus ox	(ytocin)						
Heterogeneity: Tau ² =0.1; Chi ² =	=21.39, df=5(P=0); I ² =76.62 ^o	%						
Test for overall effect: Z=5.94(I	P<0.0001)							
73.16.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol), (0 (Ergometrine plus oxytoc	in)						
Heterogeneity: Not applicable	•			1				
	Fa	vours Misoprostol	0.01	0.1	1 10	100	Favours Ergometrin	e plus oxytocin



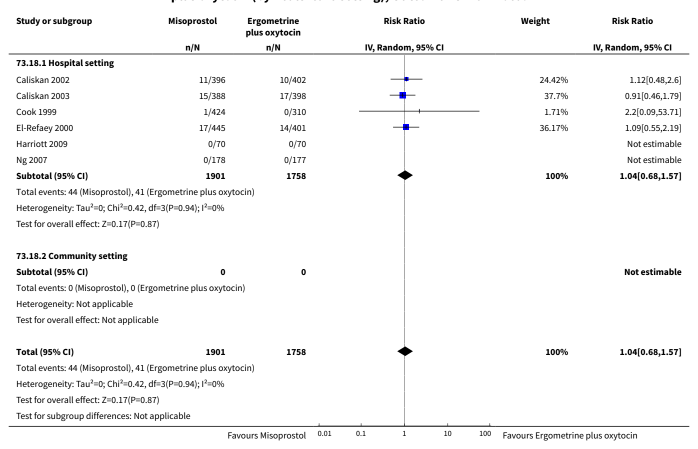


Analysis 73.17. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 17 Fever.

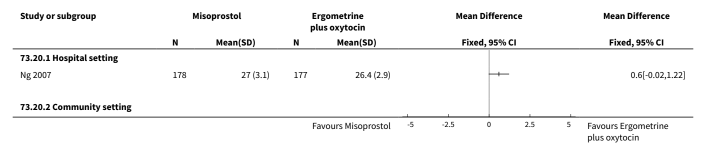
	plus oxytocin			Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
16/396	6/402		23.76%	2.71[1.07,6.85]
17/388	6/398		23.98%	2.91[1.16,7.29]
24/432	1/416		8.06%	23.11[3.14,170.06]
0/70	0/70			Not estimable
10/30	0/30		4.46%	21[1.29,342.93]
87/1026	13/1032	-	35.46%	6.73[3.78,11.98]
7/178	0/177	+	4.28%	14.92[0.86,259.21]
2520	2525	•	100%	5.33[2.87,9.88]
ິວ (Ergometrine plus oxyt	tocin)			
84, df=5(P=0.17); l ² =36.1	19%			
.0001)				
0	0			Not estimable
rgometrine plus oxytoci	n)			
ble				
2520	2525	•	100%	5.33[2.87,9.88]
6 (Ergometrine plus oxy	tocin)			
84, df=5(P=0.17); I ² =36.1	19%			
.0001)				
t applicable				
	16/396 17/388 24/432 0/70 10/30 87/1026 7/178 2520 6 (Ergometrine plus oxyt 84, df=5(P=0.17); l²=36.1 0001) 0 rgometrine plus oxyt 84, df=5(P=0.17); l²=36.1	n/N	n/N	n/N



Analysis 73.18. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 18 Diarrhoea.



Analysis 73.20. Comparison 73 Misoprostol vs Ergometrine plus oxytocin (by healthcare setting), Outcome 20 Maternal satisfaction.



Comparison 74. Misoprostol vs Misoprostol plus oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.85 [1.03, 3.33]
2.1 Hospital setting	2	1589	Risk Ratio (IV, Random, 95% CI)	1.85 [1.03, 3.33]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1589	Risk Ratio (IV, Random, 95% CI)	2.97 [1.40, 6.30]
3.1 Hospital setting	2	1589	Risk Ratio (IV, Random, 95% CI)	2.97 [1.40, 6.30]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1589	Risk Ratio (IV, Random, 95% CI)	1.93 [0.99, 3.76]
6.1 Hospital setting	2	1589	Risk Ratio (IV, Random, 95% CI)	1.93 [0.99, 3.76]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	3	1689	Risk Ratio (IV, Random, 95% CI)	1.89 [1.26, 2.83]
7.1 Hospital setting	3	1689	Risk Ratio (IV, Random, 95% CI)	1.89 [1.26, 2.83]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	792	Mean Difference (IV, Random, 95% CI)	48.0 [24.68, 71.32]
8.1 Hospital setting	1	792	Mean Difference (IV, Random, 95% CI)	48.0 [24.68, 71.32]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	3	1689	Mean Difference (IV, Random, 95% CI)	0.27 [-0.42, 0.96]
9.1 Hospital setting	3	1689	Mean Difference (IV, Random, 95% CI)	0.27 [-0.42, 0.96]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

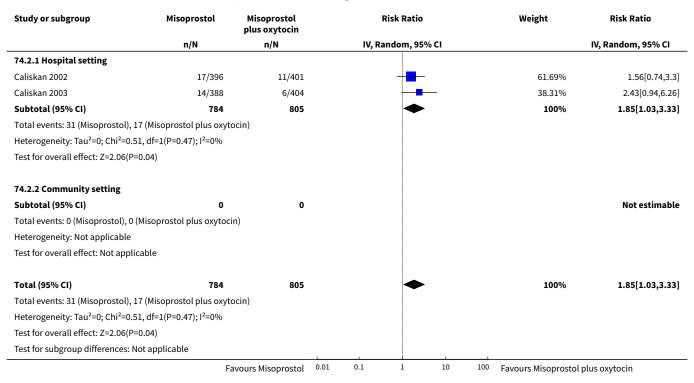


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	100	Risk Ratio (IV, Random, 95% CI)	1.83 [0.73, 4.57]
11.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	1.83 [0.73, 4.57]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	3	1689	Risk Ratio (IV, Random, 95% CI)	1.39 [0.51, 3.82]
12.1 Hospital setting	3	1689	Risk Ratio (IV, Random, 95% CI)	1.39 [0.51, 3.82]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	1689	Risk Ratio (IV, Random, 95% CI)	0.94 [0.72, 1.22]
16.1 Hospital setting	3	1689	Risk Ratio (IV, Random, 95% CI)	0.94 [0.72, 1.22]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	3	1689	Risk Ratio (IV, Random, 95% CI)	0.97 [0.62, 1.53]
17.1 Hospital setting	3	1689	Risk Ratio (IV, Random, 95% CI)	0.97 [0.62, 1.53]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	1589	Risk Ratio (IV, Random, 95% CI)	1.22 [0.70, 2.13]
18.1 Hospital setting	2	1589	Risk Ratio (IV, Random, 95% CI)	1.22 [0.70, 2.13]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

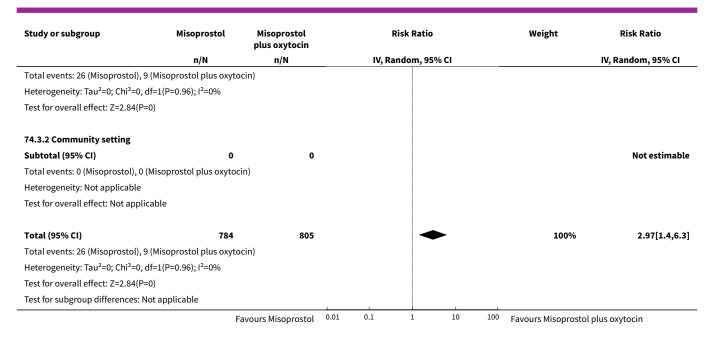
Analysis 74.2. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.



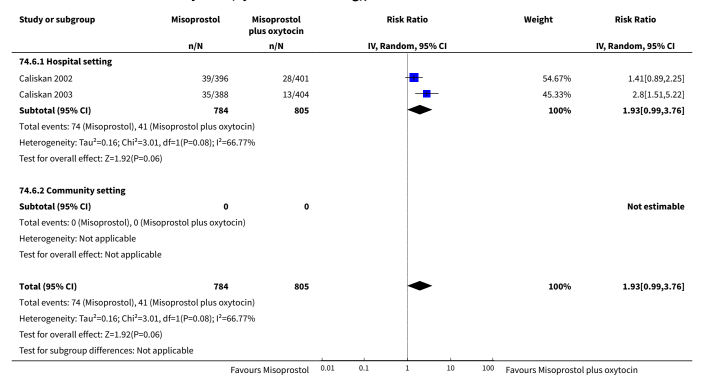
Analysis 74.3. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 3 Blood transfusion.

Study or subgroup	Misoprostol	Misoprostol plus oxytocin			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
74.3.1 Hospital setting									
Caliskan 2002	12/396	4/401			-			44.79%	3.04[0.99,9.34]
Caliskan 2003	14/388	5/404			-			55.21%	2.92[1.06,8.02]
Subtotal (95% CI)	784	805				▶ .		100%	2.97[1.4,6.3]
	Favours Misoprostol		0.01	0.1	1	10	100	Favours Misoprostol plus oxytocin	



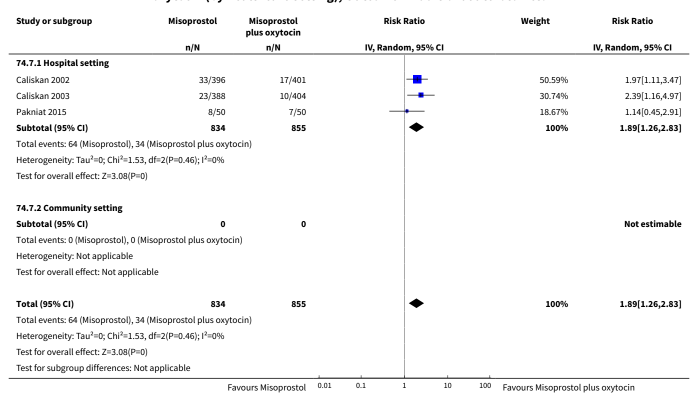


Analysis 74.6. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.





Analysis 74.7. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.



Analysis 74.8. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

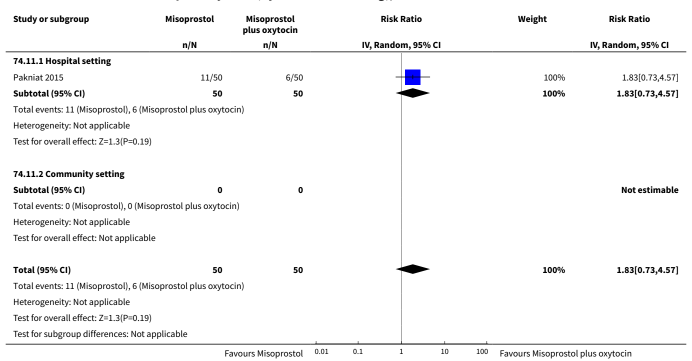
Study or subgroup	Mis	oprostol	Misoprostol plus oxytocin			Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
74.8.1 Hospital setting										
Caliskan 2003	388	328 (152)	404	280 (182)			_	_	100%	48[24.68,71.32]
Subtotal ***	388		404				•	-	100%	48[24.68,71.32]
Heterogeneity: Not applicable										
Test for overall effect: Z=4.03(P<0.0	0001)									
74.8.2 Community setting										
Subtotal ***	0		0							Not estimable
Heterogeneity: Not applicable										
Test for overall effect: Not applicab	ole									
Total ***	388		404					-	100%	48[24.68,71.32]
Heterogeneity: Not applicable										
Test for overall effect: Z=4.03(P<0.0	0001)									
Test for subgroup differences: Not	applicable									
			Favour	s Misoprostol	-100	-50	0 5	0 100	Favours Mis	oprostol plus oxytocin



Analysis 74.9. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

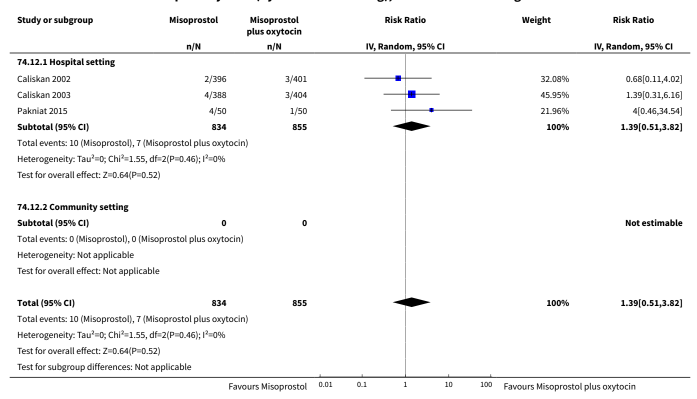
Study or subgroup	Misoprostol		Misoprostol plus oxytocin		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
74.9.1 Hospital setting							
Caliskan 2002	396	15 (12)	401	15 (13)	+	15.59%	0[-1.74,1.74]
Caliskan 2003	388	14 (11)	404	14 (13)	+	16.77%	0[-1.67,1.67]
Pakniat 2015	50	11.4 (2.9)	50	11 (0.8)		67.64%	0.4[-0.43,1.23]
Subtotal ***	834		855		•	100%	0.27[-0.42,0.96]
Heterogeneity: Tau ² =0; Chi ² =0.29, c	df=2(P=0.8	7); I ² =0%					
Test for overall effect: Z=0.77(P=0.4	4)						
74.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicab	le						
Total ***	834		855			100%	0.27[-0.42,0.96]
Heterogeneity: Tau ² =0; Chi ² =0.29, c	df=2(P=0.8	7); I ² =0%					
Test for overall effect: Z=0.77(P=0.4	4)						
Test for subgroup differences: Not a	applicable						
			Favou	rs Misoprostol -100	-50 0 50	100 Favours Mis	oprostol plus oxytocin

Analysis 74.11. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 11 Nausea.

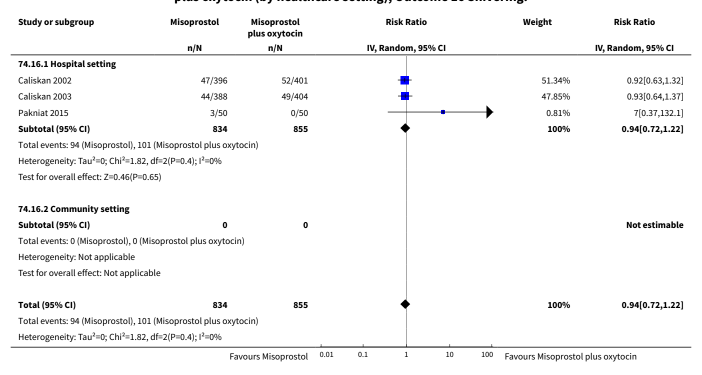




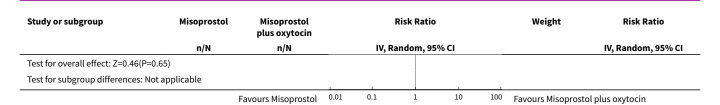
Analysis 74.12. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 12 Vomiting.



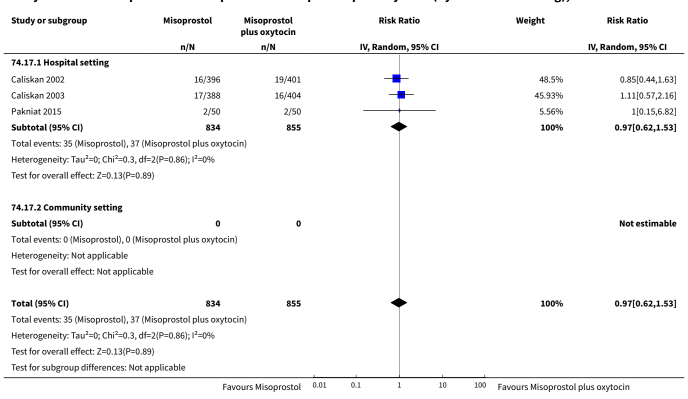
Analysis 74.16. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 16 Shivering.







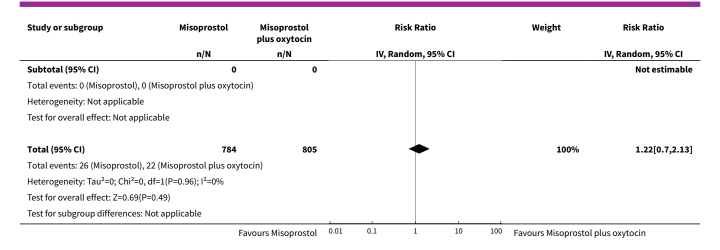
Analysis 74.17. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 17 Fever.



Analysis 74.18. Comparison 74 Misoprostol vs Misoprostol plus oxytocin (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	roup Misoprostol Misoprostol Risk Ratio plus oxytocin			Weight	Risk Ratio			
	n/N	n/N		IV, F	Random, 95% C	I		IV, Random, 95% CI
74.18.1 Hospital setting								
Caliskan 2002	11/396	9/401			-		41.29%	1.24[0.52,2.95]
Caliskan 2003	15/388	13/404			-		58.71%	1.2[0.58,2.49]
Subtotal (95% CI)	784	805			•		100%	1.22[0.7,2.13]
Total events: 26 (Misoprostol)	, 22 (Misoprostol plus oxyto	ocin)						
Heterogeneity: Tau ² =0; Chi ² =0	, df=1(P=0.96); I ² =0%							
Test for overall effect: Z=0.69(P=0.49)							
74.18.2 Community setting						i		
	Fa	vours Misoprostol	0.01	0.1	1	10 10	⁰⁰ Favours Misoprosto	l plus oxytocin





Comparison 75. Carbetocin vs Injectable prostaglandins (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 76. Injectable prostaglandins vs Ergometrine (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	100	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.1 Hospital setting	1	50	Risk Ratio (IV, Random, 95% CI)	5.0 [0.25, 99.16]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
3 Blood transfusion	4	384	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.58]
3.1 Hospital setting	4	384	Risk Ratio (IV, Random, 95% CI)	1.01 [0.04, 23.58]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	3	399	Risk Ratio (IV, Random, 95% CI)	1.42 [0.64, 3.17]
6.1 Hospital setting	3	399	Risk Ratio (IV, Random, 95% CI)	1.42 [0.64, 3.17]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	5	684	Risk Ratio (IV, Random, 95% CI)	0.78 [0.40, 1.51]
7.1 Hospital setting	5	684	Risk Ratio (IV, Random, 95% CI)	0.78 [0.40, 1.51]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	8	1195	Mean Difference (IV, Random, 95% CI)	-16.08 [-57.17, 25.00]
8.1 Hospital setting	8	1195	Mean Difference (IV, Random, 95% CI)	-16.08 [-57.17, 25.00]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	300	Mean Difference (IV, Random, 95% CI)	-1.28 [-6.58, 4.01]
9.1 Hospital setting	2	300	Mean Difference (IV, Random, 95% CI)	-1.28 [-6.58, 4.01]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	5	684	Risk Ratio (IV, Random, 95% CI)	1.70 [0.98, 2.94]
11.1 Hospital setting	5	684	Risk Ratio (IV, Random, 95% CI)	1.70 [0.98, 2.94]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	5	684	Risk Ratio (IV, Random, 95% CI)	2.70 [0.97, 7.55]
12.1 Hospital setting	5	684	Risk Ratio (IV, Random, 95% CI)	2.70 [0.97, 7.55]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	80	Risk Ratio (IV, Random, 95% CI)	2.0 [0.39, 10.31]
13.1 Hospital setting	1	80	Risk Ratio (IV, Random, 95% CI)	2.0 [0.39, 10.31]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	3	384	Risk Ratio (IV, Random, 95% CI)	1.70 [0.07, 40.44]
14.1 Hospital setting	3	384	Risk Ratio (IV, Random, 95% CI)	1.70 [0.07, 40.44]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	2	315	Risk Ratio (IV, Random, 95% CI)	0.16 [0.02, 1.37]
15.1 Hospital setting	2	315	Risk Ratio (IV, Random, 95% CI)	0.16 [0.02, 1.37]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	3	434	Risk Ratio (IV, Random, 95% CI)	0.38 [0.16, 0.91]
16.1 Hospital setting	3	434	Risk Ratio (IV, Random, 95% CI)	0.38 [0.16, 0.91]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	4	534	Risk Ratio (IV, Random, 95% CI)	4.52 [0.76, 26.80]
17.1 Hospital setting	4	534	Risk Ratio (IV, Random, 95% CI)	4.52 [0.76, 26.80]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	5	664	Risk Ratio (IV, Random, 95% CI)	10.09 [2.75, 37.00]
18.1 Hospital setting	5	664	Risk Ratio (IV, Random, 95% CI)	10.09 [2.75, 37.00]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

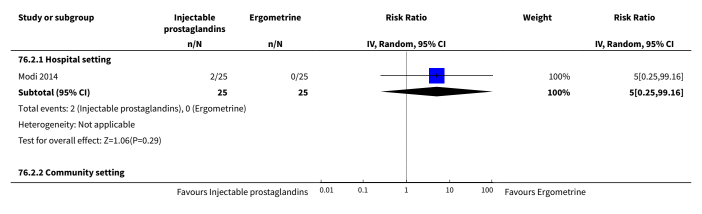


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

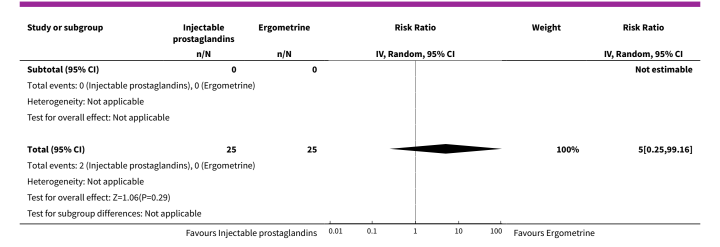
Analysis 76.1. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 1 Death.

Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
0/50	0/50			Not estimable
50	50			Not estimable
ndins), 0 (Ergometrin	e)			
0	0			Not estimable
ndins), 0 (Ergometrin	e)			
50	50			Not estimable
ndins), 0 (Ergometrin	e)			
2		İ		
plicable		İ		
	n/N 0/50 50 ndins), 0 (Ergometrin e 0 ndins), 0 (Ergometrin	n/N	n/N	n/N n/N IV, Random, 95% CI 0/50 0/50 50 50 ndins), 0 (Ergometrine) e 50 50 ndins), 0 (Ergometrine) e

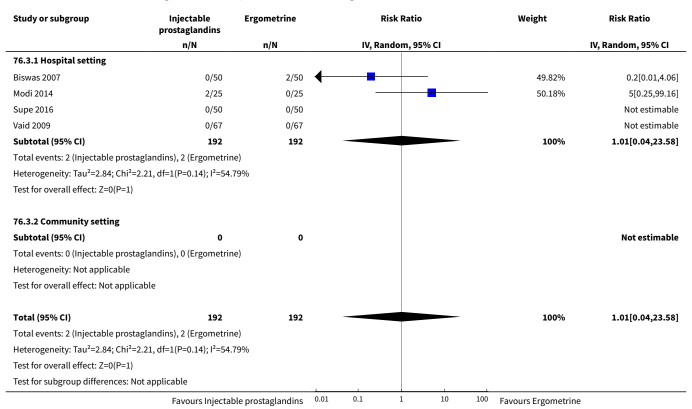
Analysis 76.2. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 2 PPH >= 1000 mL.





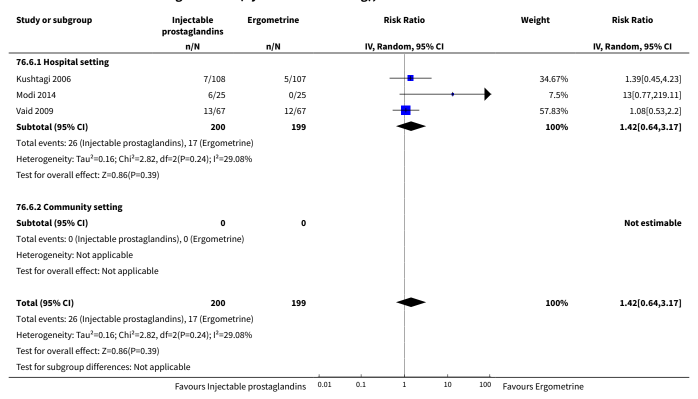


Analysis 76.3. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 3 Blood transfusion.





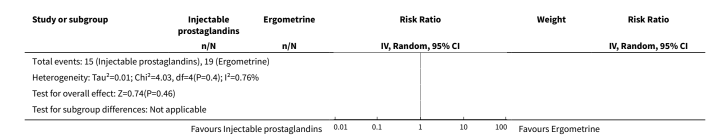
Analysis 76.6. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 6 PPH >= 500 mL.



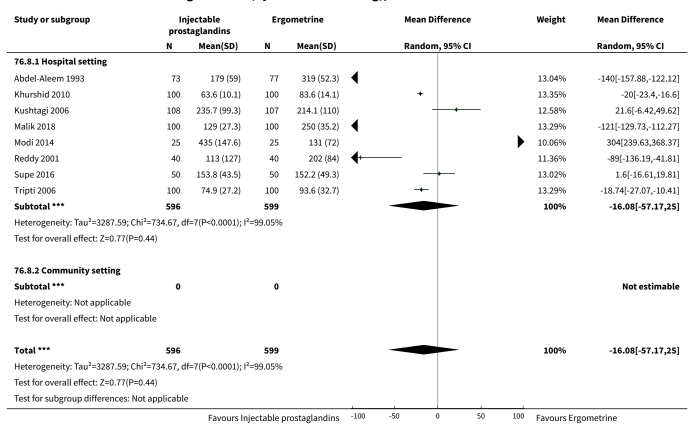
Analysis 76.7. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
76.7.1 Hospital setting					
Khurshid 2010	0/100	1/100		4.25%	0.33[0.01,8.09]
Modi 2014	2/25	0/25		4.84%	5[0.25,99.16]
Supe 2016	4/50	2/50	- •	15.71%	2[0.38,10.43]
Tripti 2006	0/100	2/100		4.72%	0.2[0.01,4.11]
Vaid 2009	9/67	14/67	-	70.48%	0.64[0.3,1.38]
Subtotal (95% CI)	342	342	•	100%	0.78[0.4,1.51]
Total events: 15 (Injectable prosta	glandins), 19 (Ergomet	rine)			
Heterogeneity: Tau ² =0.01; Chi ² =4.0	03, df=4(P=0.4); I ² =0.76	%			
Test for overall effect: Z=0.74(P=0.4	46)				
76.7.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostag	landins), 0 (Ergometrin	e)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	ole				
Total (95% CI)	342	342	•	100%	0.78[0.4,1.51]
	Favours Injectal	ole prostaglandins	0.01 0.1 1 10	100 Favours Ergometrine	2





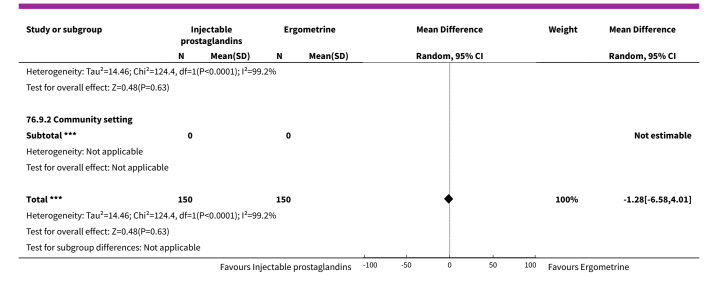
Analysis 76.8. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 8 Blood loss.



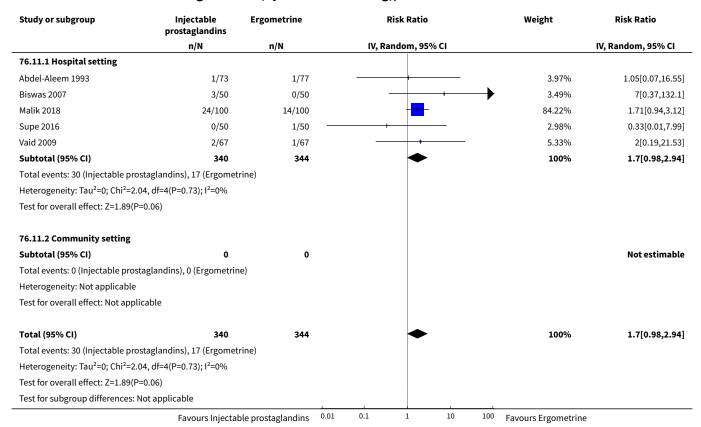
Analysis 76.9. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup		jectable taglandins	Erg	ometrine		M	lean Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		R	andom, 95%	6 CI			Random, 95% CI
76.9.1 Hospital setting											
Malik 2018	100	4.7 (3.2)	100	8.7 (3.2)						49.7%	-4[-4.89,-3.11]
Supe 2016	50	9 (1.2)	50	7.6 (0.2)			•			50.3%	1.4[1.06,1.74]
Subtotal ***	150		150				•			100%	-1.28[-6.58,4.01]
		Favours Inj	ectable p	rostaglandins	-100	-50	0	50	100	Favours Erg	ometrine



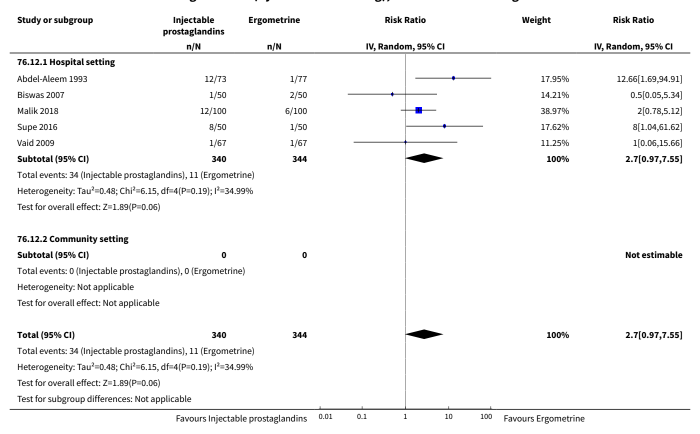


Analysis 76.11. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 11 Nausea.





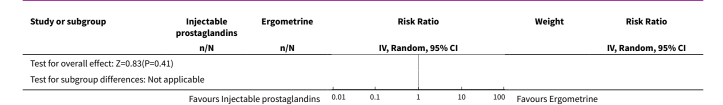
Analysis 76.12. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 12 Vomiting.



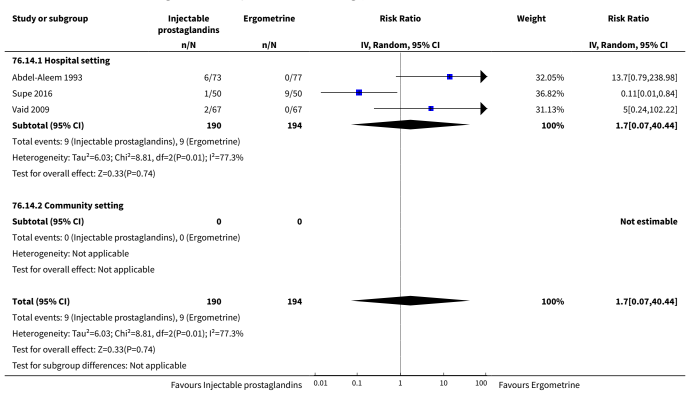
Analysis 76.13. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 13 Headache.

Study or subgroup	Injectable prostaglandins	Ergometrine		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	IV, Ra	andom, 95% CI			IV, Random, 95% CI
76.13.1 Hospital setting							
Reddy 2001	4/40	2/40				100%	2[0.39,10.31]
Subtotal (95% CI)	40	40				100%	2[0.39,10.31]
Total events: 4 (Injectable prostag	landins), 2 (Ergometrin	e)					
Heterogeneity: Not applicable							
Test for overall effect: Z=0.83(P=0.4	41)						
76.13.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Injectable prostag	landins), 0 (Ergometrin	e)					
Heterogeneity: Not applicable							
Test for overall effect: Not applical	ole						
Total (95% CI)	40	40				100%	2[0.39,10.31]
Total events: 4 (Injectable prostag	landins), 2 (Ergometrin	e)					
Heterogeneity: Not applicable							
	Favours Injectat	ole prostaglandins	0.01 0.1	1 10	100	Favours Ergometrine	

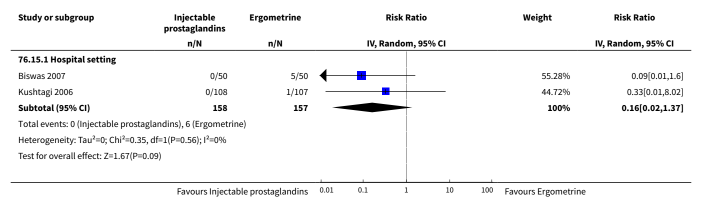




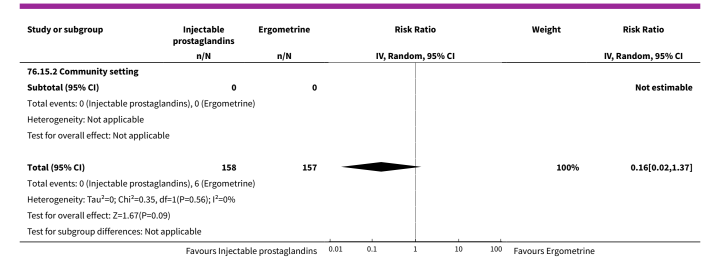
Analysis 76.14. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 14 Abdominal pain.



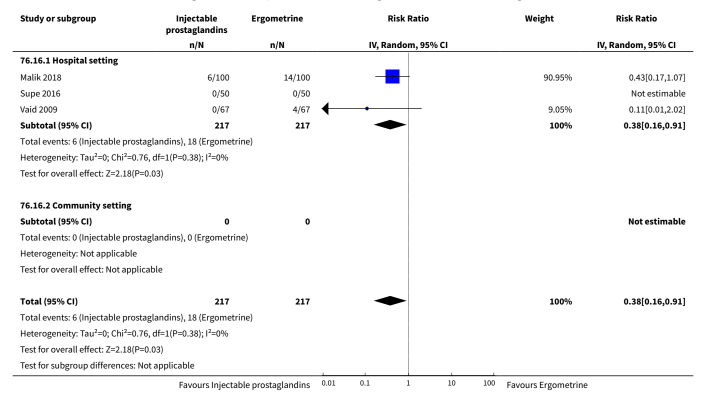
Analysis 76.15. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 15 Hypertension.







Analysis 76.16. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 16 Shivering.





Analysis 76.17. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
76.17.1 Hospital setting					
Biswas 2007	1/50	0/50		31.36%	3[0.13,71.92]
Malik 2018	4/100	0/100		37.41%	9[0.49,165]
Supe 2016	0/50	0/50			Not estimable
Vaid 2009	1/67	0/67		31.24%	3[0.12,72.35]
Subtotal (95% CI)	267	267		100%	4.52[0.76,26.8]
Total events: 6 (Injectable prostagle	andins), 0 (Ergometrine)			
Heterogeneity: Tau ² =0; Chi ² =0.34, c	df=2(P=0.84); I ² =0%				
Test for overall effect: Z=1.66(P=0.1	.)				
76.17.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagl	andins), 0 (Ergometrine)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	267	267		100%	4.52[0.76,26.8]
Total events: 6 (Injectable prostagl	andins), 0 (Ergometrine)			
Heterogeneity: Tau ² =0; Chi ² =0.34, c	df=2(P=0.84); I ² =0%				
Test for overall effect: Z=1.66(P=0.1	.)				
Test for subgroup differences: Not a	applicable				
	Favours Injectable	e prostaglandins 0.03	0.1 1 10 10	DO Favours Ergometrine	2

Analysis 76.18. Comparison 76 Injectable prostaglandins vs Ergometrine (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
76.18.1 Hospital setting					
Abdel-Aleem 1993	2/73	0/77		18.53%	5.27[0.26,107.96]
Malik 2018	2/100	0/100		18.48%	5[0.24,102.85]
Reddy 2001	7/40	0/40	 	21.1%	15[0.89,254.13]
Supe 2016	7/50	0/50	 	21%	15[0.88,255.78]
Vaid 2009	7/67	0/67	 	20.9%	15[0.87,257.48]
Subtotal (95% CI)	330	334		100%	10.09[2.75,37]
Total events: 25 (Injectable prostag	glandins), 0 (Ergometri	ine)			
Heterogeneity: Tau ² =0; Chi ² =0.61, c	df=4(P=0.96); I ² =0%				
Test for overall effect: Z=3.49(P=0)					
76.18.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Ergometrin	ie)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicab	le				
Total (95% CI)	330	334		100%	10.09[2.75,37]
	Favours Injectal	ole prostaglandins	0.01 0.1 1 10 100	Favours Ergometrine	e



Study or subgroup	Injectable prostaglandins	Ergometrine			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		IV, F	Random, 95	% CI			IV, Random, 95% CI
Total events: 25 (Injectable p	rostaglandins), 0 (Ergometr	ine)							
Heterogeneity: Tau ² =0; Chi ² =	:0.61, df=4(P=0.96); I ² =0%								
Test for overall effect: Z=3.49	(P=0)								
Test for subgroup difference	s: Not applicable								
	Favours Injecta	ble prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	

Comparison 77. Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	598	Risk Ratio (IV, Random, 95% CI)	0.80 [0.17, 3.78]
2.1 Hospital setting	2	598	Risk Ratio (IV, Random, 95% CI)	0.80 [0.17, 3.78]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	69	Risk Ratio (IV, Random, 95% CI)	0.65 [0.17, 2.53]
3.1 Hospital setting	1	69	Risk Ratio (IV, Random, 95% CI)	0.65 [0.17, 2.53]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	598	Risk Ratio (IV, Random, 95% CI)	1.29 [0.92, 1.79]
6.1 Hospital setting	2	598	Risk Ratio (IV, Random, 95% CI)	1.29 [0.92, 1.79]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	1	112	Risk Ratio (IV, Random, 95% CI)	1.07 [0.16, 7.36]

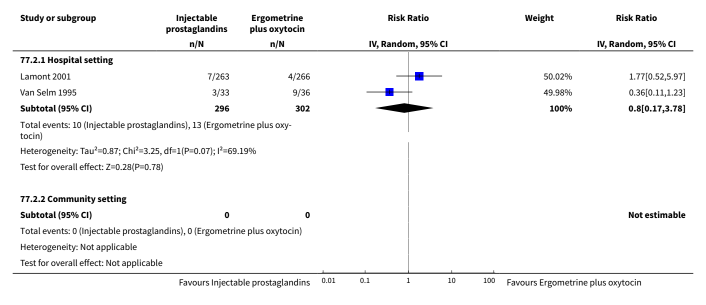


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7.1 Hospital setting	1	112	Risk Ratio (IV, Random, 95% CI)	1.07 [0.16, 7.36]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	2	598	Mean Difference (IV, Random, 95% CI)	-26.18 [-104.63, 52.27]
8.1 Hospital setting	2	598	Mean Difference (IV, Random, 95% CI)	-26.18 [-104.63, 52.27]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	1	529	Risk Ratio (IV, Random, 95% CI)	5.06 [1.12, 22.86]
11.1 Hospital setting	1	529	Risk Ratio (IV, Random, 95% CI)	5.06 [1.12, 22.86]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	1	529	Risk Ratio (IV, Random, 95% CI)	0.92 [0.40, 2.13]
12.1 Hospital setting	1	529	Risk Ratio (IV, Random, 95% CI)	0.92 [0.40, 2.13]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

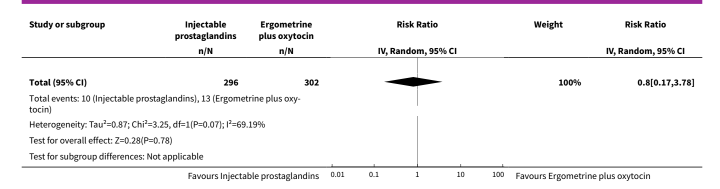


Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	641	Risk Ratio (IV, Random, 95% CI)	23.70 [7.55, 74.45]
18.1 Hospital setting	2	641	Risk Ratio (IV, Random, 95% CI)	23.70 [7.55, 74.45]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

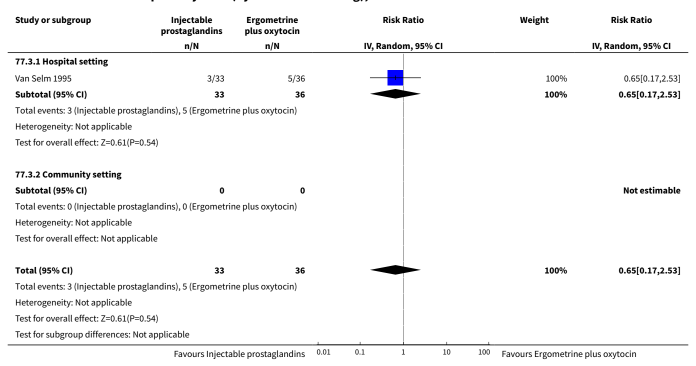
Analysis 77.2. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.







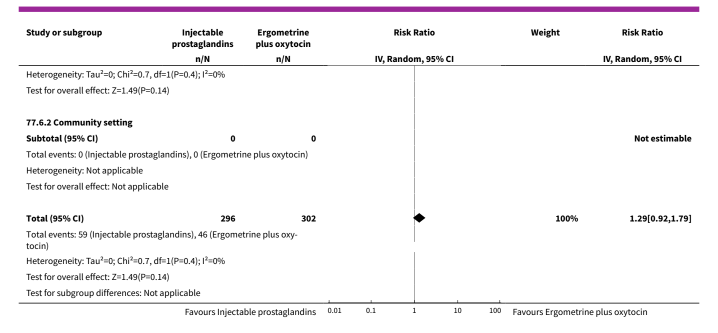
Analysis 77.3. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 3 Blood transfusion.



Analysis 77.6. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		IV, F	andom, 95% (CI			IV, Random, 95% CI
77.6.1 Hospital setting									
Lamont 2001	43/263	30/266			-			57.72%	1.45[0.94,2.24]
Van Selm 1995	16/33	16/36			-			42.28%	1.09[0.66,1.81]
Subtotal (95% CI)	296	302			•			100%	1.29[0.92,1.79]
Total events: 59 (Injectable protocin)	ostaglandins), 46 (Ergomet	rine plus oxy-							
	Favours Injecta	ble prostaglandins	0.01	0.1	1	10	100	Favours Ergometrine	e plus oxytocin



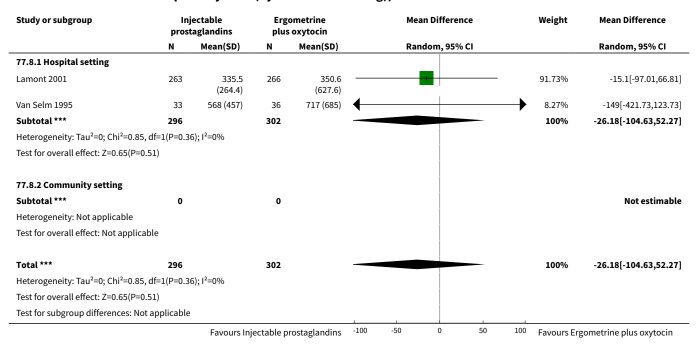


Analysis 77.7. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
77.7.1 Hospital setting					
Chua 1995	2/54	2/58		100%	1.07[0.16,7.36]
Subtotal (95% CI)	54	58		100%	1.07[0.16,7.36]
Total events: 2 (Injectable prostagla	ndins), 2 (Ergometrii	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.07(P=0.94	1)				
77.7.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Injectable prostagla	ndins), 0 (Ergometrii	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicabl	e				
Total (95% CI)	54	58		100%	1.07[0.16,7.36]
Total events: 2 (Injectable prostagla	ndins), 2 (Ergometrii	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.07(P=0.94	1)				
Test for subgroup differences: Not a	pplicable				
	Favours Injecta	ble prostaglandins 0.0	01 0.1 1 10	100 Favours Ergometrin	e plus oxytocin



Analysis 77.8. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

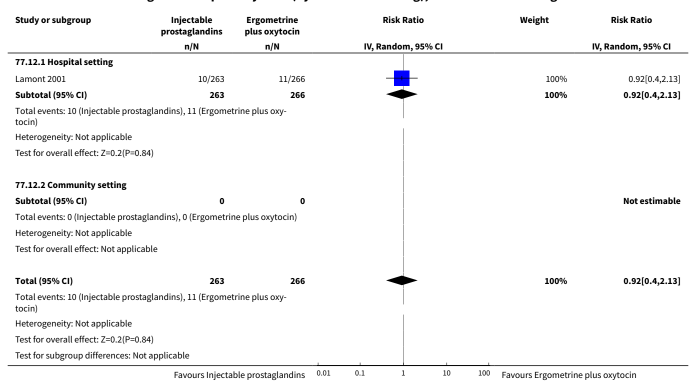


Analysis 77.11. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 11 Nausea.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin		Risk F	Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Randor	n, 95% CI			IV, Random, 95% CI
77.11.1 Hospital setting								
Lamont 2001	10/263	2/266					100%	5.06[1.12,22.86]
Subtotal (95% CI)	263	266					100%	5.06[1.12,22.86]
Total events: 10 (Injectable prostag	glandins), 2 (Ergometr	ine plus oxytocin)						
Heterogeneity: Not applicable								
Test for overall effect: Z=2.11(P=0.0	04)							
77.11.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Injectable prostagl	andins), 0 (Ergometrir	ne plus oxytocin)						
Heterogeneity: Not applicable								
Test for overall effect: Not applicab	le							
Total (95% CI)	263	266			~		100%	5.06[1.12,22.86]
Total events: 10 (Injectable prostag	glandins), 2 (Ergometr	ine plus oxytocin)						
Heterogeneity: Not applicable								
Test for overall effect: Z=2.11(P=0.0	04)							
Test for subgroup differences: Not	applicable							
	Favours Injecta	ble prostaglandins	0.01 0.	.1 1	. 10	¹⁰⁰ Fav	ours Ergometrine	e plus oxytocin



Analysis 77.12. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 12 Vomiting.



Analysis 77.18. Comparison 77 Injectable prostaglandins vs Ergometrine plus oxytocin (by healthcare setting), Outcome 18 Diarrhoea.

Study or subgroup	Injectable prostaglandins	Ergometrine plus oxytocin	Risk	Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rando	m, 95% CI		IV, Random, 95% CI
77.18.1 Hospital setting						
Chua 1995	16/54	1/58			33.21%	17.19[2.36,125.22]
Lamont 2001	55/263	2/266			66.79%	27.81[6.86,112.85]
Subtotal (95% CI)	317	324		-	100%	23.7[7.55,74.45]
Total events: 71 (Injectable prostag	landins), 3 (Ergometri	ne plus oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.15, d	If=1(P=0.7); I ² =0%					
Test for overall effect: Z=5.42(P<0.0	001)					
77.18.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Injectable prostagla	andins), 0 (Ergometrin	e plus oxytocin)				
Heterogeneity: Not applicable						
Test for overall effect: Not applicab	le					
Total (95% CI)	317	324		-	100%	23.7[7.55,74.45]
Total events: 71 (Injectable prostag	landins), 3 (Ergometri	ne plus oxytocin)				
Heterogeneity: Tau ² =0; Chi ² =0.15, d	If=1(P=0.7); I ² =0%					
Test for overall effect: Z=5.42(P<0.0	001)					
Test for subgroup differences: Not a	applicable					
	Favours Injectal	ole prostaglandins	0.01 0.1	1 10 100	Favours Ergometrine	e plus oxytocin



Comparison 78. Misoprostol plus oxytocin vs Injectable prostaglandins (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbid- ity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 79. Ergometrine vs Carbetocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of participants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 80. Carbetocin vs Ergometrine plus oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
2 PPH >= 1000 mL	4	1210	Risk Ratio (IV, Random, 95% CI)	0.34 [0.13, 0.86]
2.1 Hospital setting	4	1210	Risk Ratio (IV, Random, 95% CI)	0.34 [0.13, 0.86]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	4	1030	Risk Ratio (IV, Random, 95% CI)	1.42 [0.42, 4.82]
3.1 Hospital setting	4	1030	Risk Ratio (IV, Random, 95% CI)	1.42 [0.42, 4.82]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	2	320	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	4	1030	Risk Ratio (IV, Random, 95% CI)	0.96 [0.44, 2.09]
6.1 Hospital setting	4	1030	Risk Ratio (IV, Random, 95% CI)	0.96 [0.44, 2.09]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	6	1530	Risk Ratio (IV, Random, 95% CI)	0.54 [0.30, 1.00]
7.1 Hospital setting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.54 [0.30, 1.00]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	5	1330	Mean Difference (IV, Random, 95% CI)	-44.08 [-82.41, -5.75]
8.1 Hospital setting	5	1330	Mean Difference (IV, Random, 95% CI)	-44.08 [-82.41, -5.75]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	5	1160	Mean Difference (IV, Random, 95% CI)	-2.57 [-4.32, -0.82]
9.1 Hospital setting	5	1160	Mean Difference (IV, Random, 95% CI)	-2.57 [-4.32, -0.82]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	6	1530	Risk Ratio (IV, Random, 95% CI)	0.29 [0.19, 0.45]
11.1 Hospital setting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.29 [0.19, 0.45]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.27 [0.16, 0.46]
12.1 Hospital setting	6	1530	Risk Ratio (IV, Random, 95% CI)	0.27 [0.16, 0.46]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	5	1330	Risk Ratio (IV, Random, 95% CI)	1.41 [0.45, 4.38]
13.1 Hospital setting	5	1330	Risk Ratio (IV, Random, 95% CI)	1.41 [0.45, 4.38]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	4	930	Risk Ratio (IV, Random, 95% CI)	0.57 [0.36, 0.91]
14.1 Hospital setting	4	930	Risk Ratio (IV, Random, 95% CI)	0.57 [0.36, 0.91]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	3	660	Risk Ratio (IV, Random, 95% CI)	0.56 [0.22, 1.39]
15.1 Hospital setting	3	660	Risk Ratio (IV, Random, 95% CI)	0.56 [0.22, 1.39]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	5	1290	Risk Ratio (IV, Random, 95% CI)	0.45 [0.26, 0.80]
16.1 Hospital setting	5	1290	Risk Ratio (IV, Random, 95% CI)	0.45 [0.26, 0.80]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

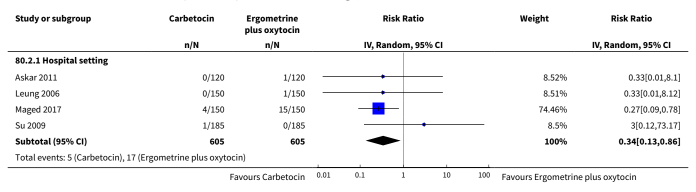


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

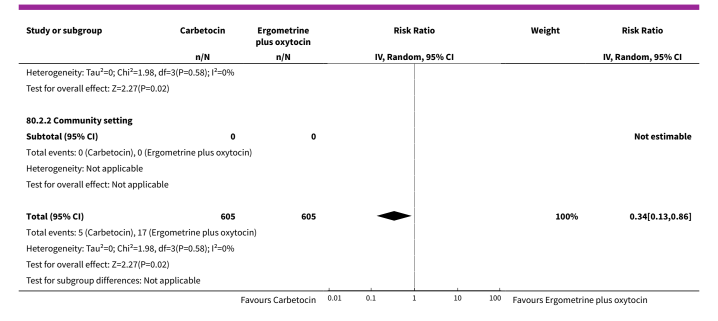
Analysis 80.1. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Risk Ratio	•	Weight	Risk Ratio
	n/N	n/N		IV, Random, 9	5% CI		IV, Random, 95% CI
80.1.1 Hospital setting							
Nirmala 2009	0/60	0/60					Not estimable
Samimi 2013	0/100	0/100					Not estimable
Subtotal (95% CI)	160	160					Not estimable
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	1)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
80.1.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	1)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Total (95% CI)	160	160					Not estimable
Total events: 0 (Carbetocin), 0 (Ergom	etrine plus oxytocir	1)					
Heterogeneity: Not applicable							
Test for overall effect: Not applicable							
Test for subgroup differences: Not app	olicable						
	Fa	vours Carbetocin	0.01	0.1 1	10 100	Favours Ergometr	ine plus oxytocin

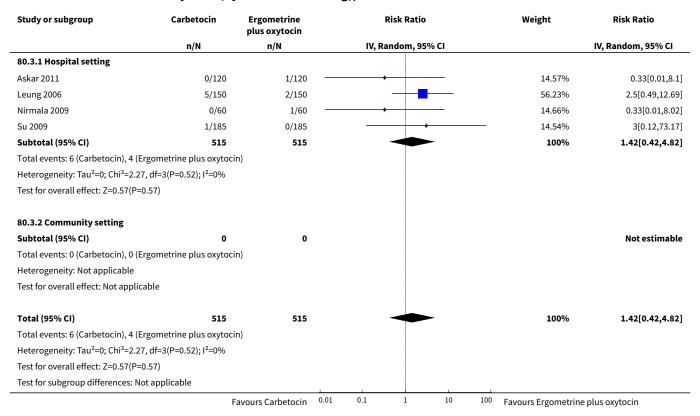
Analysis 80.2. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.







Analysis 80.3. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 3 Blood transfusion.





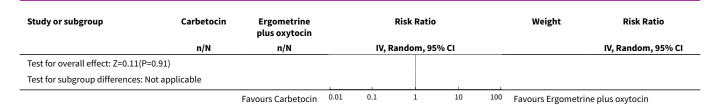
Analysis 80.4. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 4 Severe maternal morbidity: intensive care admissions.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
80.4.1 Hospital setting					
Nirmala 2009	0/60	0/60			Not estimable
Samimi 2013	0/100	0/100			Not estimable
Subtotal (95% CI)	160	160			Not estimable
Total events: 0 (Carbetocin), 0 (Ergom	netrine plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
80.4.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Ergon	netrine plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Total (95% CI)	160	160			Not estimable
Total events: 0 (Carbetocin), 0 (Ergom	netrine plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
Test for subgroup differences: Not ap	plicable				
	Fa	vours Carbetocin 0.	.01 0.1 1 10	100 Favours Ergometrin	ne plus oxytocin

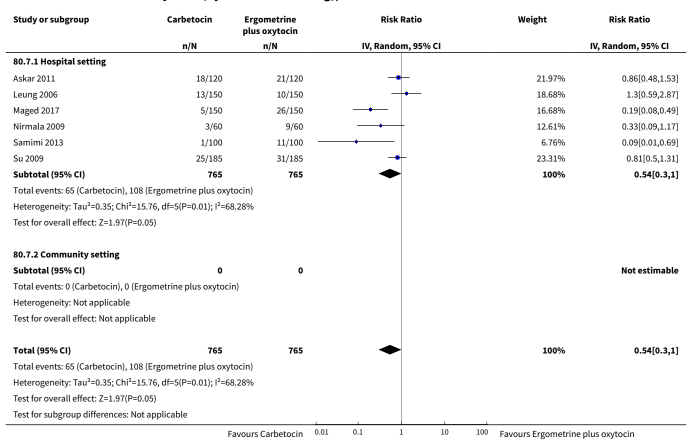
Analysis 80.6. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
80.6.1 Hospital setting					
Askar 2011	2/120	3/120		19.18%	0.67[0.11,3.92]
Leung 2006	6/150	2/150		23.88%	3[0.62,14.63]
Nirmala 2009	3/60	6/60		33.16%	0.5[0.13,1.91]
Su 2009	3/185	3/185		23.79%	1[0.2,4.89]
Subtotal (95% CI)	515	515	*	100%	0.96[0.44,2.09]
Total events: 14 (Carbetocin), 14 (Ergometrine plus oxyto	ocin)			
Heterogeneity: Tau ² =0.01; Chi ² =3.	06, df=3(P=0.38); I ² =2.0	9%			
Test for overall effect: Z=0.11(P=0.	.91)				
80.6.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Carbetocin), 0 (Erg	gometrine plus oxytoci	n)			
Heterogeneity: Not applicable					
Test for overall effect: Not applica	ble				
Total (95% CI)	515	515	•	100%	0.96[0.44,2.09]
Total events: 14 (Carbetocin), 14 (Ergometrine plus oxyto	ocin)			
Heterogeneity: Tau ² =0.01; Chi ² =3.	06, df=3(P=0.38); I ² =2.0	9%			
	F	avours Carbetocin 0.1	01 0.1 1 10 1	Favours Ergometrin	e plus oxytocin





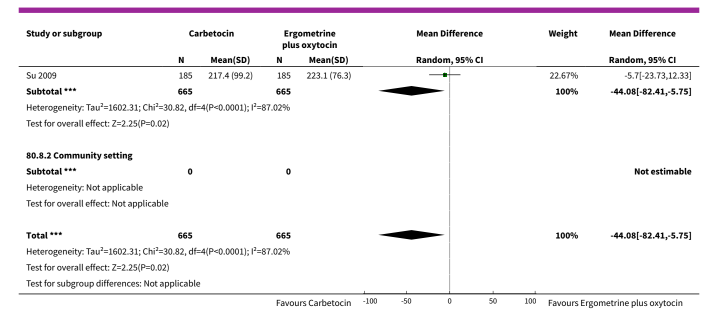
Analysis 80.7. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.



Analysis 80.8. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Car	rbetocin	_	ometrine s oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
80.8.1 Hospital setting							
Askar 2011	120	224.6 (110.6)	120	306.1 (95.7)		21.48%	-81.5[-107.66,-55.34]
Leung 2006	150	232 (122)	150	249 (175)		20.07%	-17[-51.14,17.14]
Maged 2017	150	578 (178)	150	602 (213)		18.07%	-24[-68.42,20.42]
Nirmala 2009	60	244 (114)	60	343 (143)	←	17.71%	-99[-145.27,-52.73]
			Favou	ırs Carbetocin	-100 -50 0 50 10	⁰ Favours Erg	cometrine plus oxytocin



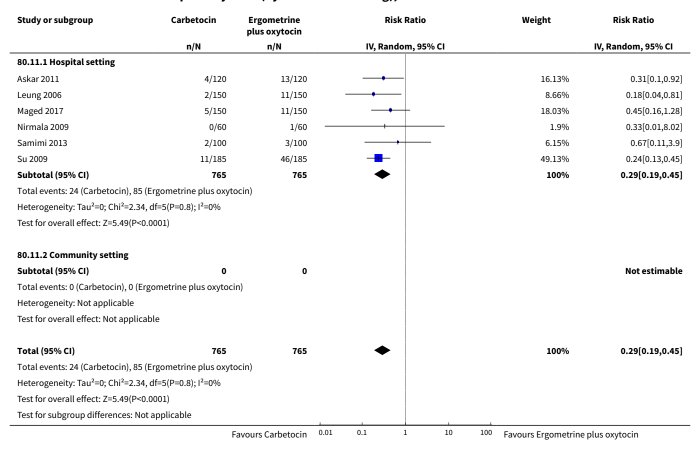


Analysis 80.9. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup	Ca	rbetocin	_	ometrine s oxytocin	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
80.9.1 Hospital setting							
Askar 2011	120	8 (2)	120	11 (3)	-	24.41%	-3[-3.65,-2.35]
Leung 2006	150	14 (11)	150	15 (13)		15.67%	-1[-3.73,1.73]
Maged 2017	150	11 (12)	150	12 (13)	+ -	15.21%	-1[-3.83,1.83]
Nirmala 2009	60	3 (2)	60	4 (2)	-#-	24.23%	-1[-1.72,-0.28]
Samimi 2013	100	4.1 (3.6)	100	10.4 (7.8)		20.47%	-6.3[-7.98,-4.62]
Subtotal ***	580		580		•	100%	-2.57[-4.32,-0.82]
Heterogeneity: Tau ² =3.16; Chi ² =40	.53, df=4(P	<0.0001); I ² =90.1	3%				
Test for overall effect: Z=2.88(P=0)							
80.9.2 Community setting							
Subtotal ***	0		0				Not estimable
Heterogeneity: Not applicable							
Test for overall effect: Not applicable	ole						
Total ***	580		580		•	100%	-2.57[-4.32,-0.82]
Heterogeneity: Tau ² =3.16; Chi ² =40	.53, df=4(P	<0.0001); I ² =90.1	3%				
Test for overall effect: Z=2.88(P=0)							
Test for subgroup differences: Not	applicable	!					
			Favou	ırs Carbetocin -	10 -5 0 5	10 Favours Erg	ometrine plus oxytocin



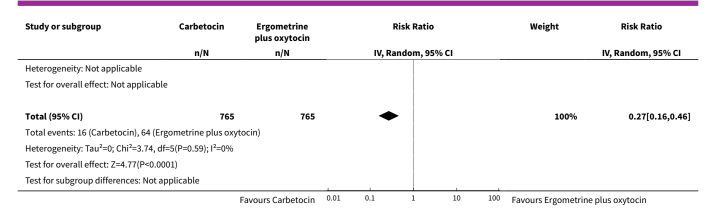
Analysis 80.11. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 11 Nausea.



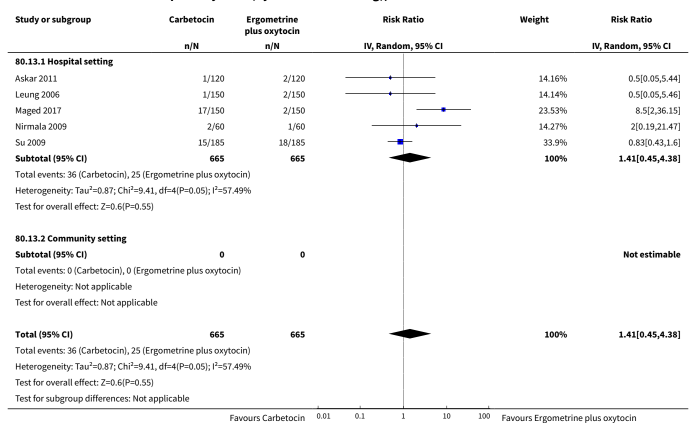
Analysis 80.12. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin	Risk	(Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Rand	om, 95% CI		IV, Random, 95% CI
80.12.1 Hospital setting						
Askar 2011	3/120	12/120		-	18.89%	0.25[0.07,0.86]
Leung 2006	1/150	10/150	+		6.95%	0.1[0.01,0.77]
Maged 2017	4/150	10/150		+	22.44%	0.4[0.13,1.25]
Nirmala 2009	0/60	2/60	+	 	3.19%	0.2[0.01,4.08]
Samimi 2013	1/100	0/100		+	2.85%	3[0.12,72.77]
Su 2009	7/185	30/185	-		45.68%	0.23[0.11,0.52]
Subtotal (95% CI)	765	765	•		100%	0.27[0.16,0.46]
Total events: 16 (Carbetocin), 64	(Ergometrine plus oxyto	cin)				
Heterogeneity: Tau ² =0; Chi ² =3.74	4, df=5(P=0.59); I ² =0%					
Test for overall effect: Z=4.77(P<0	0.0001)					
80.12.2 Community setting						
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Carbetocin), 0 (E	rgometrine plus oxytocir	n)				
	Fa	avours Carbetocin	0.01 0.1	1 10	100 Favours Ergometrine	e plus oxytocin



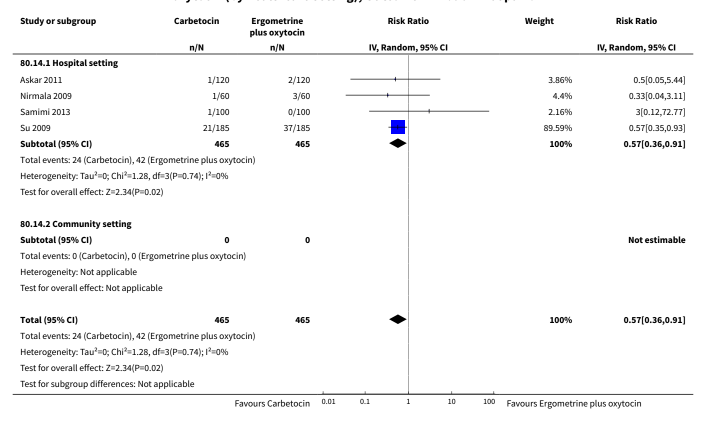


Analysis 80.13. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 13 Headache.





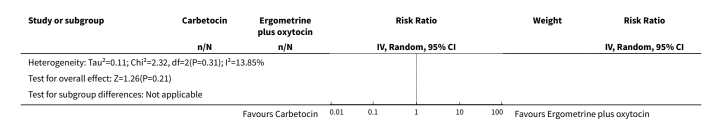
Analysis 80.14. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 14 Abdominal pain.



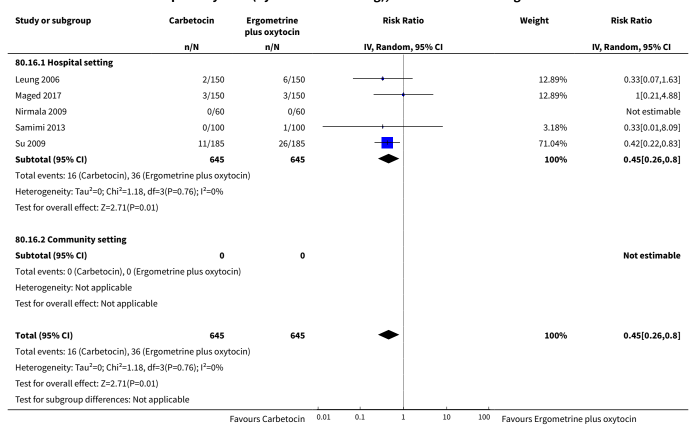
Analysis 80.15. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 15 Hypertension.

Study or subgroup	Carbetocin	Ergometrine plus oxytocin		Risk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Random, 95	% CI		IV, Random, 95% CI
80.15.1 Hospital setting							
Askar 2011	2/120	4/120				25.6%	0.5[0.09,2.68]
Leung 2006	0/150	6/150	\leftarrow	•		9.63%	0.08[0,1.35]
Nirmala 2009	7/60	9/60		-		64.77%	0.78[0.31,1.95]
Subtotal (95% CI)	330	330				100%	0.56[0.22,1.39]
Total events: 9 (Carbetocin), 19 (I	Ergometrine plus oxytoci	n)					
Heterogeneity: Tau ² =0.11; Chi ² =2	2.32, df=2(P=0.31); I ² =13.8	5%					
Test for overall effect: Z=1.26(P=0	0.21)						
80.15.2 Community setting							
Subtotal (95% CI)	0	0					Not estimable
Total events: 0 (Carbetocin), 0 (E	rgometrine plus oxytocin)					
Heterogeneity: Not applicable							
Test for overall effect: Not application	able						
Total (95% CI)	330	330		•		100%	0.56[0.22,1.39]
Total events: 9 (Carbetocin), 19 (I	Ergometrine plus oxytocii	n)					
	Fa	vours Carbetocin	0.01	0.1 1	10 100	Favours Ergometrine	e plus oxytocin





Analysis 80.16. Comparison 80 Carbetocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 16 Shivering.



Comparison 81. Misoprostol plus oxytocin vs Carbetocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	•	
1 Death	1	380	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	1	380	Risk Ratio (IV, Random, 95% CI)	4.00 [0.45, 35.46]
3.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	4.00 [0.45, 35.46]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	1	380	Risk Ratio (IV, Random, 95% CI)	1.19 [0.74, 1.93]
7.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	1.19 [0.74, 1.93]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	1	380	Mean Difference (IV, Random, 95% CI)	-1.70 [-3.72, 0.32]
9.1 Hospital setting	1	380	Mean Difference (IV, Random, 95% CI)	-1.70 [-3.72, 0.32]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
11 Nausea	1	380	Risk Ratio (IV, Random, 95% CI)	1.21 [0.62, 2.39]
11.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	1.21 [0.62, 2.39]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	1	380	Risk Ratio (IV, Random, 95% CI)	1.33 [0.58, 3.09]
12.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	1.33 [0.58, 3.09]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	1	380	Risk Ratio (IV, Random, 95% CI)	2.0 [0.37, 10.79]
13.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	2.0 [0.37, 10.79]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	1	380	Risk Ratio (IV, Random, 95% CI)	7.83 [3.43, 17.88]
16.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	7.83 [3.43, 17.88]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	1	380	Risk Ratio (IV, Random, 95% CI)	8.5 [1.99, 36.28]
17.1 Hospital setting	1	380	Risk Ratio (IV, Random, 95% CI)	8.5 [1.99, 36.28]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

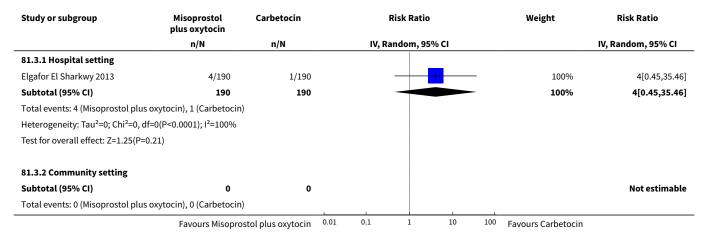


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

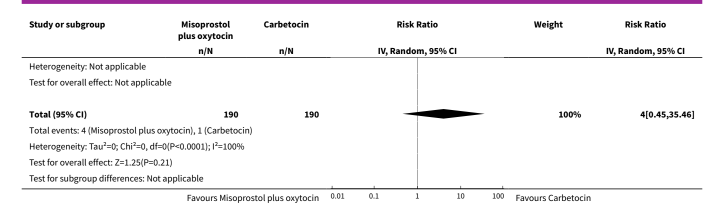
Analysis 81.1. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 1 Death.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin		Ri	sk Ratio		Weight	Risk Ratio
	n/N	n/N		IV, Rar	ıdom, 95% CI			IV, Random, 95% CI
81.1.1 Hospital setting								
Elgafor El Sharkwy 2013	0/190	0/190						Not estimable
Subtotal (95% CI)	190	190						Not estimable
Total events: 0 (Misoprostol plus oxy	rtocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
81.1.2 Community setting								
Subtotal (95% CI)	0	0						Not estimable
Total events: 0 (Misoprostol plus oxy	rtocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
Total (95% CI)	190	190						Not estimable
Total events: 0 (Misoprostol plus oxy	rtocin), 0 (Carbetocin)							
Heterogeneity: Not applicable								
Test for overall effect: Not applicable	2							
Test for subgroup differences: Not a	oplicable							
	Favours Misopros	tol plus oxytocin	0.01	0.1	1 10	100	Favours Carbetocin	

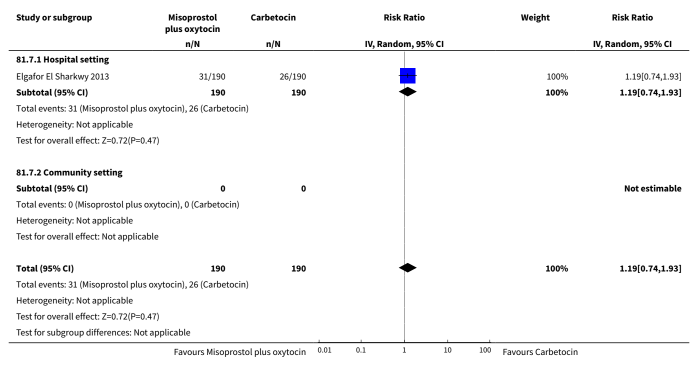
Analysis 81.3. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 3 Blood transfusion.







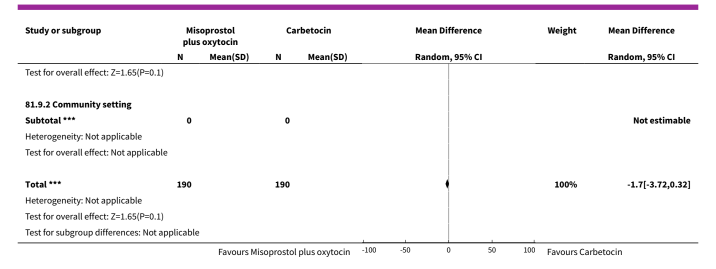
Analysis 81.7. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 7 Additional uterotonics.



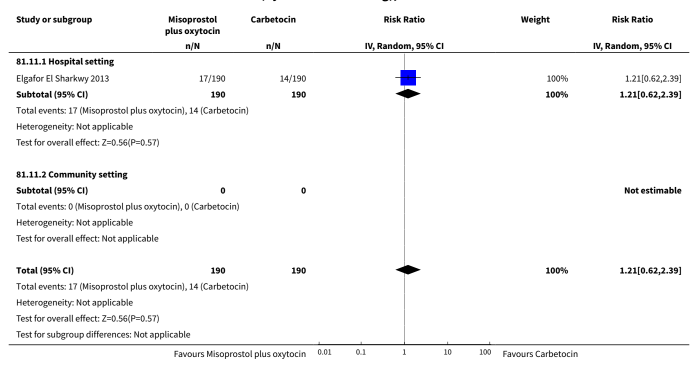
Analysis 81.9. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 9 Change in haemoglobin.

Study or subgroup		oprostol oxytocin	Ca	rbetocin		М	lean Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		R	andom, 95%	6 CI			Random, 95% CI
81.9.1 Hospital setting											
Elgafor El Sharkwy 2013	190	13.4 (8.7)	190	15.1 (11.2)			+			100%	-1.7[-3.72,0.32]
Subtotal ***	190		190				•			100%	-1.7[-3.72,0.32]
Heterogeneity: Not applicable											
		Favours Mis	oprostol	plus oxytocin	-100	-50	0	50	100	Favours Carl	petocin





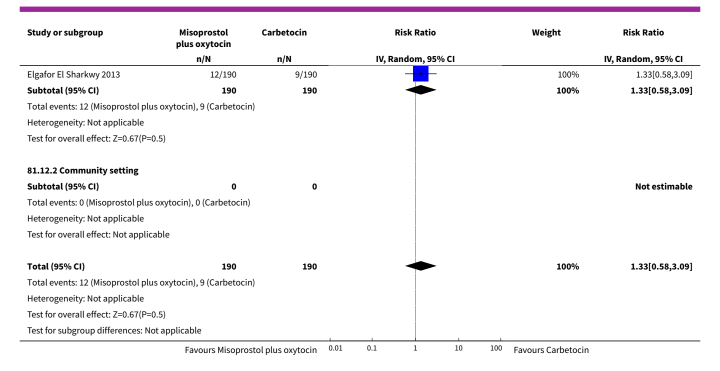
Analysis 81.11. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 11 Nausea.



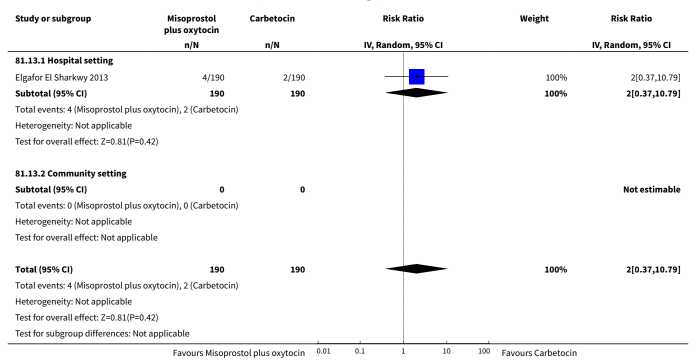
Analysis 81.12. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 12 Vomiting.

Study or subgroup	Misoprostol plus oxytocin	Carbetocin	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		IV, R	andom, 95	% CI			IV, Random, 95% CI
81.12.1 Hospital setting									
	Favours Misopro	stol plus oxytocin	0.01	0.1	1	10	100	Favours Carbetocin	



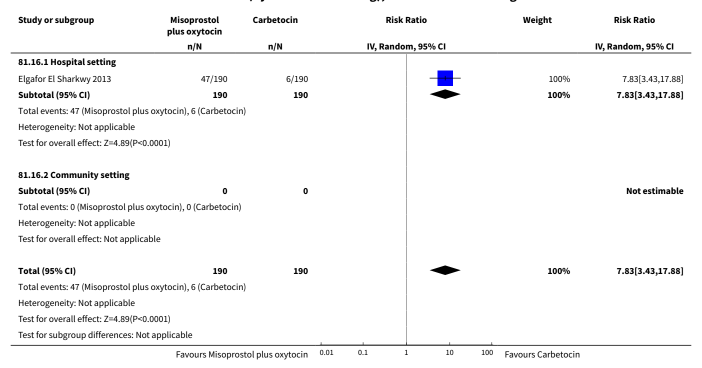


Analysis 81.13. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 13 Headache.





Analysis 81.16. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 16 Shivering.



Analysis 81.17. Comparison 81 Misoprostol plus oxytocin vs Carbetocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin			Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
81.17.1 Hospital setting					
Elgafor El Sharkwy 2013	17/190	2/190	- 	100%	8.5[1.99,36.28]
Subtotal (95% CI)	190	190		100%	8.5[1.99,36.28]
Total events: 17 (Misoprostol plus o	xytocin), 2 (Carbetocin))			
Heterogeneity: Not applicable					
Test for overall effect: Z=2.89(P=0)					
81.17.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus ox	ytocin), 0 (Carbetocin)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	e				
Total (95% CI)	190	190	-	100%	8.5[1.99,36.28]
Total events: 17 (Misoprostol plus o	xytocin), 2 (Carbetocin))			
Heterogeneity: Not applicable					
Test for overall effect: Z=2.89(P=0)					
Test for subgroup differences: Not a	pplicable				
	Favours Misopros	tol plus oxytocin 0.0	0.1 1 10 10	D Favours Carbetocin	



Comparison 82. Ergometrine vs Ergometrine plus oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbidity: intensive care admissions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	1	144	Risk Ratio (IV, Random, 95% CI)	0.17 [0.01, 4.06]
6.1 Hospital setting	1	144	Risk Ratio (IV, Random, 95% CI)	0.17 [0.01, 4.06]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	144	Mean Difference (IV, Random, 95% CI)	20.10 [5.76, 34.44]
8.1 Hospital setting	1	144	Mean Difference (IV, Random, 95% CI)	20.10 [5.76, 34.44]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

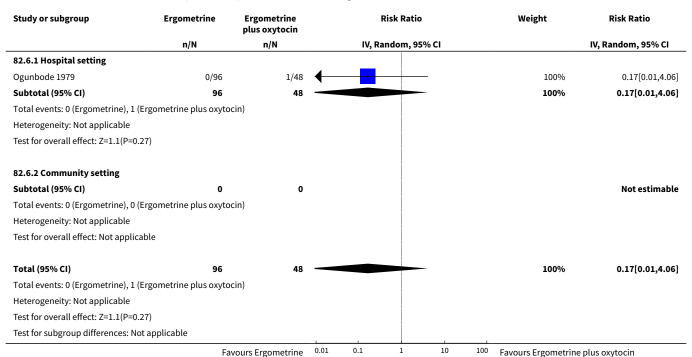


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Analysis 82.6. Comparison 82 Ergometrine vs Ergometrine plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.





Analysis 82.8. Comparison 82 Ergometrine vs Ergometrine plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

Study or subgroup	Erg	ometrine	_	ometrine oxytocin	Mea	nn Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Ran	dom, 95% CI		Random, 95% CI
32.8.1 Hospital setting								
Ogunbode 1979	96	96 (54.2)	48	75.9 (33.2)		-	100%	20.1[5.76,34.44]
Subtotal ***	96		48			•	100%	20.1[5.76,34.44]
Heterogeneity: Not applicable								
Test for overall effect: Z=2.75(P=0.01)								
32.8.2 Community setting								
Subtotal ***	0		0					Not estimable
Heterogeneity: Not applicable								
est for overall effect: Not applicable								
Fotal ***	96		48			•	100%	20.1[5.76,34.44]
Heterogeneity: Not applicable								
Test for overall effect: Z=2.75(P=0.01)								
Test for subgroup differences: Not ap	plicable							
			Favours	s Ergometrine -10) -50	0	50	50 100 Favours Erg

Comparison 83. Misoprostol plus oxytocin vs Ergometrine (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.1 Hospital setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 84. Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting)

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Death	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 PPH >= 1000 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.41 [0.68, 2.94]
2.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.41 [0.68, 2.94]
2.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 Blood transfusion	2	1605	Risk Ratio (IV, Random, 95% CI)	0.89 [0.36, 2.19]
3.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	0.89 [0.36, 2.19]
3.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 Severe maternal morbid- ity: intensive care admis- sions	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Severe maternal morbidity: shock	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 PPH >= 500 mL	2	1605	Risk Ratio (IV, Random, 95% CI)	1.39 [0.65, 2.99]
6.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.39 [0.65, 2.99]
6.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
7 Additional uterotonics	2	1605	Risk Ratio (IV, Random, 95% CI)	1.48 [0.82, 2.69]
7.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.48 [0.82, 2.69]
7.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
8 Blood loss	1	802	Mean Difference (IV, Random, 95% CI)	-16.0 [-40.24, 8.24]
8.1 Hospital setting	1	802	Mean Difference (IV, Random, 95% CI)	-16.0 [-40.24, 8.24]
8.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9 Change in haemoglobin	2	1605	Mean Difference (IV, Random, 95% CI)	-0.50 [-1.72, 0.72]
9.1 Hospital setting	2	1605	Mean Difference (IV, Random, 95% CI)	-0.50 [-1.72, 0.72]
9.2 Community setting	0	0	Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Breastfeeding	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

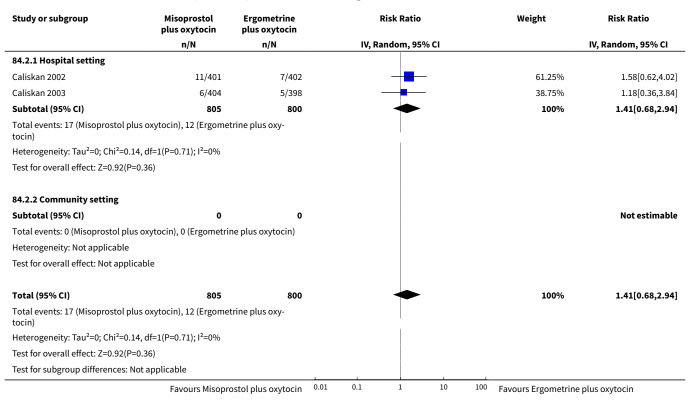


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
10.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11 Nausea	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
11.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
12 Vomiting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.04 [0.23, 4.77]
12.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	1.04 [0.23, 4.77]
12.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13 Headache	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
13.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14 Abdominal pain	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
14.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15 Hypertension	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.1 Hospital setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
15.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
16 Shivering	2	1605	Risk Ratio (IV, Random, 95% CI)	2.95 [2.02, 4.29]
16.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	2.95 [2.02, 4.29]
16.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
17 Fever	2	1605	Risk Ratio (IV, Random, 95% CI)	2.89 [1.51, 5.54]
17.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	2.89 [1.51, 5.54]
17.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
18 Diarrhoea	2	1605	Risk Ratio (IV, Random, 95% CI)	0.81 [0.46, 1.41]
18.1 Hospital setting	2	1605	Risk Ratio (IV, Random, 95% CI)	0.81 [0.46, 1.41]
18.2 Community setting	0	0	Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19 Maternal well-being	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected

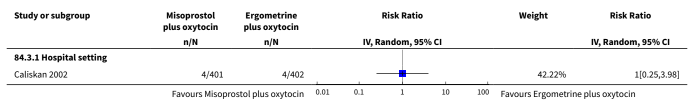


Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
19.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
19.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20 Maternal satisfaction	0		Risk Ratio (IV, Random, 95% CI)	Totals not selected
20.1 Hospital setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]
20.2 Community setting	0		Risk Ratio (IV, Random, 95% CI)	0.0 [0.0, 0.0]

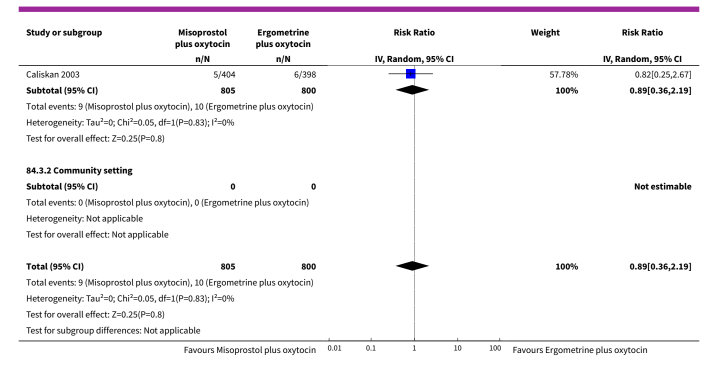
Analysis 84.2. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 2 PPH >= 1000 mL.



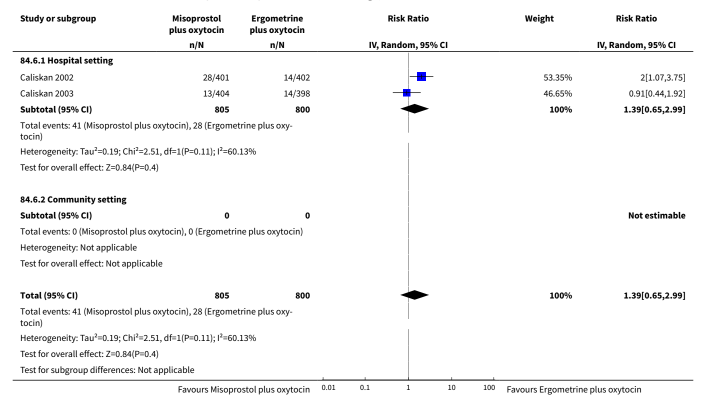
Analysis 84.3. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 3 Blood transfusion.





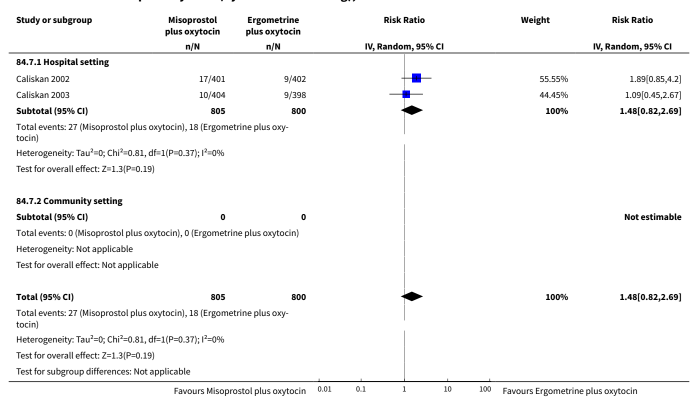


Analysis 84.6. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 6 PPH >= 500 mL.





Analysis 84.7. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 7 Additional uterotonics.

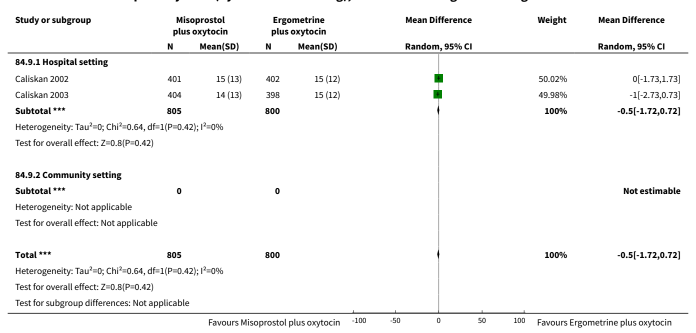


Analysis 84.8. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 8 Blood loss.

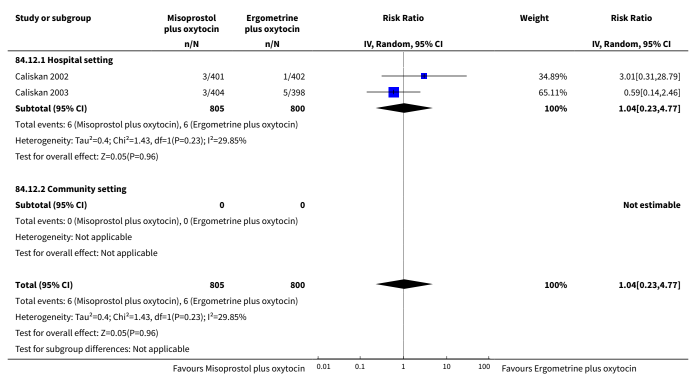
Study or subgroup		oprostol oxytocin	_	ometrine oxytocin		Mea	an Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	ndom, 95% CI			Random, 95% CI
84.8.1 Hospital setting										
Caliskan 2003	404	280 (182)	398	296 (168)		_			100%	-16[-40.24,8.24]
Subtotal ***	404		398			-			100%	-16[-40.24,8.24]
Heterogeneity: Not applicable										
Test for overall effect: Z=1.29(P=0.2))									
84.8.2 Community setting										
Subtotal ***	0		0							Not estimable
Heterogeneity: Not applicable										
Test for overall effect: Not applicable	e									
Total ***	404		398			~			100%	-16[-40.24,8.24]
Heterogeneity: Not applicable										
Test for overall effect: Z=1.29(P=0.2))									
Test for subgroup differences: Not a	pplicable									
		Favours Mis	oprostol	plus oxytocin	-100	-50	0 50	100	Favours Erg	ometrine plus oxytocin



Analysis 84.9. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 9 Change in haemoglobin.



Analysis 84.12. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 12 Vomiting.





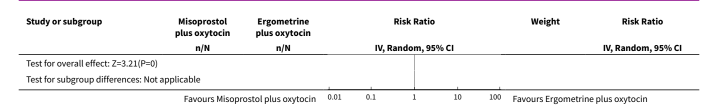
Analysis 84.16. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 16 Shivering.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
84.16.1 Hospital setting					
Caliskan 2002	52/401	19/402	-	55.1%	2.74[1.65,4.55]
Caliskan 2003	49/404	15/398	-	44.9%	3.22[1.84,5.64]
Subtotal (95% CI)	805	800	•	100%	2.95[2.02,4.29]
Total events: 101 (Misoprostol pitocin)	lus oxytocin), 34 (Ergom	etrine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.1	7, df=1(P=0.68); I ² =0%				
Test for overall effect: Z=5.63(P<	0.0001)				
84.16.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus	s oxytocin), 0 (Ergometri	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applic	cable				
Total (95% CI)	805	800	•	100%	2.95[2.02,4.29]
Total events: 101 (Misoprostol pi tocin)	lus oxytocin), 34 (Ergom	etrine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.1	7, df=1(P=0.68); I ² =0%				
Test for overall effect: Z=5.63(P<	0.0001)				
Test for subgroup differences: No	ot applicable				
	Favours Misopr	ostol plus oxytocin 0.0	1 0.1 1 10 1	.00 Favours Ergometrin	e plus oxytocin

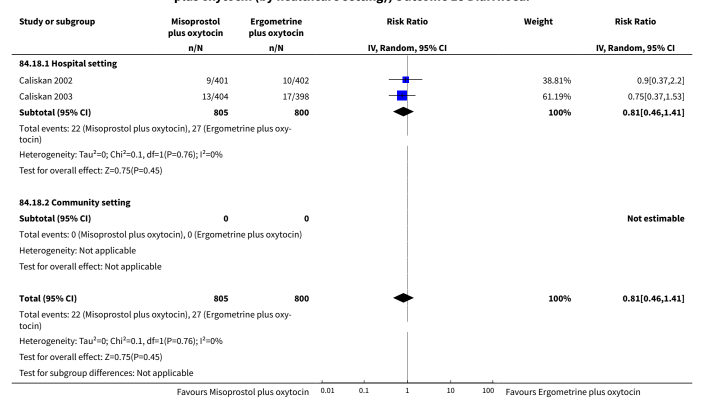
Analysis 84.17. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 17 Fever.

Study or subgroup	Misoprostol plus oxytocin	Ergometrine plus oxytocin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	IV, Random, 95% CI		IV, Random, 95% CI
84.17.1 Hospital setting					
Caliskan 2002	19/401	6/402	_ 	51.12%	3.17[1.28,7.87]
Caliskan 2003	16/404	6/398	-	48.88%	2.63[1.04,6.64]
Subtotal (95% CI)	805	800	•	100%	2.89[1.51,5.54]
Total events: 35 (Misoprostol plus or tocin)	kytocin), 12 (Ergome	rine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.08, df	f=1(P=0.77); I ² =0%				
Test for overall effect: Z=3.21(P=0)					
84.17.2 Community setting					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Misoprostol plus oxy	tocin), 0 (Ergometrii	ne plus oxytocin)			
Heterogeneity: Not applicable					
Test for overall effect: Not applicable	е				
Total (95% CI)	805	800	•	100%	2.89[1.51,5.54]
Total events: 35 (Misoprostol plus or tocin)	kytocin), 12 (Ergome	trine plus oxy-			
Heterogeneity: Tau ² =0; Chi ² =0.08, df	f=1(P=0.77); I ² =0%				
	Favours Misopro	ostol plus oxytocin 0.	01 0.1 1 10 100	Favours Ergometrin	e plus oxytocin





Analysis 84.18. Comparison 84 Misoprostol plus oxytocin vs Ergometrine plus oxytocin (by healthcare setting), Outcome 18 Diarrhoea.



APPENDICES

Appendix 1. Search terms

ClinicalTrials.gov and the WHO International Clinical Trials Registry Platform (ICTRP)

Third stage AND labo(u)r AND oxytocin

Third stage AND labo(u)r AND misoprostol

Third stage AND labo(u)r AND carbetocin

Third stage AND labo(u)r AND ergometrine

uterotonic* AND oxytocin

uterotonic* AND misoprostol



uterotonic* AND carbetocin

uterotonic* AND ergometrine

uterotonic* AND labo(u)r

uterotonic* AND h(a)emorrhage

h(a)emorrhage AND postpartum AND ergometrine

h(a)emorrhage AND postpartum AND oxytocin

h(a)emorrhage AND postpartum AND carbetocin

h(a)emorrhage AND postpartum AND misoprostol

Appendix 2. Supplementary Appendix

For all subgroup and sensitivity analyses please see Appendix 1 https://edata.bham.ac.uk/284/

For all forest plots for all outcomes and interventions see Appendix 2 https://edata.bham.ac.uk/284/

For additional data from trialists see Appendix 3 https://edata.bham.ac.uk/284/

WHAT'S NEW

Date	Event	Description
24 May 2018	New search has been performed	Search updated. We have included 56 new trials in this update involving 46,612 women. We have updated the methods - all changes are summarised in detail in 'Differences between protocol and review'. Six authors have stepped down from the team and 10 new authors have joined the review team.
24 May 2018	New citation required but conclusions have not changed	With the addition of 56 new trials (46,612 women), the update now includes a total of 196 trials (135,559 women). The conclusions remain largely the same. The results for the primary outcome of postpartum haemorrhage (PPH) \geq 500 mL were similar to the previously published review (Gallos 2018), although the quality of the evidence for carbetocin has changed from 'very low-certainly' to 'moderate-certainty evidence' for this outcome, due to the addition of data from three studies including approximately 30,000 women. For the primary outcome of PPH \geq 1000 mL, none of the agents is significantly more effective when compared with the reference uterotonic agent oxytocin. In the previous version of the review, high-quality evidence suggested that ergometrine plus oxytocin was more effective in reducing PPH \geq 1000 mL in comparison to oxytocin. For all other outcomes (blood transfusion; additional uterotonics; and side effects), the results are largely the same.

CONTRIBUTIONS OF AUTHORS

Ioannis D Gallos (IDG) and Arri Coomarasamy (AC) conceived the idea for this study. IDG, Malcolm J Price (MJP), Aurelio Tobias (AT), Olufemi T Oladapo (OTO) and AC designed the meta-analysis. IDG designed all electronic data collection forms. IDG, Argyro Papadopoulou (AP), Rebecca Man (RM), Nikos Athanasopoulos (NA) screened trials and extracted data. MJP and AT performed the statistical analysis. Myfanwy Williams (MJW), Virginia Diaz (VD), Julia Pasquale (JP), Monica Chamillard (MC), Josh Vogel (JPV) and OTO graded the evidence and created the 'summary of findings' tables. IDG drafted the protocol and all versions of the review. AP, RM, NA, MP, AT, MP, OT, MJW, VD, JP, MC, Mariana Widmer (MW), Özge Tuncalp (OT), G Justus Hofmeyr (GJH), Fernando Althabe (FA), A Metin Gulmezoglu (AMG), JPV, OTO and AC edited and revised the review.



DECLARATIONS OF INTEREST

Ioannis D Gallos (IDG): is a co-applicant to the UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis.* He has been involved in one or more previous or ongoing trials related to the use of uterotonics for the prevention of PPH that were eligible for inclusion in this review. Ferring Pharmaceuticals and Novartis supplied carbetocin and oxytocin for these studies. IDG did not participate in any decisions regarding these trials (i.e. assessment for inclusion/exclusion, trial quality, data extraction) for the purposes of this review (and will not for future updates) – these tasks were carried out by other members of the team who were not directly involved in the trials.

Argyro Papadopoulou (AP): none known.

Rebecca Man (RM): none known.

Nikolaos Athanasopoulos (NA): none known.

Malcolm J Price (MJP): is a co-applicant to the UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic* agents for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis.

Aurelio Tobias: none known.

Myfanwy Williams (MJW): is employed by the University of Liverpool as a Research Associate at Cochrane Pregnancy and Childbirth. Her role is supported by the World Health Organization.

Virginia Diaz (VD): none known.

Julia Pasquale (JP): none known.

Monica Chamillard (MC): none known.

Mariana Widmer (MW): has been involved in a trial related to the use of uterotonics for the prevention of PPH that is included in this review. Ferring Pharmaceuticals and Novartis supplied carbetocin and oxytocin for the trial and the study is supported by WHO/Merck for Mothers. MW did not participate in any decisions regarding this trial (i.e. assessment for inclusion/exclusion, trial quality, data extraction) for the purposes of this review or future updates – these tasks were carried out by other members of the team who were not directly involved in the trial.

Özge Tunçalp (OT): is a co-applicant to the UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic* agents for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis.

G Justus Hofmeyr (GJH): has been and continues to be involved in a number of studies that may be eligible for inclusion in this review, but has not been (and will not participate in) data extraction or quality assessment of the studies in which he was involved. GJH is a co-investigator on the UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis*. Neither he nor his institution receives funding from this grant.

Fernando Althabe: none known.

A Metin Gulmezoglu (AMG): was part of the central coordination unit of the large World Health Organization multicentre trial comparing carbetocin with oxytocin included in the review. He is a co-applicant to the UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic drugs for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis*.

Joshua Vogel (JPV): led the updating of WHO recommendations on uterotonics for the prevention of postpartum haemorrhage based on this review.

Olufemi T Oladapo (OTO): led the updating of WHO recommendations on uterotonics for the prevention of postpartum haemorrhage based on the findings of this review update.

Arri Coomarasamy (AC): is the Chief Investigator of UK National Institute for Health Research HTA Project Award 14/139/17 entitled *Uterotonic agents for preventing postpartum haemorrhage: a network meta-analysis and cost-effectiveness analysis*. He has been involved in one or more previous or ongoing trials related to the use of uterotonics for the prevention of PPH that were eligible for inclusion in this review. Ferring Pharmaceuticals and Novartis supplied carbetocin and oxytocin for these studies and another study is supported by WHO/ Merck for Mothers. AC did not participate in any decisions regarding these trials (i.e. assessment for inclusion/exclusion, trial quality, data extraction) for the purposes of this review or future updates – these tasks have been carried out by other members of the team who were not directly involved in the trials. AC is a member of the Executive Board of Ammalife (UK registered charity 1120236). He does not receive any payment for this relationship.



SOURCES OF SUPPORT

Internal sources

· University of Birmingham, UK.

The authors of this review are employed by the institutions indicated by their respective affiliations except where otherwise stated.

• World Health Organization, Switzerland.

The authors of this review are employed by the institutions indicated by their respective affiliations except where otherwise stated.

· University of the Witwatersrand, South Africa.

The authors of this review are employed by the institutions indicated by their respective affiliations except where otherwise stated.

External sources

· National Institute for Health Research, UK.

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· Birmingham Women's NHS Foundation Trust, UK.

Additional financial support to meet the employment costs of Helen M Willliams (HMW) and Abi Merriel (AM) is provided by Birmingham Women's NHS Foundation Trust.

· Ammalife, UK.

Additional financial support to meet the employment costs of Abi Merriel (AM) is provided by Ammalife (UK Registered Charity 1120236).

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

There are some differences between the published protocol for this review (Gallos 2015) and the full review, these are listed below.

Objectives

We have clarified the objectives of this review.

In our protocol the stated objectives were: "We aim to assess the clinical effectiveness and side-effect profile of uterotonic agents to prevent PPH, and to generate a clinically useful ranking of available uterotonics according to their effectiveness and side effects. We will explore the effects according to various key prognostic and treatment factors. The population of interest is women following a vaginal birth or a caesarean section in the hospital or the community setting. All uterotonic agents considered by the WHO are eligible and the outcomes include blood loss-related outcomes and side effects."

In the review, our objectives are listed as:

To identify the most effective uterotonic agent(s) to prevent PPH with the least side effects, and generate a ranking according to their effectiveness and side-effect profile.

Methods/types of interventions

The text in this section has been edited to add sensitivity analyses that became necessary during the review and explain how we grouped the agents for analysis.

In the protocol, this section stated:

"We will consider trials of uterotonics described by WHO (WHO 2012) (oxytocin, ergometrine, misoprostol, carbetocin, or combinations of uterotonics) administered prophylactically by healthcare professionals for preventing PPH via any systemic route (sublingual, subcutaneous, intramuscular, rectal, oral, intravenous bolus and/or infusion) compared with another uterotonic or with placebo or no treatment. If we identify in the included studies interventions that we are not aware of, we will consider them as eligible and include them in the network after assessing their comparability with those named above. We will include trials in which non-pharmacologic cointerventions such as controlled cord traction, cord clamping, or uterine massage was performed as a randomised intervention in all arms of the trial. We will stratify all agents according to mode of birth, prior risk of PPH, healthcare setting, specific dosage, regimen and route, to detect inequalities in subgroups that could affect comparative effectiveness."

Figure 1 (in the published protocol) shows the overall network of eligible comparisons in the review at the agent level.



"Multi-arm trials that compare different dosages, regimens or routes of one uterotonic agent, but also compare those versus another uterotonic agent, will be included. Intervention arms of different dosages, regimens or routes of the same uterotonic agent will be merged together for the global analysis of all outcomes and treated as separate independent comparisons only for the relevant subgroup analysis according to dosage, regimen and route of administration, while taking into account the correlation between the comparisons. We will exclude trials comparing exclusively different dosages, regimens or routes of administration of the same uterotonic agent. The review will be restricted to studies evaluating uterotonic agents administered systemically at the birth of the baby for preventing PPH. Studies considering non-uterotonic agents, uterotonic agents administered locally (for example, via intraumbilical or intrauterine routes) or at a later stage of delivery (for example, for the treatment of PPH or for retained placenta) will be excluded."

In our review this section now states:

Trials were eligible if they administered uterotonic agents of any dosage, route or regimen systemically at birth for preventing postpartum haemorrhage (PPH), and compared them with other uterotonic agents, placebo or no treatment. Trials evaluating uterotonic agents not administered systemically, such as intrauterine administration, or not immediately after birth, or exclusively comparing different dosages, routes or regimens of the same uterotonic agent were excluded. We included trials in which non-pharmacologic co-interventions such as controlled cord traction, cord clamping, or uterine massage was performed as a randomised intervention in all arms of the trial and the effects of such co-interventions were tested through a sensitivity analysis.

We classified agents into single agents including oxytocin, carbetocin, injectable prostaglandins (carboprost tromethamine, sulprostone), misoprostol, ergometrine (included also ergonovine, methylergonovine), and combination agents including ergometrine plus oxytocin (Syntometrine ® as a fixed-combination agent containing 5 international units (IU) of oxytocin and 500 mcg of ergometrine, any oxytocin dose and route when combined with any dose and route of ergometrine, ergonovine, or methylergonovine), and misoprostol plus oxytocin (any oxytocin dose and route when combined with any dose and route of misoprostol).

Methods/search methods

The search methods have been updated in line with the current standard search methods text of Cochrane Pregnancy and Childbirth.

Methods/types of outcomes/secondary outcomes

We have edited our outcome list based on the core outcome set for prevention of PPH. We have edited the composite outcome 'maternal deaths or severe morbidity events' and split that into 'severe maternal morbidity: 'intensive care admissions' and 'severe maternal morbidity: shock (as defined by the trialists)'. We have removed the outcomes 'manual removal of placenta'; 'mean durations of the third stage of labour (minutes)'; 'neonatal unit admission requirement'; 'tachycardia'; and 'hypotension'. We have added outcomes 'diarrhoea'; 'maternal sense of well-being (as defined by the trialists)'; 'maternal satisfaction (as defined by the trialists)'.

Assessment of reporting biases

We have edited the assessment of reporting biases. In the protocol, these sections stated:

We assessed potential reporting bias for the primary outcomes by assessing the sensitivity of results to exclusion of studies with fewer than 400 participants.

In our review these sections now state:

If there were 10 or more studies in the meta-analysis, we investigated reporting biases (such as publication bias) using funnel plots. The funnel plots were assessed visually for asymmetry. We also assessed potential reporting bias for the primary outcomes by assessing the sensitivity of results to exclusion of studies with fewer than 400 participants.

Methods/investigation of heterogeneity and inconsistency and also subgroup analysis

We have edited the intervention subgroups for exploring heterogeneity and inconsistency and also subgroup analysis. In the protocol, these sections stated:

Intervention: dose, regimen or route.

We assessed subgroup differences by firstly comparing the network diagram for each subgroup. Next, we performed a network metaanalysis for each subgroup and we compared their relative treatment effects and their relative treatment ranking.

In our review these sections now state:

Intervention: Dose of misoprostol (≥ 600 mcg versus < 600 mcg), and regimen of oxytocin (bolus versus bolus plus infusion versus infusion only).

We assessed subgroup differences by firstly comparing the network diagram for each subgroup. Next, we performed a pairwise and network meta-analysis for each subgroup and we compared their relative treatment effects and their relative treatment ranking. We examined



the subgroups for qualitative interactions where the direction of effect could be reversed, that is if an intervention was beneficial in one subgroup but harmful in another.

Methods/investigation of heterogeneity and inconsistency and also subgroup analysis

We have carried out additional sensitivity analyses that became necessary during the conduct of the review. These are listed below.

- 1. Trials that also randomised participants to co-interventions such as uterine massage or controlled cord traction.
- 2. Trials with more than 10% missing data.
- 3. Trials published before 1990.

Analysis

Since publication of the protocol for this review, further methods became available to perform the analysis with a frequentist approach in STATA. We changed our analysis for this reason to STATA rather than WinBUGS and a Bayesian environment.

'Summary of findings' table

We have modified our approach to assessing confidence in the evidence generated by this network meta-analysis, in line with recent guidance published by the GRADE working group (see Brignardello-Petersen 2018; Puhan 2014). This was not planned at the protocol stage, because we were not aware then of the most up-to-date guidance.

INDEX TERMS

Medical Subject Headings (MeSH)

*Network Meta-Analysis; Drug Therapy, Combination [adverse effects] [methods]; Ergonovine [adverse effects] [*therapeutic use]; Fever [chemically induced]; Hypertension [chemically induced]; Misoprostol [*therapeutic use]; Oxytocics [*therapeutic use]; Oxytocin [adverse effects] [*analogs & derivatives] [*therapeutic use]; Postpartum Hemorrhage [*prevention & control]; Prostaglandins [*therapeutic use]; Randomized Controlled Trials as Topic; Vomiting [chemically induced]

MeSH check words

Female; Humans