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Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review)

Oddie SJ, Young L, McGuire W

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TABLE OF CONTENTS

ABSTRACT 1
PLAIN LANGUAGE SUMMARY
SUMMARY OF FINDINGS
BACKGROUND
OBJECTIVES
METHODS
RESULTS
8 Figure 1.
Figure 2
Figure 3
Figure 4
DISCUSSION
AUTHORS' CONCLUSIONS
ACKNOWLEDGEMENTS
REFERENCES
CHARACTERISTICS OF STUDIES
DATA AND ANALYSES
Analysis 1.1. Comparison 1: Slow versus faster rates of feed advancement, Outcome 1: Necrotising enterocolitis - all infants 36
Analysis 1.2. Comparison 1: Slow versus faster rates of feed advancement, Outcome 2: Necrotising enterocolitis - ELBW/ 36
extremely preterm infants
Analysis 1.3. Comparison 1: Slow versus faster rates of feed advancement, Outcome 3: Necrotising enterocolitis - small for 37
gestational age or growth restricted
Analysis 1.4. Comparison 1: Slow versus faster rates of feed advancement, Outcome 4: Necrotising enterocolitis - AREDFV 37
Analysis 1.5. Comparison 1: Slow versus faster rates of feed advancement, Outcome 5: Mortality - all infants
Analysis 1.6. Comparison 1: Slow versus faster rates of feed advancement, Outcome 6: Mortality - ELBW/extremely preterm 38
infants
Analysis 1.7. Comparison 1: Slow versus faster rates of feed advancement, Outcome 7: Mortality - small for gestational age or 39 growth restricted
Analysis 1.8. Comparison 1: Slow versus faster rates of feed advancement, Outcome 8: Mortality - AREDFV
Analysis 1.9. Comparison 1: Slow versus faster rates of feed advancement, Outcome 9: Weight z-score at hospital discharge 39
Analysis 1.10. Comparison 1: Slow versus faster rates of feed advancement, Outcome 10: Head circumference z-score at hospital 39
discharge
Analysis 1.11. Comparison 1: Slow versus faster rates of feed advancement, Outcome 11: Moderate or severe disability 40
Analysis 1.12. Comparison 1: Slow versus faster rates of feed advancement, Outcome 12: Cerebral palsy
Analysis 1.13. Comparison 1: Slow versus faster rates of feed advancement, Outcome 13: Visual impairment
Analysis 1.14. Comparison 1: Slow versus faster rates of feed advancement, Outcome 14: Hearing impairment
Analysis 1.15. Comparison 1: Slow versus faster rates of feed advancement, Outcome 15: Cognitive impairment
Analysis 1.16. Comparison 1: Slow versus faster rates of feed advancement, Outcome 16: Feed intolerance (causing interruption 41 of enteral feeding)
Analysis 1.17. Comparison 1: Slow versus faster rates of feed advancement, Outcome 17: Invasive infection
APPENDICES
WHAT'S NEW
HISTORY
CONTRIBUTIONS OF AUTHORS
DECLARATIONS OF INTEREST
SOURCES OF SUPPORT
DIFFERENCES BETWEEN PROTOCOL AND REVIEW
INDEX TERMS

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants

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ABSTRACT

Background

Early enteral feeding practices are potentially modifiable risk factors for necrotising enterocolitis (NEC) in very preterm or very low birth weight (VLBW) infants. Observational studies suggest that conservative feeding regimens, including slowly advancing enteral feed volumes, reduce the risk of NEC. However, it is unclear whether slow feed advancement may delay establishment of full enteral feeding, and if it could be associated with infectious morbidities secondary to prolonged exposure to parenteral nutrition.

Objectives

To determine the effects of slow rates of enteral feed advancement on the risk of NEC, mortality, and other morbidities in very preterm or VLBW infants.

Search methods

We searched CENTRAL (2020, Issue 10), Ovid MEDLINE (1946 to October 2020), Embase via Ovid (1974 to October 2020), Maternity and Infant Care database (MIDIRS) (1971 to October 2020), CINAHL (1982 to October 2020), and clinical trials databases and reference lists of retrieved articles for eligible trials.

Selection criteria

We included randomised or quasi-randomised controlled trials that assessed effects of slow (up to 24 mL/kg/d) versus faster rates of advancement of enteral feed volumes on the risk of NEC in very preterm or VLBW infants.

Data collection and analysis

Two review authors separately evaluated trial risk of bias, extracted data, and synthesised effect estimates using risk ratio (RR), risk difference (RD), and mean difference. We used the GRADE approach to assess the certainty of evidence. Outcomes of interest were NEC, all-cause mortality, feed intolerance, and invasive infection.

Main results

We included 14 trials involving a total of 4033 infants (2804 infants participated in one large trial). None of the trials masked parents, caregivers, or investigators. Risk of bias was otherwise low. Most infants were stable very preterm or VLBW infants of birth weight appropriate for gestation. About one-third of all infants were extremely preterm or extremely low birth weight (ELBW), and about one-fifth were small for gestational age, growth-restricted, or compromised as indicated by absent or reversed end-diastolic flow velocity in the foetal umbilical artery. Trials typically defined slow advancement as daily increments of 15 to 24 mL/kg, and faster advancement as daily increments of 30 to 40 mL/kg.

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



Meta-analyses showed that slow advancement of enteral feed volumes probably has little or no effect on the risk of NEC (RR 1.06, 95% confidence interval (CI) 0.83 to 1.37; RD 0.00, 95% CI –0.01 to 0.02; 14 trials, 4026 infants; moderate-certainty evidence) or all-cause mortality prior to hospital discharge (RR 1.13, 95% CI 0.91 to 1.39; RD 0.01, 95% CI –0.01 to 0.02; 13 trials, 3860 infants; moderate-certainty evidence). Meta-analyses suggested that slow advancement may slightly increase feed intolerance (RR 1.18, 95% CI 0.95 to 1.46; RD 0.05, 95% CI –0.02 to 0.12; 9 trials, 719 infants; low-certainty evidence) and may slightly increase the risk of invasive infection (RR 1.14, 95% CI 0.99 to 1.31; RD 0.02, 95% CI –0.00 to 0.05; 11 trials, 3583 infants; low-certainty evidence).

Authors' conclusions

The available trial data indicate that advancing enteral feed volumes slowly (daily increments up to 24 mL/kg) compared with faster rates probably does not reduce the risk of NEC, death, or feed intolerance in very preterm or VLBW infants. Advancing the volume of enteral feeds at a slow rate may slightly increase the risk of invasive infection.

PLAIN LANGUAGE SUMMARY

Slowly advancing milk feeds to prevent necrotising enterocolitis in very low birth weight infants

Review question

Does limiting the rate of increase in milk feeds that very low birth weight infants receive each day during the first few weeks after birth reduce the risk of severe bowel problems?

Background

Very preterm (born more than eight weeks early) or very low birth weight (weighing < 1500 grams at birth) newborn babies are at risk of developing a severe bowel disorder called necrotising enterocolitis (where the bowel becomes inflamed and dies). It is thought that one way to prevent this condition may be to limit the milk feeds that infants receive each day for the first few weeks after birth.

Study characteristics

We searched for randomised controlled trials (a type of study where participants are randomly assigned to one of two or more treatment groups) comparing slow versus faster rates of increase in the amount of milk fed to newborn infants who were very preterm or very low birth weight. We included 14 trials involving a total of 4033 infants (2804 infants participated in one large trial). The search is up-to-date as of October 2020.

Key results

Combined analysis of the included trials showed that slow advancement of enteral feed volumes probably does not affect the risk of necrotising enterocolitis or death (moderate-certainty evidence).

Conclusions and certainty of evidence

Slowly advancing enteral feed volumes probably does not reduce the risk of necrotising enterocolitis or death before hospital discharge for very preterm or very low birth weight infants.

SUMMARY OF FINDINGS

Summary of findings 1. Slow compared with faster rates of enteral feed advancement for preventing necrotising enterocolitis in very preterm or very low birth weight infants

Slow compared with faster rates of enteral feed advancement for preventing necrotising enterocolitis in very preterm or very low birth weight infants

Patient or population: very preterm or very low birth weight infants

Setting: neonatal care facilities in North America, Colombia, Bangladesh, Iran, India, Turkey, South Africa, Ireland, and the UK

Intervention: slow rates of enteral feed advancement

Comparison: faster rates of enteral feed advancement

Outcomes	Anticipated absolute ef	ífects* (95% CI)	Relative effect (95% CI)	No. of partici- pants	Certainty of the evidence
	Risk with faster rates of enteral feed ad- vancement	Risk with slow rates of enteral feed enhancement		(studies)	(GRADE)
Necrotising enterocolitis before hospital discharge	54 per 1000	57 per 1000 (45 to 77)	RR 1.06 (0.83 to 1.37)	4026 (14 trials)	⊕⊕⊕⊝ MODERATE ^a
Mortality before hospital discharge	71 per 1000	80 per 1000 (64 to 98)	RR 1.13 (0.91 to 1.39)	3860 (13 trials)	⊕⊕⊕⊙ MODERATE ^a
Feed intolerance before hospital dis- charge	282 per 1000	333 per 1000 (268 to 412)	RR 1.18 (0.95 to 1.46	719 (9 trials)	⊕⊕⊝⊝ LOWa, b
Invasive infection before hospital dis- charge	170 per 1000	194 per 1000 (168 to 223)	RR 1.14 (0.99 to 1.31)	3583 (11 trials)	⊕⊕⊝⊝ LOWa, b

*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

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.

(Review)

4



BACKGROUND

Description of the condition

Necrotising enterocolitis (NEC), a syndrome of acute intestinal necrosis of unknown aetiology, affects about 5% of very preterm (< 32 weeks) or very low birth weight (VLBW) (< 1500 grams) infants (Horbar 2012). Infants who develop NEC experience more infections, have lower levels of nutrient intake, grow more slowly, and have longer durations of intensive care and hospital stay than gestation-comparable infants who do not develop NEC (Battersby 2018; Berrington 2012). The associated mortality rate is greater than 20%, and infants who develop NEC, especially if associated with bloodstream infections, have a higher risk of neurodevelopmental problems and disability compared with their peers (Hickey 2018; Martin 2010; Shah 2012).

In addition to low gestational age at birth, the other major perinatal risk factor for developing NEC is intrauterine growth restriction, especially if it is associated with absent or reversed end-diastolic flow velocities in Doppler studies of the foetal aorta or umbilical artery (Bernstein 2000; Dorling 2005; Garite 2004; Luig 2005; Samuels 2017). Most very preterm or VLBW infants who develop NEC have received enteral milk feeds. Feeding with human milk rather than cow's milk formula reduces the risk of NEC (Quigley 2019). Other differences in enteral feeding regimens, such as the timing of introduction of feeds and the size of daily volume increments, may also contribute to inter-unit variation in the incidence of NEC (Walsh 2019). Observational studies have suggested that delaying the introduction of enteral feeds beyond the first few days after birth, or increasing the volume of feeds by less than about 24 mL/kg body weight each day, is associated with a lower risk of developing NEC in very preterm or VLBW infants (Henderson 2009; Patole 2005).

Description of the intervention

Approaches to early enteral feeding vary by gestational age and clinical condition of the very preterm or VLBW infant (Hay 2018). Oral feeding is not usually possible because of neurological immaturity or respiratory distress challenging breathing, suck, and swallow co-ordination. Newborn infants, particularly extremely preterm (< 28 weeks) or extremely low birth weight (ELBW: < 1000 grams) infants, may not be able to tolerate enteral milk feed volumes required to match nutritional requirements because of delayed gastric emptying and immature intestinal peristalsis.

Substantial variation in practice exists with regard to early enteral feeding strategies for very preterm or VLBW infants (Klingenberg 2012). In many high-income countries, policy and practice has tended to favour the conservative approach to introducing and advancing enteral feeds for these infants because of concerns that early full enteral feeding might increase the risk of feed intolerance, gastro-oesophageal reflux and aspiration of stomach contents, and necrotising enterocolitis in very preterm or VLBW infants (de Waard 2018; Leaf 2013; Maas 2018). In low- and middle-income countries with fewer resources for neonatal care, practice is more pragmatic and tends to favour early introduction and advancement of enteral feeds (sometimes facilitated by 'kangaroo' mother care) for stable infants born at 28 weeks' gestation or greater, or with a birth weight of 1000 grams or more (Conde-Agudelo 2016; Sankar 2008).

How the intervention might work

Slow advancement of enteral milk feed volumes aims to reduce the risk of feed intolerance and NEC by limiting the physiological and metabolic stresses on the immature gastrointestinal tract during the weeks after birth. There are, however, potential disadvantages associated with conservative enteral feeding regimens (Flidel-Rimon 2004). Because gastrointestinal hormone secretion and motility are stimulated by milk feeds, slow enteral feed volume advancement may delay the functional adaptation of the gastrointestinal tract and disrupt the patterns of microbial colonisation (Embleton 2017). Intestinal dysmotility and dysbiosis might exacerbate feed intolerance and delay the establishment of enteral feeding independently of parenteral nutrition (Pammi 2017). Prolonging the duration of parenteral nutrition is associated with infectious and metabolic complications that increase mortality and morbidity, prolong hospital stay, and adversely affect growth and development (Embleton 2013). It has been argued that the risk of NEC should not be considered in isolation from these other potential clinical outcomes in the determination of feeding policies and practices for very preterm or VLBW infants (Flidel-Rimon 2006; Härtel 2009).

Why it is important to do this review

Given the potential for the rate of advancement of enteral feed volumes to affect important outcomes for very preterm or VLBW infants, it is important to identify, appraise, and synthesise the available evidence from randomised controlled trials (RCTs) to inform practice and research. This review focuses on the question of whether advancing feed volumes at slow rates compared with faster rates affects the risk of NEC, mortality, and other morbidities. Other Cochrane Reviews have addressed the questions of whether delaying the introduction of any enteral milk feeding or restricting feed volumes to trophic levels (minimal enteral nutrition) affects the risk of NEC in very preterm or VLBW infants (Morgan 2013a; Morgan 2014a; Walsh 2020).

OBJECTIVES

To determine the effects of slow rates of enteral feed advancement on the risk of NEC, mortality, and other morbidities in very preterm or VLBW infants.

METHODS

Criteria for considering studies for this review

Types of studies

We included controlled trials utilising random or quasi-random participant allocation.

Types of participants

We included enterally fed very preterm (< 32 weeks) or VLBW (< 1500 grams) newborn infants.

Types of interventions

Advancement of enteral feeds at no more than 24 mL/kg (birth weight or current body weight) per day versus faster rates of feed advancement. All infants should have received the same type of milk, and in both groups advancement of feed volume should have commenced within five days of introduction of enteral feeds.

Types of outcome measures

Infant- and parent-important outcomes likely to affect survival and quality of life.

Primary outcomes

- NEC confirmed at surgery or autopsy or using standardised clinical and radiological criteria (Walsh 1986):
 - at least one of: bilious gastric aspirate or emesis; or abdominal distention; or blood in stool; and
 - at least one of: abdominal radiograph showing pneumatosis intestinalis; or gas in the portal venous system; or free air in the abdomen.
- All-cause mortality before discharge from hospital.

Secondary outcomes

- Growth:
 - Time to regain birth weight and rates of weight gain, linear growth, head growth, or skinfold thickness growth up to six months (corrected for preterm birth).
 - Long-term growth: weight, height, or head circumference (or proportion of infants who remained below the 10th percentile for the index population's distribution) assessed at intervals from six months of age.
- Neurodevelopmental disability, defined as moderate or severe developmental delay (> two standard deviations (SD) below the mean of standardised infant developmental assessment aged > 18 months), and classifications of disability, including nonambulant cerebral palsy and auditory or visual impairment.
- Time to establish full enteral feeding independently of parenteral nutrition.
- Time to establish oral feeding (independently of parenteral nutrition or enteral tube feeding, or both).
- Feed intolerance (requirement to cease enteral feeds > four hours).
- Invasive infection confirmed by culture of bacteria or fungus from blood, cerebrospinal fluid, or another normally sterile body space.
- Duration of hospital stay (days).

Search methods for identification of studies

We used the standard methods of Cochrane and Cochrane Neonatal.

Electronic searches

We searched the Cochrane Central Register of Controlled Trials (CENTRAL; 2020, Issue 10; 19 October 2020) in the Cochrane Library, Ovid MEDLINE (1946 to October 2020), Embase via Ovid (1974 to 19 October 2020), Maternity and Infant Care database (MIDIRS; 1971 to 19 October 2020), and the Cumulative Index to Nursing and Allied Health Literature (CINAHL; 1982 to 19 October 2020) using the search terms described in Appendix 1. We did not apply language restrictions. We searched ClinicalTrials.gov (clinicaltrials.gov) for ongoing or recently completed trials.

This search updates the searches conducted for previous versions of this review (Appendix 2).

Searching other resources

We searched the reference lists of the included studies to identify any additional relevant articles.

Data collection and analysis

We used the standard methods of Cochrane Neonatal (neonatal.cochrane.org/).

Selection of studies

One review author (WM) screened the titles and abstracts of all records identified by the search, coding the records as 'order' or 'exclude'. A second review author (LY or SO) assessed all records coded as 'order' and made the final decision as to which records should be ordered as full-text articles. Two review authors read the full texts and used a checklist to assess each article's eligibility for inclusion based on the prespecified inclusion and exclusion criteria.

Data extraction and management

Two review authors (WM and LY) independently extracted data using a data collection form to aid extraction of information on design, methods, participants, interventions, outcomes, and treatment effects from each included study. We discussed disagreements until we reached consensus. If data from trial reports were insufficient, we contacted trialists to seek further information.

Assessment of risk of bias in included studies

Two review authors (WM and SO) independently assessed risk of bias (low, high, or unclear) of all the included trials using the Cochrane risk of bias tool (Higgins 2011), for the following domains (Appendix 3).

- Sequence generation (selection bias).
- Allocation concealment (selection bias).
- Blinding of participants and personnel (performance bias).
- Blinding of outcome assessment (detection bias).
- Incomplete outcome data (attrition bias).
- Selective reporting (reporting bias).
- Any other bias.

We resolved disagreements by discussion or by consultation with a third review author.

Measures of treatment effect

We calculated risk ratio (RR) and risk difference (RD) for dichotomous data, and mean difference (MD) for continuous data, with respective 95% confidence intervals (CIs). When we deemed it appropriate to combine two or more study arms, we obtained treatment effects from combined data using the methods described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2021). We determined the number needed to treat for an additional beneficial outcome (NNTB) or the number needed to treat for an additional harmful outcome (NNTH) for a statistically significant difference in RD.

Unit of analysis issues

The unit of analysis was the participating infant in individually randomised trials. Had we identified any cluster-randomised trials, we would have undertaken analyses at the level of the individual



whilst accounting for clustering in the data using the methods recommended in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2021).

Dealing with missing data

We requested additional data from trial investigators when data on important outcomes were missing or required clarification. When data remained missing, we examined the impact on effect size estimates by performing sensitivity analyses.

Assessment of heterogeneity

We examined the treatment effects in individual trials and heterogeneity between trial results by inspecting forest plots if more than one trial was included in a meta-analysis. We calculated the l^2 statistic for each analysis to quantify inconsistency across studies and to describe the percentage of variability in effect estimates that may be due to heterogeneity rather than to sampling error. If we detected moderate or high levels of heterogeneity ($l^2 > 50\%$), we explored possible causes by performing subgroup analyses.

Assessment of reporting biases

We assessed reporting bias by comparing the stated primary outcomes and secondary outcomes with the reported outcomes. Where study protocols were available, we compared these to the full publications to determine the likelihood of reporting bias. We documented studies using the interventions in a potentially eligible infant population but not reporting on any of our primary and secondary outcomes in the Characteristics of included studies tables. We used funnel plots to screen for publication bias where there were a sufficient number of studies (> 10) reporting the same outcome. If publication bias was suggested by a significant asymmetry of the funnel plot on visual assessment, we incorporated this into our assessment of the certainty of the evidence.

Data synthesis

We used a fixed-effect model inverse-variance meta-analysis for combining data where trials examined the same intervention, and the populations and methods of the trials were judged to be similar.

Subgroup analysis and investigation of heterogeneity

We prespecified subgroup analyses for our primary outcomes to compare the effects in trials in which most infants were exclusively formula-fed versus trials in which most infants were at least partially fed with human milk.

Sensitivity analysis

We undertook sensitivity analyses to explore effects on primary outcomes in:

- trials in which most participants were ELBW or extremely preterm;
- trials in which participants were infants with intrauterine growth restriction;
- infants with absent or reversed end-diastolic flow velocities (AREDFV) detected on antenatal Doppler studies of the foetal aorta or umbilical artery.

We planned to perform sensitivity analyses if:

- there was unexplained moderate to high heterogeneity (explored by removing the outlying trial or trials);
- a trial with high risk of bias (including high level of missing outcome data) was included in the meta-analysis of an outcome where the other studies had low risk of bias (removed the study with high risk of bias).

Summary of findings and assessment of the certainty of the evidence

We used the GRADE approach, as outlined in the GRADE Handbook (Schünemann 2013), to assess the certainty of evidence for the following outcomes: NEC, mortality, feed intolerance (causing interruption of enteral feeding), invasive infection.

Two review authors (WM and LY) independently assessed the certainty of the evidence for each of the outcomes above. We considered evidence from RCTs as high certainty, downgrading the evidence one level for serious (or two levels for very serious) limitations based upon the following: design (risk of bias), consistency across studies, directness of the evidence, precision of estimates, and presence of publication bias (Walsh 2021). We used GRADEpro GDT to create Summary of findings 1 to report the certainty of the evidence (GRADEpro GDT).

The GRADE approach results in an assessment of the certainty of a body of evidence as one of four grades.

- **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.

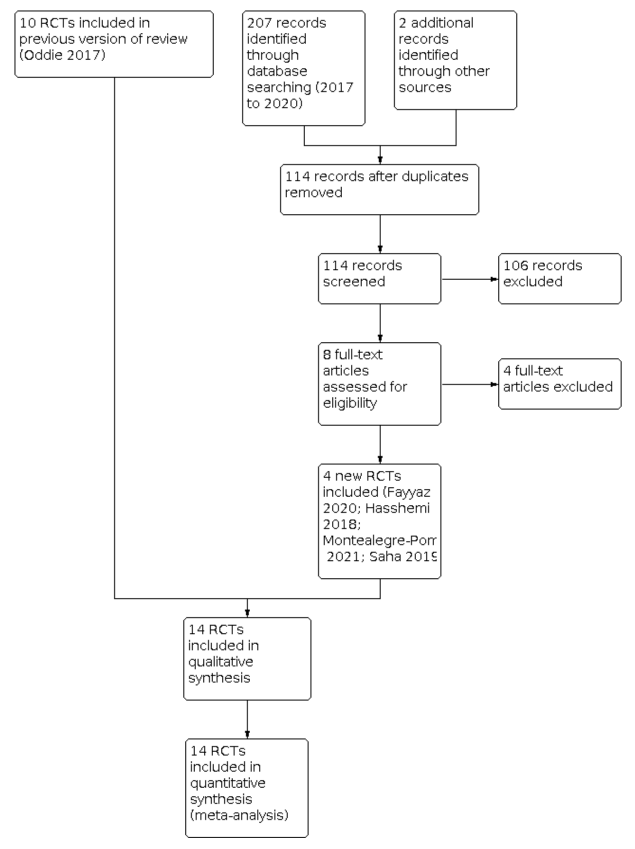
RESULTS

Description of studies

See Figure 1.



Figure 1. Study flow diagram: review update.



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Results of the search

After removal of duplicates, we screened 114 titles and abstracts, which included forward and backward citation searches, clinical trials registers, and grey literature. We re-evaluated previously included studies alongside eight articles sourced as full-text reports, therefore assessing the eligibility of 18 articles.

Included studies

We included 14 RCTs in this updated review (see Characteristics of included studies) (Caple 2004; Fayyaz 2020; Hasshemi 2018; Jain 2016; Karagol 2013; Krishnamurthy 2010; Modi 2015; Montealegre-Pomar 2021; Raban 2014a; Raban 2014b; Rayyis 1999; Saha 2019; Salhotra 2004; SIFT 2016).

Population

A total of 4033 infants participated in the 14 included trials. Almost 70% of infants were participants in a recent large, multicentre trial (SIFT 2016). Trials were undertaken at neonatal care centres in North America (Caple 2004; Rayyis 1999), Colombia (Montealegre-Pomar 2021), Bangladesh (Saha 2019), Iran (Hasshemi 2018); India (Fayyaz 2020; Jain 2016; Krishnamurthy 2010; Modi 2015; Salhotra 2004), Turkey (Karagol 2013), South Africa (Raban 2014a; Raban 2014b), and the UK and Ireland (SIFT 2016).

All but one of the trials specified participant birth weight eligibility criteria (Hasshemi 2018 specified gestational age at birth criteria).

- Caple 2004: 1000 to 2000 grams
- Fayyaz 2020: < 1500 grams
- Hasshemi 2018: 30 to 34 weeks
- Jain 2016: 1000 to 1249 grams
- Karagol 2013: 750 to 1250 grams
- Krishnamurthy 2010: 1000 to 1500 grams
- Modi 2015: 750 to 1250 grams
- Montealegre-Pomar 2021: < 1500 grams
- Raban 2014a: < 1001 grams
- Raban 2014b: < 1001 grams
- Rayyis 1999: < 1500 grams
- Saha 2019: < 1500 grams
- Salhotra 2004: < 1250 grams
- SIFT 2016: < 1500 grams

Most of the participants in Caple 2004, Hasshemi 2018, and Jain 2016 were of birth weight less than 1500 grams or gestational age less than 32 weeks, therefore we made a consensus decision to include these trials. Infants born 'small for gestational age' (birth weight < 10th percentile of the index population distribution) were not eligible to participate in Caple 2004 or Saha 2019, but were included in the other trials. More than 95% of participants in Salhotra 2004 were small for gestational age, and all participants in Jain 2016 had antenatal evidence of absent or reversed end-diastolic flow.

Interventions and comparisons

All trials commenced interval bolus intragastric feeding within the first seven days after birth. Infants were randomly allocated to one of two rates of daily increments in enteral feed volume, as follows.

- Caple 2004: 20 versus 35 mL/kg
- Fayyaz 2020: 15 to 20 versus 25 to 30 mL/kg
- Hasshemi 2018: 20 to 24 versus 30 mL/kg
- Jain 2016: 20 versus 30 mL/kg
- Krishnamurthy 2010: 20 versus 30 mL/kg
- Karagol 2013: 20 versus 30 mL/kg
- Modi 2015: 15 to 20 versus 30 to 40 mL/kg
- Montealegre-Pomar 2021: 20 versus 30 mL/kg
- Raban 2014a: 24 versus 36 mL/kg
- Raban 2014b: 24 versus 36 mL/kg
- Rayyis 1999: 15 versus 35 mL/kg
- Saha 2019: 20 versus 30 mL/kg
- Salhotra 2004: 15 versus 30 mL/kg
- SIFT 2016: 18 versus 30 mL/kg

In one trial, only formula-fed infants were eligible to participate (Rayyis 1999). Infants in eight trials received expressed breast milk or formula, or a combination of both (Caple 2004; Fayyaz 2020; Jain 2016; Karagol 2013; Krishnamurthy 2010; Modi 2015; Montealegre-Pomar 2021; SIFT 2016). In five trials, participating infants were fed exclusively with expressed human milk (Hasshemi 2018; Raban 2014a; Raban 2014b; Saha 2019; Salhotra 2004). Most trial protocols specified indications for interrupting or ceasing enteral feeding, such as residual gastric contents of more than about one-third of the previous feed volume, frequent vomiting, abdominal distension, or detection of blood in the stools (including occult blood). SIFT 2016 did not prespecify these criteria but allowed clinicians and caregivers to apply unit-specific policies and practices.

Outcomes

All 14 included trials reported NEC confirmed radiologically or at surgery or at autopsy. Other reported outcomes included all-cause mortality, time to regain birth weight, time to establish full enteral feeding, duration of hospital stay, and rates of invasive infection.

Excluded studies

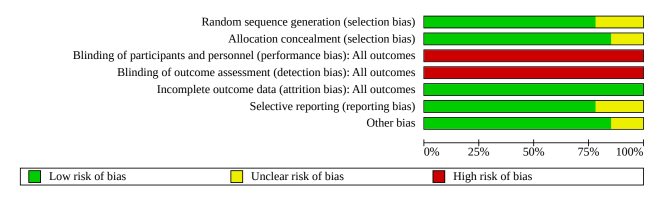
We excluded four reports after full-text review, resulting in a total fo 10 excluded reports (Ahmed 2020; Berseth 2003; Book 1976; Gray 2017; Ibrahim 2017; Jayaraman 2017; Kadam 2016; Nangia 2019; Tewari 2018; Viswanathan 2017; see Characteristics of excluded studies).

Risk of bias in included studies

The methodological quality of the included trials was generally high, but the nature of the intervention meant that parents, caregivers, or clinical investigators were aware of each infant's allocated feeding group (Figure 2).



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



Allocation

Trials employed adequate methods to generate random sequences (typically computer-generated) and to ensure adequate allocation concealment (typically using sealed, opaque envelopes). We assessed no trial as at high risk of selection bias.

Blinding

None of the included trials was able to mask feeding strategies from parents, caregivers, or clinical investigators (although there was some masked assessment of abdominal radiographs for diagnosis of NEC). We assessed all of the included trials as at high risk of performance and detection bias.

Incomplete outcome data

All trials reported complete or near-complete assessments of primary outcomes (low risk of attrition bias).

Selective reporting

Although trial protocols were not available for most trials, we did not consider selective reporting bias to be a major threat given that all relevant clinical outcomes were reported.

Other potential sources of bias

We did not find evidence of important between-group baseline differences in participant characteristics or demographics in any of the included trials.

Effects of interventions

See: **Summary of findings 1** Slow compared with faster rates of enteral feed advancement for preventing necrotising enterocolitis in very preterm or very low birth weight infants

See Summary of findings 1.

Primary outcomes

Necrotising enterocolitis

Meta-analysis of data from 14 trials (4026 infants) showed that slow advancement probably does not reduce the risk of NEC (risk ratio (RR) 1.06, 95% confidence interval (CI) 0.83 to 1.37; risk difference (RD) 0.00, 95% CI –0.01, to 0.02; Analysis 1.1). The funnel plot was not markedly asymmetrical. We assessed the certainty of evidence as moderate using GRADE methods, downgrading for serious study design limitations (lack of masking).

One trial (185 infants) restricted participation to exclusively formula-fed infants (Rayyis 1999). In the other 13 trials (3841 infants), participating infants were at least partially fed with human milk. Subgroup analysis did not show evidence of differences in effect (Figure 3).

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Figure 3. Forest plot of comparison: 1 Slow versus faster rates of feed advancement, outcome: 1.1 Necrotising enterocolitis - all infants.

	Slow	rate	Fast	rate		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.1.1 Infants at least partiall	ly fed with hu	man milk					
Caple 2004	2	84	4	74	3.9%	-0.03 [-0.09 , 0.03]	-
Fayyaz 2020	1	16	0	15	0.8%	0.06 [-0.10 , 0.22]	_ _
Hasshemi 2018	2	100	0	40	2.9%	0.02 [-0.03 , 0.07]	+
Jain 2016	1	15	2	15	0.8%	-0.07 [-0.28 , 0.15]	
Karagol 2013	5	46	4	46	2.3%	0.02 [-0.10 , 0.14]	
Krishnamurthy 2010	1	50	2	50	2.5%	-0.02 [-0.09 , 0.05]	4
Modi 2015	2	65	1	66	3.3%	0.02 [-0.04 , 0.07]	-
Montealegre - Pomar 2021	0	12	2	8	0.5%	-0.25 [-0.56 , 0.06]	
Raban 2014a	1	51	7	47	2.4%	-0.13 [-0.24 , -0.02]	
Raban 2014b	9	52	2	50	2.6%	0.13 [0.02 , 0.25]	
Saha 2019	3	55	2	38	2.3%	0.00 [-0.09 , 0.09]	<u> </u>
Salhotra 2004	0	26	2	27	1.3%	-0.07 [-0.19 , 0.04]	
SIFT 2016	78	1399	70	1394	69.9%	0.01 [-0.01 , 0.02]	•
Subtotal (95% CI)		1971		1870	95.4%	0.00 [-0.01 , 0.02]	T
Total events:	105		98				
Heterogeneity: Chi ² = 18.24, o	df = 12 (P = 0.	11); I ² = 3	4%				
Test for overall effect: $Z = 0.2$	26 (P = 0.79)						
1.1.2 Infants exclusively for	mula fed						
Rayyis 1999	13	98	8	87	4.6%	0.04 [-0.05, 0.13]	
Subtotal (95% CI)		98		87	4.6%		
Total events:	13		8				
Heterogeneity: Not applicable	2						
Test for overall effect: $Z = 0.8$							
Total (95% CI)		2069		1957	100.0%	0.00 [-0.01 , 0.02]	
Total events:	118		106			-	
Heterogeneity: Chi ² = 18.86, o	df = 13 (P = 0.	13); I ² = 3	1%				
Test for overall effect: $Z = 0.5$						I	Favours slow rates Favours faster r
Test for overall effect: $Z = 0.5$. ,	1 1 (D	0 41) 12	00/		1	ravours slow rates ravours rast

Test for subgroup differences: $Chi^2 = 0.69$, df = 1 (P = 0.41), $I^2 = 0\%$

Sensitivity analyses did not show an effect in:

- extremely preterm or ELBW infants: RR 1.01 (95% CI 0.74 to 1.38); five trials, 1299 infants (Analysis 1.2);
- infants small for gestation or with intrauterine growth restriction: RR 1.26 (95% CI 0.67 to 2.37); two trials, 639 infants (Analysis 1.3);
- infants with evidence of AREDFV: RR 1.59 (95% CI 0.74 to 3.40); two trials, 465 infants (Analysis 1.4).

Mortality

Meta-analysis of data from 13 trials (3860 infants) showed that slow advancement probably does not reduce the risk of mortality (RR

1.13, 95% CI 0.91 to 1.39; RD 0.01, 95% CI -0.01 to 0.02; Analysis 1.5). The funnel plot was not markedly asymmetrical. We assessed the certainty of evidence as moderate, downgrading for serious study design limitations (lack of masking).

One trial (185 infants) restricted participation to exclusively formula-fed infants (Rayyis 1999). In the other 12 trials (3644 infants), participating infants were at least partially fed with human milk. Subgroup analysis did not show evidence of differences in effect (Figure 4).

Figure 4. Forest plot of comparison: 1 Slow versus faster rates of feed advancement, outcome: 1.5 Mortality - all infants.

	Slow	rate	Fast	rate		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.5.1 Infants at least partiall	ly fed with hu	man milk					
Salhotra 2004	12	26	7	27	1.4%	0.20 [-0.05 , 0.46]	I <u> </u>
Krishnamurthy 2010	6	50	4	50	2.6%	0.04 [-0.08 , 0.16]	I
Karagol 2013	4	46	3	46	2.4%	0.02 [-0.09 , 0.13]	I <u>+</u>
Raban 2014b	16	52	14	50	2.7%	0.03 [-0.15 , 0.20]	I
Raban 2014a	13	51	19	47	2.6%	-0.15 [-0.33 , 0.04]	I
Modi 2015	28	65	20	66	3.4%	0.13 [-0.04 , 0.29]	Ⅰ
Jain 2016	3	15	0	15	0.8%	0.20 [-0.02 , 0.42]	I <u> </u>
SIFT 2016	65	1393	60	1392	72.7%	0.00 [-0.01 , 0.02]	I n
Hasshemi 2018	0	100	0	40	3.0%	0.00 [-0.04 , 0.04]	ı T
Saha 2019	7	55	4	38	2.3%	0.02 [-0.11 , 0.15]	I <u>+</u>
Fayyaz 2020	0	16	1	15	0.8%	-0.07 [-0.23 , 0.10]	I
Montealegre-Pomar 2021	0	12	1	8	0.5%	-0.13 [-0.39 , 0.14]	I
Subtotal (95% CI)		1881		1794	95.2%	0.01 [-0.01 , 0.03]	
Total events:	154		133				
Heterogeneity: Chi ² = 13.04, o	df = 11 (P = 0.	29); I ² = 1	6%				
Test for overall effect: $Z = 1.1$	4 (P = 0.25)						
1.5.2 Infants exclusively for	mula fed						
Rayyis 1999	0	98	0	87	4.8%	0.00 [-0.02 , 0.02]	
Subtotal (95% CI)		98		87	4.8%	0.00 [-0.02 , 0.02]	
Total events:	0		0				ľ
Heterogeneity: Not applicable	<u>.</u>						
Test for overall effect: $Z = 0.0$	00 (P = 1.00)						
Total (95% CI)		1979		1881	100.0%	0.01 [-0.01 , 0.02]	
Total events:	154		133			-	ľ
Heterogeneity: Chi ² = 13.68, o	df = 12 (P = 0.	32); I ² = 1	2%				-1 -0.5 0 0.5
Test for overall effect: $Z = 1.1$	4 (P = 0.25)	-					Favours slow rates Favours faster
Test for subgroup differences:	. ,	lf = 1 (P =	0.48), $I^2 =$	0%			

Sensitivity analyses did not show an effect in:

- extremely preterm or ELBW infants: RR 0.83 (95% CI 0.55 to 1.25); two trials, 200 infants (Analysis 1.6);*
- infants small for gestation or with intrauterine growth restriction: RR 1.78 (95% CI 0.83 to 3.81); one trial (Salhotra 2004), 53 infants (Analysis 1.7);*
- infants with evidence of AREDFV: RR 7.00 (95% CI 0.39 to 124.83); one trial (Jain 2016), 30 infants (Analysis 1.8).*

(*Subgroup data not available for SIFT 2016.)

Secondary outcomes

Growth

Ten trials reported that infants in the slow-rate-of-advancement group took a longer time to regain birth weight (meta-analysis not possible).

- Caple 2004: mean difference (MD) 2 days (95% Cl 1 to 3)
- Hasshemi 2018: data not available
- Karagol 2013: MD 3.8 days (CI not given)
- Krishnamurthy 2010: median difference 6 days
- Montealegre-Pomar 2021: median difference 0.5 days

- Raban 2014a: data not available
- Raban 2014b: data not available
- Rayyis 1999: median difference 2 days
- Saha 2019: MD 4.8 days (CI not given)
- Salhotra 2004: median difference 5 days

Three trials did not report growth parameters (Fayyaz 2020; Jain 2016; Modi 2015).

SIFT 2016 did not show effects on weight (MD 0.00, 95% CI –0.08 to 0.08; 2602 infants; Analysis 1.9) or head circumference (MD 0.00, 95% CI –0.13 to 0.13; 2286 infants; Analysis 1.10), z-scores at hospital discharge.

None of the included trials reported post-hospital discharge growth parameters.

Neurodevelopment

One trial reported neurodevelopmental outcomes assessed in children aged 18 to 24 months (SIFT 2016). Analysis suggested that slow advancement of enteral feed volumes probably does not affect the risk of moderate or severe disability (motor, visual, or hearing impairment, or cognitive or language delay) (RR 0.90, 95% CI 0.79 to 1.02; 2325 infants; Analysis 1.11). Analyses of individual domains

suggest little or no effect on visual or hearing impairment, or cognitive or language delay, but suggest that slow advancement of enteral feed volumes may reduce the risk of cerebral palsy slightly:

- cerebral palsy (clinician diagnosed): RR 0.60, 95% CI 0.39 to 0.90; 2183 infants; RD -0.02, 95% CI -0.04 to -0.00 (Analysis 1.12);
- visual impairment: RR 0.75, 95% CI 0.39 to 1.43; 2327 infants (Analysis 1.13);
- hearing impairment: RR 0.69, 95% CI 0.47 to 1.02; 2315 infants (Analysis 1.14);
- cognitive or language delay (assessed using the Parent Report of Children's Abilities – Revised): RR 0.93, 95% CI 0.81 to 1.07; 2326 infants (Analysis 1.15).

Time to establish full enteral feeding

Eleven trials reported that it took longer to establish full enteral feeds in infants in the slow-rate-of-advancement group (meta-analysis not possible).

- Caple 2004: MD 3 days (95% CI 2 to 3)
- Hasshemi 2018: data not available
- Jain 2016: MD 0.6 days (CI not given)
- Karagol 2013: MD 3.2 days (CI not given)
- Krishnamurthy 2010: median difference 2 days
- Modi 2015: MD 4 days (Cl not given)
- Montealegre-Pomar 2021: median difference 1.5 days
- Rayyis 1999: median difference 4 days
- Saha 2019: MD 5.3 days (CI not given)
- Salhotra 2004: MD 4.8 days (CI not given)
- SIFT 2016: median difference 3 days

Three trials did not report this outcome (Fayyaz 2020; Raban 2014a; Raban 2014b).

Time to establish oral feeding

None of the trials reported time to establish full oral feeding.

Feed intolerance

Meta-analysis of data from nine trials (719 infants) showed that slow advancement of enteral feed volumes may slightly increase the risk of feed intolerance (RR 1.18, 95% CI 0.95 to 1.46; RD 0.05, 95% CI –0.02 to 0.12) (Analysis 1.16). We assessed the certainty of evidence as low, downgrading for serious study design limitations (lack of masking) and imprecision.

Invasive infection

Meta-analysis of data from 11 trials (3583 infants) showed that slow advancement of enteral feed volumes may slightly increase the risk of invasive infection (RR 1.14, 95% CI 0.99 to 1.31; RD 0.02, 95% CI –0.00 to 0.05) (Analysis 1.17). We assessed the certainty of evidence as low, downgrading for serious study design limitations and imprecision.

Duration of hospital stay

Six trials reported no difference in duration of hospital stay between groups.

• Caple 2004: MD 5 days (95% CI -1 to 8)

- Hasshemi 2018: data not available
- Raban 2014a: data not available
- Raban 2014b: data not available
- Rayyis 1999: median difference 4 days
- SIFT 2016: median difference 0 days (54 versus 54 days)

Four trials reported that duration of hospital stay was longer amongst infants in the slow-rate-of-advancement group.

- Fayyaz 2020: MD 4 days (CI not given)
- Karagol 2013: MD 6 days (CI not given)
- Krishnamurthy 2010: median difference 1.5 days
- Saha 2019: MD 4 days (Cl not given)

Meta-analysis was not feasible.

The remaining four trials did not report on duration of hospital stay (Jain 2016; Modi 2015; Montealegre-Pomar 2021; Salhotra 2004).

DISCUSSION

Summary of main results

The trial data included in this review provide moderate-certainty evidence that advancing enteral feed volumes at slow rates (up to 24 mL/kg/d) compared with faster rates probably does not reduce the risk of NEC in very preterm or VLBW infants. The boundaries of the 95% CI for the estimate of effect are consistent with either two extra or one fewer cases of NEC in every 100 infants who have slow rates of feed advancement. Meta-analysis of data from these trials did not show evidence of an effect on all-cause mortality, and prespecified analyses did not show effects on risk of NEC or death amongst ELBW or extremely preterm infants, nor amongst infants with growth restriction or evidence of absent or reversed enddiastolic flow velocity. Meta-analysis of data from 11 trials showed a slightly higher risk of late-onset infection amongst infants who had slow advancement of enteral feeds. The point estimate suggests that an extra episode of late-onset infection occurs for every 50 infants who have slow rather than faster feed volume advancement, but the lower bound of the 95% CI was consistent with no effect.

Although meta-analysis suggests that slow advancement of enteral feeds may slightly increase feed intolerance, and infants who had slow advancement established full enteral feeding and regained birth weight several days later than infants who had faster rates of advancement, there was no evidence of an effect on duration of hospital stay. The clinical importance of these effects is unclear, since long-term growth outcomes were not assessed.

One large trial has now reported neurodevelopmental outcomes in participants assessed at about two years post-term (SIFT 2016). Analysis did not show an effect on the risk of moderate or severe neurodevelopmental disability. Similarly, there was no evidence of an effect on visual or hearing impairment, or on cognitive or language delay. Slow advancement of feed volumes was associated with a lower risk of cerebral palsy (clinician diagnosed). The 95% CI for the estimate was consistent with either no effect or with four fewer cases of cerebral palsy in every 100 infants who have slow rather than faster feed volume advancement. This estimate should be interpreted with caution due to possible attrition bias: complete assessment was possible for less that 80% of trial participants (similar across study groups).

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



Overall completeness and applicability of evidence

Most participants in the included trials were stable very preterm or VLBW infants of birth weight appropriate for gestational age. About one-third of all participants were extremely preterm or ELBW, and about one-fifth were small for gestational age, growth-restricted, or compromised in utero, as indicated by absent or reversed enddiastolic flow velocity in the umbilical artery. Infants who had severe respiratory distress requiring oxygen supplementation or ventilatory support were eligible to participate in all but three of the trials (Karagol 2013; Krishnamurthy 2010; Salhotra 2004). The findings of this review should therefore be applicable across these populations, who are at highest risk of developing feed intolerance or NEC (Luig 2005).

Most participating infants were at least partially fed with breast milk. Evidence indicates that artificial formula feeding increases risks of feed intolerance and NEC (Quigley 2019). The risk-benefit balance of enteral feeding strategies may differ between human milk-fed and formula-fed very preterm or VLBW infants, but available data were insufficient to show effects of different rates of feed advancement on important outcomes for infants fed exclusively with artificial formula. It is also unclear whether the review findings can be applied to infants who receive continuous infusion of intragastric feeds, as a vast majority of the infants in the included trials received enteral feeds as interval boluses. Randomised controlled trials have reported conflicting findings about the effect of continuous enteral infusion on feed tolerance in very preterm or VLBW infants (Premji 2011).

Although the finding that slow enteral feed volume advancement delays establishment of full enteral feeds may seem intuitive, it is plausible that advancing feed volumes faster could have resulted in more feed intolerance and therefore a delay in the establishment of full enteral feeding. The included trials prespecified definitions of feed intolerance that mandated interrupting or ceasing feed volume advancement, principally detection of pre-feed 'gastric residuals' (gastric content aspirated before a planned gastric tube feed) and abdominal distension. However, trial reports presented only limited data on the frequency of these outcomes. Furthermore, limited evidence suggests that the volume or colour of gastric residuals is predictive of risk of NEC for infants whose feed volumes are advanced conservatively (Bertino 2009; Cobb 2004; Mihatsch 2002). Similarly, the clinical importance of abdominal distension or bowel loops visible through the abdominal wall (without other features of intra-abdominal pathology) is unclear, especially in the modern era, when early and prolonged use of continuous positive airway pressure results in intestinal gaseous distension.

Quality of the evidence

Our GRADE assessment of the certainty of evidence for primary outcomes was moderate, downgraded from high due to lack of masking in the included trials (Summary of findings 1). Although the included trials were generally of good methodological quality, in common with other trials of feeding interventions in this population, it was not possible to mask parents, caregivers, and clinical assessors to the nature of the intervention (Figure 2). Lack of masking may have resulted in surveillance and ascertainment biases. It is more likely, however, to have caused an overestimation of feed intolerance and NEC amongst infants whose feed volumes were advanced faster. Assessment of abdominal radiographs for signs of NEC was masked in most trials to ensure that the diagnosis of severe NEC (confirmed by radiological detection of gas in the bowel wall or portal tract) was not prone to bias. However, as microbial generation of gas in the bowel wall is substratedependent, infants who received more enteral milk (substrate) may have been more likely to demonstrate this radiological sign than infants with equally severe bowel disease who had less intraluminal substrate. This 'substrate effect' is also more likely to cause overascertainment of NEC amongst infants who had faster rates of feed volume advancement (Tyson 2007).

Potential biases in the review process

The main concern with the review process is the possibility that findings are subject to publication and other reporting biases. We attempted to minimise this threat by screening the reference lists of included trials and related reviews and searching the proceedings of major international perinatal conferences to identify trial reports that were not published in full form in academic journals. Inspection of funnel plots of meta-analyses that included at least 10 data points did not show sufficient asymmetry to raise concerns about possible publication or small-study bias.

Agreements and disagreements with other studies or reviews

This review focused specifically on the comparison of slow versus faster rates of feed volume advancement. Other Cochrane Reviews have assessed how (i) enteral fasting with trophic feeding (minimal enteral nutrition), (ii) delayed versus early introduction of progressive enteral feeds, and (iii) early full enteral feeding versus gradual introduction of feeds affects important outcomes in very preterm or VLBW infants (Morgan 2013a; Morgan 2014a; Walsh 2020). These reviews, consistent with the findings of this review, found evidence that conservative feeding strategies probably do not reduce the risk of NEC, mortality, or associated morbidity.

AUTHORS' CONCLUSIONS

Implications for practice

Advancing enteral feed volumes at slow rates (up to 24 mL/kg/ d) probably does not reduce the risk of necrotising enterocolitis, death, or feed intolerance in very preterm or very low birth weight infants (moderate-certainty evidence), including extremely preterm or extremely low birth weight infants, or in infants who are growth-restricted or compromised in utero. Advancing the volume of enteral feeds at faster rates (typically 30 to 40 mL/kg/ d) shortens by several days the time taken to regain birth weight and establish full enteral feeds, and may slightly reduce the risk of late-onset invasive infection (low-certainty evidence). Clinicians, policymakers, and guideline-producers can consider how to apply this evidence to practice in their local context (Soll 2019).

Implications for research

Additional trials are unlikely to alter these effect estimates for necrotising enterocolitis or death. With regard to very preterm or very low birth weight infants who are clinically stable after birth (typically infants with birth weight more than 1000 grams or gestational age more than 27 weeks), the key research question is now whether exclusive enteral feeding from birth is better than slow introduction and advancement (Walsh 2020).



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CHARACTERISTICS OF STUDIES

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* Indicates the major publication for the study

Study characteristics	
Methods	Randomised controlled trial
Participants	Preterm infants of birth weight 1000 to 2000 grams (appropriate birth weight for gestational age) and o gestational age < 35 weeks at birth, who were starting formula feeds
	Setting: Neonatal Unit, Department of Pediatrics, University of Texas Medical School, Houston, Texas, USA (1999 to 2002)
Interventions	Feed (formula only) advancement at 20 mL/kg/d (n = 84) vs 30 mL/kg/d (n = 74)
Outcomes	 NEC Time to regain birth weight Time to achieve full enteral feeds Time to hospital discharge

Caple 2004 (Continued)

Notes

Feeds were ceased if the residual gastric aspirate was more than one-third of the previous feed volume, or if frequent vomiting, abdominal distention, or bloody stools (including occult blood) were noted.

We were unable to obtain data on all-cause mortality from the principal investigators.

Risk of bias Bias **Authors' judgement** Support for judgement Random sequence genera-Low risk Random number sequence tion (selection bias) Allocation concealment Low risk Blinded draw from envelope by caregivers not involved in the study (selection bias) Blinding of participants High risk Unmasked and personnel (performance bias) All outcomes Blinding of outcome as-High risk Unmasked (radiologists may have been masked) sessment (detection bias) All outcomes Incomplete outcome data Low risk 3 infants excluded after enrolment because of protocol violations were includ-(attrition bias) ed in this review and meta-analysis. 2 infants (1 in each group) were excluded All outcomes because they were determined ineligible for enrolment as the result of an in utero gastrointestinal perforation and foetal alcohol syndrome; these infants were not included in the meta-analysis. Unclear risk Selective reporting (re-Mortality not reported. porting bias) No evidence of baseline imbalance Other bias Low risk

Fayyaz 2020

Randomised controlled trial				
Preterm infants of birth weight < 1500 grams (gestational age 30 to 36 weeks)				
Setting: Neonatal Unit, Izzat Ali Shah Hospital, Wah Cantt, India (2019 to 2020)				
Feed advancement at 15 to 20 mL/kg/d (n = 16) vs 25 to 30 mL/kg/d (n = 15)				
Time to hospital discharge				
Information on randomisation methods and data on NEC and infection courtesy of Dr Shahzad Haider (March 2021)				
Authors' judgement Support for judgement				



Fayyaz 2020 (Continued)

Random sequence genera- tion (selection bias)	Low risk	Random numbers table
Allocation concealment (selection bias)	Low risk	Sealed, opaque envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	Complete follow-up for primary outcomes
Selective reporting (re- porting bias)	Unclear risk	Mortality, NEC, and other clinical outcomes not reported.
Other bias	Unclear risk	Baseline data not reported.

Hasshemi 2018

Study characteristics			
Methods	Randomised controlled trial		
Participants	Preterm infants of birth gestational age 30 to 34 weeks		
	Setting: Neonatal Unit,	, Tabriz Alzahra Teaching Hospital, Tabriz, Iran (2011 to 2012)	
Interventions	Feed advancement at ((i) 20 mL/kg/d (n = 60) vs (ii) 24 mL/kg/d (n = 40) vs (iii) 30 mL/kg/d (n = 40)	
	(groups (i) and (ii) com	bined for analysis in this review)	
Outcomes	 NEC Mortality Invasive infection (culture-confirmed) 		
Notes	Information on randomisation methods and data on NEC, mortality, and infection courtesy of Dr Man- izheh Mostafa Gharehbaghhi (April 2021)		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated sequence	
Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes	



Hasshemi 2018 (Continued)

Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	No losses to follow-up
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Jain 2016

Study characteristics				
Methods	Randomised controlled trial			
Participants		Preterm infants (birth weight 1000 to 1249 grams and gestational age > 30 weeks at birth) who have an- tenatal evidence of absent end-diastolic flow velocities (presumed in umbilical artery)		
	Setting: Department of garh, India (2013 to 201	f Paediatrics, Postgraduate Institute of Medical Education & Research, Chandi- 14)		
Interventions	Feed advancement at 2	Feed advancement at 20 mL/kg/d (n = 15) vs 30 mL/kg/d (n = 15)		
Outcomes	 NEC Late-onset bloodstream (culture-positive) infection In-hospital mortality Time to achieve full enteral feeds 			
Notes	Prespecified subgroup of a larger trial that enrolled infants with birth weight > 1250 grams and com- pared feed advancement at 30 mL/kg/d vs 40 mL/kg/d			
	Additional data courte	sy of Dr Mukhopadhyay (September 2014)		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Computer-generated		
Allocation concealment (selection bias)	Low risk	Sealed, opaque envelopes		
Blinding of participants and personnel (perfor- mance bias)	High risk	Unmasked		



Jain 2016 (Continued) All outcomes

Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	Complete follow-up for primary outcomes
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Karagol 2013

Study characteristics			
Methods	Randomised controlled trial		
Participants	Preterm infants < 32 weeks' gestation with birth weight of 750 to 1250 grams 32% of infants weighed < 1000 grams.		
	Exclusion criteria included major congenital malformations, severe respiratory distress, presence of umbilical vessel catheters, contraindications to enteral feeding, perinatal asphyxia, and cardiovascula compromise.		
	Setting: Division of Neonatology, Dr Sami Ulus Maternity, Children's Education and Research Hospital, Ankara, Turkey (2011 to 2012)		
Interventions	Slow advancement at 20 mL/kg/d (n = 46) vs rapid advancement at 30 mL/kg/d (n = 46)		
Outcomes	 NEC All-cause mortality Time to regain birth Time to reach full er Feed intolerance Invasive infection (Subgroup analysis) 	nteral feeds	
Notes	Feeds were ceased if any of the following occurred: gastric residuals > 5 mL/kg or > 50% of feed volume, vomiting > 3 times in 24 hours, increase in abdominal girth > 2 cm between feeds, abdominal tenderness or erythema, reduced bowel sounds, blood in the stools, or recurrent apnoea.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated sequence	
Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes	



Karagol 2013 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	No participants lost to follow-up
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Krishnamurthy 2010

Study characteristics			
Methods	Randomised controlled trial		
Participants	Preterm infants (birth v	weight 1000 to 1499 grams) and gestational age < 34 weeks at birth	
	Exclusion criteria inclu cal arterial or venous c	ded respiratory distress, mechanical ventilation, inotrope support, and umbili- atheterisation.	
	Setting: Department of	Paediatrics, University College of Medical Sciences, Delhi, India (2007 to 2009)	
Interventions	Feed advancement at 2	20 mL/kg/d (n = 50) vs 30 mL/kg/d (n = 50)	
Outcomes	• NEC		
	Incidence of invasiv	e infection	
	In-hospital mortality		
	Time to regain birth weight		
	Time to achieve full enteral feeds		
	Time to hospital dis	charge	
Notes	All feeds were delivered	d by gavage via nasogastric tube at 2-hour intervals.	
	volume (delayed if volu	ny of the following occurred: residual gastric contents > 50% of previous feed ume was 25% to 50%), > 3 episodes of apnoea in the preceding hour, abdominal ss, or bloody stools (including occult blood).	
	Parenteral nutrition was not available.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated sequence	

Krishnamurthy 2010 (Continued)

Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Modi 2015

Study characteristics

-			
Methods	Randomised controlled trial		
Participants		pirth weight of 750 to 1250 grams who commenced enteral feeds within 4 days tional age of participants was 31 weeks.	
	Exclusion criteria were tract").	(quote: "gross congenital malformation and anomalies of gastrointestinal	
	Setting: Department of Neonatology, Maulana Azad Medical College, New Delhi, India (2012 to 2014)		
Interventions	Feed advancement at 15 to 20 mL/kg/d (n = 65) vs 30 to 40 mL/kg/d (n = 66)		
Outcomes	 NEC Incidence of feed intolerance Invasive infection In-hospital (all-cause) mortality Mean daily weight gain Time to achieve full enteral feeds 		
Notes	Published as abstract only Further information available from www.ctri.nic.in/Clinicaltrials/pmaindet2.php?trialid=5289		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Unclear risk	Stratified block randomisation (computer-generated)	

tion (selection bias)



Modi 2015 (Continued)

Allocation concealment (selection bias)	Low risk	Sequentially numbered, sealed, opaque envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Montealegre-Pomar 2021

Study characteristics		
Methods	Randomised controlled trial	
Participants	Preterm infants of gestational age < 34 weeks at birth and birth weight 1000 to 1499 grams	
	Setting: Ignacio Univer	rsity Hospital, Bogata, Colombia (2014 to 2015)
Interventions	Feed advancement at 20 mL/kg/d (n = 12) vs 30 mL/kg/d (n = 8)	
Outcomes	 NEC Mortality Invasive infection Time to hospital discharge 	
Notes	Additional information courtesy of Dr Montealegre-Pomar	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Stratified block randomisation (computer-generated)
Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked



Montealegre-Pomar 2021 (Continued)

-		
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked
Incomplete outcome data (attrition bias) All outcomes	Low risk	Near complete (1 study withdrawal postrandomisation)
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Raban 2014a

Study characteristics	
Methods	Randomised controlled trial (2 × 2 factorial design with Raban 2014b)
Participants	ELBW infants
	Setting: Groote Schuur Hospital, Cape Town, South Africa (2011 to 2013)
Interventions	Feed advancement (from 12 mL/kg/d on day 2) in daily increments of 24 mL/kg (n = 51) vs 36 mL/kg (n = 47) until enteral feeds of 200 mL/kg/d were attained
Outcomes	• NEC
	Feed intolerance
	Mortality
	Invasive infection
	Time to attain 1500 grams of weight
	Time to regain birth weight
Notes	Factorial design also randomised to commencing feeds on day 1 (24 mL/kg) or day 2 (12 mL/kg).
	Infants received maternal-expressed breast milk or donor breast milk.
	Trial registration: ISRCTN96923718

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Computer-generated
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked
Blinding of outcome as- sessment (detection bias)	High risk	Unmasked



Raban 2014a (Continued) All outcomes

Incomplete outcome data (attrition bias) All outcomes	Low risk	Complete follow-up for primary outcomes
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Raban 2014b

Study characteristics			
Methods	Randomised controlled trial (2 × 2 factorial design with Raban 2014a)		
Participants	ELBW infants	ELBW infants	
	Setting: Groote Schuur	Hospital, Cape Town, South Africa (2011 to 2013)	
Interventions		om 24 mL/kg/d on day 1) in daily increments of 24 mL/kg (n = 52) vs 36 mL/kg (n = of 200 mL/kg/d were attained	
Outcomes	 NEC Feed intolerance Mortality Invasive infection Time to attain 1500 grams of weight Time to regain birth weight 		
Notes	Factorial design also randomised to commencing feeds on day 1 (24 mL/kg) or day 2 (12 mL/kg). Infants received maternal-expressed breast milk or donor breast milk. Trial registration: ISRCTN96923718		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated	
Allocation concealment (selection bias)	Unclear risk	Not described	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked	
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked	



Raban 2014b (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Complete follow-up for primary outcomes
Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Rayyis 1999

Study characteristics			
Methods	Randomised controlled trial		
Participants	VLBW infants of gestational age < 34 weeks at birth		
	Setting: Neonatal Unit, Department of Pediatrics, University of Alabama, Birmingham, Alabama, USA (dates not stated)		
Interventions	Feed advancement at 15 mL/kg/d (n = 98) vs 35 mL/kg/d (n = 87)		
Outcomes	• NEC		
	Time to regain birth weight		
	Time to achieve full enteral feeds		
	Time to hospital discharge		
Notes	Infants for whom full or partial feeding with expressed breast milk was planned were not eligible to pa ticipate.		
	Feeds were ceased if: residual gastric contents > 30% of previous feed volume, abdominal distension of tenderness, or bloody stools (including occult blood).		

Risk of bias Bias **Authors' judgement** Support for judgement Not stated Random sequence genera-Unclear risk tion (selection bias) Allocation concealment Low risk Opaque, sealed envelopes (selection bias) Blinding of participants Unmasked High risk and personnel (performance bias) All outcomes Unmasked Blinding of outcome as-High risk sessment (detection bias) All outcomes Incomplete outcome data 7 protocol violations occurred after enrolment, but all infants were included in Low risk (attrition bias) the final data analysis. All outcomes



Rayyis 1999 (Continued)

Selective reporting (re- porting bias)	Low risk	Unlikely
Other bias	Low risk	No evidence of baseline imbalance

Saha 2019

Study characteristics				
Methods	Randomised controlled trial			
Participants	n weight 1200 to 1500 grams (appropriate for gestational age)			
	Exclusion criteria included recurrent apnoea, respiratory distress requiring supplemental receipt of inotrope support.			
	Setting: Neonatal Unit, Dhaka Shishu Hospital, Dhaka, Bangladesh (2017 to 2018)			
Interventions	Feed (expressed huma	n milk only) advancement at 20 mL/kg/d (n = 55) vs 30 mL/kg/d (n = 38)		
Outcomes	 NEC Mortality pre-discharge Feed intolerance (interruption) Duration of hospital stay 			
Notes	Further information sought from Dr Saha and colleagues.			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Unclear risk	Not stated		
Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked		
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked		
Incomplete outcome data (attrition bias) All outcomes	Low risk	Complete until hospital discharge		
Selective reporting (re- porting bias)	Unclear risk	Invasive infection not reported.		
Other bias	Unclear risk	No evidence of baseline imbalance		



Salhotra 2004

Study characteristics				
Methods	Randomised controlled trial			
Participants	Preterm infants of birt	h weight < 1250 grams (> 95% of participants were "small for gestational age")		
	Exclusion criteria inclu receipt of inotrope sup	ded recurrent apnoea, respiratory distress requiring supplemental oxygen, and oport.		
	Setting: Neonatal Unit, Maulana Azad Medical College (tertiary-level teaching hospital), New Delhi, In- dia (2001 to 2003)			
Interventions	Feed advancement at 2	15 mL/kg/d (n = 26) vs 30 mL/kg/d (n = 27)		
Outcomes	 NEC Neonatal mortality Time to regain birth weight Time to achieve full enteral feeds Time to hospital discharge 			
otes Feeds were ceased if residual gastric content was > 30% of previous feed volume or if a tension was noted.				
	Mortality data courtesy of Dr Namasivayam Ambalavanan			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Computer-generated sequence		
Allocation concealment (selection bias)	Low risk	Opaque, sealed envelopes		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked		
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked		
Incomplete outcome data (attrition bias) All outcomes	Low risk	No losses to follow-up		
Selective reporting (re- porting bias)	Low risk	Unlikely		
Other bias	Low risk	No evidence of baseline imbalance		



SIFT 2016

Study characteristics				
Methods	Randomised controlled trial			
Participants	Very preterm or VLBW infants			
Interventions	Feeds advancement at	18 mL/kg/d (n = 1404) vs 30 mL/kg/d (n = 1400) (June 2013 to June 2015)		
Outcomes	 NEC Mortality Neurodisability at 24 months post-term Invasive infection Time to full milk feeds Growth (change in z-score - weight and head circumference for gestational age) from birth to discharge home Length of time in intensive care Length of hospital stay 			
Notes	See: Declarations of interest			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Computer-generated		
Allocation concealment (selection bias)	Low risk	Computer-based random allocation		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unmasked		
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unmasked (radiologists may have been masked)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	> 99% outcome data for in-hospital outcomes > 87% outcome data for neurodevelopmental outcomes at 18 to 24 months		
Selective reporting (re- porting bias)	Low risk	Adhered to protocol		
Other bias	Low risk No evidence of baseline imbalance			

ELBW: extremely low birth weight; n: number of infants; NEC: necrotising enterocolitis; VLBW: very low birth weight

Characteristics of excluded studies [ordered by study ID]

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Study	Reason for exclusion			
Ahmed 2020	RCT of early (< 48 h) vs later initiation of progressive enteral feeding in preterm infants			
Berseth 2003	Infants were randomly allocated to a stable (not progressively increased) trophic feeding volume or to feed volume advancement at 20 mL/kg/d.			
Book 1976	Enteral feeding volumes were advanced at 10 mL/kg/d vs 20 mL/kg/d (both groups received 'slow' advancement of feed volumes).			
Gray 2017	RCT of different feeding intervals (not different rates of feed volume advancement) in very preterm infants			
Ibrahim 2017	RCT of different feeding intervals (not different rates of feed volume advancement) in very preterm infants			
Jayaraman 2017	RCT examining the effect on breast milk feeding of early vs delayed kangaroo mother care in low birth weight infants (no intention to advance enteral feed volumes at different rates)			
Kadam 2016	Retrospective cohort study			
Nangia 2019	RCT of early full enteral feeding (not progressive advancement at different rates)			
Tewari 2018	RCT of early vs delayed initiation of progressive enteral feeding in very preterm infants (feeds were advanced at 10 to 15 mL/kg/d in both groups)			
Viswanathan 2017	Retrospective cohort study			

RCT: randomised controlled trial

DATA AND ANALYSES

Comparison 1. Slow versus faster rates of feed advancement

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.1 Necrotising enterocolitis - all in- fants	14	4026	Risk Difference (M-H, Fixed, 95% CI)	0.00 [-0.01, 0.02]
1.1.1 Infants at least partially fed with human milk	13	3841	Risk Difference (M-H, Fixed, 95% CI)	0.00 [-0.01, 0.02]
1.1.2 Infants exclusively formula fed	1	185	Risk Difference (M-H, Fixed, 95% CI)	0.04 [-0.05, 0.13]
1.2 Necrotising enterocolitis - ELBW/ extremely preterm infants	5		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.3 Necrotising enterocolitis - small for gestational age or growth restrict- ed	2	639	Risk Ratio (M-H, Fixed, 95% CI)	1.26 [0.67, 2.37]
1.4 Necrotising enterocolitis - AREDFV	2	465	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [0.74, 3.40]



Cochrane Database of Systematic Reviews

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.5 Mortality - all infants	13	3860	Risk Difference (M-H, Fixed, 95% CI)	0.01 [-0.01, 0.02]
1.5.1 Infants at least partially fed with human milk	12	3675	Risk Difference (M-H, Fixed, 95% CI)	0.01 [-0.01, 0.03]
1.5.2 Infants exclusively formula fed	1	185	Risk Difference (M-H, Fixed, 95% CI)	0.00 [-0.02, 0.02]
1.6 Mortality - ELBW/extremely preterm infants	2		Risk Ratio (M-H, Fixed, 95% Cl)	Subtotals only
1.7 Mortality - small for gestational age or growth restricted	1		Risk Ratio (M-H, Fixed, 95% Cl)	Subtotals only
1.8 Mortality - AREDFV	1		Risk Ratio (M-H, Fixed, 95% Cl)	Subtotals only
1.9 Weight z-score at hospital dis- charge	1	2602	Mean Difference (IV, Fixed, 95% CI)	0.00 [-0.08, 0.08]
1.10 Head circumference z-score at hospital discharge	1	2286	Mean Difference (IV, Fixed, 95% CI)	0.00 [-0.13, 0.13]
1.11 Moderate or severe disability	1	2325	Risk Ratio (M-H, Fixed, 95% Cl)	0.90 [0.79, 1.02]
1.12 Cerebral palsy	1	2183	Risk Ratio (M-H, Fixed, 95% Cl)	0.60 [0.39, 0.90]
1.13 Visual impairment	1	2327	Risk Ratio (M-H, Fixed, 95% Cl)	0.75 [0.39, 1.43]
1.14 Hearing impairment	1	2315	Risk Ratio (M-H, Fixed, 95% Cl)	0.69 [0.47, 1.02]
1.15 Cognitive impairment	1	2326	Risk Ratio (M-H, Fixed, 95% Cl)	0.93 [0.81, 1.07]
1.16 Feed intolerance (causing inter- ruption of enteral feeding)	9	719	Risk Ratio (M-H, Fixed, 95% Cl)	1.18 [0.95, 1.46]
1.17 Invasive infection	11	3583	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [0.99, 1.31]



Analysis 1.1. Comparison 1: Slow versus faster rates of feed advancement, Outcome 1: Necrotising enterocolitis - all infants

	Slow	rate	Fast	ate		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.1.1 Infants at least partially	y fed with hu	man milk					
Caple 2004	2	84	4	74	3.9%	-0.03 [-0.09 , 0.03]	
Fayyaz 2020	1	16	0	15	0.8%	0.06 [-0.10 , 0.22]	
Hasshemi 2018	2	100	0	40	2.9%	0.02 [-0.03 , 0.07]	1 -
ain 2016	1	15	2	15	0.8%	-0.07 [-0.28 , 0.15]]
Karagol 2013	5	46	4	46	2.3%	0.02 [-0.10 , 0.14]]
Krishnamurthy 2010	1	50	2	50	2.5%	-0.02 [-0.09 , 0.05]	
Modi 2015	2	65	1	66	3.3%	0.02 [-0.04 , 0.07]	
Montealegre-Pomar 2021	0	12	2	8	0.5%	-0.25 [-0.56 , 0.06]	
Raban 2014a	1	51	7	47	2.4%	-0.13 [-0.24 , -0.02]	
Raban 2014b	9	52	2	50	2.6%	0.13 [0.02 , 0.25]	
Saha 2019	3	55	2	38	2.3%	0.00 [-0.09 , 0.09]	
alhotra 2004	0	26	2	27	1.3%	-0.07 [-0.19, 0.04]	
SIFT 2016	78	1399	70	1394	69.9%	0.01 [-0.01, 0.02]	
Subtotal (95% CI)		1971		1870	95.4%	0.00 [-0.01 , 0.02]	i T
Total events:	105		98				
Heterogeneity: Chi ² = 18.24, d	lf = 12 (P = 0.)	11); I ² = 3	4%				
Cest for overall effect: $Z = 0.2$	6 (P = 0.79)						
1.1.2 Infants exclusively form	nula fed						
Rayyis 1999	13	98	8	87	4.6%	0.04 [-0.05, 0.13]	1
Subtotal (95% CI)		98		87	4.6%	0.04 [-0.05 , 0.13]	
Total events:	13		8			. , .	
Heterogeneity: Not applicable							
Test for overall effect: $Z = 0.8$							
	. ,						
Total (95% CI)		2069		1957	100.0%	0.00 [-0.01 , 0.02]	1
Total events:	118		106				
Heterogeneity: Chi ² = 18.86, d	lf = 13 (P = 0.)	13); I ² = 3	1%				-1 -0.5 0 0.5 1
Test for overall effect: Z = 0.5	1 (P = 0.61)						Favours slow rates Favours faster ra
est for subgroup differences:	$Chi^2 = 0.69$	df = 1 (P =	(0.41) , $I^2 =$	0%			

Analysis 1.2. Comparison 1: Slow versus faster rates of feed advancement, Outcome 2: Necrotising enterocolitis - ELBW/extremely preterm infants

	Slow	rate	Fast	rate	Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Karagol 2013	1	14	2	15	0.54 [0.05 , 5.28	
Raban 2014a	1	51	7	47	0.13 [0.02 , 1.03]
Raban 2014b	9	52	2	50	4.33 [0.98 , 19.05]
Rayyis 1999	9	43	3	33	2.30 [0.68 , 7.84]
SIFT 2016	53	498	56	496	0.94 [0.66 , 1.34] +
Test for subgroup differ	rences: Not a	pplicable				0.01 0.1 1 10 100 Favours slow rates Favours faster rates



Analysis 1.3. Comparison 1: Slow versus faster rates of feed advancement, Outcome 3: Necrotising enterocolitis - small for gestational age or growth restricted

	Slow	rate	Fast	rate		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
Salhotra 2004	0	26	2	27	15.0%	0.21 [0.01 , 4.12]	
SIFT 2016	20	291	14	295	85.0%	1.45 [0.75 , 2.81]	
Total (95% CI)		317		322	100.0%	1.26 [0.67 , 2.37	ı 🔶	
Total events:	20		16					
Heterogeneity: Chi ² = 1	.57, df = 1 (I	P = 0.21); I	I² = 36%				0.01 0.1 1 10 100	
Test for overall effect: 2	Z = 0.72 (P =	0.47)					Favours slow rates Favours faster ra	ites
Test for subgroup differ	ences: Not a	pplicable						

Analysis 1.4. Comparison 1: Slow versus faster rates of feed advancement, Outcome 4: Necrotising enterocolitis - AREDFV

	Slow	rate	Fast	rate		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Jain 2016	1	15	2	15	19.4%	0.50 [0.05 , 4.94]
SIFT 2016	16	226	8	209	80.6%	1.85 [0.81 , 4.23] +
Total (95% CI)		241		224	100.0%	1.59 [0.74 , 3.40	1
Total events:	17		10				
Heterogeneity: Chi ² = 1.	11, df = 1 (F	P = 0.29); I	$1^2 = 10\%$				0.01 0.1 1 10 100
Test for overall effect: Z	= 1.19 (P =	0.23)					Favours slow rates Favours faster rates
Test for subgroup differe	ences: Not a	pplicable					

Analysis 1.5. Comparison 1: Slow versus faster rates of feed advancement, Outcome 5: Mortality - all infants

	Slow	rate	Fast	rate		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.5.1 Infants at least partiall	y fed with hu	man milk	:				
Salhotra 2004	12	26	7	27	1.4%	0.20 [-0.05 , 0.46]	
Krishnamurthy 2010	6	50	4	50	2.6%	0.04 [-0.08 , 0.16]	∣
Karagol 2013	4	46	3	46	2.4%	0.02 [-0.09 , 0.13]	।
Raban 2014b	16	52	14	50	2.7%	0.03 [-0.15 , 0.20]	I
Raban 2014a	13	51	19	47	2.6%	-0.15 [-0.33 , 0.04]	∣
/Iodi 2015	28	65	20	66	3.4%	0.13 [-0.04 , 0.29]	∣ ∔⊷
Jain 2016	3	15	0	15	0.8%	0.20 [-0.02 , 0.42]	I <u>L.</u>
5IFT 2016	65	1393	60	1392	72.7%	0.00 [-0.01 , 0.02]	I 1
Hasshemi 2018	0	100	0	40	3.0%	0.00 [-0.04 , 0.04]	ı T
Saha 2019	7	55	4	38	2.3%	0.02 [-0.11 , 0.15]	∣
ayyaz 2020	0	16	1	15	0.8%	-0.07 [-0.23 , 0.10]	∣
Montealegre-Pomar 2021	0	12	1	8	0.5%	-0.13 [-0.39 , 0.14]	I
ubtotal (95% CI)		1881		1794	95.2%	0.01 [-0.01 , 0.03]	∣ ∳
otal events:	154		133				
Heterogeneity: Chi ² = 13.04, c	f = 11 (P = 0.)	29); I ² = 1	6%				
Test for overall effect: $Z = 1.1$	4 (P = 0.25)						
1.5.2 Infants exclusively forr	nula fed						
Rayyis 1999	0	98	0	87	4.8%	0.00 [-0.02 , 0.02]	ı 🖡
Subtotal (95% CI)		98		87	4.8%	0.00 [-0.02 , 0.02]	I 🖌
Total events:	0		0				ľ
Heterogeneity: Not applicable							
Test for overall effect: Z = 0.0	0 (P = 1.00)						
Fotal (95% CI)		1979		1881	100.0%	0.01 [-0.01 , 0.02]	• •
Fotal events:	154		133				
Heterogeneity: Chi ² = 13.68, c		.32); I ² = 1	2%				-1 -0.5 0 0.5 1
Test for overall effect: $Z = 1.1$,						Favours slow rates Favours faster r
est for subgroup differences:	$Chi^2 = 0.50, o$	df = 1 (P =	0.48), I ² =	0%			

Analysis 1.6. Comparison 1: Slow versus faster rates of feed advancement, Outcome 6: Mortality - ELBW/extremely preterm infants

	Slow	rate	Fast 1	rate	Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI	M-H, Fixed, 959	% CI
Raban 2014b	16	52	14	50	1.10 [0.60 , 2.01]	_	
Raban 2014a	13	51	19	47	0.63 [0.35 , 1.13]		
Test for subgroup differ	ences: Not aj	pplicable			0. Fav		10 100 vours faster rates



Analysis 1.7. Comparison 1: Slow versus faster rates of feed advancement, Outcome 7: Mortality - small for gestational age or growth restricted

Study or Subgroup	Slow Events	rate Total	Fast 1 Events	rate Total	Risk Ratio M-H, Fixed, 95% CI		Ratio d, 95% CI
Salhotra 2004	12	26	7	27	1.78 [0.83 , 3.81]	-	+
Test for subgroup differe	ences: Not a	pplicable			0. Fav	01 0.1 2 vours slow rates	10 100 Favours faster rates

Analysis 1.8. Comparison 1: Slow versus faster rates of feed advancement, Outcome 8: Mortality - AREDFV

	Slow	Slow rate Fas		rate	Risk Ratio	Risk	Ratio
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI	M-H, Fixe	ed, 95% CI
Jain 2016	3	15	0	15	5 7.00 [0.39 , 124.83]		i →
Test for subgroup differe	ences: Not a	pplicable				0.01 0.1 Favours slow rates	1 10 100 Favours faster rates

Analysis 1.9. Comparison 1: Slow versus faster rates of feed advancement, Outcome 9: Weight z-score at hospital discharge

	s	low rate		F	ast rate			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
SIFT 2016	-1.5	1.1	1295	-1.5	1.1	1307	100.0%	0.00 [-0.08 , 0.08]
Total (95% CI) Heterogeneity: Not appl Test for overall effect: Z Test for subgroup differ	z = 0.00 (P =		1295			1307	100.0%	0.00 [-0.08 , 0.08	-0.050.025 0 0.0250.05 Favours slow rates Favours faster rates

Analysis 1.10. Comparison 1: Slow versus faster rates of feed advancement, Outcome 10: Head circumference z-score at hospital discharge

Study or Subgroup	S Mean	ilow rate SD	Total	I Mean	ast rate SD	Total	Weight	Mean Difference IV, Fixed, 95% CI	Mean Difference IV, Fixed, 95% CI
SIFT 2016	-0.8	1.7	1156	-0.8	1.5	1130	100.0%	0.00 [-0.13 , 0.13]
Total (95% CI) Heterogeneity: Not appl Test for overall effect: Z Test for subgroup differ	z = 0.00 (P =	· ·	1156			1130	100.0%	0.00 [-0.13 , 0.13	-0.1 -0.05 0 0.05 0.1 Favours slow rates Favours faster rates

Analysis 1.11. Comparison 1: Slow versus faster rates of feed advancement, Outcome 11: Moderate or severe disability

Study or Subgroup	Slow Events	rate Total	Fast i Events	rate Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk Ratio M-H, Fixed, 95% CI
SIFT 2016	321	1169	354	1156	100.0%	0.90 [0.79 , 1.02]	l P
Total (95% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z Test for subgroup differe	= 1.68 (P =		354	1156	100.0%		0.01 0.1 1 10 100 Favours slow rates Favours faster rates

Analysis 1.12. Comparison 1: Slow versus faster rates of feed advancement, Outcome 12: Cerebral palsy

	Slow	rate	Fast	rate		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
SIFT 2016	35	1099	58	1084	100.0%	0.60 [0.39 , 0.90]	_ _	
Total (95% CI)		1099		1084	100.0%	0.60 [0.39 , 0.90]		
Total events:	35		58					
Heterogeneity: Not appli	icable							
Test for overall effect: Z	= 2.47 (P =	0.01)]	Favours slow rates Favours faster i	rates
Test for subgroup differences: Not applicable								

Analysis 1.13. Comparison 1: Slow versus faster rates of feed advancement, Outcome 13: Visual impairment

	Slow 1	rate	Fast	rate		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
SIFT 2016	16	1171	21	1156	100.0%	0.75 [0.39 , 1.43] -
Total (95% CI)		1171		1156	100.0%	0.75 [0.39 , 1.43	1
Total events:	16		21				•
Heterogeneity: Not app	licable						0.01 0.1 1 10 100
Test for overall effect: Z	Z = 0.87 (P =	0.39)					Favours slow rates Favours faster rates
Test for subgroup differ	ences: Not ap	pplicable					

Analysis 1.14. Comparison 1: Slow versus faster rates of feed advancement, Outcome 14: Hearing impairment

	Slow	rate	Fast	rate		Risk Ratio	Risk R	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed	l, 95% CI
SIFT 2016	41	1172	58	1143	100.0%	0.69 [0.47 , 1.02]]	
Total (95% CI)		1172		1143	100.0%	0.69 [0.47 , 1.02]	1	
Total events:	41		58				•	
Heterogeneity: Not app	olicable						0.01 0.1 1	10 100
Test for overall effect:	Z = 1.86 (P =	0.06)					Favours slow rates	Favours fast rates
Test for subgroup diffe	rences: Not aj	pplicable						

Analysis 1.15. Comparison 1: Slow versus faster rates of feed advancement, Outcome 15: Cognitive impairment

Study or Subgroup	Slow 1 Events	rate Total	Fast 1 Events	rate Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk Ratio M-H, Fixed, 95% CI
SIFT 2016	289	1170	307	1156	100.0%	0.93 [0.81 , 1.07]
Total (95% CI)		1170		1156	100.0%	0.93 [0.81 , 1.07	1
Total events:	289		307				1
Heterogeneity: Not appl	icable						0.01 0.1 1 10 100
Test for overall effect: Z	= 1.02 (P =	0.31)					Favours slow rates Favours faster rates
Test for subgroup differe	ences: Not ap	Test for subgroup differences: Not applicable					

Analysis 1.16. Comparison 1: Slow versus faster rates of feed advancement, Outcome 16: Feed intolerance (causing interruption of enteral feeding)

	Slow	rate	Fast	rate		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Salhotra 2004	17	26	14	27	13.7%	1.26 [0.80 , 1.99]]
Krishnamurthy 2010	12	50	8	50	8.0%	1.50 [0.67 , 3.35]]
Karagol 2013	13	46	11	46	10.9%	1.18 [0.59 , 2.36]]
Raban 2014a	20	51	19	47	19.7%	0.97 [0.60 , 1.58]]
Raban 2014b	24	52	19	50	19.3%	1.21 [0.77 , 1.92]	
Modi 2015	17	65	12	66	11.8%	1.44 [0.75 , 2.77]]
Jain 2016	4	15	5	15	5.0%	0.80 [0.27 , 2.41]]
Saha 2019	15	55	10	38	11.8%	1.04 [0.52 , 2.06]]
Montealegre-Pomar 2021	0	12	0	8		Not estimable	2
Total (95% CI)		372		347	100.0%	1.18 [0.95 , 1.46]	
Total events:	122		98				-
Heterogeneity: Chi ² = 2.02, df = 7 (P = 0.96); I ² = 0%							
Test for overall effect: Z = 1.4	9 (P = 0.14)						Favours slow rates Favours faster ra
T	NT . 1 . 1 1						

Test for subgroup differences: Not applicable

Analysis 1.17. Comparison 1: Slow versus faster rates of feed advancement, Outcome 17: Invasive infection

	Slow	rate	Fast	rate		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Salhotra 2004	10	26	5	27	1.6%	2.08 [0.82 , 5.26]	
Krishnamurthy 2010	5	50	4	50	1.3%	1.25 [0.36 , 4.38]]
Karagol 2013	10	46	6	46	2.0%	1.67 [0.66 , 4.21]]
Raban 2014b	10	52	4	50	1.4%	2.40 [0.81 , 7.17]]
Raban 2014a	9	51	9	47	3.1%	0.92 [0.40 , 2.12]	
Modi 2015	24	65	17	66	5.6%	1.43 [0.85 , 2.41]	1 4
SIFT 2016	267	1397	247	1389	82.5%	1.07 [0.92 , 1.26]]
Jain 2016	4	15	2	15	0.7%	2.00 [0.43 , 9.32]	
Hasshemi 2018	4	100	0	40	0.2%	3.65 [0.20 , 66.34]]
Fayyaz 2020	1	16	2	15	0.7%	0.47 [0.05 , 4.65]]
Montealegre-Pomar 2021	1	12	2	8	0.8%	0.33 [0.04 , 3.09]	I
Total (95% CI)		1830		1753	100.0%	1.14 [0.99 , 1.31]	1
Total events:	345		298				•
Heterogeneity: Chi² = 8.49, df = 10 (P = 0.58); I² = 0%							-++++++++++++++++++++++++++++++++++++
Test for overall effect: Z = 1.85	Test for overall effect: $Z = 1.85$ (P = 0.06)						Favours slow rates Favours faster rates
Test for subgroup differences:	Not applicabl	e					

APPENDICES

Appendix 1. 2020 search methods

Databases searches: CENTRAL (via Cochrane Library); CINAHL; Embase, Maternity & Infant Care (MIDIRS); MEDLINE; ClinicalTrials.gov (WHO ICTRP – currently not available)

CENTRAL via Cochrane Library

Search date 19th October 2020

#1 MeSH descriptor: [Infant, Newborn] explode all trees

#2 MeSH descriptor: [Premature Birth] explode all trees

#3 ((neonat* or neo nat*)):ti,ab,kw OR ((newborn* or new born* or newly born*)):ti,ab,kw OR ((preterm or preterms or pre term or pre terms)):ti,ab,kw OR ((preemie* or premie or premies)):ti,ab,kw OR ((prematur* NEAR/3 (birth* or born or deliver*))):ti,ab,kw (Word variations have been searched)

#4 ((low NEAR/3 (birthweight* or birth weight*))):ti,ab,kw OR ((lbw or vlbw or elbw)):ti,ab,kw OR (infan*):ti,ab,kw OR ((baby or babies)):ti,ab,kw (Word variations have been searched)

#5 #1 OR #2 OR #3 OR #4

#6 MeSH descriptor: [Enteral Nutrition] explode all trees

#7 (((enteral or enteric) NEAR/2 (nutrition or feed*))):ti,ab,kw OR (((oral or sip or tube) NEAR/2 feeding*)):ti,ab,kw OR (((nasogastric or gastrostomy or jejunostomy) NEAR/2 tube*)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteral feed*)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or speed* or slow* or volume*) NEAR/3 enteral feed*)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteric feed*)):ti,ab,kw (Word variations have been searched)

#8 (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteral intake*)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteric intake*)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteral nutrition)):ti,ab,kw OR (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) NEAR/3

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



rapid* or routine* or speed* or slow* or volume*) NEAR/3 enteric nutrition)):ti,ab,kw OR (((aggressive* or fast or rapid* or slow* or speed*) NEAR/3 feed*)):ti,ab,kw (Word variations have been searched)

#9 (((aggressive* or fast or rapid* or slow* or speed*) NEAR/3 volume*)):ti,ab,kw OR (trophic feeding*):ti,ab,kw OR (((gut or gastrointestinal) NEAR/2 priming)):ti,ab,kw (Word variations have been searched)

#10 #6 or #7 or #8 or #9

#11 #5 and #10

#12 MeSH descriptor: [Parenteral Nutrition] explode all trees and with qualifier(s): [adverse effects - AE]

#13 MeSH descriptor: [Enterocolitis, Necrotizing] explode all trees and with qualifier(s): [etiology - ET, epidemiology - EP, prevention & control - PC]

#14 MeSH descriptor: [Infections] 1 tree(s) exploded and with qualifier(s): [epidemiology - EP]

#15 (((prevent* or risk*) NEAR/3 necrotising enterocolitis)):ti,ab,kw OR (((prevent* or risk*) NEAR/3 necrotizing enterocolitis)):ti,ab,kw (Word variations have been searched)

#16 #12 or #13 or #14 or #15

#17 #5 and #16

#18 #11 or #17

CINAHL via EBSCO

Search date 19th October 2020

1062 records identified

S1 (MH "Infant, Newborn+")

S2 (MH "Childbirth, Premature")

S3 TI ((neonat* or neo nat*)) OR AB ((neonat* or neo nat*)) OR TI ((newborn* or new born* or newly born*)) OR AB ((newborn* or new born* or newly born*)) OR TI ((preterm or preterms or pre terms or pre terms)) OR AB ((preterm or preterms or pre terms))

S4 TI ((preemie* or premie or premies)) OR AB ((preemie* or premie or premies)) OR TI ((prematur* N3 (birth* or born or deliver*))) OR AB ((prematur* N3 (birth* or born or deliver*))) OR TI ((low N3 (birthweight* or birth weight*))) OR AB ((low N3 (birthweight* or birth weight*))) OR AB ((low N3 (birthweight* or birth weight*))) OR AB ((low N3 (birthweight* or birth weight*))) OR AB (low N3 (birth* or born or deliver*))) OR TI (low N3 (birth* or born or deliver*))) OR AB (low N3 (birth* or born or deliver*))) OR TI (low N3 (birth* or born or deliver*))) OR AB (low N3 (birth* or born or deliver*))) OR TI (low N3 (birth* or born or deliver*))) OR AB (low N3 (birth* or born or deliver*))) OR AB (low N3 (birth* or born or deliver*))) OR TI (low N3 (birth* or born or deliver*))) OR AB

S5 TI ((lbw or vlbw or elbw)) OR AB ((lbw or vlbw or elbw)) OR TI infan* OR AB infan* OR TI ((baby or babies)) OR AB ((baby or babies))

S6 S1 OR S2 OR S3 OR S4 OR S5

S7 (MH "Enteral Nutrition")

S8 TI (((enteral or enteric) N2 (nutrition or feed*))) OR AB (((enteral or enteric) N2 (nutrition or feed*))) OR TI (((oral or sip or tube) N2 feeding*)) OR AB (((oral or sip or tube) N2 feeding*)) OR TI (((nasogastric or gastrostomy or jejunostomy) N2 tube*)) OR AB (((nasogastric or gastrostomy or jejunostomy) N2 tube*)) OR AB (((nasogastri

S9 TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral feed*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral feed*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral feed*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric feed*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric feed*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric feed*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric feed*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric feed*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral intake*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral intake*)) OR AB (((advanc* or

S10 TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric intake*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric intake*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric intake*)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric intake*)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or molonged or rapid* or routine* or speed* or slow* or volume*) N3 enteral nutrition)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or progress* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or early or fast or full or increas* or prolonged or rapid* or progress* or prolonged or progress* or prolonged or progress* or prolonged or prapid* or progress* or



rapid* or routine* or speed* or slow* or volume*) N3 enteral nutrition)) OR TI (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric nutrition)) OR AB (((advanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or slow* or volume*) N3 enteric nutrition)) OR AB ((idvanc* or aggressive* or delay* or early or fast or full or increas* or minimal or progress* or prolonged or rapid* or routine* or speed* or slow* or volume*) N3 enteric nutrition))

S11 TI (((aggressive* or fast or rapid* or slow* or speed*) N3 feed*)) OR AB (((aggressive* or fast or rapid* or slow* or speed*) N3 feed*)) OR TI (((aggressive* or fast or rapid* or slow* or speed*) N3 volume*)) OR AB (((aggressive* or fast or rapid* or slow* or speed*) N3 volume*)) OR TI trophic feeding OR AB trophic feeding* OR TI (((gut or gastrointestinal) N2 priming)) OR AB ((gut or gastrointestinal) N2 priming)) OR AB ((gut or gastrointestinal) N2 priming)) OR AB ((gut or gastrointestinal) N2 priming)) OR AB ((gut or gastrointestinal) (gut or gastrointestinal) (gut or gastrointestinal) (gut or

S12 S7 OR S8 OR S9 OR S10 OR S11

S13 S6 AND S12

S14 (MH "Double-Blind Studies")

S15 (MH "Single-Blind Studies")

S16 (MH "Random Assignment")

S17 (MH "Pretest-Posttest Design")

S18 (MH "Cluster Sample")

S19 TI (randomized or randomised) OR AB random* OR TI trial

S20 MH "sample size" AND AB ((assigned or allocated or control))

S21 MH placebos

S22 PT randomised controlled trial OR AB control W5 group OR MH "crossover design" OR MH "comparative studies" OR AB cluster W3 RCT

S23 S14 OR S15 OR S16 OR S17 OR S18 OR S19 OR S20 OR S21 OR S22

S24 S13 AND S23

S25 (MH "Parenteral Nutrition/AE")

S26 (MH "Enterocolitis, Necrotizing/CO/ET/EP/PC")

S27 (MH "Infection/EP")

S28 TI (((prevent* or risk*) N3 necrotising enterocolitis)) OR AB (((prevent* or risk*) N3 necrotising enterocolitis)) OR TI (((prevent* or risk*) N3 necrotizing enterocolitis)) OR AB (((prevent* or risk*) N3 necrotizing enterocolitis))

S29 S25 OR S26 OR S27 OR S28

S30 S6 AND S23 AND S29

S31 S24 OR S30

View Results (1,062)

Embase via OVID

Search date 19th October 2020

1 Newborn/ (533014)

2 Prematurity/ (103974)

3 (neonat\$ or neo nat\$).ti,ab. (347799)

4 (newborn\$ or new born\$ or newly born\$).ti,ab. (196226)

5 (preterm or preterms or pre term or pre terms).ti,ab. (107483)

6 (preemie\$ or premie or premies).ti,ab. (272)

- 7 (prematur\$ adj3 (birth\$ or born or deliver\$)).ti,ab. (21963)
- 8 (low adj3 (birthweight\$ or birth weight\$)).ti,ab. (44434)
- 9 (lbw or vlbw or elbw).ti,ab. (11748)
- 10 infan\$.ti,ab. (505921)
- 11 (baby or babies).ti,ab. (98472)
- 12 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 (1150651)
- 13 Enteric Nutrition/ (32699)
- 14 ((enteral or enteric) adj2 (nutrition or feed\$)).ti,ab. (23164)
- 15 ((oral or sip or tube) adj2 feeding\$).ti,ab. (15554)
- 16 ((nasogastric or gastrostomy or jejunostomy) adj2 tube\$).ti,ab. (13591)

17 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteral feed\$).ti,ab. (1697)

18 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteric feed\$).ti,ab. (10)

19 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteral nutrition).ti,ab. (2109)

20 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteric nutrition).ti,ab. (7)

21 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteral intake).ti,ab. (40)

22 ((advanc\$ or aggressive\$ or delay\$ or early or fast or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed\$ or slow\$ or volume\$) adj3 enteric intake).ti,ab. (0)

- 23 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 feed\$).ti,ab. (3431)
- 24 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 volume\$).ti,ab. (5320)
- 25 trophic feeding\$.ti,ab. (113)
- 26 ((gut or gastrointestinal) adj2 priming).ti,ab. (41)
- 27 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 (67117)
- 28 12 and 27 (10261)
- 29 Randomized controlled trial/ (626485)
- 30 Controlled clinical study/ (465422)
- 31 Random\$.ti,ab. (1589685)
- 32 randomization/ (88577)
- 33 intermethod comparison/ (266507)
- 34 placebo.ti,ab. (313152)
- 35 (compare or compared or comparison).ti. (521918)

36 ((evaluated or evaluate or evaluating or assessed or assess) and (compare or compared or comparing or comparison)).ab. (2190407)

Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



37 (open adj label).ti,ab. (82341)

38 ((double or single or doubly or singly) adj (blind or blinded or blindly)).ti,ab. (237163)

39 double blind procedure/ (177360)

40 parallel group\$1.ti,ab. (26293)

41 (crossover or cross over).ti,ab. (107555)

42 ((assign\$ or match or matched or allocation) adj5 (alternate or group\$1 or intervention\$1 or patient\$1 or subject\$1 or participant \$1)).ti,ab. (339974)

- 43 (assigned or allocated).ti,ab. (400614)
- 44 (controlled adj7 (study or design or trial)).ti,ab. (360663)
- 45 (volunteer or volunteers).ti,ab. (252353)
- 46 human experiment/ (519930)

47 trial.ti. (311954)

48 or/29-47 (5173763)

49 (random\$ adj sampl\$ adj7 ("cross section\$" or questionnaire\$1 or survey\$ or database\$1)).ti,ab. not (comparative study/ or controlled study/ or randomi?ed controlled.ti,ab. or randomly assigned.ti,ab.) (8271)

50 Cross-sectional study/ not (randomized controlled trial/ or controlled clinical study/ or controlled study/ or randomi?ed controlled.ti,ab. or control group\$1.ti,ab.) (249866)

- 51 (((case adj control\$) and random\$) not randomi?ed controlled).ti,ab. (17728)
- 52 (Systematic review not (trial or study)).ti. (155685)
- 53 (nonrandom\$ not random\$).ti,ab. (16497)
- 54 "Random field\$".ti,ab. (2428)
- 55 (random cluster adj3 sampl\$).ti,ab. (1312)
- 56 (review.ab. and review.pt.) not trial.ti. (836864)
- 57 "we searched".ab. and (review.ti. or review.pt.) (33674)
- 58 "update review".ab. (110)
- 59 (databases adj4 searched).ab. (38341)

60 (rat or rats or mouse or mice or swine or porcine or murine or sheep or lambs or pigs or piglets or rabbit or rabbits or cat or cats or dog or dogs or cattle or bovine or monkey or monkeys or trout or marmoset\$1).ti. and animal experiment/ (1084361)

61 Animal experiment/ not (human experiment/ or human/) (2283901)

62 or/49-61 (3568186)

63 48 not 62 (4602096)

64 28 and 63 (2237)

65 Parenteral Nutrition/ (29655)

66 complication/ (221706)

67 safety/ or patient safety/ (371167)

68 (adverse\$ adj2 (effect\$ or event\$ or impact\$ or outcome\$)).ti,ab. (650603)

69 (complication\$ or risk\$ or safe or safety or safety or sequaela or side effect\$ or tolerated or toxicities or toxicity).ti,ab. (5830131)

70 65 and (66 or 67 or 68 or 69) (10653)

- 71 Necrotizing Enterocolitis/co, ep, et, pc [Complication, Epidemiology, Etiology, Prevention] (2682)
- 72 ((prevent\$ or risk\$) adj3 necrotising enterocolitis).ti,ab. (204)
- 73 ((prevent\$ or risk\$) adj3 necrotizing enterocolitis).ti,ab. (800)
- 74 Infection/ep [Epidemiology] (4639)
- 75 70 or 71 or 72 or 73 or 74 (18499)
- 76 12 and 63 and 75 (1149)
- 77 64 or 76 (3050)

Maternity & Infant Care Via OVID

Search date 19th October 2020

- 1 (neonat\$ or neo nat\$).ti,ab. (48117)
- 2 (newborn\$ or new born\$ or newly born\$).ti,ab. (21553)
- 3 (preterm or preterms or pre term or pre terms).ti,ab. (28518)
- 4 (preemie\$ or premie or premies).ti,ab. (57)
- 5 (prematur\$ adj3 (birth\$ or born or deliver\$)).ti,ab. (4262)
- 6 (low adj3 (birthweight\$ or birth weight\$)).ti,ab. (11416)
- 7 (lbw or vlbw or elbw).ti,ab. (3290)
- 8 infan\$.ti,ab. (68527)
- 9 (baby or babies).ti,ab. (30608)
- 10 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 (127398)
- 11 (Infant premature or Infant very low birth weight or Infant newborn).de. (30913)
- 12 10 or 11 (130585)
- 13 Enteral nutrition.de. (287)
- 14 ((enteral or enteric) adj2 (nutrition or feed\$)).ti,ab. (779)
- 15 ((oral or sip or tube) adj2 feeding\$).ti,ab. (480)
- 16 ((nasogastric or gastrostomy or jejunostomy) adj2 tube\$).ti,ab. (157)

17 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral feed\$).ti,ab. (324)

18 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteric feed\$).ti,ab. (1)

19 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral intake\$).ti,ab. (9)

20 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteric intake\$).ti,ab. (0)

21 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral nutrition).ti,ab. (71)





\$ or slow\$ or volume\$) adj3 enteric nutrition).ti,ab. (0)
23 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 feed\$).ti,ab. (76)
24 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 volume\$).ti,ab. (31)
25 trophic feeding\$.ti,ab. (23)

22 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed

- 26 ((gut or gastrointestinal) adj2 priming).ti,ab. (5)
- 27 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 (1471)
- 28 12 and 27 (1365)
- 29 limit 28 to randomised controlled trial (103)
- 30 Parenteral nutrition.de. (206)
- 31 Enterocolitis necrotizing.de. (1)
- 32 (adverse\$ adj2 (effect\$ or event\$ or impact\$ or outcome\$)).ti,ab. (13711)
- 33 (complication\$ or risk\$ or safe or safely or safer or safety or sequaela or side effect\$ or tolerated or toxicities or toxicity).ti,ab. (94379)
- 34 Complications.de. (199)
- 35 safety.de. (1953)
- 36 (30 or 31) and (32 or 33 or 34 or 35) (96)
- 37 ((prevent\$ or risk\$) adj3 necrotising enterocolitis).ti,ab. (71)
- 38 ((prevent\$ or risk\$) adj3 necrotizing enterocolitis).ti,ab. (231)
- 39 36 or 37 or 38 (392)
- 40 limit 39 to randomised controlled trial (17)
- 41 29 or 40 (110)

MEDLINE via OVID

Search date 19th October 2020 [Ovid MEDLINE(R) ALL <1946 to October 16, 2020>]

- 1 exp Infant, Newborn/ (611659)
- 2 Premature Birth/ (14154)
- 3 (neonat\$ or neo nat\$).ti,ab. (267306)
- 4 (newborn\$ or new born\$ or newly born\$).ti,ab. (167592)
- 5 (preterm or preterms or pre term or pre terms).ti,ab. (76438)
- 6 (preemie\$ or premie or premies).ti,ab. (174)
- 7 (prematur\$ adj3 (birth\$ or born or deliver\$)).ti,ab. (15880)
- 8 (low adj3 (birthweight\$ or birth weight\$)).ti,ab. (35103)
- 9 (lbw or vlbw or elbw).ti,ab. (8568)
- 10 infan\$.ti,ab. (441307)
- 11 (baby or babies).ti,ab. (70876)
- 12 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 (1067192)

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13 Enteral Nutrition/ (19843)

- 14 ((enteral or enteric) adj2 (nutrition or feed\$)).ti,ab. (14402)
- 15 ((oral or sip or tube) adj2 feeding\$).ti,ab. (10325)

16 ((nasogastric or gastrostomy or jejunostomy) adj2 tube\$).ti,ab. (8813)

17 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral feed\$).ti,ab. (1664)

18 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteric feed\$).ti,ab. (14)

19 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral intake\$).ti,ab. (41)

20 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteric intake\$).ti,ab. (0)

21 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteral nutrition).ti,ab. (1450)

22 ((advanc\$ or aggressive\$ or delay\$ or early or fast or full or increas\$ or minimal or progress\$ or prolonged or rapid\$ or routine\$ or speed \$ or slow\$ or volume\$) adj3 enteric nutrition).ti,ab. (5)

- 23 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 feed\$).ti,ab. (2933)
- 24 ((aggressive\$ or fast or rapid\$ or slow\$ or speed\$) adj3 volume\$).ti,ab. (3947)
- 25 trophic feeding\$.ti,ab. (89)
- 26 ((gut or gastrointestinal) adj2 priming).ti,ab. (33)
- 27 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 (43937)
- 28 12 and 27 (7156)
- 29 Parenteral Nutrition/ae [Adverse Effects] (2782)
- 30 Enterocolitis, Necrotizing/ep, et, pc [Epidemiology, Etiology, Prevention & Control] (1662)
- 31 ((prevent\$ or risk\$) adj3 necrotising enterocolitis).ti,ab. (143)
- 32 ((prevent\$ or risk\$) adj3 necrotizing enterocolitis).ti,ab. (610)
- 33 Infection/ep [Epidemiology] (4070)
- 34 29 or 30 or 31 or 32 or 33 (8854)
- 35 12 and 34 (3319)
- 36 28 or 35 (9891)
- 37 randomized controlled trial.pt. (515341)
- 38 controlled clinical trial.pt. (93888)
- 39 randomized.ab. (495059)
- 40 placebo.ab. (211516)
- 41 drug therapy.fs. (2243399)
- 42 randomly.ab. (342541)
- 43 trial.ab. (523032)



44 groups.ab. (2102194)

45 37 or 38 or 39 or 40 or 41 or 42 or 43 or 44 (4813149)

46 exp animals/ not humans.sh. (4744580)

47 45 not 46 (4179397)

48 36 and 47 (2836)

Appendix 2. Previous search methods

Electronic searches

We searched the Cochrane Central Register of Controlled Trials (CENTRAL; 2017, Issue 5), MEDLINE via PubMed (2015 to June 2017), Embase (2015 to June 2017), and the Cumulative Index to Nursing and Allied Health Literature (CINAHL; 2015 to June 2017) using search terms adapted for individual databases: ("Infant-Nutrition"/all subheadings OR Infant Formula OR milk OR formula OR trophic feeding OR minimal enteral nutrition OR gut priming), plus database-specific limiters for randomised controlled trials (RCTs) and neonates (see Appendix 1). We did not apply language restrictions.

We searched clinical trials registries for ongoing or recently completed trials (ClinicalTrials.gov (clinicaltrials.gov); the World Health Organization International Clinical Trials Registry Platform (www.who.int/clinical-trials-registry-platform/); and the ISRCTN registry (www.isrctn.com/)).

Searching other resources

We searched abstracts from annual meetings of the Pediatric Academic Societies (1993 to 2017), the European Society for Paediatric Research (1995 to 2016), the UK Royal College of Paediatrics and Child Health (2000 to 2017), and the Perinatal Society of Australia and New Zealand (2000 to 2016). Trials reported only as abstracts were eligible if sufficient information was available from the report or through contact with study authors to fulfil the inclusion criteria.

Standard search methods

PubMed: ((infant, newborn[MeSH] OR newborn OR neonate OR neonatal OR premature OR low birth weight OR VLBW OR LBW or infan* or neonat*) AND (randomized controlled trial [pt] OR controlled clinical trial [pt] OR randomized [tiab] OR placebo [tiab] OR drug therapy [sh] OR randomly [tiab] OR trial [tiab] OR groups [tiab]) NOT (animals [mh] NOT humans [mh]))

Embase: (infant, newborn or newborn or neonate or neonatal or premature or very low birth weight or low birth weight or VLBW or LBW or Newborn or infan* or neonat*) AND (human not animal) AND (randomized controlled trial or controlled clinical trial or randomized or placebo or clinical trials as topic or randomly or trial or clinical trial)

CINAHL: (infant, newborn OR newborn OR neonate OR neonatal OR premature OR low birth weight OR VLBW OR LBW or Newborn or infan* or neonat*) AND (randomized controlled trial OR controlled clinical trial OR randomized OR placebo OR clinical trials as topic OR randomly OR trial OR PT clinical trial)

Cochrane Library: (infant or newborn or neonate or neonatal or premature or preterm or very low birth weight or low birth weight or VLBW or LBW)

Appendix 3. Risk of bias tool

Sequence generation (checking for possible selection bias). Was the allocation sequence adequately generated?

For each included study, we categorised the method used to generate the allocation sequence as:

- low risk (any truly random process, e.g. random number table; computer random number generator);
- high risk (any non-random process, e.g. odd or even date of birth; hospital or clinic record number); or
- unclear risk.

Allocation concealment (checking for possible selection bias). Was allocation adequately concealed?

For each included study, we categorised the method used to conceal the allocation sequence as:

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

Blinding of participants and personnel (checking for possible performance bias). Was knowledge of the allocated intervention adequately prevented during the study?

For each included study, we categorised the methods used to blind study participants and personnel from knowledge of which intervention a participant received. We assessed blinding separately for different outcomes or class of outcomes. We categorised the methods as:

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

Blinding of outcome assessment (checking for possible detection bias). Was knowledge of the allocated intervention adequately prevented at the time of outcome assessment?

For each included study, we categorised the methods used to blind outcome assessment. We assessed blinding separately for different outcomes or class of outcomes. We categorised the methods as:

- low risk for outcome assessors;
- high risk for outcome assessors; or
- unclear risk for outcome assessors.

Incomplete outcome data (checking for possible attrition bias through withdrawals, dropouts, protocol deviations). Were incomplete outcome data adequately addressed?

For each included study and for each outcome, we described the completeness of data, including attrition and exclusions from the analysis. We noted whether attrition and exclusions were reported; 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 categorised the methods as:

- low risk (< 20% missing data);
- high risk (≥ 20% missing data); or
- unclear risk.

Selective reporting bias. Are reports of the study free of the suggestion of selective outcome reporting?

For each included study, we described how we investigated the possibility of selective outcome reporting bias and what we found. For studies for which study protocols were published in advance, we compared the prespecified outcomes versus outcomes reported in the published results. If the study protocol was not published in advance, we contacted the study authors to gain access to the study protocol. We assessed the methods as:

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

Other sources of bias. Was the study apparently free of other problems that could put it at a high risk of bias?

For each included study, we described any important concerns we had about other possible sources of bias (e.g. whether there was a potential source of bias related to the specific study design, or if the trial was stopped early due to some data-dependent process). We assessed whether each study was free of other problems that could put it at risk of bias as:

- low risk;
- high risk; or
- unclear risk.

If necessary, we explored the impact of the level of bias through the undertaking of sensitivity analyses.

WHAT'S NEW



Date	Event	Description
20 October 2020	New search has been performed	This updates the review 'Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants' (Oddie 2017).
20 October 2020	New citation required but conclusions have not changed	The updated search identified four new trials for inclusion (Fayyaz 2020; Hasshemi 2018; Montealegre-Pomar 2021; Saha 2019).

HISTORY

Protocol first published: Issue 4, 1998 Review first published: Issue 4, 1998

Date	Event	Description
11 January 2011	New citation required and conclusions have changed	New data and an increased total number of participating infants (to 496) narrowed confidence intervals for the estimates of effect and modified the implications for practice and research.
15 December 2010	New search has been performed	This updates the review 'Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants', which was published in the Cochrane Database of Systematic Reviews, Issue 2, 2008 (McGuire 2008).
		We updated the search in December 2010 and included one new trial (Krishnamurthy 2010).
		We included the following new co-authors on the review team: Jessie Morgan and Lauren Young.
13 February 2008	New citation required but conclusions have not changed	We added the following new review authors: Sarah Bombell and William McGuire.
2 February 2008	New search has been performed	This updates the review 'Rapid versus slow rate of advancement of feedings for promoting growth and preventing necrotizing enterocolitis in parenterally fed low-birth-weight infants', by Kennedy and Tyson, which was published in the Cochrane Data- base of Systematic Reviews, Issue 2, 2000 (Kennedy 2000).
		We modified the title to read 'Slow advancement of enteral feed volumes to prevent necrotising enterocolitis in very low birth weight infants', and added the following new review authors: Sarah Bombell and William McGuire. We made the following changes to the original protocol.
		 We defined 'slow' rate of feed advancement as daily increments up to 24 mL/kg (body weight). We restricted the population to very low birth weight and very preterm infants. We added mortality, adverse neurodevelopment, growth parameters, and infection rates as outcomes of interest.
		We updated the search in December 2007. We included one new trial (Salhotra 2004), and excluded one previously included trial (Book 1976).



Date	Event	Description
		The findings and implications for practice and research of this re- view have not changed overall.
11 January 2008	Amended	We converted the review to new review format.

CONTRIBUTIONS OF AUTHORS

Drs Oddie, Young, and McGuire updated the search, independently determined the eligibility of identified studies, assessed the methodological quality of the included trials, and extracted relevant information and data.

DECLARATIONS OF INTEREST

SO has no conflict of interest to declare.

LY has no conflict of interest to declare.

WM was a clinical investigator for SIFT 2016 (LY and SO extracted data for this study). SIFT 2016 was funded by the National Institute for Health Research (NIHR) (UK).

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- Bradford Neonatal, Bradford Royal Infirmary, UK

Logistical

External sources

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• Vermont Oxford Network, USA

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• The Gerber Foundation, USA

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We made the following changes to the previous publication of the review (Oddie 2017).

- We updated the certainty of evidence and the risk of bias tool in Appendix 2.
- We revised the text in the following sections:
- How the intervention might work
- Assessment of reporting biases
- Sensitivity analysis

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- Allocation (selection bias)
- Blinding (performance bias and detection bias)
- Incomplete outcome data (attrition bias)
- Selective reporting (reporting bias)
- Other potential sources of bias
- We added new external sources of support.
- We developed a new search strategy. The previous search methods are shown in Appendix 2.

INDEX TERMS

Medical Subject Headings (MeSH)

*Enteral Nutrition; *Enterocolitis, Necrotizing [epidemiology] [prevention & control]; Infant, Premature; Infant, Very Low Birth Weight; Parenteral Nutrition [adverse effects]

MeSH check words

Female; Humans; Infant, Newborn; Pregnancy