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Single induction dose of etomidate versus other induction agents for endotracheal intubation in critically ill patients (Review)

Bruder EA, Ball IM, Ridi S, Pickett W, Hohl C

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[Intervention Review]

Single induction dose of etomidate versus other induction agents for endotracheal intubation in critically ill patients

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ABSTRACT

Background

The use of etomidate for emergency airway interventions in critically ill patients is very common. In one large registry trial, etomidate was the most commonly used agent for this indication. Etomidate is known to suppress adrenal gland function, but it remains unclear whether or not this adrenal gland dysfunction affects mortality.

Objectives

The primary objective was to assess, in populations of critically ill patients, whether a single induction dose of etomidate for emergency airway intervention affects mortality.

The secondary objectives were to address, in populations of critically ill patients, whether a single induction dose of etomidate for emergency airway intervention affects adrenal gland function, organ dysfunction, or health services utilization (as measured by intensive care unit (ICU) length of stay (LOS), duration of mechanical ventilation, or vasopressor requirements).

We repeated analyses within subgroups defined by the aetiologies of critical illness, timing of adrenal gland function measurement, and the type of comparator drug used.

Search methods

We searched the Cochrane Central Register of Controlled Trials (CENTRAL); MEDLINE; CINAHL; EMBASE; LILACS; International Pharmaceutical Abstracts; Web of Science; the Database of Abstracts of Reviews of Effects (DARE); and ISI BIOSIS Citation indexSM on 8 February 2013. We reran the searches in August 2014. We will deal with any studies of interest when we update the review.

We also searched the Scopus database of dissertations and conference proceedings and the US Food and Drug Administration Database. We handsearched major emergency medicine, critical care, and anaesthesiology journals.

We handsearched the conference proceedings of major emergency medicine, anaesthesia, and critical care conferences from 1990 to current, and performed a grey literature search of the following: Current Controlled Trials; National Health Service – The National Research Register; ClinicalTrials.gov; NEAR website.

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Selection criteria

We included randomized controlled trials in patients undergoing emergency endotracheal intubation for critical illness, including but not limited to trauma, stroke, myocardial infarction, arrhythmia, septic shock, hypovolaemic or haemorrhagic shock, and undifferentiated shock states. We included single (bolus) dose etomidate for emergency airway intervention compared to any other rapid-acting intravenous bolus single-dose induction agent.

Data collection and analysis

Refinement of our initial search results by title review, and then by abstract review was carried out by three review authors. Full-text review of potential studies was based on their adherence to our inclusion and exclusion criteria. This was decided by three independent review authors. We reported the decisions regarding inclusion and exclusion in accordance with the PRISMA statement.

Electronic database searching yielded 1635 potential titles, and our grey literature search yielded an additional 31 potential titles. Duplicate titles were filtered leaving 1395 titles which underwent review of their titles and abstracts by three review authors. Sixty seven titles were judged to be relevant to our review, however only eight met our inclusion criteria and seven were included in our analysis.

Main results

We included eight studies in the review and seven in the meta-analysis. Of those seven studies, only two were judged to be at low risk of bias. Overall, no strong evidence exists that etomidate increases mortality in critically ill patients when compared to other bolus dose induction agents (odds ratio (OR) 1.17; 95% confidence interval (CI) 0.86 to 1.60, 6 studies, 772 participants, moderate quality evidence). Due to a large number of participants lost to follow-up, we performed a post hoc sensitivity analysis. This gave a similar result (OR 1.15; 95% CI 0.86 to 1.53). There was evidence that the use of etomidate in critically ill patients was associated with a positive adrenocorticotropic hormone (ACTH) stimulation test, and this difference was more pronounced at between 4 to 6 hours (OR 19.98; 95% CI 3.95 to 101.11) than after 12 hours (OR 2.37; 95% CI 1.61 to 3.47) post-dosing. Etomidate's use in critically ill patients was associated with a small increase in SOFA score, indicating a higher risk of multisystem organ failure (mean difference (MD) 0.70; 95% CI 0.01 to 1.39, 2 studies, 591 participants, high quality evidence), but this difference was not clinically meaningful. Etomidate use did not have an effect on ICU LOS (MD 1.70 days; 95% CI -2.00 to 5.40, 4 studies, 621 participants, moderate quality evidence), hospital LOS (MD 2.41 days; 95% CI -7.08 to 11.91, 3 studies, 152 participants, moderate quality evidence), duration of mechanical ventilation (MD 2.14 days; 95% CI -1.67 to 5.95, 3 studies, 621 participants, moderate quality evidence), or duration of vasopressor use (MD 1.00 day; 95% CI -0.53 to 2.53, 1 study, 469 participants).

Authors' conclusions

Although we have not found conclusive evidence that etomidate increases mortality or healthcare resource utilization in critically ill patients, it does seem to increase the risk of adrenal gland dysfunction and multi-organ system dysfunction by a small amount. The clinical significance of this finding is unknown. This evidence is judged to be of moderate quality, owing mainly to significant attrition bias in some of the smaller studies, and new research may influence the outcomes of our review. The applicability of these data may be limited by the fact that 42% of the patients in our review were intubated for "being comatose", a population less likely to benefit from the haemodynamic stability inherent in etomidate use, and less at risk from its potential negative downstream effects of adrenal suppression.

PLAIN LANGUAGE SUMMARY

Etomidate for sedating critically ill people during emergency endotracheal intubation

Review question

Does a single dose of etomidate increase mortality or complications in people who are critically ill and undergoing emergency endotracheal intubation?

Background

People who are critically ill often need help breathing. One way to do this is called endotracheal intubation. This involves placing a tube into the windpipe (trachea) and having a ventilator (breathing machine) help the patient breathe.

People are often given sedative agents during endotracheal intubation to make them unaware of the procedure. Many sedative agents cause a potentially harmful drop in blood pressure.

Etomidate is commonly used to sedate patients before endotracheal intubation because it has minimal effects on blood pressure. However, when someone is given etomidate their adrenal glands do not function as well. This may be harmful to them.

Study characteristics

We looked at the evidence up to February 2013 and found 1666 studies. We included eight studies in our review and seven studies (involving 772 patients) in our meta-analysis. The studies involved people who were in an unstable condition and critically ill. They were given one



dose of etomidate or another sedative agent for endotracheal intubation. We reran the search in August 2014. We will deal with any studies of interest when we update the review.

Results

No strong evidence exists to suggest that etomidate, when compared to other bolus dose induction agents, increases mortality in critically ill patients. We must be careful in interpreting this finding because only large studies would be able to show a difference in mortality. So far, no such study has been completed.

Etomidate does seem to impair adrenal gland functioning. Functioning is impaired most between four and six hours after etomidate is given.

Sequential Organ Failure Assessment (SOFA) scores are used to find out how badly someone's organs are failing. Using etomidate results in worse SOFA scores but this difference is small and not clinically meaningful.

The effects of impaired adrenal gland functioning and higher SOFA scores on people's health is unknown. Using etomidate does not seem to increase the length of time someone is in hospital (including an intensive care unit), the length of time a person is connected to a mechanical ventilator (a machine to assist with breathing), or the use of vasopressors (medicines to increase blood pressure).

Quality of the evidence

Most of the evidence was moderate quality. This is mainly because some small studies we looked at did not check up on people adequately after they were intubated.

Most people that were involved in one study were intubated because they were in a coma. These people comprise 42% of those involved in the studies we looked at. People in a coma are unlike other critically ill people because they may not benefit to the same extent from having stable blood pressure during endotracheal intubation, which etomidate provides, nor are they at high risk from impaired adrenal gland function compared to other critically ill patients, for example those with severe infection.

SUMMARY OF FINDINGS

Summary of findings for the main comparison. Etomidate versus all other induction agents for endotracheal intubation in critically ill patients

Etomidate versus all other induction agents for endotracheal intubation in critically ill patients

Patient or population: patients with endotracheal intubation in critically ill patients **Settings:**

Intervention: etomidate versus all other induction agents

Outcomes	Outcomes Illustrative comparative risks* (95% CI)			No of partici-	Quality of the Commer	nts
	Assumed risk	Corresponding risk	(3370 CI)	(studies)	(GRADE)	
	Control	Etomidate versus all other induc- tion agents				
Mortality	Study population		OR 1.17	772 (6 studies)	⊕⊕⊕⊝ moderate1	
	296 per 1000	330 per 1000 (265 to 402)				
	Moderate	rate				
	241 per 1000	271 per 1000 (214 to 337)				
ACTH stimulation test	Study population	OR 2.37	519 (3 studies)	⊕⊕⊕⊙ moderate1		
is considered positive if the change in serum cor- tisol level was less than	228 per 1000	411 per 1000 (322 to 506)	()	(000000)	inouclute	
9 μg/dL (248 nmol/L) af- ter the administration of	Moderate					
250 μg of cosyntropin Follow-up: 24 hours	167 per 1000	322 per 1000 (244 to 410)				
Random serum cortisol levels (µg/dL) after re- ceiving intervention	The mean random serum cortisol levels (μg/dl) after receiving intervention in the control groups was 21 to 28 μg/dL	The mean random serum cortisol levels (μg/dl) after receiving inter- vention in the intervention groups was 4.96 lower (8.06 to 1.86 lower)		105 (3 studies)	⊕⊕⊕⊝ moderate ¹	

Organ system dysfunc- tion Sequential Organ Fail- ure Assessment (SOFA) Score. Scale from: 1 to 24	The mean organ system dysfunction in the control groups was 9.6	The mean organ system dysfunction in the intervention groups was 0.7 higher (0.01 to 1.39 higher)		469 (1 study)	⊕⊕⊕⊕ high
ICU length of stay (days)	The mean ICU length of stay (days) in the control groups was 3 to 22 days	The mean ICU length of stay (days) in the intervention groups was 1.7 higher (2 lower to 5.4 higher)		621 (3 studies)	⊕⊕⊕⊝ moderate ¹
Hospital length of stay (days)	The mean hospital length of stay (days) in the con- trol groups was 6.4 to 10 days	The mean hospital length of stay (days) in the intervention groups was 2.41 higher (7.08 lower to 11.91 higher)		152 (2 studies)	⊕⊕⊕⊝ moderate ¹
Duration of mechanical ventilation (days)	The mean duration of me- chanical ventilation (days) in the control groups was 1.5 to 13 days	The mean duration of mechanical ventilation (days) in the intervention groups was 2.14 higher (1.67 lower to 5.95 higher)		621 (3 studies)	⊕⊕⊕⊝ moderate ¹
*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (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; OR: Odds ratio					

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹ A significant number of patients were lost to follow-up

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BACKGROUND

More than 60% of all emergency airway interventions in the United States (US) use etomidate as the bolus induction agent (Sivilotti 2003) owing to its favourable haemodynamic properties and ease of dosing. Data from the National Emergency Airway Registry (NEAR) show that etomidate is the most commonly used induction agent for emergency airway intervention (Sivilotti 2003). The NEAR registry currently collects data on emergency department airway interventions from 25 hospitals in five countries. At the time of Sivilotti's publication in 2003, the database included 20 hospitals in the US, one in Canada, and one in Asia. Etomidate was used in 62% of all rapid sequence intubations (RSI) (1468 of 2380 patients) (Sivilotti 2003). Benzodiazepines were used 18% of the time and were the next most common agents used. In the NEAR database, 63% of emergency physicians and 26% of anaesthesiologists used etomidate (Sivilotti 2003). In the Corticosteroid Therapy of Septic Shock (CORTICUS) trial examining steroid use in septic intensive care unit (ICU) patients, 19% of these critically ill patients received etomidate (Cuthbertson 2009).

Etomidate suppresses the normal cortisol production of the adrenal glands through inhibition of 11-beta-hydroxylase. In one trial, 94% of patients receiving etomidate failed to respond to a corticotropin stimulation test (Absalom 1999), but whether this suppression leads to clinically relevant outcomes is uncertain. Specifically, it is unknown if a single dose of etomidate affects the mortality of critically ill patients. Given its widespread popularity, there is potential for harm if mortality is increased through its use. Should there not be a negative effect on mortality, then many clinicians can be reassured that this medication is safe for use in the sickest patients. Either outcome of this review would be practice changing for the large number of physicians caring for critically ill patients.

Description of the condition

Many critically ill patients require airway control, where an endotracheal tube is placed within the trachea. This procedure is painful and difficult to tolerate when awake, so most clinicians sedate patients for the procedure. Indications for airway control in critical illness are numerous but broadly include airway protection when airway patency is threatened by distorted anatomy or the patient's level of consciousness. This therapy is also used to support respiratory failure and to allow mechanical ventilation in patients with haemodynamic instability, including patients with septic shock and other critical illnesses.

Patients requiring this therapy typically have abnormal vital signs including hypotension, tachycardia, hypoxia, or an altered level of consciousness. Insults to the patient's central nervous system (CNS) and other vital organs may be exacerbated if the induction agents worsen hypotension. Rapidly acting induction agents decrease critically ill patients' blood pressure further through vasodilatory effects or direct myocardial suppression, or both.

Description of the intervention

When faced with the decision to obtain airway control in critically ill patients, clinicians must weigh the benefits and potential harms of a multitude of pharmacological agents. They must then apply this decision to a complex and dynamic physiological state in patients sensitive to further physiological insults. Several classes of agents are used to sedate critically ill patients, each with their own benefits and weaknesses.

Etomidate is a short-acting intravenous (IV) medication used for anaesthesia induction and sedation. Single-dose etomidate is commonly used to facilitate endotracheal intubation in critically ill patients because etomidate is less likely to cause a harmful drop in blood pressure than other induction agents, after an induction dose of 0.3 mg/kg IV. After this dose, there are minimal changes in heart rate, stroke volume, or cardiac output; and mean arterial blood pressure may decrease up to 15% because of decreases in systemic vascular resistance. Etomidate (1-(1-phenylethyl)-1H-imidazole-5carboxylic acid ethyl ester) is a carboxylated imidazole derivative used for the induction of general anaesthesia. Following a standard dose (0.3 mg/kg), hypnosis occurs in less than one minute and is maintained for 4 to 10 minutes by producing gamma-aminobutyric acid (GABA)-like effects on the CNS.

Benzodiazepines (midazolam, lorazepam, diazepam, etc.) induce CNS depression through GABA effects. Midazolam is a commonly used rapid-acting benzodiazepine for RSI. Midazolam, when administered as an IV bolus (0.05 to 0.15 mg/kg) for induction of anaesthesia has an onset of action of one to two minutes. Duration of action after an induction dose of 0.15 mg/kg IV to young healthy volunteers was 17 minutes to awakening. The clinical effects of midazolam can be prolonged in elderly patients or patients with impaired renal or hepatic function.

Propofol (2,6-diisopropylphenol) is presumed to exert its sedativehypnotic effects through a GABA receptor interaction. At a standard dose of 1.5 to 2.5 mg/kg IV, anaesthesia is induced in less than one minute (10 to 50 seconds) and is maintained for five minutes. Propofol produces decreases in systemic blood pressure and has a negative inotropic effect. Bradycardia and asystole have been observed after induction of anaesthesia with propofol, potentially owing to a decrease in sympathetic nervous system activity that results in a predominance of parasympathetic activity.

Opiate derivatives (morphine, fentanyl, remifentanil, etc.) in large doses have been used as the sole anaesthetic in critically ill patients. Morphine and hydromorphone can cause histamine release, which causes hypotension owing to peripheral vasodilation. Fentanyl and remifentanil do not cause release of histamine. Dose, metabolism, elimination, and side effects vary depending on the opioid administered. Morphine, hydromorphone, and remifentanil may produce mild decreases in systemic blood pressure and heart rate. Fentanyl may produce bradycardia.

Ketamine is another induction agent with favourable haemodynamic and kinetic profiles. At IV bolus doses of 1 to 2 mg/kg for induction, dissociation occurs within 30 to 60 seconds with a duration of action of 10 to 20 minutes. Ketamine produces cardiovascular effects that resemble sympathetic nervous system stimulation. The mechanisms for these ketamine-induced cardiovascular effects are complex. Direct stimulation of the CNS leading to increased sympathetic nervous system outflow seems to the most important mechanism for cardiovascular stimulation (Wong 1974). This may result in an increase in systemic and pulmonary arterial blood pressure, heart rate, cardiac output, cardiac workload, and therefore myocardial oxygen demand. Ketamine also has a direct negative cardiac inotropic effect. This effect is usually overshadowed by central sympathetic stimulation



How the intervention might work

The use of etomidate infusions in the ICU setting has been largely abandoned secondary to adrenal suppression and increased mortality. This is thought to be mediated by etomidate's transient, reversible suppression of 11-beta-hydroxylase, an enzyme responsible for the production of active steroids from the adrenal glands (de Jong 1984). There is a growing body of literature supporting the hypothesis that even single-dose etomidate causes adrenal suppression (de Jong 1984; Hildreth 2008; Jabre 2009; Zed 2006). The association of adrenal dysfunction in septic shock patients, and the possible survival benefit from exogenous steroids, has been debated and investigated in the critical care literature (Annane 2002; Annane 2004; Cronin 1995; Sprung 2008).

Whether the adrenal dysfunction has a causal role in mortality, or is simply another indicator of organ dysfunction, remains unclear.

Why it is important to do this review

Debate remains regarding the clinical effects of single dose etomidate in critically ill patients. Systematic reviews on the subject have led to conflicting results.

Hohl et al pooled data from seven studies examining the effects of etomidate in critically ill patients. None of the individual studies were powered to detect a mortality difference, and a pooled odds ratio (OR) estimate of mortality showed no statistical difference (Hohl 2010).

Albert et al published a systematic review of 19 etomidate trials (Albert 2011). The authors concluded that strong evidence exists for an increased relative risk for etomidate-induced adrenal suppression. They also stated that weak evidence exists for any association between etomidate and mortality. The authors very correctly assert that the mortality conclusions are weak based on: "a preponderance of non-randomized trials and heterogeneity of studies". In their review, Albert et al combined clinically heterogeneous data from 15 retrospective, observational, and non-randomized trials with four prospective randomized trials. The conclusion that etomidate use is associated with greater mortality in critically ill patients must be interpreted with caution (Albert 2011).

Chan and colleagues also published a meta-analysis of randomized controlled trials and observational studies examining the effects of etomidate on adrenal insufficiency and all-cause mortality in septic patients. In this meta-analysis, they report a pooled relative risk of 1.20 (95% confidence interval (CI) 1.02 to 1.42) for mortality, and a pooled relative risk of 1.33 (95% CI 1.22 to 1.46) for the development of adrenal insufficiency (Chan 2012). The practice of combining observational and randomized data is methodologically questionable, and the results must be interpreted with caution.

While it is clear that etomidate is associated with transient adrenal suppression, the literature has yet to answer whether or not this effect is clinically meaningful. It is also unclear whether the immediate haemodynamic safety benefits of etomidate outweigh its potential harm from transient adrenal suppression.

Given the vast number of doses administered on an annual basis to critically ill patients, there exists a risk of harm if mortality is affected by etomidate-induced adrenal suppression. Should no harm be identified, then physicians can be reassured that the use of an otherwise favourable drug is safe.

OBJECTIVES

The primary objective was to assess, in populations of critically ill patients, whether a single induction dose of etomidate for emergency airway intervention affects mortality.

The secondary objectives were to address, in populations of critically ill patients, whether a single induction dose of etomidate for emergency airway intervention affects adrenal gland function, organ dysfunction, or health services utilization (as measured by ICU length of stay (LOS), duration of mechanical ventilation, or vasopressor requirements).

We repeated analyses within subgroups defined by the aetiologies of critical illness, timing of adrenal gland function measurement, and the type of comparator drug used.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomized controlled trials (RCTs).

We excluded non-randomized or quasi-randomized trials.

Types of participants

We included adult and paediatric patients undergoing emergency endotracheal intubation (defined as endotracheal intubation for an unstable clinical condition) for critical illness, including but not limited to: trauma, stroke, myocardial infarction, arrhythmia, septic shock, hypovolaemic or haemorrhagic shock, and undifferentiated shock states.

We excluded elective anaesthesia induction in stable patients.

Types of interventions

We included a single (bolus) dose of etomidate for emergency airway intervention compared to any other rapid-acting IV bolus single-dose induction agent (ketamine, midazolam, propofol, thiopental, etc.).

We excluded etomidate infusions and etomidate use for indications other than airway intervention (for example procedural sedation).

Types of outcome measures

Primary outcomes

All-cause mortality. Mortality data at 30 days (including sensitivity analysis of death before 24 hours, up to 7 days, and 28 days) will be reported if available.



Secondary outcomes

- 1. Mortality at 30 days within groups of patients with adrenal gland dysfunction, as available in published reports.
- Adrenal gland dysfunction (at times < 4 hours, between 4 to 6 hours, between 6 to 12 hours, and > 12 hours from etomidate dose) as described by Marik 2008, defined as:
 - a. random cortisol level < 10 μg/dL (276 nmol/L);
 - b. failed adrenocorticotropic hormone (ACTH) stimulation tests where the delta cortisol is < 9 μ g/dL (248 nmol/L) after a 250 μ g cosyntropin administration (or body surface-area appropriate dose in the paediatric population).
- 3. Organ dysfunction:
 - a. Sequential Organ Failure Assessment (SOFA) score;
 - b. other validated systems for reporting organ dysfunction.
- 4. ICU LOS (sensitivity analysis stratified by patients who died before 24 hours, within 7 days, and within 28 days).
- 5. Duration of mechanical ventilation (sensitivity analysis stratified by patients who died before 24 hours, within 7 days, and within 28 days).
- 6. Vasopressor requirements (duration in days of any vasoactive medication infusion).

Search methods for identification of studies

Electronic searches

We searched the Cochrane Central Register of Controlled Trials (CENTRAL) (2012, Issue 12); MEDLINE via OvidSP (1950 to 8 February 2013); CINAHL (EBSCOhost) (1982 to 8 February 2013); EMBASE via OvidSP (1980 to 8 February 2013); LILACS (BIREME) (1982 to 8 February 2013); International Pharmaceutical Abstracts (1970 to 8 February 2013); Web of Science (1980 to 8 February 2013); the Database of Abstracts of Reviews of Effects (DARE) (8 February 2013); and ISI BIOSIS Citation indexSM (1969 to 8 February 2013).

We also searched the Scopus database of dissertations and conference proceedings (1980 to 8 February 2013) and the US Food and Drug Administration (FDA) Database (1980 to 8 February 2013).

We did not limit the selection of studies by region of publication or by language.

We adopted the MEDLINE search strategy for searching all other databases (see Appendix 1 for detailed search strategies).

We reran the search in August, 2014. We will deal with any studies of interest when we update the review.

Searching other resources

We handsearched the following medical journals from 2000 to February 2013:

- Annals of Emergency Medicine;
- Academic Emergency Medicine;
- Canadian Journal of Emergency Medicine;

- Emergency Medicine Clinics of North America;
- Journal of Emergency Medicine;
- Anesthesiology;
- Canadian Journal of Anesthesia;
- Anesthesia and Analgesia;
- British Journal of Anaesthesia;
- Journal of Trauma;
- Intensive Care Medicine;
- Critical Care Medicine;
- Chest:
- American Journal of Respiratory and Critical Care Medicine.

We searched the conference proceedings of major emergency medicine, anaesthesia, and critical care conferences from 1990 to February 2013 to identify data published in abstract form only.

A grey literature search included electronic searches of the following clinical trial registry websites:

- Current Controlled Trials;
- National Health Service The National Research Register;
- ClinicalTrials.gov;
- NEAR website.

We contacted authors of all ongoing trials for unreported data. We contacted drug manufacturers and asked them to provide any published or unpublished data, however these publications were not provided to us, citing that, "Clinical studies not already published are proprietary information. Regretfully, for this reason, this data cannot be provided" (Lloyd 2013 [pers comm]).

The bibliographies of all relevant retrieved articles identified in the search above were handsearched for any missed studies.

Data collection and analysis

Selection of studies

Refinement of our initial search results by title review and then by abstract review was carried out by three review authors (EB, IB, SR). We decided the final inclusion for full-text review by majority vote (two out of three).

Full-text review of potential studies was based on their adherence to our inclusion and exclusion criteria. This was decided by three independent review authors (EB, IB, SR). We resolved disagreements by open discussion and consensus agreement, with the principle review author (EB) making the final decision. If required, we contacted the authors of studies to clarify their eligibility for inclusion and whether their publication was a duplicate report of a single study. If any doubt remained, the default was inclusion for data extraction.

We reported the decisions regarding inclusion and exclusion in accordance with the PRISMA statement (Figure 1). We analysed multiple reports of a single study as a single study.



Figure 1. Study flow diagram.



Figure 1. (Continued)

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We were not blinded to the authors or journals of publication.

Data extraction and management

Three review authors (EB, IB, SR) independently extracted the data from studies using a data extraction form (see Appendix 2). We (EB, IB, SR) pilot tested the data extraction form on 10 articles selected at random from our initial search strategy and applied it to studies that both met, and did not meet, our inclusion criteria. We did not modify the data extraction methodology and form after this pilot test.

We extracted the following data, according to outcome:

- all-cause mortality;
- Adrenal gland dysfunction:
- random serum cortisol levels,
- positive ACTH stimulation tests;
- Health services utilization:
 - hospital LOS in days (mean, standard deviation (SD), numbers (n); as well as median and interquartile range (IQR) as reported),
 - ICU LOS in days (mean, SD, n; as well as median and IQR as reported),
 - duration of mechanical ventilation in days (mean, SD, n; as well as median and IQR as reported),
 - vasopressor requirements,

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 duration of any vasopressor requirement in days (mean, SD, n).

Assessment of risk of bias in included studies

We performed risk of bias assessments using the 'Risk of bias' tool described in Chapter 8 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011).

We assessed each trial according to the quality domains of random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting, and any other potential threats to validity. We judged each criterion regarding its risk of bias and recorded an assessment of the magnitude and direction of each source of bias. Data points for the risk of bias assessment are part of our data extraction form, attached as Appendix 2.

We considered a trial to have a low risk of bias if all domains were assessed as adequate. We considered a trial to have a high risk of bias if one or more domain was assessed as inadequate or unclear.

We report the 'Risk of bias' table (Figure 2) as part of the table 'Characteristics of included studies' and present a 'Risk of bias summary' figure (Figure 3), which details all of the judgements made for all included studies in the review.



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





	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Incomplete outcome data (attrition bias): Mortality	Incomplete outcome data (attrition bias): Adrenal Gland Dysfunction	Incomplete outcome data (attrition bias): Hospital LOS	Incomplete outcome data (attrition bias): ICU LOS	Incomplete outcome data (attrition bias): Duration of mechanical Ventillation	Incomplete outcome data (attrition bias): Vasopressor requirements	Incomplete outcome data (attrition bias): Organ Dysfunction	Incomplete outcome data (attrition bias): All outcomes	Selective reporting (reporting bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)
Absalom 1999	?	?	•	•	•	•	•	•	•	•	•	?	?
Cinar 2010	?	?	?	?	•	?	?	•	•	?	•	•	•
Hildreth 2008	?	•	•	•	•	•	•	•	•	•	•	●	?
Jabre 2009	•	•	•	•	•	•	•	•	•	•	•	•	•
Jacoby 2006	•	•	•	•	•	•	•	•	•	•	•	•	•
Koksal 2013	•	?	?	?	•	•	•	•	?	?	•	?	•
Schenarts 2001	•	•	•	•	•	•	•	•	•	•	•	•	•
Tekwani 2010	•	•	•	•	•	•	•	•	•	•	•	•	•

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

We assessed the risk of bias for each outcome independently. Studies were not weighted by risk of bias for our analysis.

Measures of treatment effect

We reported the odds ratio (OR) for dichotomous data (mortality, ACTH stimulation tests). We reported the mean difference (MD) and SD for continuous data with the same unit of measure, and the standardized mean difference (SMD) for continuous data that were reported using different units of measure (hospital and ICU LOS, duration of mechanical ventilation, duration of vasopressor requirements, serum cortisol levels). Serum cortisol levels reported in nmol/L were converted to μ g/dL, where possible, using the 'SIU Conversion Calculator' embedded within the Micromedex 2.0 system by Truven Health Analytics Inc (Truven 2013).

Variables with non-normal distributions, reported as medians with IQRs, were assumed to be normally distributed for the purposes of this analysis. The median was assumed to be an acceptable estimation of the mean, and the IQR was considered to be 1.35 times the SD. This process is described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011) and has been used by other review authors in the past (Zacharias 2013).

Unit of analysis issues

The unit of analysis was the patient.

In studies with multiple treatment arms, only interventions relevant to our review were included for analysis, and a description of these studies is included in the 'Characteristics of included studies' table. We combined all control groups (for example propofol, ketamine, and benzodiazepines) for comparison with the intervention (etomidate). We analysed each pair-wise comparison separately as subgroup analyses for mortality. For dichotomous outcomes, such as mortality, we divided the number of events and the total number of patients in proportions similar to the proportions of the total number of participants per experimental group.

The nature of the intervention precluded case cross-over trial designs. Therefore, we did not encounter any of these study designs. Cluster-randomized trials were included in our analysis if they reported the outcome data ignoring the cluster design for the total number of individuals.

Dealing with missing data

Whenever possible, we contacted the authors of studies and asked for primary data (Cinar 2010; Hildreth 2008; Koksal 2013).

Where this was not possible, we performed an intention-to-treat (ITT) analysis on mortality data where we assumed that all missing data represented patient deaths in both the intervention and control groups.

Assessment of heterogeneity

We only pooled data meta-analytically for clinically homogenous data reporting the same outcome measures. For all pooled data, we used the Chi² statistic, degrees of freedom, and the l² statistic to assess the degree of statistical heterogeneity across the studies. If the l² was < 40%, we reported the results of pooled data using meta-analytic techniques. If the l² was > 40%, we used sensitivity analysis to explore potential causes for the heterogeneity.

Assessment of reporting biases

We did not employ a funnel plot of included studies to ensure that reporting bias did not significantly affect our findings as fewer than 10 trials were included (Egger 1997) in this review.

Data synthesis

We conducted a meta-analysis, using a random-effects model, for studies with similar design and interventions.

We report ORs with 95% confidence intervals (CIs) for dichotomous variables such as mortality, and MDs with 95% CIs for continuous variables. When different scales or units were used to report continuous data, we calculated and reported the SMDs.

Subgroup analysis and investigation of heterogeneity

We performed subgroup analysis on the following:

- comparator drug;
- different aetiologies of critical illness (i.e. septic, cardiogenic, trauma, undifferentiated).

We had planned to perform a subgroup analysis on paediatric patients, but no reports of paediatric patients were identified.

Sensitivity analysis

We planned to conduct sensitivity analyses on the following data:

- mortality timing, less than 24 hours versus 7 days, 28 days, and all;
- ICU LOS stratified by timing of mortality;
- duration of mechanical ventilation stratified by timing of mortality;
- sensitivity analysis where studies assessed as being at high risk of bias were excluded.

We were unable to conduct these sensitivity analyses due to unavailable data.

We conducted sensitivity analysis to assess the impact of excluding poor quality studies, assessed as being at high risk of bias. Only two studies were judged to be at low risk of bias (Jabre 2009; Tekwani 2010).

Summary of findings tables

We used the principles of the GRADE system (Guyatt 2008) to assess the quality of the body of evidence associated with specific outcomes (mortality, adrenal gland dysfunction, organ dysfunction, ICU LOS) in our review and constructed a summary of findings (SoF) table using the GRADE software. The GRADE approach appraises the quality of a body of evidence based on the extent to which we can be confident that an estimate of effect or association reflects the item being assessed. The quality of a body of evidence considers within study risk of bias (methodological quality), the directness of the evidence, heterogeneity of the data, precision of effect estimates, and risk of publication bias.



RESULTS

Description of studies

See the 'Characteristics of included studies' and 'Characteristics of excluded studies' tables below.

Results of the search

Electronic database searching yielded 1635 potential titles, and our grey literature search yielded an additional 31 potential tiles. Duplicate titles were filtered leaving 1395 titles which underwent review of their titles and abstracts by three review authors (EB, IB, SR). We judged 68 titles to be relevant to our review, however only eight met our inclusion criteria. One title (Cinar 2010) met our inclusion criteria but could not be included in our meta-analysis because it was a poster at a research meeting that did not present its numerical data. This paper presented results like "Groups' mean serum cortisol and 11b-deoxycortisol concentrations were similar before and five minutes after intubation (P < 0.05). The groups were not significantly different with regards to intubation conditions, length of ICU stay, duration of mechanical ventilation, and mortality (P < 0.05)" (Cinar 2010). Multiple attempts were made to contact both the primary and supervising authors, but no reply was received. As a result we included seven studies in our metaanalysis (Figure 1).

We reran our search on 22 August 2014 and identified an additional three studies of interest (Driver 2014; Freund 2014; Punt 2014). We will deal with these studies of interest when we update the review.

Included studies

We included eight studies in our review and seven studies in our meta-analysis. All seven were RCTs of critically ill patients requiring emergency airway intervention, with a combined census of 772 patients. The studies were conducted in North American (Hildreth 2008; Schenarts 2001; Tekwani 2010) or European (Absalom 1999; Jabre 2009; Koksal 2013) tertiary care centres, or in pre-hospital settings in the United States (Jacoby 2006). All studies randomized patients to receive etomidate or other single-dose induction agents including thiopentone (Absalom 1999), fentanyl and midazolam (Hildreth 2008), ketamine (Jabre 2009), or midazolam alone (Jacoby 2006; Koksal 2013; Schenarts 2001; Tekwani 2010).

There were differences in the overall mortality rates reported within the studies, which may suggest clinical heterogeneity. Most studies reported an overall mortality rate of 23% to 38% (Absalom 1999; Jabre 2009; Jacoby 2006; Tekwani 2010), however two studies reported overall mortality rates of 3% (Schenarts 2001) and 7% (Hildreth 2008).

Excluded studies

We excluded 17 studies. The majority of these studies were excluded because of their observational design or they were retrospective reviews of charts or databases. Please refer to 'Characteristics of excluded studies'. Three studies warranted specific discussion (Asehnoune 2012; Cherfan 2011; Cuthbertson 2009).

Asehnoune reported a substudy of the HYPOLYTE trial, which intended to report the impact of etomidate on the rate of hospital acquired pneumonia in trauma patients intubated for more than 48 hours. They also performed ACTH stimulation tests. While the use of etomidate was prospectively collected, the participants in this trial were randomized, in a double blind, placebo controlled manner, to receive either hydrocortisone or placebo. These patients were not randomized to receive etomidate or other induction agents (Asehnoune 2012).

Cherfan evaluated the effect of low-dose steroids on septic patients intubated with etomidate. This was a nested cohort study within a RCT, which evaluated the use of low-dose hydrocortisone in septic cirrhotic patients. Mortality was compared between groups who did, and did not, receive etomidate (Cherfan 2011).

Cuthbertson's study received a lot of attention when published. It concluded that etomidate use was associated with a statistically significant increase in mortality (61.0% versus 44.6%) and the authors cautioned against using etomidate. This study was an a priori substudy of the CORTICUS trial, a multi-centre, double blind, placebo controlled RCT comparing patients who received either hydrocortisone or placebo. Cuthbertson's report compared mortality in patients who received etomidate within 72 hours of enrolment to those who did not. It may stand to reason that the higher mortality rate seen in the etomidate group was due to their more severe underlying illness and was not an effect of the drug. Using multivariate analysis, the authors controlled for the severity of illness and still showed an increase in mortality (42.7% versus 30.5%) in the patients who received etomidate. This study was excluded from our analysis because individual patients were randomized to receive either placebo or hydrocortisone, and the data relating to the exposure to etomidate was observational (Cuthbertson 2009).

Studies awaiting classification

Three studies (Driver 2014; Freund 2014; Punt 2014) are awaiting classification (see 'Characteristics of studies awaiting classification'). We will deal with these studies of interest when we update the review.

Risk of bias in included studies

Please refer to our 'Characteristics of included studies', risk of bias graph (Figure 2), and risk of bias summary (Figure 3) for our judgements regarding each study's risk of bias. Overall, only two studies (Jabre 2009; Tekwani 2010) were judged to be at low risk of bias.

Allocation

In only three of the eight trials (Cinar 2010; Jacoby 2006; Tekwani 2010) were the healthcare worker(s) administering the experimental treatment blinded. In two trials (Absalom 1999; Koksal 2013) it is unclear, whereas in one trial (Jabre 2009), despite there being an appropriate randomization process in place, the physicians administering the study medication were unblinded. This raised significant concerns as unblinded healthcare workers with preconceived beliefs about the safety and efficacy of the study drugs may have altered the randomization sequence. The concealment methodology utilized would make it unlikely that the allocation sequence of the participants could have been altered. Cinar 2010 described their methodology as a "prospective, randomized, double blinded study", but did not elaborate on their methods of sequence generation or concealment. This was likely adequate, but the risk of bias remained unclear in the absence of this description.



Blinding

In three of the eight trials (Cinar 2010; Jacoby 2006; Tekwani 2010) the healthcare worker(s) administering the experimental treatment were blinded. In one trial (Jabre 2009), although the workers administering the study drug were unblinded, the teams providing all subsequent care were blinded to treatment allocation. It was unclear to what degree knowledge of treatment allocation may have altered patient care. An obvious possibility was the administration of steroids to patients who received etomidate. Differences in fluid resuscitation, administration of vasoactive medications, or other critical care unit treatments were unlikely but cannot be ruled out.

Virtually all eight included trials' outcomes were objective (ACTH stimulation test results, serum cortisol levels, mortality, etc) and the trials (Cinar 2010; Jacoby 2006) that did use more subjective outcomes (ease of intubation, vocal cord visualization, etc.) blinded the assessors. On this basis, detection bias was not a concern when interpreting the conclusions from this review.

Incomplete outcome data

One of the advantages of critical care research is the ability to obtain excellent patient follow-up (as patients are intubated and ventilated). Yet, many of these trials are threatened by the significant proportions of patients lost to follow-up. In one unblinded trial (Hildreth 2008) over 50% of eligible patients were not enrolled (a significant proportion of which were because of "protocol violations"), and no additional explanation was provided. In one trial (Jacoby 2006), close to 20% of enrolled patients were lost to follow-up. In another trial (Schenarts 2001), 9/31 patients were excluded.

In two of the larger trials (Jabre 2009; Tekwani 2010), follow-up was excellent with only two patients lost to follow-up. One of the smaller trials (Absalom 1999) also had very good follow-up, with only one patient lost to follow-up.

In six of the eight trials, follow-up was either poor or not described. Fortunately the large trials did demonstrate excellent followup, thereby reducing concern about attrition bias.

Selective reporting

In the current controversy regarding the haemodynamic effect of etomidate, the adrenal suppression, and ultimately any effect it may or may not have on mortality, any finding is noteworthy and publishable. It was unlikely that negative studies would not be submitted for publication (and we did not identify any examples of this during our handsearch of conference abstracts), or that negative findings within a study (so-called within study reporting bias) would not be described.

Other potential sources of bias

Designing a trial to compare single-dose etomidate to another agent(s) is fraught with challenges. The heterogeneity of the patient populations that require emergency intubation, managing informed consent for an emergent treatment, and blinding the healthcare team to medications that require emergent dose titration are impediments to bias free studies.

Another important source of bias is the characteristics of the study population. The concern of negative consequences from adrenal

suppression is widely believed to be most important in critically ill patients in shock, particularly those with sepsis. In one recent trial (Jabre 2009), the majority of patients were intubated for airway protection due to neurologic compromise. It was not surprising that etomidate-induced adrenal suppression did not affect SOFA score as these patients were neither septic nor in shock.

Effects of interventions

See: Summary of findings for the main comparison Etomidate versus all other induction agents for endotracheal intubation in critically ill patients

Primary outcome: mortality

None of the included studies reported mortality as their primary outcome, and none were powered to detect a mortality difference. No individual trial showed a significant difference in mortality. There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 2.80, df = 5 (P = 0.73); l² = 0%). We employed a randomeffects model for meta-analysis, describing the OR and 95% CI. The pooled result of 390 patients receiving etomidate, compared to 382 patients receiving other induction agents, showed no difference in mortality (OR 1.17; 95% CI 0.86 to 1.60) (see Analysis 1.1).

We performed a subgroup analysis comparing etomidate to individual comparator agents. Only one study (Absalom 1999) compared etomidate to thiopentone. We employed a randomeffects model for meta-analysis, describing the OR and 95% CI. When 17 patients receiving etomidate were compared to 17 patients receiving thiopentone, there was no significant difference in mortality (OR 1.94; 95% CI 0.38 to 9.88). Four studies compared etomidate to midazolam (with or without fentanyl) (Hildreth 2008; Jacoby 2006; Schenarts 2001, Tekwani 2010). There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 2.28, df = 3 (P = 0.52); $I^2 = 0\%$). We employed a randomeffects model for meta-analysis, describing the OR and 95% CI. The pooled result of 139 patients receiving etomidate, compared to 130 patients receiving midazolam, showed no difference in mortality (OR 1.06; 95% CI 0.61 to 1.83). Only one study (Jabre 2009) compared etomidate to ketamine. We employed a random-effects model for meta-analysis, describing the OR and 95% CI. When 234 patients receiving etomidate were compared to 235 patients receiving ketamine, there was no statistically significant difference in mortality (OR 1.20; 95% CI 0.81 to 1.76) (see Analysis 1.2).

We performed a subgroup analysis comparing the effects of etomidate on different etiologies of critical illness. Only one study (Tekwani 2010) studied etomidate specifically in patients with septic shock. We employed a random-effects model for metaanalysis, describing the OR and 95% CI. When 61 septic patients receiving etomidate were compared to 59 septic patients receiving other agents, there was no significant difference in mortality (OR 1.34; 95% CI 0.64 to 2.81). Only one study (Hildreth 2008) studied etomidate specifically in traumatized patients. We employed a random-effects model for meta-analysis, describing the OR and 95% CI. When 18 trauma patients receiving etomidate were compared to 12 trauma patients receiving other agents, there was no statistically significant difference in mortality (OR 3.79; 95% CI 0.17 to 86.13). No studies involving patients in cardiogenic shock were identified. Four studies reported results for patients with undifferentiated, or unreported, etiologies of critical illness (Absalom 1999; Jabre 2009; Jacoby 2006; Schenarts 2001). There

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was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 2.07, df = 3 (P = 0.56); I² = 0%). We employed a randomeffects model for meta-analysis, describing the OR and 95% CI. The pooled result of 311 undifferentiated patients receiving etomidate, compared to 311 undifferentiated patients receiving other induction agents, showed no difference in mortality (OR 1.12; 95% CI 0.79 to 1.58) (see Analysis 1.3).

Only two studies were judged to be at low risk of bias (Jabre 2009; Tekwani 2010). We analysed mortality when only these two studies were included. There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 0.07, df = 1 (P = 0.79); I² = 0%). We employed a random-effects model for meta-analysis, describing the OR and 95% CI. The pooled result of 295 patients receiving etomidate, compared to 294 patients receiving other induction agents, showed no difference in mortality (OR 1.23; 95% CI 0.87 to 1.73) (see Analysis 1.4).

Due to a large proportion of patients lost to follow-up, we performed a post hoc sensitivity analysis where all missing patients were assumed to have died. There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 2.64, df = 5 (P = 0.76); l² = 0%). We employed a random-effects model for meta-analysis, describing the OR and 95% CI. The pooled result of 425 patients receiving etomidate, compared to 408 patients receiving other induction agents, showed no difference in mortality (OR 1.15; 95% CI 0.86 to 1.53) (see Analysis 1.5).

The 30 day mortality data were not available. We reported mortality data at the individual study conclusion.

Secondary outcomes

ACTH stimulation test

Four studies reported dichotomous data regarding ACTH stimulation tests (Absalom 1999; Hildreth 2008; Jabre 2009; Schenarts 2001). ACTH stimulation tests were considered to be positive or negative according to standardized criteria (Marik 2008). There was some statistical heterogeneity in this comparison (Tau² = 0.42; Chi^2 = 6.55, df = 4 (P = 0.16); I^2 = 39%). We employed a random-effects model for meta-analysis, describing the OR and 95% CI. No studies reported the results of ACTH stimulation tests performed less than four hours after receiving etomidate, or the comparator. Two studies (Hildreth 2008; Schenarts 2001) reported the results of ACTH stimulation tests performed between four and six hours after induction. There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 0.20, df = 1 (P = 0.66); $I^2 = 0\%$). We employed a random-effects model for meta-analysis, describing the OR and 95% CI. The pooled result of 28 patients receiving etomidate, compared to 20 patients receiving other induction agents, showed a significant difference in the proportion of positive ACTH stimulation tests (OR 19.98; 95% CI 3.95 to 101.11). Only one study (Schenarts 2001) reported ACTH stimulation test results performed between 6 and 12 hours post-induction. None of the patients in either treatment arm demonstrated a positive ACTH stimulation test (n etomidate = 7, n other = 7), so the treatment effect could not be estimated. Three studies (Absalom 1999; Jabre 2009; Schenarts 2001) reported results of ACTH stimulation tests performed more than 12 hours after induction. There was no statistical heterogeneity in this comparison (Tau² = 0.00; Chi² = 0.03, df = 2 (P = 0.99); $I^2 = 0\%$). We employed a random-effects model for meta-analysis, describing the OR and 95% CI. The pooled result of 260 patients receiving etomidate, compared to 259 patients receiving other induction agents, showed a significant difference in the proportion of positive ACTH stimulation tests (OR 2.37; 95% CI 1.61 to 3.47) (see Analysis 1.6).

Random serum cortisol levels

Three studies (Absalom 1999; Hildreth 2008; Koksal 2013) reported continuous data regarding random serum cortisol levels. There was statistical heterogeneity in this comparison (Tau² = 2.38; Chi² = 4.46, df = 3 (P = 0.22); l² = 33%). We employed a random-effects model for meta-analysis, describing the MD and 95% CI. The pooled result of 75 patients receiving etomidate, compared to 70 patients receiving other induction agents, showed a significant difference in the random serum cortisol level (MD -4.96; 95% CI -8.06 to -1.86) (see Analysis 1.7). For Analysis 1.7.2 and Analysis 1.7.4, the sample sizes in the Koksal 2013 paper were adjusted from 20 to 10 in each of the treatment and control groups in order to avoid double counting the sample size in the overall estimate of effect. The mean and SD values were left unchanged.

SOFA score

One study (Jabre 2009) reported the effects of etomidate on SOFA score. Statistical heterogeneity was not calculated. We employed a random-effects model for meta-analysis, describing the MD and 95% CI. The pooled result of 234 patients receiving etomidate, compared to 235 patients receiving other induction agents, showed a significant difference in the SOFA score (MD 0.70; 95% CI 0.01 to 1.39) favouring other induction agents over etomidate (see Analysis 1.8).

ICU length of stay (LOS)

Three studies (Hildreth 2008; Jabre 2009; Tekwani 2010) reported the effects of etomidate on ICU LOS. There was significant statistical heterogeneity in this comparison (Tau² = 8.82; Chi² = 12.89, df = 2 (P = 0.002); I² = 84%). We employed a random-effects model for metaanalysis, describing the MD and 95% CI. The pooled result of 315 patients receiving etomidate, compared to 306 patients receiving other induction agents, showed no significant difference in ICU LOS (MD 1.70; 95% CI -2.00 to 5.40) (see Analysis 1.9).

Hosptial length of stay (LOS)

Two studies (Hildreth 2008; Tekwani 2010) reported the effects of etomidate on hospital LOS. There was significant statistical heterogeneity in this comparison (Tau² = 42.71; Chi² = 10.85, df = 1 (P = 0.0010); I² = 91%). We employed a random-effects model for meta-analysis, describing the MD and 95% CI. The pooled result of 81 patients receiving etomidate, compared to 71 patients receiving other induction agents, showed no statistically significant difference in hospital LOS (MD 2.41; 95% CI -7.08 to 11.91) (see Analysis 1.10).

Duration of mechanical ventilation

Three studies (Hildreth 2008; Jabre 2009; Tekwani 2010) reported the effects of etomidate on duration of mechanical ventilation. There was significant statistical heterogeneity in this comparison (Tau² = 9.59; Chi² = 14.75, df = 2 (P = 0.0006); l² = 86%). We employed a random-effects model for meta-analysis, describing the MD and 95% CI. The pooled result of 315 patients receiving etomidate, compared to 306 patients receiving other induction agents, showed no significant difference in the duration of mechanical ventilation (MD 2.14; 95% CI -1.67 to 5.95) (see Analysis 1.11).



Duration of vasopressor support

Only one study (Jabre 2009) reported the effects of etomidate on the duration of vasopressor support. We employed a randomeffects model for meta-analysis, describing the MD and 95% CI. The pooled result of 234 patients receiving etomidate, compared to 235 patients receiving other induction agents, showed no significant difference in the duration of vasopressor support (MD 1.00; 95% CI -0.53 to 2.53) (see Analysis 1.12).

DISCUSSION

Summary of main results

There are insufficient data to conclude that etomidate use is associated with an increased risk of harm, or to conclude that it is safe.

Mortality

A non-significant trend towards increased mortality was observed. Overall, no strong evidence exists to suggest that etomidate, when compared to other bolus dose induction agents, increases mortality in critically ill patients (OR 1.17; 95% CI 0.86 to 1.60).

No strong evidence exists to suggest that etomidate increases mortality when compared to thiopentone (OR 1.94; 95% CI 0.38 to 9.88), midazolam with or without fentanyl (OR 1.06; 95% CI 0.61 to 1.83), or ketamine (OR 1.20; 95% CI 0.81 to 1.76).

No strong evidence exists to suggest that etomidate increases mortality in the subset of critically ill patients with septic shock (OR 1.34; 95% CI 0.64 to 2.81), trauma (OR 3.79; 95% CI 0.17 to 86.13), cardiogenic shock (no data), or undifferentiated critical illness (OR 1.12; 95% CI 0.79 to 1.58).

This lack of strong evidence for increased mortality due to the use of etomidate remains when only studies judged to be at low risk of bias are included in the analysis (OR 1.23; 95% CI 0.87 to 1.73). Due to a large number of participants lost to follow-up, we performed a post hoc sensitivity analysis where all patients lost to follow up were assumed to have died. Also, in this scenario no strong evidence of increased mortality associated with etomidate use was identified (OR 1.15; 95% CI 0.86 to 1.53).

Therefore, with the existing evidence to date, despite a trend towards harm, etomidate use does not seem to cause an increase in mortality when used for emergency endotracheal intubation in patients with critical illness. This evidence is of moderate quality.

Adrenal function

Overall, strong evidence of moderate quality suggests that the use of etomidate in critically ill patients increases the likelihood of a positive ACTH stimulation test, and this difference is greater at four to six hours (OR 19.98; 95% CI 3.95 to 101.11) than after 12 hours post-induction (OR 2.37; 95% CI 1.61 to 3.47). Likewise, the use of etomidate in critically ill patients appears to be associated with statistically significant reductions in random serum cortisol levels (MD -4.96; 95% CI -8.06 to -1.86). Again, this difference is more pronounced at four to six hours (MD -5.73; 95% CI -10.24 to -1.23) after induction compared to greater than 12 hours (MD -3.78; 95% CI -10.01 to 2.44) when it is no longer statistically significant. This evidence is of moderate quality.

Organ system dysfunction (SOFA score)

The use of etomidate in critically ill patients is associated with a statistically significant increase in SOFA score, indicating a higher risk of multi-system organ failure when compared to other induction agents (MD 0.70; 95% CI 0.01 to 1.39). This evidence is of high quality. It should be noted that Jabre (Jabre 2009) concluded that no difference in SOFA score could be attributed to the use of etomidate, while we have concluded that a difference does exist. This is due to the fact that Jabre 2009 utilized one decimal place in their data analysis and concluded that the mean difference in SOFA score of 0.7 (95% CI 0.0 to 1.4) was insignificant because the confidence interval included 0.0. When the raw data were entered into RevMan for analysis, data were calculated to the second decimal place, which reached statistical significance because the lower confidence interval was 0.01. The SOFA score is a numerical surrogate score for organ system dysfunction that ranges from 0 (good organ function) to 24 (worse organ function). The difference in the maximum SOFA score in the etomidate group was 10.3 compared to 9.6 in the ketamine group, with a difference of 0.7 between groups (Jabre 2009). Regardless of whether this difference was statistically significant it is not clinically meaningful.

Health services utilization

Non-significant trends towards increased health services utilization were observed. However, when compared to other induction agents, no strong evidence exists to suggest that etomidate increases ICU length of stay (MD 1.70; 95% CI -2.00 to 5.40), hospital length of stay (MD 2.41; 95% CI -7.08 to 11.91), duration of mechanical ventilation (MD 2.14; 95% CI -1.67 to 5.95), or duration of vasopressor use (MD 1.00; 95% CI -0.53 to 2.53). This evidence is of moderate quality.

Overall completeness and applicability of evidence

We searched for randomized trials that compared etomidate to another induction agent or a combination of agents for endotracheal intubation of critically ill patients. We identified only seven published papers and one unpublished paper. For an agent as commonly used as etomidate this is a very small number of studies. Despite this small number of studies we are confident that all eligible reports were identified. The total number of patients from all studies was only 772.

In the largest trial included in our analysis (Jabre 2009) 69% of patients in both arms were intubated for being comatose, while the minority were intubated for 'shock', 'acute respiratory failure', or 'other' reasons. These obtunded patients (n = 324) account for 42% of all patients in our review. Although it is accepted that patients obtunded secondary to intracranial catastrophes and drug intoxications are critically ill, this is not the group most likely to benefit from the haemodynamic stability inherent in etomidate use, nor are they the patients most at risk from its potential negative downstream effects of adrenal suppression. Applying conclusions from a small trial with a high preponderance of obtunded patients to patients in shock should be done with caution. Larger studies of patients in shock are still required to assess the appropriateness of etomidate's use.

Each of the individual trials included in our review had point estimates of effect suggesting increased mortality with the use of etomidate, but none were powered sufficiently to detect a difference. All confidence intervals crossed the line of no effect. The

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fact that all of the studies, and our pooled estimate, cross the line of no effect is likely because all studies were underpowered. It is possible that with more studies the 95% confidence intervals will no longer cross the line of no effect.

Quality of the evidence

Using the principals of the GRADE system, the quality of the data from randomized controlled trials (with the exception of SOFA score) were downgraded to 'moderate'. This is due mainly to methodological limitations and the high proportions of patients lost to follow-up in the smaller studies (Hildreth 2008; Jacoby 2006; Schenarts 2001). Other methodological limitations involving randomization and allocation concealment supported this downgrade as well. In particular, the authors of one study did not describe their sequence generation or concealment (Absalom 1999); another unblinded study (Hildreth 2008) only enrolled half of the eligible patients, with almost half of the excluded patients excluded due to 'protocol violations'. A third study (Schenarts 2001) was unblinded and did not describe their allocation concealment procedures. In addition, one third of the patients in this unblinded study were excluded from the analysis, raising a significant concern in relation to bias (Schenarts 2001).

We did not downgrade our quality of evidence based on directness as all trials directly compared etomidate to a comparator agent in directly generalizable populations.

We did not downgrade our quality of evidence based on heterogeneity or inconsistency as we demonstrated very little statistical heterogeneity, and all studies reported similar results. There were differences in the overall mortality rates reported within the studies, which may suggest clinical heterogeneity. Most studies reported an overall mortality rate of 23% to 38% (Absalom 1999; Jabre 2009; Jacoby 2006; Tekwani 2010), however two studies reported overall mortality rates of 3% (Schenarts 2001) and 7% (Hildreth 2008).

We did not downgrade our quality of evidence based on imprecision as the reported confidence intervals were sufficiently narrow.

We did not downgrade our quality of evidence based on the potential for publication bias. While we did not employ a funnel plot (we did not identify 10 studies for inclusion), all studies reported similar findings and the risk of publication bias was judged to be low.

Potential biases in the review process

We believe that our search methodology was sufficient to minimize the chances that any study was missed. We did not employ a funnel plot because fewer than 10 studies were included in our analysis. We believe that our three investigator, data abstraction process also minimized the chances for individual beliefs to influence our results. Some of our statistical analysis has led to transformed data. Specifically, we were unable to enter non-normally distributed data into RevMan as they were reported as medians and IQRs. Instead, we replicated the data transformation process of other investigators (Zacharias 2013) and roughly transformed these data into means and standard deviations. While all data were transformed in the same manner, the exact precision of the results may be altered by this process. If future editions of RevMan allow median and IQR data to be analysed primarily, we will report the data as such in future review updates. This data transformation affects data regarding hospital and ICU length of stay, duration of ventilator support, and duration of vasopressor support. It does not affect data on adrenal gland dysfunction or mortality.

Agreements and disagreements with other studies or reviews

Three other systematic reviews have been published on the topic of etomidate use in critically ill patients. Our results were similar to Hohl et al, where the pooled data were from seven studies examining the effects of etomidate in critically ill patients. None of the individual studies were powered to detect a mortality difference, and a pooled odds ratio estimate of mortality showed no statistical difference (Hohl 2010).

Albert et al published a systematic review of 19 etomidate trials (Albert 2011). The authors concluded that strong evidence exists for an increased relative risk for etomidate-induced adrenal suppression. They also stated that weak evidence exists for any association between etomidate and mortality. The authors very correctly assert that the mortality conclusions are weak based on "a preponderance of non-randomized trials and heterogeneity of studies". In their review, Albert et al combined clinically heterogeneous data from 15 retrospective, observational, and non-randomized trials with four prospective randomized trials. Conclusions drawn from combining such heterogeneous trials should be interpreted with caution.

Chan and colleagues also published a meta-analysis of randomized controlled trials and observational studies examining the effects of etomidate on adrenal insufficiency and all-cause mortality in septic patients. In this meta-analysis, they reported a pooled relative risk of 1.20 (95% CI 1.02 to 1.42) for mortality, and a pooled relative risk of 1.33 (95% CI 1.22 to 1.46) for the development of adrenal insufficiency (Chan 2012). The combination of observational and randomized data may be subject to bias and the results must be interpreted appropriately.

AUTHORS' CONCLUSIONS

Implications for practice

Based on the currently available evidence, the use of etomidate in critically ill patients does not seem to increase mortality, organ system dysfunction, or healthcare resource utilization. This observation must be interpreted with caution owing to the moderate quality of evidence, the patient population described above, and the fact that no randomized trial to date has been adequately powered to detect a mortality difference. As in non-critically ill patients, it appears that etomidate's use does negatively affect adrenal gland function but it is unclear whether or not this adrenal gland dysfunction influences patient outcomes in the first four to six hours (more so than after 12 hours). Again, this must be interpreted with caution, acknowledging the moderate quality of evidence. The use of etomidate is associated with an increase in SOFA score indicating an increased risk of organ system dysfunction. This increase in SOFA score is not clinically meaningful. With respect to healthcare resource utilization, no strong evidence exists to suggest that etomidate increases ICU length of stay, hospital length of stay, or duration of mechanical ventilation.



Implications for research

Additional randomized trials of high quality are still required to confirm the safety of etomidate's use in critically ill patients. Due to the moderate quality of the existing evidence, new research may influence the outcomes of our review upon updating the review.

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CHARACTERISTICS OF STUDIES

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* Indicates the major publication for the study

Absalom 1999	
Methods	RCT comparing etomidate (n = 17) to thiopentone (n = 18)
Participants	35 patients: ASA ¹ \geq III, with 2 or more organ systems failure, and requiring admission to ICU



Absalom 1999 (Continued)

Induction with etomidate or thiopentone (unreported doses)

Interventions Outcomes

Adrenal gland function using ACTH stimulation test 24 hours post-intervention

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	"patients were randomly allocated to receive either etomidate or thiopen- tone" (p861). Sequence generation not described
Allocation concealment (selection bias)	Unclear risk	Sequence generation and concealment not described
Incomplete outcome data (attrition bias) Mortality	Low risk	One patient was lost from the thiopentone group. This is unlikely to affect re- sults
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Low risk	One patient was lost from the thiopentone group. This is unlikely to affect re- sults
Incomplete outcome data (attrition bias) Hospital LOS	Low risk	N/A
Incomplete outcome data (attrition bias) ICU LOS	Low risk	N/A
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	Low risk	N/A
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	N/A
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A
Incomplete outcome data (attrition bias) All outcomes	Low risk	One patient was lost from the thiopentone group. This is unlikely to affect re- sults
Selective reporting (re- porting bias)	Low risk	All relevant data reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No description of blinding reported, however the outcome is entirely objective



Absalom 1999 (Continued)

Blinding of outcome assessment (detection bias) All outcomes No description of blinding reported, however the outcome is entirely objective

Cinar 2010						
Methods	RCT comparing etomid	RCT comparing etomidate (n = 12) and ketamine (n = 10)				
Participants	22 ICU patients					
Interventions	Etomidate 0.3 mg/kg ve	ersus ketamine 2 mg/kg				
Outcomes	Intubating conditions, l tion, duration of mecha	haemodynamic response to intubation, total serum cortisol 5 min post-induc- anical ventilation, ICU length of stay, mortality				
Notes	Need more information. This was conference poster and no actual data is presented. Primary author and supervising author contacted by e-mail. Waiting on response. Unable to include in analysis due to lack of data					
Risk of bias						
Bias	Authors' judgement	Support for judgement				
Random sequence genera- tion (selection bias)	Unclear risk	"prospective, randomized, double blinded study" reported, but sequence gen- eration and concealment not described				
Allocation concealment (selection bias)	Unclear risk	"prospective, randomized, double blinded study" reported, but sequence gen- eration and concealment not described				
Incomplete outcome data (attrition bias) Mortality	Unclear risk	No description of incomplete data. Impacts on risk estimates not possible to describe, but bias possible				
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Unclear risk	No description of incomplete data. Impacts on risk estimates not possible to describe, but bias possible				
Incomplete outcome data (attrition bias)	Low risk	N/A				

Hospital LOS		
Incomplete outcome data (attrition bias) ICU LOS	Unclear risk	No description of incomplete data. Impacts on risk estimates not possible to describe, but bias possible
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	Unclear risk	No description of incomplete data. Impacts on risk estimates not possible to describe, but bias possible
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	N/A



Cinar 2010	(Continued)
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Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No description of incomplete data. Impacts on risk estimates not possible to describe, but bias possible
Selective reporting (re- porting bias)	Low risk	all relevant outcomes addressed
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	"prospective, randomized, double blinded study" reported, but no further de- scription
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	"prospective, randomized, double blinded study" reported. Exact nature of double blinding not fully specified, so effects on measurement remain un- known

Hildreth 2008

Methods	RCT comparing etomidate (n = 18) to fentanyl and midazolam combination (n = 12)	
Participants	Adult trauma patients requiring intubation < 48 hours post-injury	
Interventions	Etomidate 0.3 mg/kg versus fentanyl 100 μg and midazolam 5 mg	
Outcomes	Primary: Adrenal function (ACTH Stimulation test and random serum cortisol levels at 4 to 6 h after in- tervention). Secondary: vasopressor requirements, post-intubation haemodynamics, transfusion re- quirements, mortality, hospital LOS ² , ICU LOS ² , ventilator days	
Notes	Clarification about sequence generation, allocation concealment, and details about the protocol viola- tions have been sought from the author. Raw data for ACTH stimulation tests were provided by the au- thor. Risk of bias judgments may change once this information is received	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Reports "randomization packets" but no description of sequence genera- tion. "Medical personnel who participated in randomization procedures were trained". Effects on randomization unclear
Allocation concealment (selection bias)	Low risk	"Envelopes were half [etomidate] and half [Fentanyul/Midazolam] and sealed. Opened in ED or helicopter or ICU by person intubating. No way it could have been predicted prior to enrolment. Envelopes were identical." (Hildreth 2013 [pers comm])
Incomplete outcome data (attrition bias) Mortality	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias)	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In



Hildreth 2008 (Continued) Adrenal Gland Dysfunc- tion		an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias) Hospital LOS	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias) ICU LOS	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias) Vasopressor requirements	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A
Incomplete outcome data (attrition bias) All outcomes	High risk	This study reported 61 eligible patients, but only enrolled 30. Of the 31 pa- tients who were excluded, 14 (45%) were excluded for protocol violations. In an unblinded study, this raises significant concerns for the reviewers, and this study is judged at high risk of bias for attrition bias
Selective reporting (re- porting bias)	Low risk	All relevant data reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Non-blinded study. Unlikely to affect empiric outcomes relevant to this review, however, when considered in the context of the high rate of patient exclusion for protocol violation, the risk of bias due to unblinded assessors and person- nel is high
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Non-blinded study. Unlikely to affect empiric outcomes relevant to this review

Jabre 2009	
Methods	RCT comparing etomidate (n = 234) and ketamine (n = 235)
Participants	Adult pre-hospital, emergency department, or ICU patients requiring intubation for being comatose, shock, acute respiratory failure, or 'other'
Interventions	Etomidate 0.3 mg/kg versus ketamine 2 mg/kg
Outcomes	Primary: maximum Sequential Organ Failure Assessment (SOFA) score. Secondary: change in SOFA score, 28 day mortality, ICU LOS ² , ventilation duration, vasopressor use, ACTH stimulation test performed 0 to 48 h after intervention

Jabre 2009 (Continued)

Notes

ICU length of stay reported as "ICU free days at day 28", Vasopressor use reported as "Catecholamine free days until day 28", ventilation duration reported as "Mechanical ventilation free days at day 28". These variables were transformed by subtracting these results from 28 days to estimate the duration of each of these variables. This analysis assumes a single ICU visit, ventilation treatment, vasopressor use; 69% of patients in both arms were intubated for being comatose, while the minority were intubated for shock, acute respiratory failure, or 'other' reasons

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Utilized a computerized random number generator
Allocation concealment (selection bias)	Low risk	Study drugs were sealed in sequentially numbered, identical boxes containing the entire treatment for each patient. "The emergency physician enrolling pa- tients was aware of study group assignment. However nurses and intensivists in the intensive care unit were masked to the treatment assigned" (Jabre 2009. p294). While this single blind methodology raises concern, the allocation con- cealment was adequate to prevent enrolling physicians from altering the allo- cation sequence
Incomplete outcome data (attrition bias) Mortality	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) Hospital LOS	Low risk	N/A
Incomplete outcome data (attrition bias) ICU LOS	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Incomplete outcome data (attrition bias) All outcomes	Low risk	Missing data on 2 patients from each group, and one patient in the etomidate withdrew consent. Unlikely to affect results, given the small proportion of the sample size. Effect on risk estimates likely to be negligible
Selective reporting (re- porting bias)	Low risk	All relevant data reported



Jabre 2009 (Continued)

Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Emergency physicians giving the drugs were not blinded, but nurses and ICU physicians providing care for the duration of the study were blinded. Unlikely to affect the performance of participants and personnel, hence the effects of the intervention
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Emergency physicians giving the drugs were not blinded, but nurses and ICU physicians providing care for the duration of the study were blinded. Unlikely to affect the outcome assessment, and hence opportunity for information bias minimized

Jacoby 2	2006
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Methods	RCT comparing etomidate (n = 55) and midazolam (n = 55)	
Participants	Pre-hospital (EMS) intubation for respiratory distress or altered level of consciousness	
Interventions	Etomidate (20mg) versus midazolam (7 mg)	
Outcomes	Primary: intubation success. Secondary: vocal cord visualization, number of attempts or intubation dif- ficulty, post-intubation hypotension, vomiting, fasciculations, survival to discharge	

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Used a random number table
Allocation concealment (selection bias)	Low risk	Numbered syringes, identical to each other, volume corrected.
Incomplete outcome data (attrition bias) Mortality	High risk	115 patients enrolled in the study and 110 (55 per group) patients completed. Mortality data available for 88 (44 per group). Equal proportion in each group of lost patients (11 per group), but a description of lost patients with regards to mortality was not reported. Effects on risk estimates unclear due to this loss to follow-up. Selection bias possible, but direction of bias not predictable
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Low risk	N/A
Incomplete outcome data (attrition bias) Hospital LOS	Low risk	N/A
Incomplete outcome data (attrition bias) ICU LOS	Low risk	N/A
Incomplete outcome data (attrition bias)	Low risk	N/A



Jacoby 2006 (Continued) Duration of mechanical

Ventillation		
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	N/A
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A
Incomplete outcome data (attrition bias) All outcomes	Low risk	115 patients enrolled in the study, and 110 patients completed. Patients ac- counted for, and small proportion of lost patients unlikely to affect results
Selective reporting (re- porting bias)	Low risk	All relevant data reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinded. Master allocation lists were never accessed so blinding was not com- promised
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Blinded. Master allocation lists were never accessed so blinding was not com- promised

Koksal 2013

Methods	RCT comparing etomidate (n = 20), etomidate + methylprednisolone (n = 20), and midazolam (n = 20)	
Participants	Adult patients, ASA III-IV, requiring intubation in the ED or ICU	
Interventions	Group 1: etomidate 0.3 mg/kg, Group 2: etomidate 0.3 mg/kg AND methylprednisolone 2 mg/kg, Group 3: midazolam 0.5 mg/kg	
Outcomes	Primary: Random serum cortisol levels at 4 and 24 h. Secondary: post-intubation haemodynamics	
Notes	Group 2 was excluded from our analysis because methylprednisolone was given. All results from this study exclude the outcomes for the 20 patients in group 2. This is an unpublished study identified in our grey literature search. The author has kindly provided us with a copy of the manuscript for inclusion. The study is to be submitted for publication	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Computer generated randomization
Allocation concealment (selection bias)	Unclear risk	Awaiting details from author
Incomplete outcome data (attrition bias) Mortality	Unclear risk	Awaiting details from author

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Koksal 2013 (Continued)		
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Unclear risk	Awaiting details from author
Incomplete outcome data (attrition bias) Hospital LOS	Low risk	N/A
Incomplete outcome data (attrition bias) ICU LOS	Low risk	N/A
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	Low risk	N/A
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	N/A
Incomplete outcome data (attrition bias) Organ Dysfunction	Unclear risk	N/A
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Awaiting details from author
Selective reporting (re- porting bias)	Low risk	All outcomes from trial registration were reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	single blind study. Only outcome assessor was blind. Awaiting more details from author regarding allocation concealment
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	single blind study. Only outcome assessor was blind

Schenarts 2001	
Methods	RCT comparing etomidate (n = 16) and midazolam (n = 15)
Participants	Adult patients requiring intubation in a tertiary care emergency department
Interventions	Etomidate 0.3 mg/kg versus midazolam 0.05 to 0.1 mg/kg
Outcomes	Primary: ACTH Stimulation tests at 4, 12 and 24 hours after intervention. Secondary: mortality, random serum cortisol levels (at 4, 12, and 24 h), hospital LOS ² , ICU LOS ² , ventilator duration
Notes	



Schenarts 2001 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Computer generated randomization table
Allocation concealment (selection bias)	High risk	No description of allocation concealment reported. However, with the lack of blinding, in the setting of significant incomplete data (9 of 31 patients), the reviewers have considerable concern regarding the allocation concealment
Incomplete outcome data (attrition bias) Mortality	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Incomplete outcome data (attrition bias) Hospital LOS	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Incomplete outcome data (attrition bias) ICU LOS	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	N/A
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A
Incomplete outcome data (attrition bias) All outcomes	High risk	9 of 31 enrolled patients have incomplete data, and were excluded from analy- sis. Effects on risk estimates unclear, although could be substantial; risk of bias high
Selective reporting (re- porting bias)	Low risk	All relevant data reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded, and in the setting of a significant proportion of patients excluded due to incomplete data, the reviewers have considerable concern that the un- blinded nature of this study has led to bias in risk estimates
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Unblinded, and in the setting of a significant proportion of patients excluded due to incomplete data, the reviewers have considerable concern that the un- blinded nature of this study has led to bias in risk estimates



Tekwani 2010

Methods	RCT comparing Etomidate (n=63) and midazolam (n = 59)			
Participants	Septic adults requiring intubation in a tertiary care emergency department			
Interventions	Etomidate 0.3 mg/kg v	Etomidate 0.3 mg/kg versus midazolam 0.1 mg/kg		
Outcomes	Primary: Hospital LOS ² haemodynamics, vasoj	. Secondary: inhospital mortality, ICL LOS ² , ventilator duration, post-intubation pressor requirements		
Notes				
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Computer generated blocks of 10		
Allocation concealment (selection bias)	Low risk	Kits numbered and dispensed by an automated medication dispensing cabi- net. No description of the blocking factor were provided, but given the com- puter generated sequence generation and automated medication dispensing, we believe that participant anticipation of subsequent allocation would be dif- ficult, and that the subsequent risk of bias is low		
Incomplete outcome data (attrition bias) Mortality	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		
Incomplete outcome data (attrition bias) Adrenal Gland Dysfunc- tion	Low risk	N/A		
Incomplete outcome data (attrition bias) Hospital LOS	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		
Incomplete outcome data (attrition bias) ICU LOS	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		
Incomplete outcome data (attrition bias) Duration of mechanical Ventillation	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		
Incomplete outcome data (attrition bias) Vasopressor requirements	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		
Incomplete outcome data (attrition bias) Organ Dysfunction	Low risk	N/A		
Incomplete outcome data (attrition bias)	Low risk	Only 2 patients had incomplete data. Likely to have minimal effects on the strength and direction of risk estimates		



Tekwani 2010 (Continued) All outcomes

Selective reporting (re- porting bias)	Low risk	All relevant data reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Identical vials with volume-equivalent concentrations. Double blinded study
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Identical vials with volume-equivalent concentrations. Data collectors were blinded

1. American Society of Anesthesiologists (ASA) physical status

2. LOS = Length of stay

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Asehnoune 2012	This is a substudy of the HYPOLYTE (Roquilly 2011) multi-centred double blind RCT in which trauma patients were randomized to receive hydrocortisone or placebo
Baird 2009	Observational study
Borner 1985	Elective anaesthesia induction
Bramwell 2002	Not RCT
Cherfan 2011	This was a nested cohort study within an RCT evaluating hydrocortisone use in septic cirrhotic pa- tients
Cotton 2008	Retrospective registry study
Cuthbertson 2009	This was an a priori substudy of the CORTICUS trial (Sprung 2008). In this double blind RCT, patients were randomized to receive hydrocortisone or placebo
den Brinker 2008	Retrospective review
Dmello 2010	Retrospecitve study
Ehrman 2010	Retrospective cohort study
McPhee 2013	Retrospective cohort study
Mohammad 2006	Retrospective study
Morel 2011	Elective cardiac surgery induction
Ray 2007	Retrospective chart review
Vinclair 2008	Observational cohort study
Warner Kier 2009	Not RCT



Study

Reason for exclusion

Zed 2006

Observational study

Characteristics of studies awaiting assessment [ordered by study ID]

Driver 2014	
Methods	Randomized clinical trial. Single blind where ICU physicians but not ER physicians were blinded
Participants	n = 98. All patients receiving RSI
Interventions	Ketamine 2 mg/kg versus etomidate 0.3 mg/kg
Outcomes	Primary: 30 day mortality, or discharge, whichever occurred first. Secondary: first pass intuba- tion success, number of intubation attempts, number of post-intubation sedative boluses within 6 hours, post-intubation hypotension within 6 hours, post-intubation hypoxaemia within 2 hours
Notes	This is an interim report of their study. 98 patients have been enrolled. This reports the outcomes of 58 patients with available data

Freund 2014

Methods	Ancillary study. Off-shoot of KETASED study which was an RCT comparing ketamine and etomidate
Participants	n = 310. Patients who underwent pre-hospital RSI, and who had ACTH stimulation test within 24 h or ICU admission
Interventions	Patients in the KETASED study were randomized to ketamine or etomidate. This study groups pa- tients as having RAI or not, and then identifying risk factors based on this grouping, not on whether they received etomidate or ketamine
Outcomes	The objective of this study was to assess the effect of relative adrenal insufficiency (RAI) on prog- nostic outcomes after RSI, and factors associated with the onset of RAI
Notes	This was the same population as the Jabre 2009 study. This sample size is larger because this re- port includes the patients who were excluded by Jabre who had died before arrival to hospital, or who were discharged from the ICU before the 3 day mark

Punt 2014	
Methods	Single centre prospective open label study. Block randomized by ICU. Each ICU delivered the same agent for 10 months, and they switched to the other agent for an additional 10 months
Participants	n = 322. ICU patients requiring tracheal intubation
Interventions	Etomidate 0.2 to 0.3 mg/kg versus (S-ketamine 0.5 mg/kg + midazolam 2.5 mg)
Outcomes	Primary: 28 day mortality. Secondary: length of stay, usage of norepinephrine, and cortisol concen- trations
Notes	



DATA AND ANALYSES

Comparison 1. Etomidate versus all other induction agents

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Mortality: Data as reported	6	772	Odds Ratio (M-H, Random, 95% CI)	1.17 [0.86, 1.60]
2 Mortality: Data as reported - Sub- group analysis of comparator drugs	6	772	Odds Ratio (M-H, Random, 95% CI)	1.17 [0.86, 1.60]
2.1 Studies comparing etomidate and Thiopentone	1	34	Odds Ratio (M-H, Random, 95% CI)	1.94 [0.38, 9.88]
2.2 Studies comparing etomidate and midazolam (+/- fentanyl)	4	269	Odds Ratio (M-H, Random, 95% CI)	1.06 [0.61, 1.83]
2.3 Studies comparing etomidate and ketamine	1	469	Odds Ratio (M-H, Random, 95% CI)	1.20 [0.81, 1.76]
3 Mortality: Data as reported - Sub- group analysis of etiology of shock	6	772	Odds Ratio (M-H, Random, 95% CI)	1.17 [0.86, 1.60]
3.1 Septic Shock	1	120	Odds Ratio (M-H, Random, 95% CI)	1.34 [0.64, 2.81]
3.2 Trauma	1	30	Odds Ratio (M-H, Random, 95% CI)	3.79 [0.17, 86.13]
3.3 Cardiogenic Shock	0	0	Odds Ratio (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
3.4 Undifferentiated Critical Illness	4	622	Odds Ratio (M-H, Random, 95% CI)	1.12 [0.79, 1.58]
4 Mortality: Data as reported - Stud- ies judged to be at low risk of bias	2	589	Odds Ratio (M-H, Random, 95% CI)	1.23 [0.87, 1.73]
5 Mortality: Post Hoc ITT Analysis accounting for missing subjects	6	833	Odds Ratio (M-H, Random, 95% CI)	1.15 [0.86, 1.53]
6 ACTH Stimulation Test	4		Odds Ratio (M-H, Random, 95% CI)	Subtotals only
6.1 ACTH Stimulation test per- formed 4-6 h	2	48	Odds Ratio (M-H, Random, 95% CI)	19.98 [3.95, 101.11]
6.2 ACTH Stimulation Test Per- formed 6-12 h	1	14	Odds Ratio (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 ACTH Stimulation test per- formed >12h	3	519	Odds Ratio (M-H, Random, 95% CI)	2.37 [1.61, 3.47]



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7 Random Serum Cortisol levels (μg/dL) after receiving intervention	3	105	Mean Difference (IV, Random, 95% CI)	-4.96 [-8.06, -1.86]
7.1 Random Serum Cortisol Levels 4-6 h	2	50	Mean Difference (IV, Random, 95% CI)	-5.73 [-10.24, -1.23]
7.2 Random Serum Cortisol Levels >12h	2	55	Mean Difference (IV, Random, 95% CI)	-3.78 [-10.01, 2.44]
8 Organ System Dysfunction (SOFA Score)	1	469	Mean Difference (IV, Random, 95% CI)	0.70 [0.01, 1.39]
9 ICU Length of Stay	3	621	Mean Difference (IV, Random, 95% CI)	1.70 [0.00, 5.40]
10 Hospital Length of Stay	2	152	Mean Difference (IV, Random, 95% CI)	2.41 [-7.08, 11.91]
11 Duration of Mechanical Ventila- tion	3	621	Mean Difference (IV, Random, 95% CI)	2.14 [-1.67, 5.95]
12 Duration of Vasopressor Support	1	469	Mean Difference (IV, Random, 95% CI)	1.0 [-0.53, 2.53]

Analysis 1.1. Comparison 1 Etomidate versus all other induction agents, Outcome 1 Mortality: Data as reported.

Study or subgroup	Etomidate	Other Induc- tion Agents		Odds Ratio		Weight	Odds Ratio
	n/N	n/N	I	M-H, Random, 95%	CI		M-H, Random, 95% Cl
Absalom 1999	5/17	3/17			_	3.65%	1.94[0.38,9.88]
Hildreth 2008	2/18	0/12				0.99%	3.79[0.17,86.13]
Jabre 2009	81/234	72/235		 		64.54%	1.2[0.81,1.76]
Jacoby 2006	13/44	16/44		-+		12.11%	0.73[0.3,1.79]
Schenarts 2001	0/16	1/15			-	0.9%	0.29[0.01,7.76]
Tekwani 2010	26/61	21/59		+		17.81%	1.34[0.64,2.81]
Total (95% CI)	390	382		•		100%	1.17[0.86,1.6]
Total events: 127 (Etomidate), 113 (O	ther Induction Agen	ts)					
Heterogeneity: Tau ² =0; Chi ² =2.8, df=5	5(P=0.73); I ² =0%						
Test for overall effect: Z=1(P=0.32)							
	F	avours etomidate	0.01 0	.1 1	10 100	Favours other agents	

Analysis 1.2. Comparison 1 Etomidate versus all other induction agents, Outcome 2 Mortality: Data as reported - Subgroup analysis of comparator drugs.

Study or subgroup	Etomidate	Other Induc- tion Agents	Odds Ratio	Weight	Odds Ratio		
	n/N	n/N	M-H, Random, 95% Cl		M-H, Random, 95% CI		
1.2.1 Studies comparing etomidate a	and Thiopentone						
Absalom 1999	5/17	3/17		3.65%	1.94[0.38,9.88]		
Subtotal (95% CI)	17	17		3.65%	1.94[0.38,9.88]		
Total events: 5 (Etomidate), 3 (Other Ir	nduction Agents)						
Heterogeneity: Tau ² =0; Chi ² =0, df=0(Pe	<0.0001); l ² =100%						
Test for overall effect: Z=0.8(P=0.42)							
1.2.2 Studies comparing etomidate a	and midazolam (+/	- fentanyl)					
Hildreth 2008	2/18	0/12		- 0.99%	3.79[0.17,86.13]		
Jacoby 2006	13/44	16/44	+	12.11%	0.73[0.3,1.79]		
Schenarts 2001	0/16	1/15		0.9%	0.29[0.01,7.76]		
Tekwani 2010	26/61	21/59	-++	17.81%	1.34[0.64,2.81]		
Subtotal (95% CI)	139	130	•	31.81%	1.06[0.61,1.83]		
Total events: 41 (Etomidate), 38 (Othe	r Induction Agents)						
Heterogeneity: Tau ² =0; Chi ² =2.28, df=3	(P=0.52); I ² =0%						
Test for overall effect: Z=0.19(P=0.85)							
1.2.3 Studies comparing etomidate a	and ketamine						
Jabre 2009	81/234	72/235	<mark>₩</mark>	64.54%	1.2[0.81,1.76]		
Subtotal (95% CI)	234	235	•	64.54%	1.2[0.81,1.76]		
Total events: 81 (Etomidate), 72 (Othe	r Induction Agents)						
Heterogeneity: Not applicable							
Test for overall effect: Z=0.92(P=0.36)							
Total (95% CI)	390	382	•	100%	1.17[0.86,1.6]		
Total events: 127 (Etomidate), 113 (Ot	ner Induction Agent	ts)					
Heterogeneity: Tau ² =0; Chi ² =2.8, df=5(P=0.73); l ² =0%						
Test for overall effect: Z=1(P=0.32)							
Test for subgroup differences: Chi ² =0.5	52, df=1 (P=0.77), I ² =	=0%					
	F	avours Etomidate	0.01 0.1 1 10 10	⁰ Favours Other Agen	ts		

Analysis 1.3. Comparison 1 Etomidate versus all other induction agents, Outcome 3 Mortality: Data as reported - Subgroup analysis of etiology of shock.

Study or subgroup	Etomidate	Other Induc- tion Agents	Odds	Ratio	Weight	Odds Ratio
	n/N	n/N	M-H, Rando	om, 95% Cl		M-H, Random, 95% CI
1.3.1 Septic Shock						
Tekwani 2010	26/61	21/59	_	•	17.81%	1.34[0.64,2.81]
Subtotal (95% CI)	61	59	•		17.81%	1.34[0.64,2.81]
Total events: 26 (Etomidate), 21 (Othe	er Induction Agents)					
Heterogeneity: Tau ² =0; Chi ² =0, df=0(F	e<0.0001); l ² =100%					
Test for overall effect: Z=0.79(P=0.43)						
1.3.2 Trauma						
Hildreth 2008	2/18	0/12			0.99%	3.79[0.17,86.13]
	Fa	avours Etomidate	0.01 0.1	10 1	¹⁰⁰ Favours Other Agent	5



Study or subgroup	Etomidate	Other Induc- tion Agents	Odds Ratio	Weight	Odds Ratio
	n/N	n/N	M-H, Random, 95%	CI	M-H, Random, 95% Cl
Subtotal (95% CI)	18	12		0.99%	3.79[0.17,86.13]
Total events: 2 (Etomidate), 0 (Other I	nduction Agents)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.84(P=0.4)					
1.3.3 Cardiogenic Shock					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Etomidate), 0 (Other I	nduction Agents)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
1.3.4 Undifferentiated Critical Illnes	is				
Absalom 1999	5/17	3/17		- 3.65%	1.94[0.38,9.88]
Jabre 2009	81/234	72/235	<u>+</u> -	64.54%	1.2[0.81,1.76]
Jacoby 2006	13/44	16/44	+	12.11%	0.73[0.3,1.79]
Schenarts 2001	0/16	1/15		- 0.9%	0.29[0.01,7.76]
Subtotal (95% CI)	311	311	•	81.2%	1.12[0.79,1.58]
Total events: 99 (Etomidate), 92 (Othe	r Induction Agents)				
Heterogeneity: Tau ² =0; Chi ² =2.07, df=	3(P=0.56); I ² =0%				
Test for overall effect: Z=0.65(P=0.52)					
Total (95% CI)	390	382	•	100%	1.17[0.86,1.6]
Total events: 127 (Etomidate), 113 (Ot	her Induction Agent	s)			
Heterogeneity: Tau ² =0; Chi ² =2.8, df=5	(P=0.73); I ² =0%				
Test for overall effect: Z=1(P=0.32)					
Test for subgroup differences: Chi ² =0.	74, df=1 (P=0.69), I ² =	0%			
	Fa	avours Etomidate	0.01 0.1 1	10 100 Eavours Other Age	nts

Analysis 1.4. Comparison 1 Etomidate versus all other induction agents, Outcome 4 Mortality: Data as reported - Studies judged to be at low risk of bias.

Study or subgroup	Etomidate	Other Induc- tion Agents	Odds Ratio					Weight	Odds Ratio
	n/N	n/N		М-Н, Р	andom, 959	% CI			M-H, Random, 95% CI
Jabre 2009	81/234	72/235						78.37%	1.2[0.81,1.76]
Tekwani 2010	26/61	21/59			+			21.63%	1.34[0.64,2.81]
Total (95% CI)	295	294			•			100%	1.23[0.87,1.73]
Total events: 107 (Etomidate), 93 (C	Other Induction Agents	s)							
Heterogeneity: Tau ² =0; Chi ² =0.07, d	f=1(P=0.79); I ² =0%								
Test for overall effect: Z=1.18(P=0.24	4)								
	F	avours etomidate	0.01	0.1	1	10	100	Favours other agents	



Analysis 1.5. Comparison 1 Etomidate versus all other induction agents, Outcome 5 Mortality: Post Hoc ITT Analysis accounting for missing subjects.

Study or subgroup	Etomidate	Other Induc- tion Agents		Odds Ratio			Weight	Odds Ratio	
	n/N	n/N		М-Н, Б	andom, 95°	% CI			M-H, Random, 95% Cl
Absalom 1999	5/17	4/18		-	+	_		3.49%	1.46[0.32,6.7]
Hildreth 2008	21/37	12/24						7.62%	1.31[0.47,3.68]
Jabre 2009	84/237	74/237						55.34%	1.21[0.82,1.77]
Jacoby 2006	24/55	27/55			-+			14.37%	0.8[0.38,1.7]
Schenarts 2001	6/16	8/15			+			3.94%	0.53[0.13,2.2]
Tekwani 2010	28/63	21/59			+-			15.25%	1.45[0.7,3]
Total (95% CI)	425	408			•			100%	1.15[0.86,1.53]
Total events: 168 (Etomidate), 146	(Other Induction Agen	its)							
Heterogeneity: Tau ² =0; Chi ² =2.64, c	lf=5(P=0.76); I ² =0%								
Test for overall effect: Z=0.95(P=0.3	4)						I		
	F	avours Etomidate	0.01	0.1	1	10	100	Favours Other Agents	;

Analysis 1.6. Comparison 1 Etomidate versus all other induction agents, Outcome 6 ACTH Stimulation Test.

Study or subgroup	Etomidate	Other Induc- tion Agents	Odds Ratio	Weight	Odds Ratio
	n/N	n/N	M-H, Random, 95% Cl		M-H, Random, 95% CI
1.6.1 ACTH Stimulation test perform	med 4-6 h				
Hildreth 2008	16/18	4/12		73.02%	16[2.4,106.73]
Schenarts 2001	7/10	0/8		26.98%	36.43[1.61,826.33]
Subtotal (95% CI)	28	20		100%	19.98[3.95,101.11]
Total events: 23 (Etomidate), 4 (Othe	r Induction Agents)				
Heterogeneity: Tau ² =0; Chi ² =0.2, df=1	L(P=0.66); I ² =0%				
Test for overall effect: Z=3.62(P=0)					
1.6.2 ACTH Stimulation Test Perform	med 6-12 h				
Schenarts 2001	0/7	0/7			Not estimable
Subtotal (95% CI)	7	7			Not estimable
Total events: 0 (Etomidate), 0 (Other	Induction Agents)				
Heterogeneity: Not applicable					
Test for overall effect: Not applicable					
1.6.3 ACTH Stimulation test perform	med >12h				
Absalom 1999	5/17	3/18		5.57%	2.08[0.41,10.53]
Jabre 2009	100/234	56/235		93.13%	2.39[1.6,3.55]
Schenarts 2001	1/9	0/6		1.3%	2.29[0.08,66.02]
Subtotal (95% CI)	260	259	•	100%	2.37[1.61,3.47]
Total events: 106 (Etomidate), 59 (Ot	her Induction Agent	s)			
Heterogeneity: Tau ² =0; Chi ² =0.03, df=	=2(P=0.99); I ² =0%				
Test for overall effect: Z=4.41(P<0.000	01)				
	F	Favours Etomidate	0.001 0.1 1 10	¹⁰⁰⁰ Favours Other Agen	ts

Analysis 1.7. Comparison 1 Etomidate versus all other induction agents, Outcome 7 Random Serum Cortisol levels (μ g/dL) after receiving intervention.

Study or subgroup	Etomidate		Other Induc- tion Agents		Mean Diffe	rence	Weight	Mean Difference
	Ν	Mean(SD)	N	Mean(SD)	Random, 9	5% CI		Random, 95% Cl
1.7.1 Random Serum Cortisol Level	s 4-6 h							
Hildreth 2008	18	18.2 (8.4)	12	27.9 (13.4)			11.75%	-9.7[-18.22,-1.18]
Koksal 2013	10	6.1 (3.1)	10	10.5 (6.3)			34.04%	-4.4[-8.75,-0.05]
Subtotal ***	28		22				45.79%	-5.73[-10.24,-1.23]
Heterogeneity: Tau ² =2.14; Chi ² =1.18, o	df=1(P=0	.28); l ² =15.22%						
Test for overall effect: Z=2.49(P=0.01)								
1.7.2 Random Serum Cortisol Level	s >12h							
Absalom 1999	17	21 (8.3)	18	21 (11.2)			18.65%	-0.07[-6.57,6.43]
Koksal 2013	10	9.3 (4.6)	10	15.8 (5)			35.56%	-6.5[-10.71,-2.29]
Subtotal ***	27		28				54.21%	-3.78[-10.01,2.44]
Heterogeneity: Tau ² =12.86; Chi ² =2.65	, df=1(P=	0.1); l ² =62.23%						
Test for overall effect: Z=1.19(P=0.23)								
Total ***	55		50		•		100%	-4.96[-8.06,-1.86]
Heterogeneity: Tau ² =2.43; Chi ² =3.94, o	df=3(P=0	.27); I ² =23.85%						
Test for overall effect: Z=3.14(P=0)								
Test for subgroup differences: Chi ² =0.	25, df=1	(P=0.62), I ² =0%						
			Favours	Other Agents	-20 -10 0	10 20	Favours Etc	omidate

Analysis 1.8. Comparison 1 Etomidate versus all other induction agents, Outcome 8 Organ System Dysfunction (SOFA Score).

Study or subgroup	Etomidate		Other Induc- tion Agents			Меа	n Differer	nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		Ran	dom, 95%	CI			Random, 95% Cl
Jabre 2009	234	10.3 (3.7)	235	9.6 (3.9)						100%	0.7[0.01,1.39]
Total ***	234		235							100%	0.7[0.01,1.39]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.99(P=0.05)											
			Favou	rs Etomidate	-2	-1	0	1	2	Favours Othe	r Agents

Analysis 1.9. Comparison 1 Etomidate versus all other induction agents, Outcome 9 ICU Length of Stay.

Study or subgroup	Eto	omidate	Oth tio	er Induc- n Agents	Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% Cl		Random, 95% CI
Hildreth 2008	18	8.1 (7.2)	12	3 (2.4)		29.23%	5.1[1.51,8.69]
Jabre 2009	234	24 (16.3)	235	22 (17)		31.83%	2[-1.01,5.01]
Tekwani 2010	63	3.1 (2.7)	59	4.2 (3.5)		38.94%	-1.1[-2.21,0.01]
Total *** Hotorogonality: $T_{2}U^{2}=0.92$; $Chi^{2}=12.90$	315	-01.12-24 4004	306			100%	1.7[-2,5.4]
Heterogeneity: Tau =8.82; Cill =12.83	9, ui-2(P-	-0);1 -84.49%					
			Favou	urs Etomidate	-5 -2.5 0 2.5 5	Favours Oth	ner Agents



Study or subgroup	Et	tomidate Other Induc- tion Agents		м	lean l	Diffe	rence		Weight Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% CI			Random, 95% CI		
Test for overall effect: Z=0.9(P=0.37)								1		
			Favo	urs Etomidate	5-	-2.5	0	2.5	5	Favours Other Agents

Analysis 1.10. Comparison 1 Etomidate versus all other induction agents, Outcome 10 Hospital Length of Stay.

Study or subgroup	Etomidate		Other Induc- tion Agents		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% CI		Random, 95% Cl
Hildreth 2008	18	13.9 (9.5)	12	6.4 (4.4)	_	47.56%	7.5[2.45,12.55]
Tekwani 2010	63	7.3 (7.3)	59	9.5 (8.4)		52.44%	-2.2[-5,0.6]
Total ***	81		71			100%	2.41[-7.08,11.91]
Heterogeneity: Tau ² =42.71; Chi ² =10.8	5, df=1(F	P=0); I ² =90.79%					
Test for overall effect: Z=0.5(P=0.62)							
			Favou	Irs Etomidate	-10 -5 0 5 10	Favours Oth	er Agents

Analysis 1.11. Comparison 1 Etomidate versus all other induction agents, Outcome 11 Duration of Mechanical Ventilation.

Study or subgroup	Ete	omidate	Oth tio	er Induc- n Agents	Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% CI		Random, 95% CI
Hildreth 2008	18	6.3 (6.5)	12	1.5 (0.8)		31.54%	4.8[1.76,7.84]
Jabre 2009	234	16 (18.5)	235	13 (19.3)		29.92%	3[-0.42,6.42]
Tekwani 2010	63	2.1 (2.1)	59	2.8 (3)		38.54%	-0.7[-1.62,0.22]
Total ***	315		306			100%	2.14[-1.67,5.95]
Heterogeneity: Tau ² =9.59; Chi ² =14.	75, df=2(P	=0); I ² =86.44%					
Test for overall effect: Z=1.1(P=0.27	.)						
			Favou	Irs Etomidate	-5 -2.5 0 2.5 5	 Favours Oth	er Agents

Analysis 1.12. Comparison 1 Etomidate versus all other induction agents, Outcome 12 Duration of Vasopressor Support.

Study or subgroup	Eto	omidate	Oth tio	er Induc- 1 Agents		Меа	n Differe	nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		Rane	dom, 95%	6 CI			Random, 95% Cl
Jabre 2009	234	1 (10.4)	235	0 (5.9)				_		100%	1[-0.53,2.53]
Total ***	234		235							100%	1[-0.53,2.53]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.28(P=0.2)											
			Favou	ırs Etomidate	-5	-2.5	0	2.5	5	Favours Othe	er Agents



APPENDICES

Appendix 1. Search strategies

CENTRAL (The Cochrane Library)

#1MeSH descriptor: [Imidazoles] this term only #2MeSH descriptor: [Benzvl Compounds] explode all trees #3MeSH descriptor: [Etomidate] explode all trees #4hypnomidate or amidate or et?omidat* or r?26?490 or R?16659 or radenar?on #5#1 or #2 or #3 or #4 #6MeSH descriptor: [Anesthesia, Intravenous] this term only #7MeSH descriptor: [Anesthesia] this term only #8MeSH descriptor: [Intubation] explode all trees #9MeSH descriptor: [Intubation, Intratracheal] explode all trees #10MeSH descriptor: [Anesthesia, General] explode all trees #11MeSH descriptor: [Anesthetics] explode all trees #12(airway near protect*) or laryngoscop* or sedat*:ti,ab or hypnotic or (intubat* or an?esthe*):ti,ab #13MeSH descriptor: [Laryngoscopy] explode all trees #14MeSH descriptor: [Hypnotics and Sedatives] explode all trees #15(#6 or #7 or #8 o #9 or #10 or #11 or #12 or #13) and #14 #16MeSH descriptor: [Deep Sedation] explode all trees #17MeSH descriptor: [Conscious Sedation] explode all trees #18MeSH descriptor: [Intensive Care Units] explode all trees #19MeSH descriptor: [Burn Units] explode all trees #20MeSH descriptor: [Coronary Care Units] explode all trees #21MeSH descriptor: [Recovery Room] explode all trees #22MeSH descriptor: [Respiratory Care Units] explode all trees #23MeSH descriptor: [Intensive Care] explode all trees #24MeSH descriptor: [Critical Care] explode all trees #25MeSH descriptor: [Emergencies] explode all trees #26MeSH descriptor: [Emergency Treatment] explode all trees #27MeSH descriptor: [Emergency Service, Hospital] explode all trees #28MeSH descriptor: [Trauma Centers] explode all trees #29MeSH descriptor: [Emergency Medical Services] explode all trees #30MeSH descriptor: [Critical Illness] explode all trees

#31(single bolus dose or induction or emergenc* or ambulanc* or trauma* or ((intensive or critical* or serious*) near (ill* or care or sick*)) or ICU):ti,ab or shock:ti,ab

#32#15 or #16 or #17 or #18 or #19 or #20 or #21 or #22 or #23 or #24 or #25 or #26 or #27 or #28 or #29 or #30 or #31 #33#5 and #32

MEDLINE (OvidSP)

1. Imidazoles/ or Benzyl Compounds/ or hypnomidate.mp. or amidate.af. or exp Etomidate/ or et?omidat\$.af. or (r?26?490 or R?16659).mp. or radenar?on.mp.

2. ((Intubation, Intratracheal/ or Intubation/ or intubat\$.ti,ab. or anesthesia/ or anesthesia, intravenous/ or an?esthesia.ti,ab. or anesthesia, general/ or anesthetics/ or an?esthetic\$.ti,ab. or (airway adj5 protect\$).mp. or Laryngoscopy/ or laryngoscop\$.mp. or sedat \$.mp. or Hypnotics.mp.) and Sedatives/) or Deep Sedation/ or single bolus dose.mp. or induction.mp. or Conscious Sedation/ or intensive care units/ or burn units/ or coronary care units/ or recovery room/ or respiratory care units/ or Intensive Care/ or Critical Care/ or Emergencies/ or emergenc\$.mp. or Emergency Treatment/ or ambulanc\$.mp. or emergency service, hospital/ or trauma centers/ or Emergency medical services/ or trauma.ti,ab. or Critical Illness/ or ((intensive or critical\$ or serious\$) adj5 (ill\$ or care or sick\$)).mp. or ICU.mp. or shock\$.ti,ab.

3.1 and 2

4. ((randomized controlled trial or controlled clinical trial).pt. or randomized.ab. or placebo.ab. or drug therapy.fs. or randomly.ab. or trial.ab. or groups.ab.) not (animals not (humans and animals)).sh. 5. 3 and 4

EMBASE (OvidSP)

1. imidazole derivative/ or benzyl derivative/ or amidate.af. or et?omidat\$.af. or (r?26?490 or R?16659).mp. or radenar?on.mp. or hypnomidate.mp.

2. ((endotracheal intubation/ or intubation/ or anesthesia/ or intravenous anesthesia/ or general anesthesia/ or anesthetic agent/ or intubat\$.ti,ab. or an?esthe\$.ti,ab. or (airway adj3 protect\$).ti,ab. or laryngoscopy/ or laryngoscop\$.ti,ab. or sedat\$.ti,ab. or hypnotics.ti,ab.)



and sedative agent/) or deep sedation/ or conscious sedation/ or intensive care unit/ or burn/ or coronary care unit/ or recovery room/ or intensive care unit/ or intensive care/ or emergency/ or emergency treatment/ or emergency health service/ or critical illness/ or (emergenc \$ or single bolus doseor inductionor or ambulanc\$ or trauma or ((intensive or critical\$ or serious\$) adj3 (ill\$ or care or sick\$)) or ICU or shock\$).ti,ab.

3.1 and 2

4. (randomized-controlled-trial/ or randomization/ or controlled-study/ or multicenter-study/ or phase-3-clinical-trial/ or phase-4-clinical-trial/ or double-blind-procedure/ or single-blind-procedure/ or (random* or cross?over* or multicenter* or factorial* or placebo* or volunteer*).mp. or ((singl* or doubl* or trebl* or tripl*) adj3 (blind* or mask*)).ti,ab. or (latin adj square).mp.) not (animals not (humans and animals)).sh.

5.3 and 4

CINAHL (EBSCOhost)

S1 ((MM "Imidazoles") OR (MM "Benzyl Compounds") OR (MM "Etomidate")) OR (etomidate or radenaron or hypnomidate)

S2 ((MH "Intubation, Intratracheal") OR (MH "Anesthesia") OR (MH "Anesthetics")) OR (intubat* or an?esthe* or (airway and protect*) or laryngoscop* or sedat* or hypnotic*)

S3 (MM "Hypnotics and Sedatives")

S4 S2 AND S3

S5 ((MH "Sedation") OR (MH "Intensive Care Units") OR (MH "Burn Units") OR (MH "Coronary Care Units") OR (MH "Respiratory Care Units") OR (MH "Critical Care") OR (MH "Emergencies") OR (MH "Emergency Treatment (Non-Cinahl)") OR (MH "Trauma Centers") OR (MH "Emergency Medical Services") OR (MH "Critical Illness")) OR (emergenc* or ambulanc* or trauma) OR (((intensive or critical* or serious*) and (ill* or care or sick*))) OR (ICU or shock*)

S6 S4 OR S5

S7 random* or (trial* and (clinical or control*)) or placebo* or multicenter or prospective or ((blind* or mask*) and (single or double or triple or treble))

S8 S1 AND S6 AND S7

ISI Web of Science

#1 TS=(imidazoles or benzyl compounds or hypnomidate or amidate or etomidate or (r?26?490 or r?16659) or radenaron)

#2 TS=((intubation, intratracheal or intubation or intubat* or anesthesia or anesthesia, intravenous or an?esthesia or anesthesia, general or anesthetics or an?esthetic* or (airway same protect*) or laryngoscopy or laryngoscop* or sedat* or hypnotics) and sedatives) or TS=(deep sedation or single bolus dose or induction or conscious sedation or intensive care units or burn units or coronary care units or recovery room or respiratory care units or intensive care or critical care or emergencies or emergenc* or emergency treatment or ambulanc* or emergency service, hospital or trauma centers or emergency medical services or trauma or critical illness or ((intensive or critical* or serious*) same (ill* or care or sick*)) or icu or shock*) #3 #1 and #2

LILACS (BIREME)

(hypnomidate or amidate or etomidate or radenaron)

BIOSIS Citation IndexSM

#1 TS=(hypnomidate or amidate or etomidate or radenaron)

#2 TS=((intubation, intratracheal or intubation or intubat* or anesthesia or anesthesia, intravenous or an?esthesia or anesthesia, general or anesthetics or an?esthetic* or (airway same protect*) or laryngoscopy or laryngoscop* or sedat* or hypnotics) and sedatives) or TS=(deep sedation or single bolus dose or induction or conscious sedation or intensive care units or burn units or coronary care units or recovery room or respiratory care units or intensive care or critical care or emergencies or emergenc* or emergency treatment or ambulanc* or emergency service, hospital or trauma centers or emergency medical services or trauma or critical illness or ((intensive or critical* or serious*) same (ill* or care or sick*)) or icu or shock*)

#3 TS=(random* or (trial* SAME (clinical or control*)) or placebo* or multicenter or prospective or ((blind* or mask*) SAME (single or double or triple or treble)))

#4 #1 and #2 and #3

Appendix 2. Data extraction form

Data Extraction Form Reviewer: Bruder, Ball, Ridi

Study Identifier	Author	Year

Single induction dose of etomidate versus other induction agents for endotracheal intubation in critically ill patients (Review)
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(Continued)

Citation: Authors. Title. Journal. Year; Volume (Issue): pages

Eligibility for review

Inclusion criteria	Exclusion criteria
p Comparison of a single bolus dose of etomidate to any	p Patients were extubated within 24 h
other rapid-acting, iv botus dose induction agent	p Etomidate infusion
p RCI	p Non-randomized
p Human data	p Exogenous steroid replacement
p Emergency airway intervention	p Etomidate use for indications other than airway intervention
p Critical illness	p Elective anaesthesia induction

Methods

Objectives

Participants

<u>Primary</u>

<u>Secondary</u>

Inclusion criteria

Exclusion criteria

Indication for Intubation



(Continued)	
Aetiology of Shock (circle)	Sepsis, Trauma, Cardiac, Undifferentiated, Unknown
Experimental intervention	Etomidate, Dose, Route
Control intervention	Drug, Dose, Route
Co-interventions	
Study duration	
Study location	
Sample size calculation	Yes, No
Funding source	
Analysis	Intention-to-treat? Appropriate statistics? Other

Outcomes

	Measured	Definition, Units & Timeframe
Mortality	Y/N	
Cortisol levels	Y/N	
ACTH stimulation test	Y/N	
Hospital LOS	Y/N	
ICU LOS	Y/N	
Post-intubation haemodynamics	Y/N	
Ventilator days	Y/N	
Vasopressor requirements	Y/N	

Risk of bias assessment

Domain		Description	Judgment
Random sequence generation	Describe the method used to generate the allocation sequence in sufficient detail to allow an assessment of whether it should produce comparable groups		Was the allocation se- quence adequately gener- ated?



(Continued)

		Low risk/High risk/Un- clear
Allocation conceal- ment	Describe the method used to conceal the allocation se- quence in sufficient detail to determine whether inter- vention allocations could have been foreseen in advance of, or during, enrolment	Was allocation adequately concealed?
		Low risk/High risk/Un- clear
Blinding	Describe all measures used, if any, to blind study partici- pants and personnel from knowledge of which interven- tion a participant received. Provide any information re- lating to whether the intended blinding was effective	Was knowledge of the al- located intervention ade- quately prevented during the study?
	Assess blinding for each outcome	Low risk/High risk/Un- clear
Incomplete out- come data	Describe the completeness of outcome data for each main outcome, including attrition and exclusions from the analysis. State whether attrition and exclusions were reported, the numbers in each intervention group (com- pared with total randomized participants), reasons for	Were incomplete out- come data adequately ad- dressed?
	attrition/exclusions where reported, and any re-inclu- sions in analyses performed by the review authors	Low risk/High risk/Unclear
Selective reporting	State how the possibility of selective outcome report- ing was examined by the review authors, and what was found	Are reports of the study free of suggestion of selec- tive outcome reporting?
		Low risk/High risk/Un- clear
Other bias	Selection bias (differences in comparison groups)	Was the study apparent- ly free of other problems that could put it at a high
	Performance bias (differences in the care provided other than the experimental intervention)	risk of blas?
	Attrition bias (differences in withdrawals, loss to fol- low-up, or protocol deviations)	Low risk/High risk/Unclear
	Detection bias (differences in outcome assessment be- tween groups)	
	Other	

Results



Baseline characteristics	Etomidate	Comparator	P value
Participants (n)			
Age (mean, SD)			
% Male (mean, SD)			
Lost (n)			
Disease severity			
Comorbidities			
Baseline heart rate (bpm)			
Baseline blood pressure (MAP)			
Main results			
Dichotomous			
Mortality (numbers, 2 x 2 table) Yes			
No			_
ACTH Stim (numbers, 2 x 2 table) Pos			
Neg			_
Continuous			
Cortisol levels (#, mean, SD)			
Hospital LOS (days) (#, mean, SD)			
ICU LOS (days) (#, mean, SD)			
Post-intubation haemodynamics			
Ventilator days (days) (#, mean, SD)			
Vasopressor requirements			
Other			

NOTES

WHAT'S NEW



Date	Event	Description
20 December 2018	Amended	Editorial team changed to Cochrane Emergency and Critical Care

CONTRIBUTIONS OF AUTHORS

Conceiving the review: Eric A Bruder (EB), Ian Ball (IB), Corinne Hohl (CH) Co-ordinating the review: EB Undertaking manual searches: EB, IB, Stacy Ridi (SR) Screening search results: EB, IB, SR Organizing retrieval of papers: EB Screening retrieved papers against inclusion criteria: EB, IB, SR Appraising quality of papers: EB, IB, SR Abstracting data from papers: EB, IB, SR Writing to authors of papers for additional information: EB Providing additional data about papers: EB Obtaining and screening data on unpublished studies: EB Data management for the review: EB, William Pickett (WP) Entering data into Review Manager (RevMan 5.2): EB, IB, SR RevMan statistical data: EB, IB, WP Other statistical analysis not using RevMan: WP Double entry of data: (data entered by person one: EB; data entered by person two: IB) Interpretation of data: EB, IB, WP Statistical inferences: WP Writing the review: EB, IB Securing funding for the review: N/A Performing previous work that was the foundation of the present study: CH Guarantor for the review (one author): EB Person responsible for reading and checking review before submission: EB

DECLARATIONS OF INTEREST

Eric Bruder: none known.

Ian Ball: none known.

Stacy Ridi: none known.

William Pickett: none known.



Corinne Hohl: Dr Hohl is a full-time faculty member with the Department of Emergency Medicine at the University of British Columbia. She has received reimbursement for travel expenses by the Canadian Association of Emergency Physicians to present on this topic in Montreal, 2010.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The secondary outcome, duration of vasopressor support, was proposed to be measured in hours in the protocol (Bruder 2012). The data were reported in the literature in days, so we also reported these data in days.

Our protocol (Bruder 2012) stated that we would report both the OR and RR. This review has only reported the OR. We chose to do this in order to simplify our results and minimize information clutter.

Our protocol (Bruder 2012) stated, "We will exclude non-randomized or quasi-randomized trials". We chose to include one study that block randomized patients in blocks of four (Jabre 2009). While this deviates from our protocol, we chose to include this study because it was the single largest study on the topic with otherwise sound methodology and low risk of bias. This study also had a very low rate of attrition. We judged the small blocks of four in conjunction with computer generated randomization and identically sealed and sequentially numbered drug boxes to mitigate the shortfalls of block randomization. We feel that this deviation from our protocol did not introduce bias into our review.

Our review included a post hoc sensitivity analysis of our mortality data (Analysis 1.5) that was not described in our protocol. In designing our protocol, we did not anticipate the large number of patients lost to follow-up. This sensitivity analysis was added to ensure our mortality outcome results were true. We feel that this deviation from our protocol did not introduce bias into our review.

INDEX TERMS

Medical Subject Headings (MeSH)

*Critical Illness; *Intubation, Intratracheal; Adrenal Insufficiency [*chemically induced] [mortality]; Etomidate [*administration & dosage] [*adverse effects]; Health Services Needs and Demand [statistics & numerical data]; Hypnotics and Sedatives [*administration & dosage] [*adverse effects]; Intensive Care Units [statistics & numerical data]; Length of Stay; Organ Dysfunction Scores; Randomized Controlled Trials as Topic; Respiration, Artificial [statistics & numerical data]

MeSH check words

Humans