INTRODUCTION

Description of the problem

Traumatic injuries caused over 5 million deaths worldwide in 2010 and casualties are projected to exceed 8 million by 2020. Haemorrhagic shock is the leading cause of preventable death following trauma. Fluids have traditionally been used to support organ perfusion in the setting of haemorrhage, but recent investigations have highlighted the risks of excessive fluid administration. In a landmark trial, hypotensive patients with penetrating torso trauma were more likely to be discharged alive from hospital when fluid resuscitation was withheld until arrival to the operating theatre. These results are concordant with data from a subsequent 90-patient trial on fluid use during trauma surgery, where a mean arterial pressure target of 50 vs 65 mm Hg significantly decreased blood product use without increasing 30-day mortality. Recent trauma guidelines have incorporated restrictive fluid strategies, referred to as permissive hypotension, into their recommendations. However, as pointed out in two systematic reviews, the safety of permissive hypotension remains uncertain. This may be particularly true among patients with concomitant traumatic brain injury in whom hypotension is associated with increased mortality.

METHODS AND ANALYSIS

We will identify randomised control trials comparing early resuscitation with vasopressors versus placebo or standard care in adults following traumatic injury. Data sources will include MEDLINE, EMBASE, CENTRAL, clinical trial registries and conference proceedings. Two reviewers will independently determine trial eligibility. For each included trial, we will conduct duplicate independent data extraction and risk of bias assessment. We will assess the overall quality of the data for each individual outcome using the GRADE approach.

Ethics and dissemination: We will report this review in accordance with the PRISMA statement. We will disseminate our findings at critical care and trauma conferences and through a publication in a peer-reviewed journal. We will also use this systematic review to create clinical guidelines (http://www.magicapp.org), which will be disseminated in a standalone publication.

Trial registration number: CRD42016033437.

ABSTRACT

Introduction: Worldwide, traumatic casualties are projected to exceed 8 million by year 2020. Haemorrhagic shock and brain injury are the leading causes of death following trauma. While intravenous fluids have traditionally been used to support organ perfusion in the setting of haemorrhage, recent investigations have suggested that restricting fluid therapy by tolerating more severe hypotension may improve survival. However, the safety of permissive hypotension remains uncertain, particularly among patients who have suffered a traumatic brain injury. Vasopressors preferentially vasoconstrict blood vessels that supply non-vital organs and capacitance vessels, thereby mobilising the unstressed blood volume. Used as fluid-sparing adjuncts, these drugs can complement resuscitative measures by correcting hypotension without diluting clotting factors or increasing the risk for tissue oedema.

Methods and analysis: We will identify randomised control trials comparing early resuscitation with vasopressors versus placebo or standard care in adults following traumatic injury. Data sources will include MEDLINE, EMBASE, CENTRAL, clinical trial registries and conference proceedings. Two reviewers will independently determine trial eligibility. For each included trial, we will conduct duplicate independent data extraction and risk of bias assessment. We will assess the overall quality of the data for each individual outcome using the GRADE approach.

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Vasopressors are vasoactive agents that induce vasoconstriction and increase arterial pressure. They also vasoconstrict capacitance vessels, thereby mobilising the unstressed blood volume and increasing venous return. Vasopressors can rapidly correct hypotension in an effort to maintain end-organ perfusion in the face of hypovolaemic shock. Used as fluid-sparing adjuncts to resuscitation, vasopressors can complement resuscitative measures by correcting hypotension without diluting clotting factors or increasing the risk for tissue oedema.

In spite of this potential role as fluid-sparing adjuncts, vasopressors potentiate vasoconstriction and may therefore worsen hypoperfusion, despite high blood pressure values. Thus, trauma guidelines restrict vasopressor use to cases of severe hypotension refractory to fluid therapy, but wide practice variations exist. In the absence of high-quality evidence, experts recommend limiting vasopressors to patients unresponsive to fluid therapy, while recognising that a significant knowledge gap exists.

Why is it important to conduct this review? Given the well-recognised risks of excessive fluid administration, healthcare providers are left with few alternatives for unstable trauma patients in whom hypotension may be deleterious. Vasopressors are increasingly perceived as a complementary strategy, but their associated harms and benefits have never been conclusively examined. No systematic review has yet focused on the efficacy or safety of vasopressors in trauma.

Research question
Does the early use of vasopressors improve the survival of victims of traumatic injury, compared with standard care which incorporates vasopressors only once other modalities of support have failed?

METHODS AND ANALYSIS

Criteria for selecting studies for this review

Types of studies
We will include all randomised controlled trials and controlled observational studies (case-control or cohort). We will exclude case reports and case series. We will not use restrictions based on language, methodological quality, publication status or year of publication.

Types of participants
Our population of interest consists of adult victims of acute, non-iatrogenic traumatic injury (blunt or penetrating). We will include studies reporting diverse populations, such as severe burns, if there are extractable data on the trauma subgroup or if the non-trauma subgroup constitutes <10% of the total study population. We will exclude paediatric (<16 years) and animal studies.

Types of interventions
The interventions of interest are the administration of any vasopressor (epinephrine, norepinephrine, dopamine, phenylephrine, ephedrine vasopressin or vasopressin analogues) during early trauma resuscitation. We will include studies that consider cardiac inotropes (e.g., milrinone) if these account for <10% of the vasopressor group. We will exclude studies that report vasopressor use exclusively during the postoperative phase, after arrival in the intensive care unit or >24 hours after arrival at the trauma bay. We will also exclude studies that rely on cointerventions not available in the early phases of care, such as cerebral perfusion pressure monitoring.

Types of outcome measures
We will exclude studies where follow-up was <24 hours. The primary outcome will be short-term mortality at longest follow-up up to 90 days. Other outcomes will be long-term mortality beyond 90 days, fluid and blood product requirements during the early resuscitation period, requirements for acute (up to 90 days) or chronic (beyond 90 days) renal replacement therapy, duration of renal replacement therapy, duration of mechanical ventilation, incidence of acute kidney injury, incidence of vasopressor-associated adverse events (new-onset cardiac arrhythmia, digit, limb or skin ischaemia, mesenteric ischaemia and myocardial ischaemia), neurological outcome and long-term quality of life.

Search methods for the identification of studies
We will perform a search of the following databases for relevant studies: MEDLINE, EMBASE and the Cochrane Central Register of Controlled Trials (CENTRAL). The search strategy specific to MEDLINE is included as online supplementary appendix 1. We will perform similar searches, with keywords adapted to specific database dictionaries.

Additional search methods (grey literature)
We will screen reference lists of included studies and relevant reviews for eligible articles. We will also manually screen conference proceedings from 2005 for the following scientific meetings: Society of Critical Care Medicine, European Society of Intensive Care Medicine, International Society of Intensive Care and Emergency Medicine, American Thoracic Society, American Association for the Surgery of Trauma, Eastern Association for the Surgery of Trauma, European Society for Trauma and Emergency Surgery, Shock Society, European Shock Society and the American College of Chest Physicians. We will also search ClinicalTrials.gov for any relevant ongoing or unpublished trials. Whenever possible, we will contact authors to obtain additional data.

Study records
Pairs of reviewers will independently screen titles and abstracts using a pretested electronic screening form...
(Covidence web platform: http://www.COVIDENCE.org). We will include articles for full-text review unless both reviewers deem them irrelevant. Pairs of reviewers will then independently screen all full-text articles using specific eligibility criteria via pretested electronic screening tools (Covidence). We will resolve disagreements by consensus or third-reviewer adjudication if necessary. We will report chance-corrected agreement using Cohen’s k for full-text eligibility screening.

**Data collection**

Teams of two reviewers will perform data extraction independently and in duplicate using pretested data collection forms (Covidence). We will collect information pertaining to study design, patient baseline characteristics, intervention and comparator, clinical outcomes and risk of bias. Conflicts will be resolved by consensus or third-abstractor adjudication if necessary.

**Risk of bias assessment**

For randomised controlled trials, we used a modified version of the Cochrane Collaboration tool to assess the risk of bias of individual studies. This tool addresses the following domains: randomisation, allocation concealment, blinding of patients, healthcare providers, data collectors, outcome assessors and data analysts, loss to follow-up, selective outcome reporting and other risks of bias.

For observational studies, we will use the risk of bias tools developed by the ‘Clinical Advances through Research and Information Technology’ (CLARITY) group at McMaster university (https://distillercer.com/resources/). These tools evaluate the selection of the intervention and control groups, the adequacy of assessment of prognostic factors, exposure and clinical outcomes, statistical adjustment and/or matching, follow-up, similarity of co-interventions between groups and other risks of bias.

Studies with one or more domain assessed as a potential source of bias will be considered overall at high risk of bias. We will assess the overall quality of the data for each individual outcome using the GRADE approach.

**Summarising data and treatment effect**

We will include the results of clinically homogeneous studies in a meta-analysis using the Review Manager software (Review Manager 5.3). We will use a random-effects model and the inverse variance method to calculate individual study weights. Randomised controlled trials and observational studies will be meta-analysed separately and presented as forest plots. For dichotomous outcomes, summary effect measures will be pooled from individual study ORs and presented as risk ratios with 95% CIs. For continuous outcomes, summary effect measures will be presented as mean differences with 95% CIs. Health-related quality-of-life outcomes may be reported using a variety of different scales. In this situation, we will use the methods of reporting suggested by Thorlund et al. Their recommendations encourage the reporting of summary measures as a number needed to treat using at least two complementary methods, in order to improve the interpretability of results.

We will provide qualitative summaries of outcomes for which quantitative summaries are deemed inappropriate and justify our rationale. We will use trial sequential analysis to assess the risk of random errors.

**Assessment of heterogeneity**

We will assess and report heterogeneity quantitatively the I² statistic and perform a χ² test for homogeneity. Irrespective of the degree of heterogeneity, we will perform prespecified exploratory subgroup analyses.

**Subgroup analysis and investigation of heterogeneity**

We will perform the following comparisons to identify potential subgroup effects. (1) Patients with severe traumatic brain injury (Glasgow Coma Scale ≤8) versus those without (Glasgow Coma Scale >8), hypothesising that patients with severe traumatic brain injury benefit more from vasopressors, given their vulnerability to hypotension and the risk of harm associated with fluid resuscitation and permissive hypotension. (2) Patients with blunt trauma versus those with penetrating trauma, hypothesising that there may be a greater benefit for blunt trauma victims, as they may be less likely to benefit from transient hypotension to control a discrete source of bleeding as in penetrating trauma. (3) Patients aged ≤45 years versus those aged >45 years (if within study subgroup data available) or studies where the median age is >45 versus ≤45 years hypothesising that older patients may be more vulnerable to hypotension and therefore receive supplemental benefit from vasopressor therapy. (4) Academic trauma centre versus community setting, hypothesising that vasopressors may be more beneficial in community settings where transport times and in-hospital delays likely expose patients to resuscitative measures for prolonged periods of time, thereby increasing the risks associated with transient hypotension. (5) High or unclear risk of bias versus low risk of bias, hypothesising that studies with high or unclear risk of bias may overestimate the benefits of early vasopressor therapy. (6) Early studies versus more recent studies (within the last 10 years) hypothesising that recent studies report greater benefit of early vasopressor use.

For between-study comparisons, we will require at least five studies, with each subgroup represented by at least two studies, to undertake a subgroup analysis. A minimum of two studies will be required to conduct a within-study subgroup analysis.

**Assessment of reporting bias**

If we include 10 or more studies in a meta-analysis, we will assess the potential for publication bias visually using a funnel plot and statistically using Egger’s test for continuous outcomes and the arcsine test for dichotomous outcomes.
Assessment of confidence in estimates of effect
We will use the Grading of Recommendations, Assessment, Development and Evaluation (GRADE) framework to report our overall confidence in estimates of effect. This method considers the overall risk of bias, imprecision, inconsistency, indirectness and likelihood of publication bias to judge the overall quality of evidence for each outcome. Quality of evidence is rated ‘very low’, ‘low’, ‘moderate’ or ‘high’. Randomised controlled trials provide high-quality evidence, while observational studies yield low-quality evidence. Trials can be rated down according to the above-mentioned criteria. Observational studies can be rated up in the presence of a large magnitude of the association, a dose–response gradient or if all unaccounted confounders increase confidence in estimates of effect.

The findings of this review will be summarised and presented with a summary of findings table with an explicit judgement of quality of evidence for each outcome across studies.

DISCUSSION
Traumatic injuries remain one of the leading causes of mortality worldwide. For many patients, healthcare providers are left with few safe options for haemodynamic support. Although restrictive fluid strategies may be beneficial for victims of penetrating torso injuries,4 this strategy is unsafe for patients with signs of neurological injury. Vaspressors could provide a much-needed complementary means of haemodynamic support in this vulnerable population. Given the theoretical risks associated with these agents, a rigorous evaluation of the published evidence is required to guide clinical practice.

This methodologically rigorous systematic review will summarise the existing evidence on the efficacy and safety of early vaspressor use following traumatic injury. Strengths of this review include duplicate risk of bias assessment, evaluation of the quality of evidence using the GRADE approach, a detailed search of published studies and grey literature, predefined study eligibility criteria and a priori subgroup hypotheses.

DISSEMINATION
We will report this review in accordance with the PRISMA statement. This protocol is reported in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols (PRISMA-P) guidelines. We will disseminate our findings by emitting clinical guidelines using the MagiApp (http://www.magicapp.org) as well as conference presentations and publication in a peer-reviewed journal.

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Competing interests None declared.

Ethics approval Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement The data and analyses of the systematic review will be available to the public.

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