RESEARCH ARTICLE

Vasopressors for the Treatment of Septic Shock: Systematic Review and Meta-Analysis

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Abstract

Objective

International guidelines recommend dopamine or norepinephrine as first-line vasopressor agents in septic shock. Phenylephrine, epinephrine, vasopressin and terlipressin are considered second-line agents. Our objective was to assess the evidence for the efficiency and safety of all vasopressors in septic shock.

Methods

Systematic review and meta-analysis. We searched electronic database of MEDLINE, CENTRAL, LILACS and conference proceedings up to June 2014. We included randomized controlled trials comparing different vasopressors for the treatment of adult patients with septic shock. Primary outcome was all-cause mortality. Other clinical and hemodynamic measurements were extracted as secondary outcomes. Risk ratios (RR) and mean differences with 95% confidence intervals (CI) were pooled.

Results

Thirty-two trials (3,544 patients) were included. Compared to dopamine (866 patients, 450 events), norepinephrine (832 patients, 376 events) was associated with decreased all-cause mortality, RR 0.89 (95% CI 0.81-0.98), corresponding to an absolute risk reduction of 11% and number needed to treat of 9. Norepinephrine was associated with lower risk for major adverse events and cardiac arrhythmias compared to dopamine. No other mortality benefit was demonstrated for the comparisons of norepinephrine to epinephrine, phenylephrine and vasopressin / terlipressin. Hemodynamic data were similar between the different vasopressors, with some advantage for norepinephrine in central venous pressure, urinary output and blood lactate levels.
Conclusions
Evidence suggests a survival benefit, better hemodynamic profile and reduced adverse events rate for norepinephrine over dopamine. Norepinephrine should be regarded as the first line vasopressor in the treatment of septic shock.

Introduction
Septic shock and severe sepsis are a grave consequence of infection. Septic shock accounts for about 9% of admissions, and is the most common cause of death in intensive care units (ICUs) [1,2]. The mortality rate reported is 40 to 60% [1,3]. Septic shock is defined by the ACCP/SCCM as the need for vasopressors to reverse sepsis-induced hypotension [4]. The concept of “early goal directed therapy” was developed in order to set early hemodynamic goals. The goals include central venous or mixed venous saturation (ScvO2) higher than 70%, mean arterial pressure (MAP) ≥ 65 mmHg, central venous pressure (CVP) > 8–12 mmHg, and urine output > 0.5 ml/kg/hr within 6 hours of initiation of therapy [5]. Later the concept of lactate clearance was added to the goals as an alternative or addition to the ScvO2 [6]. Thus, hemodynamic outcomes are regarded as surrogate markers for survival, the ultimate goal of treatment.

No study to date has demonstrated a statistically significant survival benefit of one vasopressor over another. Therefore, the choice of vasopressor in septic shock is rather empiric. The Surviving Sepsis Campaign recommends norepinephrine or dopamine as the first-choice vasopressor agent followed by epinephrine in patients who respond poorly to dopamine or norepinephrine [7,8].

In this systematic review we aimed to examine the evidence on the relative clinical, hemodynamics and safety of any vasopressors in the treatment of septic shock, in order to guide future trials and treatment guidelines.

Methods
Inclusion criteria
Randomized controlled trials (RCTs) and randomized crossover trials regardless of publication status were included. We included trials assessing adult patients with septic shock and sepsis with multiple organ dysfunction syndrome diagnosed using established criteria [4], or an accepted definition of severe sepsis and end organ damage caused by hypoperfusion. In studies that recruited patients with shock due to many causes, we extracted data for septic patients where possible. The intervention assessed was vasopressor versus a different vasopressor, a combination of vasopressors, placebo or no vasopressor. The following vasopressors were included: dopamine, norepinephrine, epinephrine, phenylephrine, vasopressin and terlipressin.

We regarded vasopressin and terlipressin as non-adrenergic vasopressors. Trials that added open-label vasopressor(s) were included if it was applied to both arms of intervention. Studies comparing interventions limited to inotrope agents were excluded. We excluded studies that assessed different dosages or schedules of the same vasopressors.

Outcomes assessed
The primary outcome was all-cause mortality at 28 days following randomization, or when lacking these data, all-cause mortality as reported by the authors. Secondary outcomes included length of ICU stay and / or hospital stay in patients discharged alive; ventilator free days;
vasopressor free days; hemodynamic profiles of patients at the first hour (or first measuring point), and at the 6th hour (or second measuring point); and adverse events (AEs) (major AEs, arrhythmias, myocardial infarction, stroke, internal organ ischemic damage, and local extravasation or skin necrosis). The hemodynamic profile consisted of CVP, MAP, ScvO2, urinary output, blood lactate levels, cardiac index (CIX), systemic vascular resistance index (SVRI), heart rate, and oxygen delivery index (VIO2); we also collected measurements of splanchnic blood flow, oxygen delivery, oxygen consumption and visceral CO2 difference as measured by catheter sampling or tonometry. In studies that allowed crossover between arms, we included data regarding the first randomization only, if available. Mortality was not extracted from crossover studies. Outcome measures were collected on an intention-to-treat basis. Where such data was not presented, per-protocol results were used. We also compared results for adrenergic vasopressors versus non-adrenergic vasopressors.

Search methods
We searched MEDLINE, CENTRAL, LILACS and conference proceedings (International Symposium on Intensive Care and Emergency Medicine), up to June 2014. We also hand-searched all references of included studies, and previous meta-analyses for more trials. The words "vasopressor" and vasopressor names and their MESH terms were crossed with the terms "hypotension", "circulatory failure", "shock", "sepsis" or "bacteremia" and with the Cochrane highly sensitive filter for RCTs [9]. No language restrictions were used. Authors were contacted to complement data by e-mail and phone calls.

Data collection
Two reviewers independently inspected each reference identified by the search, scanned full-texts of relevant studies, applied the inclusion criteria and extracted the data. Disagreements in data extraction were resolved by discussion with a third reviewer. Risk of bias was assessed in duplicate using domain-based evaluation, classifying studies primarily according to the risk of non-random allocation of patients to the intervention arm, i.e. allocation concealment and sequence generation. These were graded as adequate, unclear and inadequate as recommended in The Cochrane Handbook [9]. Additional domains assessed included blinding, incomplete outcome data reporting, ethics committee, patient consent and industrial sponsorship.

Data analysis
For binary data, individual study results are expressed as RR with 95% CI. For continuous outcomes we used end-value means with standard deviations (SD), as data for change from baseline was unavailable. In studies that reported median with interquartile ranges, we converted the reported values to means assuming a normal distribution [10]. RRs and mean differences were pooled using a fixed effect model (Mantel-Haenszel method) (Review Manager [RevMan], version 5.3 for Windows, The Cochrane Collaboration, Oxford, UK). Heterogeneity was defined by a chi-square test of heterogeneity <0.1 or an I² measure of inconsistency >40%. If significant heterogeneity was identified, we used random effect model (REM). We explored potential sources of heterogeneity: trials published before versus after the year 2004 (surviving sepsis campaign), trials conducted in developed versus developing countries, and the adequacy of allocation concealment and blinding.
Results

The search yielded 2,380 publications of which 61 were potentially relevant. Thirty-six studies were excluded (Fig 1), and 7 studies were added from reference hand searching. Altogether, 32 studies [11–42] were included (all published in peer-review journals). The trials were published in the years 1989–2012 and recruited 3,544 patients (median 50 patients, Table 1 for trials’ characteristics). Five trials were crossover [16–18,37,41] and the remaining were parallel RCTs. The comparisons assessed in included trials are summarized in Fig 2. The main comparisons were between norepinephrine and dopamine (14 trials, including 2 crossover trials) and norepinephrine and epinephrine (7 trials including 3 crossover trials). One trial compared vasopressin to placebo [24]; Additional open-label vasopressors could be added to both study arms equally in 16/32 trials (Fig 2). Eight trials were designed to measure clinical endpoints as the primary outcome, while 26 trials, including all crossover trials, measured hemodynamic data (crossover time at 45–220 minutes). The follow up duration was not stated in 21/32 trials; When stated, follow-up ranged from length of the ICU stay (3 trials) to 90 days (median 28 days). The common sources of infection were respiratory (34.6% of included patients) and intra-abdominal (25.1%). Steroid administration was reported in 5 studies, and given to 31–100% of all patients (median 74%), (Table 1). The amount of fluids that was given for the resuscitation was reported in 7/32 trials and varied considerably (median 150 ml/hr, range 45–300 ml/hr). Another 10 trials reported a protocol based fluid resuscitation and titration by CIX. The weighted mean all-cause mortality was 45.0% (SD 16.2%, range 16.7–88.7%) and was not significantly correlated with publication year (Spearman’s rho -0.260, p = 0.24, 32 studies, Table 1).

Risk of bias assessment

Risk of bias assessment is detailed in Table 2. Low risk sequence generation and allocation concealment were reported in 16/32 (50%) and 13/32 (40.6%) trials, respectively. One study was
| Study ID | Country, SC / MC | Intervention | Primary outcome | N patients | Mean age | Mean MAP at baseline (mmHg) | Mean lactate at baseline (mmol/l) | Mean cardiac index (l/min/m²) | APACHE II / SAPS-2 / SOFA score at baseline¹ | 28 days, All-cause mortality | Mortality in longest follow up, follow-up duration |
|----------|-----------------|--------------|----------------|------------|---------|-----------------------------|---------------------------------|----------------------------------|---------------------------------|-----------------------------|-----------------------------------------------|
| Agrawal 2011 [11] | India, SC | Norepinephrine | hemodynamic | 25 | 52 | NS | NS | 5.45 | 25 (A) | NS | NS |
| | | Dopamine | | 25 | 54 | NS | NS | 5.28 | 24 (A) | NS | NS |
| Albanese 2005 [12] | France, SC | Norepinephrine | hemodynamic | 10 | 65 | 54 | NS | 5.1 | 29 (A) | 40% | NS |
| | | Terlipressin | | 10 | 66 | 54 | NS | 5.0 | 28 (A) | 50% | NS |
| Annane 2007 [13] | France, MC | Norepinephrine + Dobutamine | survival | 169 | 60 | 68 | 3.3 | NS | 52 (S) | 34% | 50% |
| | | Epinephrine | | 161 | 65 | 70 | 2.9 | NS | 54 (S) | 39% | 52% |
| Chen 2012 [14] | China, SC | Norepinephrine | hemodynamic | 40 | NS | NS | NS | NS | NS | 22% | NS |
| | | Dopamine | | 40 | NS | NS | NS | NS | NS | 27% | NS |
| De Backer 2003 [16] | Belgium, SC | Norepinephrine crossover | hemodynamic | 5 | NS | NS | NS | NS | NS | 19 (A) | 80% | NS |
| | | Epinephrine crossover | | 5 | NS | NS | NS | NS | NS | 19 (A) | 80% | NS |
| De Backer 2010 [15] | Belgium, Austria, Spain, MC | Norepinephrine | survival | 502 | NS | NS | NS | NS | NS | 49% | NS |
| | | Dopamine | | 542 | NS | NS | NS | NS | NS | 53% | NS |
| Duranteau 1999 [17] | France, SC | Norepinephrine crossover | hemodynamic | 4 | 54 | NS | NS | NS | NS | 58% | NS, NS |
| | | Norepinephrine + Dobutamine crossover | | 4 | 54 | NS | NS | NS | NS | 58% | NS, NS |
| | | Epinephrine crossover | | 4 | 54 | NS | NS | NS | NS | 16% | NS, NS |

(Continued)
| Study ID | Country, SC / MC | Intervention | Primary outcome | N patients | Mean age | Mean MAP at baseline (mmHg) | Mean lactate at baseline (mmol/l) | Mean cardiac index (l/min/m²) | APACHE II / SAPS-2 / SOFA score at baseline | 28 days, All-cause mortality | Mortality in longest follow up, follow-up duration |
|----------|-----------------|-------------|----------------|------------|---------|----------------------------|-----------------------------------|---------------------------------|---------------------------------------------|------------------------------------------|-------------------------------------------------|
| France, SC | Norepinephrine | crossover | 6 | 40 | 56.8 | 2.3 | 4.07 | 14 (A) | | |
| Dopamine | crossover | 6 | 40 | 56.8 | 2.3 | 4.07 | 14 (A) | | |
| High 2008 | hemodynamic | | 56 | | | | | | NS | |
| China, SC | Norepinephrine | 23 | NS | NS | NS | NS | NS | 39% | NS | |
| Dopamine | 21 | NS | NS | NS | NS | NS | 38% | NS | |
| Jain 2010 | hemodynamic | | | | | | | | NS | |
| India, SC | Norepinephrine | 27 | 42 | 68 | NS | 5.0 | 17.6 (A) | 55% | NS | |
| Phenylephrine | 27 | 45 | 69 | NS | 4.8 | 19.1 (A) | 59% | NS | |
| Lauzier 2006 | hemodynamic | | | | | | | | NS | |
| France, | Norepinephrine | 10 | 58.1 | 68 | 3.33 | 4.4 | 23.5 (A) | 33% | NS | |
| Canada, MC | Vasopressin | 13 | 51.2 | 72 | 2.87 | 4.6 | 22.8 (A) | 23% | NS | |
| Levy 1997 | hemodynamic | | | | | | | | NS | |
| France, SC | Norepinephrine | 15 | 56 | 60 | 3.1 | 4.0 | 25.6 (A) | 53% | NS | |
| + Dobutamine | 15 | 54 | 60 | 3.1 | 4.0 | 24.5 (A) | 60% | NS | |
| Epinephrine | 15 | 54 | 60 | 3.1 | 4.0 | 24.5 (A) | 60% | NS | |
| Liu 2010 | hemodynamic | | 43 | | | | | | NS | |
| China, SC | Norepinephrine | 25 | NS | 4.9 | NS | 16 (A) | 32% | NS | |
| Dopamine | 25 | NS | 4.6 | NS | 17 (A) | 48% | NS | |
| Malay 1999 | hemodynamic | | | | | | | | NS | |
| US, SC | Placebo | 5 | 56 | 64 | NS | 5.6 | 26 (A) | 40% | NS | |
| Vasopressin | 5 | 53 | 66 | NS | 4.3 | 27 (A) | 0 | NS | |
| Marik 1994 | hemodynamic | | | | | | | | NS | |
| US, SC | Norepinephrine | 10 | 46 | 65 | 1.8 | 4.2 | 18 (A) | 50% | NS | |
| Dopamine | 10 | 46 | 63 | 2.2 | 4.2 | 17 (A) | 60% | NS | |
| Marthur 2007 | hemodynamic | | | | | | | | NS | |
| India, SC | Norepinephrine | 25 | 52 | NS | NS | 5.4 | 25.6 (A) | 56% | NS | |
| Dopamine | 25 | 54 | NS | NS | 5.2 | 24.5 (A) | 76% | NS | |
| Martin 1993 | hemodynamic | | | | | | | | NS | |

(Continued)
| Study ID | Country, SC / MC | Intervention | Primary outcome | N patients | Mean age | Mean MAP at baseline (mmHg) | Mean lactate at baseline (mmol/l) | Mean cardiac index (l/min/m²) | APACHE II / SAPS-2 / SOFA score at baseline | 28 days, All-cause mortality | Mortality in longest follow up, follow-up duration |
|----------|------------------|--------------|-----------------|------------|-----------|----------------------------|----------------------------------|-----------------------------|---------------------------------|-----------------------------|-----------------------------------------------|
| France, SC | Norepinephrine | 16 | 52 | 54 | 4.8 | 5.4 | 31 (A) | 43% | NS |
| Dopamine | 16 | 53 | 53 | 4.8 | 5.3 | 30 (A) | 62% | NS |
| Morelli 2008 [30] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Italy, SC | Norepinephrine | 16 | 70 | 55 | 2.8 | 4.4 | 55 (S) | 56% | NS |
| Phenylephrine | 16 | 70 | 54 | 2.8 | 4.3 | 57 (S) | 62% | NS |
| Morelli 2009 [29] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Italy, SC | Norepinephrine | 15 | 64 | 54 | 3.1 | 4.0 | 58 (S) | 66% | NS |
| Phenylephrine | 15 | 66 | 53 | 3.0 | 4.0 | 60 (S) | 53% | NS |
| Terlipressin | 15 | 67 | 53 | 3.1 | 4.0 | 62 (S) | 46% | |
| Morelli 2011[28] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Italy, SC | Norepinephrine | 20 | 66 | 71 | 2.5 | 4.0 | 54 (S) | NS | NS |
| Vasopressin | 20 | 61 | 72 | 2.3 | 4.0 | 53 (S) | NS | NS |
| Terlipressin | 20 | 65 | 71 | 1.8 | 3.8 | 50 (S) | NS | NS |
| Myburgh 2008 [31] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Australia, MC | Norepinephrine | 82 | 60 | 66 | 2.4 | NS | 22.7 (A) | 29% | 36.5% |
| Epinephrine | 76 | 58 | 65 | 2.7 | NS | 23.4 (A) | 22% | 31% |
| Patel 2002 [32] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Canada, SC | Norepinephrine | 11 | 68 | 68 | NS | 5.0 | 24 (A) | NS | NS |
| Vasopressin | 13 | 68 | 69 | NS | 4.8 | 22 (A) | NS | NS |
| Patel 2010 [33] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| US, MC | Norepinephrine | 118 | NS | NS | NS | NS | 27 (A) | 43% | 43% |
| Dopamine | 134 | NS | NS | NS | NS | 28 (A) | 50% | 50% |
| Plotkin 2007 [34] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Russia, SC | Norepinephrine | 33 | 40 | NS | 3.2 | 2.8 | 24 (A) | 90% | NS |
| Terlipressin | 41 | 40 | NS | 3.6 | 2.8 | 24 (A) | 73% | NS |
| Ruokonen 1993 [35] | | | | | | | | | |
| Dopamine | | | | | | | | | |
| Finland, SC | Norepinephrine | 5 | 44 | 55 | 2.1 | 4.4 | 13.3 (A) | 80% | NS |
| Dopamine | 5 | 46 | 59 | 1.3 | 4.1 | 13.3 (A) | 60% | NS |
| Russell 2008 [36] | | | | | | | | | |
| Dopamine | | | | | | | | | |

Table 1. (Continued)
| Study ID | Country, SC / Intervention | Primary outcome | N | Mean patients age | Mean MAP at baseline (mmHg) | Mean lactate at baseline (mmol/l) | Mean cardiac index (l/min/m²) | APACHE II / SAPS-2 / SOFA score at baseline ² | 28 days, All-cause mortality | Mortality in longest follow-up, follow-up duration |
|----------|-----------------------------|-----------------|---|-------------------|-----------------------------|---------------------------------|-------------------------------|-----------------------------------|---------------------------------|-----------------------------------------------|
| US, Australia, Canada, MC | Norepinephrine | hemodynamic | 395 | 61 73 | 3.5 | NS | 27.1 (A) | 39% | 49% | 49% |
| Schreuder 1989 [37] | Norepinephrine | hemodynamic | 395 | 61 73 | 3.5 | NS | 27.1 (A) | 39% | 49% | 49% |
| Schreuder 2002 [38] | Norepinephrine crossover | hemodynamic | 5 | 57 56 | 2.3 | 4.07 | NS | 49% | 59% | 59% |
| Seguin 2012 [39] | Norepinephrine + Dobutamine | hemodynamic | 11 | 70 51 | 5.3 | 3.0 | 62 (S) | 45% | 56% | 56% |
| Svoboda 2012 [39] | Norepinephrine | hemodynamic | 11 | 65 54 | 3.9 | 3.2 | 57 (S) | 36% | 50% | 50% |
| Wu 2010 [40] | Norepinephrine | hemodynamic | 11 | 70 74 | 5.3 | 3.0 | 62 (S) | 45% | 56% | 56% |
| Zhou 2002 [41] | Norepinephrine crossover | hemodynamic | 4 | 59 60 | 2.3 | 4.07 | NS | 56% | 56% | 56% |
| Zhuangyu 2011 [42] | Norepinephrine | hemodynamic | 45 | 54 71 | 3.5 | 3.5 | 21 (A) | 30% | 40% | 40% |
| All results present in mean (or median if not reported).  
¹ MC—multicenter, SC—single-center.  
² (A)—APACHE II, (S)—SAPS-2, (SO)—SOFA score.

Table 1. (Continued)
triple blinded [15], 8 trials were double blinded [13, 24, 26, 27, 30–32, 36], 2 were single blinded [20, 34] and the others were open-labeled or not stated. Results were analysed by intention-to-treat in all trials. Informed consent and ethical committee approval were described in 26/32 and 25/32 trials, respectively. Industrial sponsorship was stated in one trial [39]. Definitions of sepsis and septic shock for inclusion criteria, were described as suggested by the ACCP/SCCM consensus or similar by 30/32 (93.7%) of all studies.

**Primary outcome**

Mortality was reported in eleven trials that compared norepinephrine to dopamine. Funnel plot did not detect high risk of publication bias (S1 Fig). Norepinephrine was associated with reduced all-cause mortality (RR 0.89, 95% CI 0.81–0.98, I² = 0%), corresponding to an absolute risk reduction of 11% and number needed to treat of 9 (Fig 3). Excluding De Backer’s study from 2010 [15] which contributed 64% of the weight, did not alter the results, RR 0.84 (95% CI 0.70–0.99, I² = 0%, n = 10), as the exclusion of other trials. Sensitivity analysis restricted to trials from the developed world, trials published after 2004, trials with adequate allocation concealment and generation and blinding did not alter this result. Analysis based on trials whose primary outcome was clinical [15, 33, 42] revealed the same trend for reduced mortality with norepinephrine that was not statistically significant (RR 0.91 95% CI 0.82–1.02).

There was no statistically significant mortality benefit with norepinephrine over epinephrine (RR 0.96, 95% CI 0.77–1.21, I² = 0%, n = 4). Excluding one trial that did not add dobutamine to norepinephrine did not alter the results (RR 0.89, 95% CI 0.69–1.14, I² = 0%, n = 3). There was no statistically significant mortality benefit with norepinephrine over vasopressin / terlipressin RR 1.07 (95% CI 0.91–1.26, I² = 0%, n = 5), and over phenylephrine RR 0.92 (95% CI 0.64–1.32, I² = 0%, n = 2). Results for the comparison of dopamine to terlipressin, and vasopressin to placebo were not pooled (one trial for each comparison, no change in mortality ratio within each trial). When compared to all other treatments, norepinephrine and vasopressin were not associated with a benefit in mortality (RR 0.96, 95% CI 0.86–1.04, I² = 0%, n = 16, and RR 1.01, 95% CI 0.88–1.15, I² = 14%, n = 8, respectively). Comparing adrenergic to
Table 2. Risk of bias assessment.

| Study ID          | Sequence generation | Allocation concealment | Blinding | Ethical committee / informed consent | ITT analysis of primary outcome | Incomplete data reporting | Valid definition of septic shock |
|-------------------|---------------------|------------------------|----------|-------------------------------------|-------------------------------|---------------------------|---------------------------------|
| Agrawal 2011 [11] | Unclear             | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Albanese 2005 [12]| Low                 | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Annane 2007 [13] | Low                 | Low                    | Double   | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Chen 2012 [14]   | Unclear             | Unclear                | Open     | Unclear / Unclear                   | Yes                           | Unclear                    | Yes                             |
| De Backer 2003 [16]| Unclear            | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| De Backer 2010 [15]| Low                | Low                    | Triple   | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Duranteau 1999 [17]| Unclear           | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Guérin 2005 [18] | Unclear             | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| High 2008 [19]   | Unclear             | Unclear                | Open     | Unclear / Unclear                   | Yes                           | Unclear                    | No                              |
| Jain 2010 [20]   | Low                 | Low                    | Single   | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Lauzier 2006 [21]| Low                 | Low                    | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Levy 1997 [22]   | Low                 | Low                    | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Liu 2010 [23]    | Unclear             | Unclear                | Open     | Unclear / Unclear                   | Yes                           | Unclear                    | Yes                             |
| Malay 1999 [24]  | Low                 | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Marik 1994 [25]  | Low                 | Low                    | Open     | Yes / Unclear                       | Yes                           | Unclear                    | Yes                             |
| Marthur 2007 [26]| Unclear             | Unclear                | Double   | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Martin 1993 [27] | Low                 | Low                    | Double   | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Morelli 2008 [29]| Low                 | Low                    | Double   | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Morelli 2009 [29]| Low                 | Low                    | Open     | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Morelli 2011 [28]| Unclear             | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Myburgh 2008 [31]| Low                 | Low                    | Double   | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Patel 2002 [32]  | Low                 | Unclear                | Double   | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Patel 2010 [33]  | High                | Unclear                | Open     | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Plotkin 2007 [34]| Unclear             | Unclear                | Single   | Unclear / Unclear                   | Yes                           | Unclear                    | Yes                             |
| Ruokonen 1993 [35]| Unclear            | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Russell 2008 [36]| Low                 | Low                    | Double   | Yes / Yes                           | Yes                           | No                         | Yes                             |
| Schreuder 1989 [37]| Unclear          | Unclear                | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
| Seguin 2002 [38] | Low                 | Low                    | Open     | Yes / Yes                           | Yes                           | Unclear                    | Yes                             |
non-adrenergic vasopressors resulted in no benefit in mortality for either groups (RR 1.17, 95% CI 0.90–1.51, I² = 10%, n = 7). The pre-defined sensitivity analyses did not alter these results.

Secondary outcomes

Length of ICU stay, days free of vasopressors / ventilation and AEs.  ICU stay was reported in 6 trials, all compared norepinephrine to other vasopressors (median 13 days range 7–25). No advantage was found for the use of norepinephrine over other vasopressors (mean difference 1.01 days, 95% CI -0.65–2.66, I² = 0%) (S2 Fig). Hospital stay in patients discharged alive was reported in 3 trials, all compare norepinephrine to other vasopressors (median 15 days, range 7–52). No advantage was found for the use of norepinephrine. Only two trials reported data for days free of ventilation and days free of vasopressor support, therefore we did not pool these results. Eight trials reported major AEs, which included life threatening tachyarrhythmias (37 events, 3 trials), any arrhythmias (475 events, 7 trials), myocardial ischemia or arrest (80 events, 5 trials), internal organ or limb ischemia (126 events, 4 trials) and other (9 events, 1 trial). Major AEs were decreased with norepinephrine in comparison to dopamine, RR 0.34 (95% CI 0.14–0.84, I² = 0%, n = 3, Fig 4). Arrhythmias, were significantly decreased with norepinephrine compared to dopamine, RR 0.48 (95% CI 0.40–0.58, I² = 30%, n = 4, Fig 4). Compared with vasopressin / terlipressin, norepinephrine was not associated with a statistically significant change in major AEs and arrhythmias. Due to lack of reported data, we did not pool other AEs separately. Data for AEs with epinephrine and phenylephrine was sparse and did not allow pooled meta-analysis.

Hemodynamic data—ScvO2, urinary output, lactate clearance, MAP, CVP.  Forest plots of the presented data are available as Supporting Information (S2 Fig). Clinical and hemodynamic measurements at the beginning of the interventions, varied significantly between the trials (S1 Table). No vasopressor had a statistically significant effect of the MAP at any measurement point, compared to other vasopressor. CVP was higher with norepinephrine at the
first measuring point compared to all other vasopressors by a mean of 0.84 mmHg (95% CI 0.16–1.51, I² = 0%, n = 7). No vasopressor had a statistically significant effect on ScvO₂ levels compared to other vasopressors, at any measurement point. Lactate levels in the first measuring point were lower with norepinephrine compared to vasopressin / terlipressin by a mean 0.23 mmol/l (95% CI 0.13–0.34, I² = 0, n = 6). Compared to all other treatment, norepinephrine was not associated with decreased lactate levels at the first measuring point. No other effect on lactate levels was demonstrated with either comparison and by the pre-defined sensitivity analysis. Norepinephrine was associated with an increase in urine output compared to dopamine by a mean of 0.31 ml/kg/min (95% CI 0.12–0.49, I² = 0%, n = 4). There were no other statistically significant results comparing urine output between other vasopressors.

**Hemodynamic data**—other measurements

Forest plots of the presented data are available at Supporting Information (S2 Fig). Fourteen trials (43.5%) included only patients with high baseline CIX (defined as CIX > 3.5 l/min/m² by authors). At the first measuring point, CIX decreased with norepinephrine compared to dopamine by a mean of 0.35 l/min/m² (95% CI 0.22–0.48, I² = 18%, n = 7), to epinephrine by a mean of 0.87 l/min/m² (95% CI 0.57–1.16, I² = 0%, n = 4), and to all other vasopressors by a mean 0.7 l/min/m² (95% CI 0.42–0.97, I² = 42%, REM, n = 14). Heart rate was decreased with norepinephrine compared to dopamine in the first measuring point by a mean difference of 18.76 beats per minute (95% CI 9.76–27.76. I² = 96%, REM, n = 8). When comparing norepinephrine to all other interventions, a decreased heart rate at the first measurement point of 8.92 beats per minute was noted (95% CI 1.6–16.23, I² = 97%, REM, n = 19). One-hour SVRI measurements were higher with norepinephrine compared to dopamine and vasopressin / terlipressin by a mean difference of 192.82, 95% CI 60–325, I² = 86%, REM, n = 8 and 196.56 (95% CI 4.4–3.88, I² = 0%, n = 3, respectively).

Oxygen delivery index in the first measuring point were decreased with norepinephrine compared to other vasopressors by a mean of 14.06 ml/min/m² (95% CI 0.16–27.9, I² = 59%, REM, n = 10). First measuring point splanchnic CO₂ difference was lower with norepinephrine in comparison to epinephrine by a mean difference of 3.74 (95% CI 1.82–5.66, I² = 42%, REM, n = 4). There were insufficient data to compare vasopressors with regard to splanchnic blood flow and splanchnic oxygen delivery index.

**Discussion**

The pooled evidence summarized in this meta-analysis shows absolute reduction of 11% in 28-days all-cause mortality with norepinephrine compared with dopamine corresponding to
number needed to treat of 9. Dopamine resulted in more than twice the risk for major AEs including a twofold increase in the risk for cardiac arrhythmias. The hemodynamic profile of norepinephrine was also more favorable than the other vasopressors, resulting in decreased lactate levels, increased CVP and urine output in comparison to the other vasopressors. Further benefits of norepinephrine included reduced CIX and heart rate, elevated SVRI and reduced VIO₂ and splanchnic CO₂ difference.

We did not demonstrate mortality benefit with norepinephrine over epinephrine, phenylephrine and vasopressin / terlipressin or between the other comparisons, although a trend towards reduced mortality with norepinephrine was seen in all comparisons. Clinical outcomes other than mortality were seldom reported and therefore it was not possible to get strong evidence for length of ICU / hospital stay and ventilator / vasopressor free days. AEs were also reported only by 15/32 trials. The lower rates of cardiac arrhythmia and major AEs with norepinephrine compared to dopamine might account for the reduced mortality rates observed with norepinephrine.

Data regarding clinical and hemodynamic measurement were sparse and inconsistent in methodology and time of report. Although considered as targets for early goal directed therapy, most trials did not report on the variables included in the algorithm (including both studies that were published before and after 2004 with the publication of the surviving sepsis campaign [8]). The concept of early goal directed therapy was challenged recently in a large RCT [43], in which no mortality benefit with protocol based resuscitation was observed compared with usual care. Thus, obtaining the predefined hemodynamic goal with either vasopressor may not be translated to any clinical outcome change.

No statistically significant changes were observed between vasopressin / terlipressin and all adrenergic vasopressors (norepinephrine, dopamine) regarding mortality and AEs. One of the rationales for the use of non-adrenergic vasopressors is relative vasopressin deficiency in patients with septic shock [44] and the hypothesis that vasopressin can reduce the need for catecholamines and AEs associated with adrenergic stimulation. We did not observe a reduction in AEs, but most patients in the vasopressin arm received norepinephrine at randomization and at least for a part of the trial period as open-label vasopressor.

The use of vasopressors and inotropes for several conditions were addressed previously in systematic reviews and meta-analysis. A Cochrane systematic review [45] assessed the efficacy of vasopressors for the treatment of any circulatory failure in RCTs. The authors did not pool results for studies assessing septic patients alone. No mortality benefit was demonstrated for all direct comparisons between different vasopressors or vasopressor combinations. Dopamine was associated with more arrhythmias. Another review [46], focused only on the comparison of norepinephrine and dopamine in septic shock, but included observational studies as well as randomized controlled trials. It showed an advantage of norepinephrine over dopamine with regard to 28 days, all-cause mortality, RR for increased mortality with dopamine 1.12 (95% CI 1.01–1.20, I² = 0%, n = 6). Vasu et al. [47] assessed norepinephrine and dopamine in septic shock, however the data quoted from the largest trial were inaccurate [15], thus introducing further bias.

Several limitations of this review should be noted. Included trials span a long period, starting before 1989 and the last published in 2012. During this period advances in support and early goal directed treatment had been introduced to improve the quality of care in ICU. However, mortality rates were unchanged from early to recent trials (weighted mean crude mortality rate 45.0%). We could not perform a meta-analysis of differences from baseline for the hemodynamic measures, which might be more easily interpreted than end values, because results were not reported as such. An individual patient data meta-analysis might provide a better understanding of the evidence accrued to date. To note, most of the trials were
methodologically good or very good, however the treatment algorithm were different in regard to dosages, time to events and monitoring. The trials were diverse in reporting and statistical methods.

Future trials should use a uniformly acceptable protocol consisting of adequate methodological design, a common sepsis management algorithm (including the use of fluid resuscitation, stress dose steroids and initial hemodynamic targets). The reported outcomes should include all-cause mortality and, for patients discharged alive, length of hospital or ICU stay, length of ventilation, length of vasopressor support and AEs.

Conclusion

These data supply further support for the use of norepinephrine over dopamine for the treatment of patients with septic shock in ICUs, given consistent reduced all-cause mortality at 28 days with supporting hemodynamic data and lower rate of major AEs and cardiac arrhythmias. This recommendation is based on data from all the RCTs published up to date, with no heterogeneity. Trials to guide recommendations for the use other vasopressors, especially non-adrenergic, are needed.

Supporting Information

S1 Fig. Funnel plot of norepinephrine vs dopamine primary outcome, Begg and Mazumdar rank correlation, Egger’s regression intercept. (DOCX)

S2 Fig. Forest plots (a-r). a, Norepinephrine vs epinephrine, Mortality primary. b, Norepinephrine vs vasopressin OR terlipressin, Mortality primary. c, Norepinephrine vs phenylephrine, Mortality primary. d, Norepinephrine vs other, Mortality primary. e, Vasopressin / terlipressin vs other, Mortality Primary. f, Norepinephrine vs other, ICU (or hospital) stay. g, Norepinephrine vs other, 1st measurement point CVP. h, Norepinephrine vs vasopressor or terlipressin, 1st measurement point lactate. i, Norepinephrine vs dopamine, 1st measurement point urine output. j, Norepinephrine vs dopamine, 1st measurement point CIX. k, Norepinephrine vs epinephrine, 1st measurement point CIX. l, Norepinephrine vs other, 1st measurement point CIX. m, Norepinephrine vs dopamine, 1st measurement point heart rate. n, Norepinephrine vs other, 1st measurement point heart rate. o, Norepinephrine vs dopamine, 1st measurement point SVRI. p, Norepinephrine vs vasopressor or terlipressin, 1st measurement point SVRI. q, Norepinephrine vs other, 1st measurement point Vlo2. r, Norepinephrine vs Epinephrine, 1st measurement point splanchnic CO2 difference. (DOCX)

S1 Table. Hemodynamic data reported and baseline, first and second measurement points. (DOCX)

S2 Table. List of excluded studies and reason for exclusion. (DOCX)

S3 Table. PRISMA 2009 checklist. (DOCX)
Author Contributions
Conceived and designed the experiments: TA. Performed the experiments: TA AL SL LL MP AG. Analyzed the data: TA AL. Contributed reagents/materials/analysis tools: TA. Wrote the paper: TA.

References
1. Annane D, Aegerter P, Jars-Guincestre MC, Guidet B (2003) Current epidemiology of septic shock: the CUB-Rea Network. Am J Respir Crit Care Med 168: 165–172. PMID:12851245
2. Angus DC, Linde-Zwirble WT, Lidicker J, Clermont G, Carcillo J, Pinsky MR (2001) Epidemiology of severe sepsis in the United States: analysis of incidence, outcome, and associated costs of care. Crit Care Med 29: 1303–1310. PMID:11445675
3. Friedman G, Silva E, Vincent JL (1998) Has the mortality of septic shock changed with time. Crit Care Med 26: 2078–2086. PMID:9875924
4. (1992) American College of Chest Physicians/Society of Critical Care Medicine Consensus Conference: definitions for sepsis and organ failure and guidelines for the use of innovative therapies in sepsis. Crit Care Med 20: 864–874. PMID:1597042
5. Rivers E, Nguyen B, Havstad S, Ressler J, Muzzin A, Knobilch B et al. (2001) Early goal-directed therapy in the treatment of severe sepsis and septic shock. N Engl J Med 345: 1368–1377. PMID:11794169
6. Arnold RC, Shapiro NI, Jones AE, Schorr C, Pope J, Casner E et al. (2009) Multicenter study of early lactate clearance as a determinant of survival in patients with presumed sepsis. Shock 32: 35–39. PMID:19533847
7. Hollenberg SM, Ahrens TS, Annane D, Astiz ME, Chalfin DB, Heard SO et al. (2004) Practice parameters for hemodynamic support of sepsis in adult patients: 2004 update. Crit Care Med 32: 1928–1948. PMID:15343024
8. Dellinger RP, Carlet JM, Masur H, Gerlach H, Calandra T, Cohen J et al. (2004) Surviving Sepsis Campaign guidelines for management of severe sepsis and septic shock. Crit Care Med 32: 858–873. PMID:15090974
9. Higgins JPT, Green S (editors) (2008) Cochrane Handbook for Systematic Reviews of Interventions Version 5.0.1 [updated September 2008]. The Cochrane Collaboration. Available:www.cochrane-handbook.org.
10. Hozo SP, Djulbegovic B, Hozo I (2005) Estimating the mean and variance from the median, range, and the size of a sample. BMC Med Res Methodol 5: 13. PMID:15840177
11. Agrawal A, Gupta A, Consul S, Shastri P (2011) Comparative study of dopamine and norepinephrine in the management of septic shock. Saudi J Anaesth 5: 162–166. doi:10.4103/1658-354X.82784 PMID:21804796
12. Albanese J, Leone M, Delmas A, Martin C (2005) Terlipressin or norepinephrine in hyperdynamic septic shock: a prospective, randomized study. Crit Care Med 33: 1897–1902. PMID:16148457
13. Annane D, Vignon P, Renault A, Bollaert PE, Charpentier C, Martin C et al. (2007) Norepinephrine plus dobutamine versus epinephrine alone for management of septic shock: a randomised trial. Lancet 370: 676–684. PMID:17720019
14. Chen H, Zeng Z (2012) Comparison of the effect and complication between dopamine and norepinephrine on treatment of the septic shock. Jiangxi Medical Journal 47: 565–567.
15. De Backer D, Biston P, Devisendt J, Madi C, Chochrad D, Aldecoa C et al. (2010) Comparison of dopamine and norepinephrine in the treatment of shock. N Engl J Med 362: 779–789. doi: 10.1056/NEJMoa0907118 PMID:20200382
16. De Backer D, Cretier J, Silva E, Vincent J (2003) Effects of dopamine, norepinephrine, and epinephrine on the splanchnic circulation in septic shock: which is best? Crit Care Med 31: 1659–1667. PMID:12794401
17. Duranteau J, Sitbon P, Teboul JL, Vicaut E, Anguel N, Samii K (1999) Effects of epinephrine, norepinephrine, or the combination of norepinephrine and dobutamine on gastric mucosa in septic shock. Crit Care Med 27: 893–900. PMID:10362410
18. Guerin JP, Levrat J, Samat-Long C, Leverve X, Grimaud D, Ichai C (2005) Effects of dopamine and norepinephrine on systemic and hepatosplanchnic hemodynamics, oxygen exchange, and energy balance in vasoplegic septic patients. Shock 23: 18–24. PMID:15614126
19. High K (2008) Impact of Dopamine and norepinephrine on renal perfusion in patients with septic shock. Journal of Hebei Medicine 30: 1188–1121.
20. Jain G, Singh DK (2010) Comparison of phenylephrine and norepinephrine in the management of dopamine-resistant septic shock. Indian J Crit Care Med 14: 29–34. doi: 10.4103/0972-5229.63033
PMID: 2060906

21. Lauzier F, Levy B, Lamarre P, Lesur O (2006) Vasopressin or norepinephrine in early hyperdynamic septic shock: a randomized clinical trial. Intensive Care Med 32: 1782–1789. PMID: 17019548

22. Levy B, Bollaert PE, Charpentier C, Nace L, Audibert G, Bauer P et al. (1997) Comparison of norepinephrine and dobutamine to epinephrine for hemodynamics, lactate metabolism, and gastric tonometric variables in septic shock: a prospective, randomized study. Intensive Care Med 23: 282–287. PMID: 9083230

23. Liu P, Chen T, Zhang Y (2010) Comparison evaluation of resuscitation effect of norepinephrine and dopamine on the treatment of septic shock. Clinical Education of General Practice 8: 265–267.

24. Malay MB, Ashton RC Jr., Landry DW, Townsend RN (1999) Low-dose vasopressin in the treatment of vasodilatory septic shock. J Trauma 47: 699–703; discussion 703–695. PMID: 10528604

25. Marik PE, Mohedin M (1994) The contrasting effects of dopamine and norepinephrine on systemic and splanchic oxygen utilization in hyperdynamic sepsis. JAMA 272: 1354–1357. PMID: 7933396

26. Marthur K, Dhunna R, Chakraborty A (2007) Comparison of norepinephrine and dopamine in the management of septic shock using impedance cardiography. Indian J Crit Care Med 11: 186–191.

27. Martin C, Papazian L, Perrin G, Saux P, Gouin F (1993) Norepinephrine or dopamine for the treatment of hyperdynamic septic shock? Chest 103: 1826–1831. PMID: 8404107

28. Morelli A, Donati A, Ertem C, Rehberg S, Kampmeier T, Orecchioni A et al. (2011) Effects of vasopressinergic receptor agonists on sublingual microcirculation in norepinephrine-dependent septic shock. Crit Care 15: R217. doi: 10.1186/cc10453 PMID: 21929764

29. Morelli A, Ertem C, Rehberg S, Lange M, Orecchioni A, Cecchini V et al. (2009) Continuous terlipressin versus vasopressin infusion in septic shock (TERLIVAP): a randomized, controlled pilot study. Crit Care 13: R130. doi: 10.1186/cc7990 PMID: 19664253

30. Morelli A, Ertem C, Rehberg S, Lange M, Orecchioni A, Laderchi A et al. (2008) Phenylenephrine versus norepinephrine for initial hemodynamic support of patients with septic shock: a randomized, controlled trial. Crit Care 12: R143. doi: 10.1186/cc7121 PMID: 19017409

31. Myburgh JA, Higgins A, Jovanovska A, Lipman J, Ramakrishnan N, Santamaria J (2008) A comparison of epinephrine and norepinephrine in critically ill patients. Intensive Care Med 34: 2225–2234. doi: 10.1007/s00134-008-1219-0 PMID: 18654759

32. Patel GP, Chittock DR, Russell JA, Walley KR (2002) Beneficial effects of short-term vasopressin infusion during severe septic shock. Anesthesiology 96: 576–582. PMID: 11873030

33. Patel GP, Grahe JS, Sperry M, Singla S, Elpern E, Lateef O et al. (2010) Efficacy and safety of dopamine versus norepinephrine in the management of septic shock. Shock 33: 375–380. doi: 10.1097/SHK.0b013e3181c6ba8f PMID: 19951126

34. Plotkin LL (2007) [Use of vasopressin to correct hemodynamic disorders in patients with abdominal sepsis]. Anestesiol Reanimatol: 47–49.

35. Ruokonen E, Takala J, Kari A, Saxen H, Mertsola J, Hansen EJ (1993) Regional blood flow and oxygen transport in septic shock. Crit Care Med 21: 1296–1303. PMID: 8370292

36. Russell JA, Walley KR, Singer J, Gordon AC, Hebert PC, Cooper DJ et al. (2008) Vasopressin versus norepinephrine infusion in patients with septic shock. N Engl J Med 358: 877–887. doi: 10.1056/NEJMoa067373 PMID: 18305265

37. Schreuder WG, Schneider AJ, Gruenerwold AB, Thijs LG (1989) Effect of dopamine vs norepinephrine on hemodynamics in septic shock. Emphasis on right ventricular performance. Chest 95: 1282–1288. PMID: 2721267

38. Seguin P, Belissant E, Le Tulzo Y, Laviolle B, Lessard Y, Thomas R et al. (2002) Effects of epinephrine compared with the combination of dobutamine and norepinephrine on gastric perfusion in septic shock. Clin Pharmacol Ther 71: 381–388. PMID: 12011824

39. Svboda P, Scheer P, Kantorova I, Doubek J, Dudra J, Radvan M et al. (2012) Terlipressin in the treatment of late phase catecholamine-resistant septic shock. Hepatogastroenterology 59: 1043–1047. doi: 10.5755/hge10550 PMID: 22580654

40. Wu J, Chen J, Ou Y, Yang C, Chen M, Hwang SW et al. (2010) Effect of dopamine and norepinephrine on hemodynamics and oxygen metabolism of tissue in patients with septic shock. Chin Arch Gen Surg (Electronic Edition) 4: 117–121.

41. Zhou SX, Qiu HB, Huang YZ, Yang Y, Zheng RQ (2002) Effects of norepinephrine, epinephrine, and norepinephrine-dobutamine on systemic and gastric mucosal oxygenation in septic shock. Acta Pharmacol Sin 23: 654–658. PMID: 12100762
42. Zhuangyu Y (2011) Affect of norepinephrine and dopamine on infectious tissue oxygen metabolism and hemodynamics in patients with shock. Shandong Medicine Journal 51: 93–94.

43. Pro CI, Yealy DM, Kellum JA, Huang DT, Barnato AE, Weisfeld LA et al. (2014) A randomized trial of protocol-based care for early septic shock. N Engl J Med 370: 1683–1693. doi: 10.1056/NEJMoa1401602 PMID: 24635773

44. Sharshar T, Blanchard A, Paillard M, Raphael JC, Gajdos P, Annane D (2003) Circulating vasopressin levels in septic shock. Crit Care Med 31: 1752–1758. PMID: 12794416

45. Havel C, Arrich J, Losert H, Gamper G, Mullner M, Herkner H (2011) Vaspressors for hypotensive shock. Cochrane Database Syst Rev 5: CD003709.

46. De Backer D, Aldecoa C, Njimi H, Vincent JL (2012) Dopamine versus norepinephrine in the treatment of septic shock: a meta-analysis*. Crit Care Med. 40: 725–730. doi: 10.1097/CCM.0b013e31823778ee PMID: 22036860

47. Vasu TS, Cavallazzi R, Hirani A, Kaplan G, Leiby B, Marik PE (2011) Norephinephrine or Dopamine for Septic Shock: A Systematic Review of Randomized Clinical Trials. J Intensive Care Med. 27: 172–8. doi: 10.1177/0885066610396312 PMID: 21436167