The clinical outcomes of selenium supplementation on critically ill patients
A meta-analysis of randomized controlled trials

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Abstract

Purpose: Selenium supplementation is a potentially promising adjunctive therapy for critically ill patients, but the results are controversial among studies. Accordingly, we performed this meta-analysis to more clearly detect the efficacy and safety of selenium supplementation on critically ill patients.

Methods: Systematic literature retrieval was carried out to obtain RCTs on selenium supplementation for critically ill patients up to August 2017. Data extraction and quality evaluation of these studies were performed by 2 investigators. Statistical analyses were performed by RevMan 5.3. Trial sequential analysis (TSA) was conducted to control the risks of type I and type II errors and calculate required information size (RIS).

Results: Totally 19 RCTs involving 3341 critically ill patients were carried out in which 1694 participants were in the selenium supplementation group, and 1647 in the control. The aggregated results suggested that compared with the control, intravenous selenium supplement as a single therapy could decrease the total mortality (RR = 0.86, 95% CI: 0.78–0.95, P = .002, TSA-adjusted 95% CI: 0.77–0.96, RIS = 4108, n = 3297) and may shorten the length of stay in hospital (MD = 2.30, 95% CI: 1.15–3.45, P = .002, TSA-adjusted 95% CI: 1.13–3.35, RIS = 3482, n = 3297) and may shorten the length of ICU stay (MD = –1.68, 95% CI: –1.88 to –1.48, P = .84) in critically ill patients. Our results also showed that selenium supplementation did not increase incidence of drug-induced side effect compared with the control (RR 1.04, 95% CI 0.83 to 1.30, P = .73).

Conclusions: The current evidence suggests that the use of selenium could reduce the total mortality, and TSA results showed that our outcome is reliable and no more randomized controlled trials are needed. But selenium supplementation might have no effect on reducing 28-days mortality as well as the incidence of new infections, or on length of stay in ICU or mechanical ventilation. However, the results should be used carefully because of potential limitations.

Abbreviations: CIs = confidence intervals, D² = diversity, MDs = mean differences, RCTs = randomised controlled trials, RIS = required information size, RRs = risk ratios, SIRS = systemic inflammatory response syndrome, TSA = Trial sequential analysis.

Keywords: meta-analysis, randomized controlled trials, selenium supplementation, trial sequential analysis (TSA)

1. Introduction

Endoplasmic reticulum stress, oxidative stress and inflammatory response are increasingly being recognized as the central pathophysiology for critically ill patients. Especially the development of sepsis, septic shock, and multiple organ failure is responsible for a longer hospitalization period and increased risk of mortality.[1,2] Previous studies indicated that the circulating antioxidant and anti-inflammatory levels would decrease rapidly after injury, sepsis, or surgery and would remain below the normal level for several days or even weeks.[3] The severer the trauma, the systemic inflammatory response syndrome (SIRS), or the sepsis, the larger the depletion of antioxidants appears to be.[4] These changes are associated with an increase in the free radical generation, an augmentation of the inflammatory response, and are playing a direct role in cell death, increased morbidity, and even higher mortality in the critically ill patients.[13–15] Also, studies have proved that special enzymes such as superoxide dismutase, catalase, and glutathione peroxidase (including their cofactors such as selenium, zinc, iron, and manganese), sulphydryl group donors (glutathione), and vitamins (vitamins C, E, and β-carotene) can form a functional
network to protect physiological body from the above injury mechanisms. Current studies all focus on nutrition support with these compositions that may play a critical role in the recovery of the critically ill patients.

Selenium, a trace element, is one of the essential nutrients with regulatory, immunologic, and antioxidant functions. It may play an important role as an antioxidant as well as an anti-inflammatory in the glutathione peroxidase system. Supplementation of selenium is a promising adjunctive therapy for patients with SIRS, sepsis, or septic shock. Up to now, many clinical trials have studied the effect of selenium, being administered intravenously as a monotherapy, on clinical outcomes of critically ill patients (such as mortality, the length of ICU stay, the length of hospital stay, new infections). However, most of these current studies were performed in relatively small patient populations with trauma, SIRS, or sepsis, which are underpowered to detect the treatment effect on clinically outcomes. More importantly, the results are controversial between each other. More recently, several meta-analyses have been performed about selenium supplement on critically ill patients. In 2015, the meta-analysis of Allingstrup et al demonstrated that selenium supplement can reduce the overall mortality of critically ill patients. However, in 2016, Manzanares et al reported that selenium therapy could not reduce the mortality and improve other clinical outcomes of critically ill patients. In consideration of these inconsistencies, we carried out this meta-analysis of the randomized controlled trials (RCTs), aiming to detect the efficacy and safety of selenium supplementation on critically ill patients more clearly.

2. Materials and methods

2.1. Protocol and registration

This meta-analysis of randomized controlled trials was performed according to the PRISMA (Preferred Reporting Items for Systematic reviews and meta-analyses) recommendations. A protocol for this meta-analysis has been registered on PROSPERO (http://www.crd.york.ac.uk/prospero) and the registration number is: CRD42017079365.

2.2. Literature search

Three search engines, namely PubMed (1966–2017.8), Embase (1974–2017.8), and Cochrane library (Issue 8, 2017) were retrieved. The following key words were used: 'selenium', 'selenium derivative', 'seleniumic acid', 'sodium selenite', 'antioxidant cocktails', 'selenium compounds', 'randomized controlled trial', 'randomized', 'randomly', 'trial', 'clinical trials', 'controlled clinical', 'selenious acid', 'sodium selenite', 'selenium compounds', 'antioxidant cocktails'. Each study assessed the methodological qualities of trials by 2 investigators (Yan Zhao and Hongjun Kang) independently, and 3. interventions: The patients were randomly allocated to the selenium supplementation group or the control according to the telephone computer system or computerized randomization or random number table. For the selenium supplementation group they were given parenteral selenium supplementation singly at different doses (not in combination with other antioxidant micronutrients), while the control were given placebo or maintenance dose selenium or no intervention. In addition, critical patients in the 2 groups could receive other treatment.

4. outcomes: Primary end points: mortality at day 28 and total mortality (regardless of the follow-up period). Secondary end points: new infection, length of stay in ICU, length of stay in hospital and length of mechanical ventilation during follow-up.

2.4. Data extraction

According to Table 1, 2 investigators (Yan Zhao and Hongjun Kang) independently read the titles, abstracts and full texts with the following procedures:

1. examining titles and abstracts to remove obviously irrelevant studies,
2. retrieving the full texts of potentially relevant trials,
3. examining full texts for compliance of studies with eligibility criteria, and
4. making final decisions on data entry and proceeding to data collection.

Patient’s baseline information (treatment strategy, dose, and duration of supplementation) and detailed methods of research design (publication year, research settings, designs, methods of randomization, allocation concealment, blinding) were extracted from the selected studies. Disagreement was solved by discussion with the third investigator (Feihu Zhou).

2.5. Quality evaluation

Each study assessed the methodological qualities of trials by 2 investigators (Yan Zhao and Hongjun Kang) independently. The criterion was based on criteria described in Cochrane Reviewer’s Handbook 5.1.0, including the following risk of selection, performance, detection, attrition and reporting bias domains: random sequence generation, allocation concealment, blinding, incomplete outcome data, intention to treat analysis.

2.6. Data synthesis and statistical analysis

Differences were calculated as risk ratios (RRs) and expressed with 95% confidence intervals (CIs) for dichotomous outcomes and mean differences (MDs) with 95% CIs for continuous outcomes. Heterogeneity across analysis was done using I² statistic, which is a quantitative measure of the inconsistency of the across analysis. Studies with an I² statistic of 25% to 50%, 50% to 75%, and >75% are considered as low heterogeneity, moderate heterogeneity, and high heterogeneity, respectively. An I² value greater than 50% indicates a significant heterogeneity. A random-effects model was used in the case of significant heterogeneity (I² > 50%), otherwise, a fixed-effects model was used. We conducted sensitivity analyses to explore possible explanations for the heterogeneity on the overall pooled estimate and to examine the influence of various exclusion criteria on
| Study | Disease | Study population | Male/female | Interventions | Treatment duration | No. of patients | Outcomes |
|-------|---------|------------------|-------------|---------------|-------------------|----------------|----------|
| Kuklinski, 1991 | Acute pancreatitis | 28–65 | 17/0 | Selenium Control Administration | 1000 µg/day, loading bolus, thereafter 1000 µg/day for 3 days, 250 µg for 3 days, and 155 µg for 3 days, then after 35 µg per day infusion 2000 µg for 1 day, 1000 µg/day for 4 days, 300 µg/day until discharged 474 µg for 3 days, followed by 31.6 µg for 3 days, and 15.6 µg for 3 days and thereafter 31.6 µg per day 1000 µg/day loading bolus, followed by 1000 µg/day continuous infusion | 8 | 9 | Hospital fatality |
| Zimmermann, 1997 | SIRS, sepsis and organ failure | Not given | Not given | Selenium Control Administration | 35 µg per day infusion throughout the trial Treatment period 0.9% sodium chloride placebo Standard dose of 31.6 µg sodium selenite | 20 | 20 | Hospital fatality |
| Lindner, 2004 | Acute pancreatitis | Median 50–52 | 39/28 (complete) | Selenium Control Administration | 14 days | 116 | 122 | Hospital fatality, APACHE III score, length of ICU, new infections |
| Mishra, 2007 | Septic ICU patients | 66 | 29/13 | Selenium Control Administration | 316 g for 3 days, followed by 316 g per day 0.9% sodium chloride for 1 day, 1000 g/day for 4 days, 300 g/day until discharge | 21 | 21 | SOFA, hospital fatality |
| Angstwurm, 1999 | SIRS and sepsis | 64.6 (14.0) | 162/76 | Selenium Control Administration | 9 days | 122 | 116 | Hospital fatality, APACHE III score, length of ICU, new infections |
| Forceville, 2007 | Severe septic shock patients with documented infection | 66 (14)/69 (12) | 38/22 | Selenium Control Administration | 4.1 days for selenium, 4.7 days for control | 34 | 34 | SOFA, hospital fatality |
| Montoya, 2009 | ICU admission with a diagnosis of sepsis | 66 | 38/30 | Selenium Control Administration | 10 days | 31 | 29 | Fatality, |
| Andrews, 2011 | Sepsis | 58 (17)/54 (17) | 15/16 | Selenium Control Administration | 10 days | 122 | 127 | New infection, fatality, length of ICU |
| Manzanares, 2011 | SIRS | 60 (16)/60 (15) | 97/53 | Selenium Control Administration | 14 days | 75 | 75 | Mortality |
| Valenta, 2011 | sepsis or SIRS | 53 (23–79) | 49/23 | Selenium Control Administration | 10 days | 35 | 37 | 28-day mortality APACHE II score |
| Janka, 2013 | sepsis, severe sepsis or septic shock | 62 (54–76)/66 (57–78) | 23/17 | Selenium Control Administration | 14 days | 21 | 19 | Mortality |
| Wohl 2014 | severe septic patients with multiple organ failure | 35 (17–82)/41 (19–82) | 44/10 | Selenium Control Administration | 14 days | 29 | 29 | 28-day mortality, ICU length of stay, length of mechanical ventilation, SOFA score, renal failure |
| Chelkeba, 2015 | severe sepsis and septic shock | 65.7 (13.7) | 691/98 | Selenium Control Administration | 14 days | 543 | 546 | 28-day mortality, 90-day mortality, ICU length of stay, hospital length of stay, new infections |
| Bloos, 2016 | severe sepsis and septic shock in last 24 h | 35 (17–82)/41 (19–82) | 44/10 | Selenium Control Administration | 14 days | 29 | 29 | 28-day mortality, ICU length of stay, hospital length of stay, new infections |
| Chelkeba, 2017 | severe sepsis and septic shock | 35 (36.53) | 270/37 | Selenium Control Administration | 14 days | 125 | 128 | Mortality, ICU length of stay, mortality, ICU length of stay, hospital length of stay, side effect, hospital length of stay, mechanical ventilation |
| Khalili, 2017 | traumatic brain injury | 40.07 (17.82)/42.93 (17.19) | 90/23 | Selenium Control Administration | 14 days | 57 | 56 | Mortality, ICU length of stay, hospital length of stay, side effect, mechanical ventilation |
| Moghaddam, 2017 | acute traumatic brain injury | 66 (11)/68 (10) | 302/109 | Selenium Control Administration | 0.9% sodium chloride placebo | 206 | 205 | 28-days mortality, ICU length of stay, hospital length of stay, mechanical ventilation |

**Table 1**

**Characteristics of included trials.**
the overall pooled estimate. We further conducted Begg funnel plots to identify the existence of publication bias. Differences are considered statistically significant at $P < .05$. Statistical analyses were performed by RevMan version 5.3 (Cochrane Collaboration, Oxford, UK), and sensitivity analysis and funnel plots were conducted by STATA STATA 12.0 (StatCorp, College Station, TX, USA).

2.7. Trial sequential analysis (TSA)

The same as clinical trial, systematic review and meta-analysis also need to estimate sample size to reduce the risks of random errors and ensure the reliability of results. TSA is a method which could control the risks of type I and type II errors and calculate required information size (RIS) needed by systematic review and meta-analysis. When the cumulative Z curve crosses the trial sequential monitoring boundaries with or without the achievement of RIS, we think the anticipated intervention effect may have been reached and no further trials are needed. If RIS has been reached, but the cumulative Z curve crosses neither the trial sequential monitoring boundaries nor conventional boundaries, we think there is no statistical difference between 2 groups and no more trials are needed. If the cumulative Z curve crosses the futility boundaries, we can also think no difference exists between two groups. However, if the cumulative Z curve does not cross the trial sequential monitoring boundaries, at the same time, the RIS has not been reached, we conclude that more trials are needed.

We adopted a method of constant continuity correction for handing zero-event trials, and added a continuity correction factor of 0.5 to the number of events and non-events in each group.

Two-sided tests, a type I error of 5% and a type II error of 20% (a power of 80%) were used for calculating the RIS. For dichotomous data, incidence in the control was derived from the results of our meta-analysis, and a relative risk reduction or increase was estimated according to the information from related areas.

3. Results

3.1. Process for included trials

As shown in Figure 1, a total of 2827 potentially relevant studies were identified and screened for retrieval. Totally 389 studies were excluded because of duplications and 2400 studies were excluded after the titles and abstracts had been read. Thus 37 studies were assessed for eligibility. Because 15 studies of them included other positive antioxidants, and 3 studies selected oral route for administration, finally 19 RCTs were included in our review.

3.2. Characteristics of included trials

The main characteristics of the trials included in our meta-analysis were shown in Table 1. There were totally 3341 critically ill patients of which 1694 participants were in the selenium supplementation group, and 1647 in the control. Diseases in most of studies included SIRS, sepsis, septic shock and multiple organ failure. The doses of selenium supplement on the first day varied from 500 mg to 4000 mg in different studies, and patients in the selenium supplementation group from 13 RCTs received loading bolus on the first day varied from 1000 mg to 4000 mg. In three studies (500 mg/day) and Zimmermann research (1000 mg/day) the patients were given the same dose duration the treatment, while in the rest studies the patients were given a dynamic dose duration the treatment. In the control, patients in 5 RCTs were given a low-dose selenium from 31.6 mg/day to 100 mg/day, and in 7 RCTs were given 0.9% sodium chloride placebo, and in 3 studies were given standard therapy, and in 4 studies such interventions were not reported. The total treating period was reported in 17 trials. Thus, the total dose amount could be calculated by subtracting the control from the selenium supplement group, that is, by subtracting 2050 mg from 28,000 mg. The number of patients in

![Figure 1. Process for included trials.](image-url)
these studies varied from 17 to 1089 and hospital fatality was reported in all studies.

3.3. Risk of bias and quality of evidence

Figure 2 showed risk of bias in the included trials. The GRADE evidence quality for outcomes was summarized in Table 2.

3.4. Meta-analysis results

3.4.1. Primary end points

3.4.1.1. Overall mortality. We included nineteen trials with 3297 participants reporting overall mortality in 2 groups. The result indicated that selenium supplement could reduce the overall mortality compared with placebo or no intervention in critically ill patients (RR 0.86, 95% CI 0.78–0.95, P = .002) using a fixed effects model ($I^2$ = 24%, $P = .17$) (Fig. 3A).

Trial sequential analysis was conducted in the light of overall mortality in the control of 30%, a relative risk reduction in experimental group of 18%, and diversity ($D^2$) of 48%. The required information size was 4108 participants, 80.3% of which were accrued in our meta-analysis. The cumulative Z curve (blue line) crossed the trial sequential monitoring boundaries (red inward slash) before the RIS has been reached (Fig. 3B). The TSA-adjusted 95% CI of RR was 0.77 to 0.96.

3.4.1.2. Twenty eight days all causes mortality. We included ten trials with 2510 participants reporting 28-day all causes mortality in 2 groups. No significant difference was found between selenium supplement and placebo or no intervention (RR 0.96, 95% CI 0.85 to 1.09, $P = .54$) using a fixed effects model ($I^2$ = 31%, $P = .16$) (Fig. 4).

3.4.2. Secondary end points

3.4.2.1. Length of stay in ICU. We included nine trials with 1491 participants reporting length of stay in ICU in 2 groups. The result showed that selenium supplement could not shorten the length of stay compared with placebo or no intervention in critically ill patients (MD $\bar{Y}/\bar{X}$ 0.15, 95% CI 1.68 to 1.38, $P = .84$) using a random effects model ($I^2$ = 70%, $P = .0008$) (Fig. 5). The result of sensitivity analysis found that no single study had a significant influence on pooled MD (Additional file, http://links.lww.com/MD/C984).

3.4.2.2. Length of stay in hospital. We included seven trials with 1250 participants reporting length of stay in hospital in 2 groups. The result showed that selenium supplement may shorten the length of stay in hospital compared with placebo or no intervention in critically ill patients (MD $\bar{Y}/\bar{X}$ 2.30, 95% CI 4.03 to 0.57, $P = .006$) using a random effects model ($I^2$ = 67%, $P = .006$) (Fig. 6). The result of sensitivity analysis found that no single study had a significant influence on pooled MD (Additional file, http://links.lww.com/MD/C984).

3.4.2.3. Length of mechanical ventilation during follow-up. We included 6 trials with 368 participants reporting Length of mechanical ventilation during follow-up in 2 groups. The result showed that selenium supplement could not shorten the length of stay compared with placebo or no intervention in critically ill patients (MD $\bar{Y}/\bar{X}$ 0.98, 95% CI −3.38 to 1.41, $P = .42$) using a random effects model ($I^2$ = 82%, $P < .0001$) (Fig. 7). The result of sensitivity analysis found that no single study had a significant influence on pooled MD (Additional file, http://links.lww.com/MD/C984).

3.4.2.4. New infection. We included 6 trials with 1990 participants reporting number of new infected participants in critically ill patients (RR 0.97, 95%
CI 0.89–1.05, \( P = .43 \) using a fixed effects model (\( I^2 = 27\% \), \( P = .23 \)) (Fig. 8).

### 3.4.2.5. Drug-induced side effects

We included seven trials with 1038 participants reporting drug-induced side effects in 2 groups. The result showed that selenium supplement did not increase incidence of drug-induced side effect compared with placebo or no intervention in critically ill patients (RR 1.04, 95% CI 0.83–1.30, \( P = .73 \)) using a random effects model (\( I^2 = 50\% \), \( P = .06 \)) (Fig. 9). The result of sensitivity analysis found that no single study had a significant influence on pooled RR (Additional file, http://links.lww.com/MD/C984).

### 3.4.3. Publication bias

Begg funnel plot showed no publication bias (Additional file, http://links.lww.com/MD/C984).

### 4. Discussion

The pooled results from 19 RCTs using a fixed effects model suggest that selenium supplement could cause decrease in the total mortality in hospital but could not reduce the mortality at day 28. We conduct subgroup analysis such as loading bolus and no loading bolus; high total dose and low total dose; duration ≤ 9 days, duration > 9 days, and unknown duration, no significant subgroup difference was found. For the complications, results indicate selenium supplement did not increase incidence of drug-induced side effect, but it did not yet cause reduction in the new infections. Data also show that selenium have no influence on the length of stay in ICU or the length of mechanical ventilation. Overall, the clinical heterogeneity is low among these RCTs, and most of the studies are of moderate quality and little differences are found in characteristics of the populations, regimen, and study designs. Sensitivity analysis suggests that the results are relatively stable.

Mortality in critically illness is the primary end point. Our meta-analysis shows that there is significant difference between the selenium supplement group and the control in the total mortality in hospital and the TSA result shows that our conclusion is reliable and no more trials are needed to confirm it, although there is no beneficial effect on the mortality at day 28. Total mortality in our meta-analysis refers to mortality regardless of the follow-up period, however, the longest follow-up period of our included studies is 3 months. According to our results, we suppose that selenium supplement may have a beneficial effect on the clinical outcome of long-term follow-up mortality.

To the best of our knowledge, this is not the first meta-analysis to explore the role of selenium supplement on the outcome of critically ill patients. Our partial results are different from the last meta-analysis.[9] Manzanares et al[9] including 21 studies reported that the use of high-dose selenium supplementation had no beneficial effect on overall mortality and the length of stay in hospital in critically ill patients. They did not use TSA to control the risks of type I and type II errors and calculate RIS. However, TSA was used in present article and the result of TSA demonstrated that our conclusion selenium could cause reduction in overall mortality is reliable and no more studies are needed. In the meta-analysis Manzanares et al[9] selenium as a combined therapy is also included, and the test subgroup difference between selenium as a monotherapy and combined therapy was not significant. Manzanares et al also analyzed mechanical ventilation and the incidence of new infections, and get similar results with our study.

Complications are also assessed. Although selenium supplement is generally regarded as safe and well tolerated in most populations, it should be with cautious that high dose of selenium may lead to toxicity, which is most likely resulted from their prooxidant properties.[33]

The meta-analysis has several potential limitations that should be taken into account. Firstly, even though we analyzed selenium supplement in different subgroups, the characteristics of them are different and the effect may be unequal. In the included studies, the characteristics of critically ill patients are not on a unified level, which vary from SIRS to severe multiple injuries. These factors may have a potential influence on our results. Secondly, follow-up varies from 28 days to 12 months, and the outcomes will be uncertain in mutative follow-ups. Thirdly, the route, dose and administration of selenium supplement are varying, so we are not sure to assess the impact of selenium supplement based on clinically meaningful end points. In addition, our study provides additional interesting clues that may be useful for future research on this topic. Remarkably, route of selenium supplement is by continuously intravenous infusion in all studies. Thus, one clue is to focus on route of selenium supplement and to compare the enteral selenium supplement with parenteral selenium supplement to testify the efficacy on critically illness.

In conclusion, the current evidence suggests that the use of selenium could cause reduction in overall mortality and may shorten the length of stay in hospital in critically ill patients, but could not reduce 28-days all causes mortality or shorten length of stay in ICU. Also it has no influence on mechanical ventilation or the incidence of new infections. However, the results should be used carefully because of potential limitations. Further well-designed RCTs on this topic are needed to carry out to provide more evidence to clearly answer the clinical question.
Figure 3. Figure 3A Forest plot for overall mortality. CI = confidence intervals, Fixed = a fixed effects model, M–H = Mantel–Haenszel test, Figure 3B. TSA for overall mortality. TSA = Trial sequential analysis.
Figure 4. Forest plot for 28-day-all cause mortality. CI=confidence intervals, Fixed=a fixed effects model, M-H=Mantel-Haenszel test.

Figure 5. Forest plot for ICU length of stay. CI=confidence intervals, IV=inverse variance, Random=a random effects model.

Figure 6. Forest plot for hospital length of stay. CI=confidence intervals, IV=inverse variance, Random=a random effects model.

Figure 7. Forest plot for mechanical ventilation time. CI=confidence intervals, IV=inverse variance, Random=a random effects model.
Author contributions
Hongjun Kang and Feihu Zhou designed the research. Yan Zhao and Xin Hu conducted the research. Li Wang and Zhi Mao analyzed the data. Hongjun Kang and Yan Zhao wrote the manuscript. Feihu Zhou had primary responsibility for the final content. All authors read and approved the final manuscript.

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