Therapeutic effect of intensive glycemic control therapy in patients with traumatic brain injury: A systematic review and meta-analysis of randomized controlled trials

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Abstract
Background: Hyperglycemia is associated with dismal outcomes in patients with traumatic brain injury (TBI), which is frequently treated with insulin therapy. In this study, a systematic review and meta-analysis of the published randomized controlled trials (RCTs) was performed to assess the safety and efficacy of intensive glycemic control (IGC) versus conventional glycemic control (CGC) for patients following TBI.

Methods: Databases, including PubMed, Embase, and the Cochran database, were retrieved up to January 2018. The outcomes evaluated in this study included mortality, neurological outcome, infection rate, hypoglycemia episode, and length of stay (LOS) in intensive care unit (ICU). The enrolled trials were analyzed using the Review Manager 5.3 software.

Results: A total of 7 randomized controlled trials involving 1013 cases were enrolled in this study, and the results indicated no significant difference in 6-month mortality (risk ratio [RR], 0.92; 95% confidence interval [CI] 0.76–1.10; P = .34). Subsequently, IGC was associated with a better neurological outcome (RR, 1.22; 95% CI 1.05–1.43; P = .01), lower infection rate (RR, 0.65; 95% CI 0.51–0.82; P = .0003) and shorter LOS in ICU (mean difference [MD] = −1.37; 95% CI = −2.11, −0.63; P = .0003). In addition, IGC would also increase the risk of hypoglycemia episode (RR, 4.53; 95% CI 2.18–9.42; P < .001).

Conclusions: IGC plays a protective role in improving neurological outcome, decreasing infection rate and reducing the LOS in ICU. However, IGC therapy can also remarkably increase the risk of hypoglycemia, but it will not affect the mortality in TBI patients.

Abbreviations: CGC = conventional glycemic control, CI = confidence intervals, eGOS = extended Glasgow Outcome Scale, GOS = Glasgow Outcome Scale, ICU = intensive care unit, IGC = intensive glycemic control, MD = mean difference, mRS = modified Rankin Scale, RCTs = randomized controlled trials, RR = risk ratio, TBI = traumatic brain injury.

Keywords: conventional glycemic control, intensive glycemic control, meta-analysis, traumatic brain injury

1. Introduction

Hyperglycemia frequently occurs in critically ill patients, which is also linked with increased morbidity and mortality. These observations can be found in general patients as well as those with traumatic brain injury (TBI). Typically, TBI will lead to profoundly increased glucose utilization (also known as hyperglycolysis), which can persist for up to 1 week, finally altering the ability to use ketone bodies as energetic substrates. It is suggested that hyperglycemia can exacerbate secondary brain injury and independently predict the dismal neurological outcome in severe TBI patients.

Conventional glucose control (CGC), the traditional treatment for hyperglycemia, administers insulin at the glucose level of >200 to 220 mg/dL. In the light of reports on worse outcomes of hyperglycemia, a new therapeutic approach, intensive glucose control (IGC), has been sought to maintain the glucose level within the range of 80 to 110 mg/dL. Specifically, patients treated with IGC have distinctly lower morbidity and mortality compared with those undergoing CGC. Additional studies also reveal the benefits of IGC in lowering mortality and incidence of infections among different groups of critically ill patients. However, strict glycemic control with low target ranges will inevitably carry a risk of inadvertent hypoglycemic episodes. On the contrary, other studies show that IGC has no benefits and fails to achieve glycemic control; what’s worse, patients under strict glycemic control suffer from a potentially higher incidence of hypoglycemia.

Many studies and randomized controlled trials (RCTs) have addressed the question of whether IGC can result in better outcomes for TBI patients than CGC, but no consensus has been reached yet. Therefore, the current meta-analysis of RCTs comparing IGC with CGC in patients was conducted, with...
an aim to evaluate the effect of IGC on mortality, neurological outcome, and other clinical outcomes in severe blunt TBI patients.

2. Methods

2.1. Retrieval strategy

The following electronic databases were retrieved until January 2018, including PubMed, Embase, and the Cochrane database, using retrieval terms of traumatic brain injury, subarachnoid hemorrhage, subdural hematoma, insulin therapy, intensive glucose control, glycemic control, conventional glucose control, and randomized controlled trial. Each step of pooled analysis was conducted by 2 investigators independently, and any disagreement was settled by mutual discussion. The current systematic review and meta-analysis of RCTs was conducted according to Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. The current systematic review was not registered.

2.2. Selection criteria

The inclusion criteria were: comparative study (RCT), study investigating only TBI patients, and study comparing IGC with CGC. The exclusion criteria were: review article, meta-analysis, and guideline, non-RCT, study with no medical treatment control group, and study with no CGC arm, for example, IGC versus non-insulin treatment or IGC versus saline with no insulin treatment.

2.3. Data extraction

Data were extracted by 2 reviewers independently, and any disagreement was resolved by consulting with a third reviewer. The following information was extracted from the RCTs, including name of first author, country of origin, patient characteristics (such as mean age and sex), operational definitions, and outcomes. Moreover, means, standard deviations or medians, and interquartile ranges for each treatment group, together with the numbers assessed in each group, were recorded to evaluate the continuous outcomes. Besides, the primary author was reached by email to seek the clarification for the missing information.

2.4. Study outcomes

Primary outcomes of clinical importance included: 6-month mortality, and the available time frame closest to 6 months was used when it was not specifically presented, and good neurological recovery, as defined in individual studies. To be specific, a Glasgow Outcome Scale (GOS) score of 4 to 5, a modified Rankin Scale (mRS) score of 1 to 3, or an extended Glasgow Outcome Scale (eGOS) score of 5 to 8, were considered to represent good outcomes when a full range of outcomes were presented. Secondary outcomes included number of hypoglycemia episodes, length of stay (LOS) in intensive care unit (ICU), and the incidence of infections. Specifically, the major infections included wound infections, pneumonia, urinary infections, and sepsis.

2.5. Quality assessment

The risk of bias was independently evaluated by 2 reviewers using the domain-based evaluation described in the Cochrane Handbook for Systematic Reviews of Interventions. Typically, the following domains were assessed, including selection bias (random sequence generation and allocation concealment), attrition bias (incomplete outcome data), performance and detection bias (blinding of participants, personnel and outcome assessment), reporting bias (selective reporting), and other biases (other sources of bias).

2.6. Statistical analysis

Dichotomous and continuous variables were analyzed by risk ratio (RR) and mean difference (MD), respectively. The heterogeneity between studies was accessed using Cochran Q-statistic test, and tested using I² (P < .05 stood for significant heterogeneity). In the presence of evidence for heterogeneity between studies, the random effects model was used, since it could provide a more conservative effect than that of the fixed-effects model. Meanwhile, sensitivity analysis was performed in the presence of heterogeneity through eliminating one study at a time to check the resolution of heterogeneity. Publication bias was assessed using the visual funnel plot. Data were analyzed by the Review Manager (RevMan version 5.3; Cochrane Collaboration, Oxford, UK).

2.7. Ethical consideration

This is a meta-analysis article, does not involve ethical review, and ethical approval is not necessary after inquiring the ethical review committee in our hospital.

3. Results

3.1. Study selection

A diagram summarizing the study selection process was shown in Fig. 1. As could be seen, a total of 3004 potential trials were identified by the first retrieval strategy, and 7 RCTs were identified after careful full-text evaluation in the final analysis.

3.2. Trial characteristics

Together, a total of 1013 patients were enrolled in the identified RCTs published from 2007 to 2017, including adults diagnosed with TBI. The follow-up period ranged from 6 to 24 months, and the detail information was summarized in Table 1. Five out of the 7 studies had an intensive target range of 80 to 108 (or 110)mg/dL, and only 2 studies by Bilotta et al had an intensive target range of 80 to 120mg/dL. In comparison, the target range for CGC was more variable, among which, 2 studies had an intensive target range of <220mg/dL, 2 of 180–200mg/dL, 2 of <180mg/dL, and 1 of <150mg/dL. Moreover, most studies had displayed a low risk of bias; however, a few studies had an unclear risk (Fig. 2). In the meantime, the risk of bias among studies was assessed to be low.

3.3. Primary outcomes

3.3.1. Mortality. A meta-analysis was performed to calculate the RR of mortality related to IGC versus CGC following TBI. Typically, 6 studies had reported mortality as the primary outcome. A total of 286 related deaths were reported, including 139 in the IGC arm and 147 in the CGC arm. Besides, the results of pooled analysis demonstrated no difference in the risk between
IGC and CGC (RR, 0.92; 95% CI 0.76 to 1.10; P = .34). No significant heterogeneity was observed among the identified studies after evaluating mortality ($I^2=0$, $P = .98$). (Fig. 3A)

### 3.3.2. Favorable neurological outcomes

Results of pooled analysis on the 7 studies reporting neurological recovery showed that, IGC therapy was superior to CGC therapy among patients (RR, 1.22; 95% CI 1.05–1.43; $P = .01$). At the same time, no significant heterogeneity was found ($I^2 = 0\%$, $P = .84$). (Fig. 3B)

### 3.4. Secondary outcomes

#### 3.4.1. Infection rate

Five trials had reported that, the incidence of overall infectious complications in the IGC group and CGC group was 36.6% (107/292) and 50% (145/290), respectively.\[8,9,11,18\] Meanwhile, pneumonia, urinary infection, wound infection, and sepsis were the 4 most common infections, which were included in our meta-analysis. Our results suggested that
We evaluated the clinical efficacy of IGC among the selected studies when evaluating the difference between these 2 groups was statistically significant (MD = -1.37; 95% CI = -2.11, -0.63; P = .0003) (Fig. 4E).

3.5. Sensitivity analysis and publication bias

Sensitivity analysis was performed by randomly excluding one trial and interchanging the fixed-effects model and random-effects model based on the pooled analysis, and the outcomes were confirmed to be stable. Publication bias was assessed using funnel plots and Egger test. Typically, mortality was used as an exemplary indicator for publication bias assessment. The shape of funnel plot revealed no indication of obvious asymmetry. In addition, Egger test was also employed to provide statistical evidence of funnel plot symmetry, which revealed no proof of publication bias. (Fig. 5)

4. Discussion

The present meta-analysis demonstrates that TBI patients undergoing IGC therapy are associated with improved neurological outcome, decreased infection rate, and reduced LOS in ICU compared with those receiving CGC therapy. However, IGC does not demonstrate a mortality benefit compared with CGC. Moreover, IGC therapy will also dramatically increase the risk of hypoglycemia episode.

TBI is related to a stress response including hyperglycemia, which has been shown to worsen the neurological outcome during cerebral ischemia and hypoxia. Studies on moderate to severe TBI patients indicate that, higher initial and postoperative glucose levels will lead to higher intracerebral lactate levels and worse outcome, especially for those with the glucose levels of greater than 160 to 200 mg/dL. The mechanisms by which hyperglycemia exerts the harmful effect are complex. Concretely, the contributing factors may include free radical formation and oxidative injury, activation of Nmethyl-D-aspartate receptors, raised intracellular calcium, triggering of inflammatory and apoptotic pathways, and alterations in lactate metabolism with reduced tissue pH. Therefore, the impact of interventions, such as more aggressive control over blood glucose with insulin control therapy, has been studied.

Notably, the target range of conventional glucose control in our meta-analysis is more variable, among which, 2 studies have the target range of <220 mg/dL, 2 of 180 to 200 mg/dL, 2 of <180 mg/dL, and 1 of <150 mg/dL. It is proposed in existing guidelines to initiate insulin therapy when the blood glucose level exceeds 180 mg/dL to trigger a blood glucose concentration of <180 mg/dL. Moreover, the previous guidelines propose that glycemic control infusions should aim at maintaining the blood glucose levels of <200 mg/dL in neurocritically ill patients with hyperglycaemia. In contrast, Jacob et al suggested that patients with the blood glucose levels of ≥150 mg/dL should be initiated the insulin therapy, so as to keep the blood glucose levels of <150 mg/dL in most adult trauma patients and to maintain the absolute blood glucose values of <180 mg/dL. Meanwhile, a protocol that could achieve a low rate of hypoglycemia (blood glucose level of ≤70 mg/dL) should be used to achieve the lower infection rates and shorter LOS in ICU in trauma patients. In 2001, van den Berghe et al had compared 783 patients receiving CGC therapy with 765 undergoing IGC therapy. Their results indicated that patients treated with IGC had dramatically lower morbidity and mortality compared with those receiving
CGC therapy. Additional studies also show the benefits of IGC in reducing mortality and incidence of infections among different groups of critically ill patients. Nonetheless, other studies have revealed no benefit of IGC, which fails to achieve glycemic control and potentially results in higher incidence of hypoglycemia in patients. In our meta-analysis, no differences are observed in reducing the mortality and infection rate between IGC and CGC in TBI patients. Such observation is consistent with the results from recent large-scale and multi-center RCTs performed in critically ill patients that displayed a higher degree of heterogeneity in the neurological and diagnostic categories. On the contrary, Zafar et al had demonstrated the inconsistent results in 2011. However, they not only analyzed TBI; instead, all kinds of brain injuries were included for analysis, including tumor and stroke. Moreover, our results show that IGC is beneficial for lowering the infection rates. Nevertheless, future RCTs on infection would be needed due to the small sample size.

It should be noted that, strict glycemic control with a low target range will invariably carry a risk of inadvertent hypoglycemia episode. As found in the current meta-analysis, the IGC therapy will markedly increase the incidence of hypoglycemia (RR, 4.53; 95% CI 2.18–9.42; *P* < .001). RCTs examining the effect of IGC have consistently reported an increased risk of both moderate and severe hypoglycemia among patients randomly assigned to IGC group; besides, the occurrence of hypoglycemia is strongly associated with dismal outcomes. On this account, an important question occurs, which is whether hypoglycemia episode will contribute to the worsened long-term neurological outcome following TBI. Several studies have suggested that hypoglycemia is an independent mortality factor in ICU, but it is not observed in the Computerized Glucose Control in Critically Ill Patients (CGAO-REA) study. Most studies enrolled in our analysis reveal that, hypoglycemia episode is not associated with dismal outcomes. For instance, in the study by Bilotta et al involving 97 TBI patients, IGC therapy was reported to increase the risk of hypoglycemia but would not markedly affect the 6-month mortality or neurological disability. In the study by Coester et al enrolling 88 severe TBI patients, the authors reported that IGC therapy would increase the risk of hypoglycemia but would not distinctly affect the 6-month mortality or neurological disability. In the study from Yang et al recruiting 240 patients, the authors reported no difference in mortality, but an increase in the proportion of patients with favorable neurological recovery at 6 months, which showed no statistical significance.

Contrast, our results suggest that IGC can promote the occurrence of favorable neurological outcomes, which are

**Figure 3.** Forest plot of all included trials examining the effect of IGC versus CGC on mortality and favorable neurological outcomes in TBI patients. (A) Mortality; and (B) favorable neurological outcomes. CGC = conventional glycemic control; IGC = intensive glycemic control; M-H, Random = Mantel-Haenszel, Fixed = Mantel-Haenszel, Fixed-effects model; TBI = traumatic brain injury.
Figure 4. Comparison of the infection rates, hypoglycemia episodes and LOS in ICU among TBI patients. (A) Infection rates; (B) infection rates excluding the study by Coester; (C) hypoglycemia episodes; (D) hypoglycemia episodes excluding the study by Bilotta; and (E) LOS in ICU. CGC = conventional glycemic control; IGC = intensive glycemic control; IV, Random = inverse variance, random-effects model; LOS = length of stay; M-H, Fixed = Mantel-Haenszel, Fixed-effects model; M-H, Random = Mantel-Haenszel, Random-effects model; TBI, traumatic brain injury.
consistent with those from Kramer in 2012.\(^{152}\) Such effect on neurological outcome can be explained by the central nervous system (CNS) protection of IGC. IGC therapy can protect the CNS through reducing the mean and maximal intracranial pressures in patients with isolated brain injury.\(^{15,153,154}\) The beneficial effect of IGC on intracranial pressure can be attained in the presence of similar cerebral perfusion pressures achieved with notably less norepinephrine as a vasopressor. Not only normoglycemia, but also insulin itself, has been reported to improve critically ill patients, which can be attributable to its metabolic and anti-inflammatory effects.\(^{55}\) Experimental data suggest that insulin can increase glucose uptake in astrocyte\(^{56}\) and can play a role in cerebral cortical glucose regulation.\(^{17}\) In addition, our meta-analysis also reveals that the LOS in ICU is shorter in IGC group than in CGC group.

Nonetheless, our analysis is inevitably associated with several potential limitations. Therefore, firstly, any future RCTs on IGC therapy should carefully pilot their protocol to ensure the minimized hypoglycemia. Secondly, the number of included studies is not sufficient enough to make a convincing conclusion for the secondary endpoints. Thirdly, the neurological outcomes reported in this meta-analysis are relatively crude, and it remains possible that glycemic control can have greater influence on more subtle neurocognition or indices of quality of life. Fourthly, the follow-up period of the enrolled trials is short to moderate, making it impossible to evaluate the long-term complications. Lastly, the description of allocation concealment, difficulties in the binding of participants, and outcome assessors is lacking in these included studies, which can be attributed to the nature of intervention.

5. Conclusions
This meta-analysis suggests that IGC generally has comparable efficacy to CGC in reducing mortality following TBI. Moreover, IGC plays a protective role in improving neurological outcome, decreasing infection rate and reducing the LOS in ICU.

Author contributions
Conceptualization: Tao Xu.
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