Association of vasopressor use with mortality in severe traumatic brain injury: a nationwide retrospective cohort study in Japan

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Abstract

Background: Maintaining of cerebral perfusion pressure (CPP) is frequently incorporated in severe traumatic brain injury management algorithms. However, there is limited evidence on prevalent clinical practices regarding a preferred method for achieving such CPP aims. We conducted a nationwide retrospective cohort study to determine the association between the use of vasopressors and mortality following hospital discharge in patients with severe traumatic brain injury, and to determine whether the use of vasopressors affects emergency department mortality or the occurrence of cognitive dysfunction.

Methods: Data were collected between January 2004 and December 2018 from the Japanese Trauma Data Bank, which includes data from 272 emergency hospitals in Japan. Adults aged ≥16 years with severe traumatic brain injury, without other major injuries, were examined. A severe traumatic brain injury was defined based on the Abbreviated Injury Scale code and a Glasgow Coma Scale score of 3–8 on admission. Multivariable analysis and propensity score matching were performed. Statistical significance was assessed using 95% confidence intervals (CIs).

Results: In total, 10,284 patients were eligible for analysis, with 650 patients (6.32%) included in the vasopressor group and 9,634 patients (93.68%) included in the non-vasopressor group. The proportion of deaths on hospital discharge was higher in the vasopressor group than in the non-vasopressor group (81.69% [531/650] vs. 40.21% [3,874/9,634]). This finding was confirmed by multivariable logistic regression analysis (adjusted odds ratio [OR], 5.71; 95% CI: 4.56–7.16). Regarding propensity score-matched patients, the proportion of deaths on hospital discharge remained higher in the vasopressor group than in the non-vasopressor group (81.66% [530/649] vs. 50.69% [329/649]) (OR, 4.33; 95% CI: 3.37–5.57). The vasopressor group had a higher emergency department mortality rate than the non-vasopressor group (8.01% [52/649] vs. 2.77% [18/649]) (OR, 3.05; 95% CI: 1.77–5.28). There was no reduction in complications of cognitive disorders in the vasopressor group (5.39% [35/649] vs. 5.55% [36/649]) (OR, 0.97; 95% CI: 0.60–1.57).

Conclusions: In this population, the use of vasopressors for severe traumatic brain injury was associated with higher mortality on hospital discharge. Our results could not reach the conclusion that vasopressor therapy is beneficial in improving clinical outcomes in patients with severe TBI.

Background

Severe traumatic brain injury (TBI) is a leading cause of death and disability in people of all generations worldwide [1-3] and is associated with high economic and social costs [4]. Severe TBI is characterized by a primary insult that involves the initial mechanical force of impact and results in loss of brain tissue and neuronal cell death. Secondary injury occurs in the minutes and hours following the primary injury. Although little can be done to reverse the primary brain damage, secondary brain injury due to dysregulation of cerebral blood flow is potentially preventable. Therefore, acute management of patients
after severe TBI aims to minimize secondary brain injury by maintaining cerebral perfusion pressure, defined as the difference between mean blood pressure and intracranial pressure [5-8].

Blood pressure control is the mainstay of therapy for several acute cerebral disorders, both in the early emergent phase immediately after brain injury and in the subsequent intensive care unit phase [9]. Hypotension in the acute phase of severe TBI is a key factor associated with poor clinical outcome [7-9]. Although vasopressors are considered to be harmful to trauma patients, in cases of a hemorrhagic shock state [10], The most recent international guidelines on the management of severe TBI suggests that maintaining systolic BP at $\geq 100$ mm Hg for patients 50 to 69 years old or at $\geq 110$ mmHg or above for patients 15 to 49 or $>70$ years old may decrease mortality and improve outcomes [5]. However, this is based on level $e$ evidence, and it cannot specify a preferred method for achieving such systolic BP goals.

Therefore, the aim of this study was to assess the possible association of vasopressor use with mortality in patients with severe TBI, using data from the Japanese Trauma Data Bank (JTDB) registry, which represents the biggest trauma data bank in Japan. Survivors of severe TBI often have cognitive control function deficits [11, 12]. Therefore, we also assess association of vasopressor use with the occurrence of cognitive dysfunction.

Methods

Study design, population, and setting

This was a nationwide retrospective cohort study conducted using the JTDB database. We included cases registered in the database from January 2004 to December 2018 and patients aged $\geq 16$ years who had a TBI and were transported to a JTDB-participating hospital. We segregated patients with severe TBI based on the Abbreviated Injury Scale (AIS) code (TBI = 140000) and a Glasgow Coma Scale score of 3–8 on admission. We excluded cases with a maximum head AIS score of 6 (unsurvivable injury) or 9 (unspecified injury), with other critical injuries (AIS scores of $\geq 3$), in cardiopulmonary arrest on or before hospital arrival, that underwent cardiopulmonary resuscitation (i.e., use of adrenaline during cardiopulmonary resuscitation), or that required interhospital transport. We also excluded cases with missing outcome data or variables required for propensity score (PS) matching [13, 14]. This study defined patients in cardiopulmonary arrest as those whose systolic blood pressure was 0 mmHg and/or heart rate was 0 bpm on or before hospital arrival [15].

Japanese Trauma Data Bank

The JTDB was launched in 2003 by the Japanese Association for the Surgery of Trauma (Trauma Surgery Committee) and the Japanese Association for Acute Medicine (Committee for Clinical Care Evaluation) [15, 16], similar to the trauma databases in North America, Europe, and Oceania. By 2018, 272 major emergency medical institutions across Japan had been registered in the JTDB database [16]. The included hospitals have service levels similar to Level I trauma centers in the United States. Data were collected via the Internet from participating institutions. The physicians and medical assistants who
attended the AIS coding course were the main inputters of the data [15, 16]. The JTDB records trauma patient data including age; sex; mechanism of injury; AIS code (1998 version); Injury Severity Score (ISS); vital signs on hospital arrival; date and time series from hospital arrival to discharge; medical treatments such as interventional radiology, surgery, and computed tomography; complications; and mortality on discharge [15, 16]. The ISS was calculated from the top three AIS scores in nine sites classified using the AIS codes. The data used in this study were the most recent data available in this registry.

Study endpoints

The primary outcome of this investigation was death on hospital discharge. The secondary outcomes were emergency department (ED) mortality and in-hospital cognitive dysfunction, which diagnosed by medical team in hospital.

Propensity score matching

In this study, we selected matched PS analysis as the use of vasopressors was not randomly assigned. A logistic regression analysis was performed to estimate a PS for the prediction of vasopressor use from the available predictors. Confounders were carefully selected from previous reports [9, 10, 17-20], and clinically important confounders were included to estimate PSs. The probability of receiving vasopressors (PS) for each patient was calculated using multivariable logistic regression analysis based on the following 14 variables: age (continuous variable), sex (male/female), year of onset (2004–2006, 2007–2009, 2010–2012, 2013–2015, 2016–2018), Glasgow Coma Scale score (continuous variable), systolic blood pressure (continuous variable) on admission to the ED, surgery indicated for TBI (no/yes), type of injury (blunt/non-blunt), cause of trauma (motor vehicle accident, fall, sports, other), type of TBI (isolated TBI, TBI with skull fracture, TBI with intracranial vessel injury, TBI with other head injuries), prehospital intravenous fluid administration (no/yes), use of transfusion in the first 24 hours (no/yes), past medical history of stroke (no/yes), use of anticoagulant or antiplatelet drugs (no/yes), and ISS (continuous variable). We performed a receiver operating characteristic curve analysis with the area under the curve predicting the use of vasopressors in patients with severe TBI. One-to-one pair matching between the non-vasopressor and vasopressor groups was performed by nearest-neighbor matching without replacement, using a caliper width of 0.02 as the standard deviation of the PS. Covariate balances before and after matching were checked by comparing standardized mean differences (SMDs). An SMD <10% was considered a negligible imbalance between the two groups. In the PS-matched cohort, univariable logistic regression analysis was performed to assess the association between the use of vasopressors and outcomes.

Statistical analysis

We divided patients into two groups (vasopressor and non-vasopressor). Descriptive data are presented as counts and percentages (categorical variables) or medians and interquartile ranges (continuous variables). Outcomes were evaluated using univariable and multivariable logistic regression analyses to assess the independent effect of vasopressor use. Based on these analyses, we calculated the odds
ratios (ORs) and 95% confidence intervals (CIs). Based on previous reports [9, 10, 17-20], we selected confounders for multivariable logistic regression analysis on the assumption that none were directly affected by the use of vasopressors. In the multivariable logistic regression model, we adjusted for the aforementioned 14 variables used in the PS calculation. In addition, subgroup analyses were performed to identify the potential benefits and drawbacks of the use of vasopressors. In each subgroup, a multivariable logistic regression analysis adjusted for the aforementioned variables was performed to assess the independent effect of vasopressor use on mortality on hospital discharge. P for interaction was calculated using the multivariable logistic regression model.

All statistical analyses were conducted using STATA (version 16) (StataCorp LP, 4905 Lakeway Drive, College Station, Texas, USA). Statistical significance was defined as a two-sided p value <0.05 or assessed using 95% CIs. This manuscript was written based on the STROBE statement for the reporting of cohort and cross-sectional studies [21].

Results

A total of 10,284 patients were included in the study. Of those, 650 patients (6.32%) received vasopressors and 9,634 patients (93.68%) did not. Fig. 1 depicts the flowchart of patients included in this study. The median (interquartile range) age of the included patients was 67 (51–79) years, and most were men (69.6% [7,159/10,284]). Blunt trauma was the most common type of injury (95.5% [9,819/10,284]). The patient characteristics are summarized in Table 1. There were no significant differences in age, sex, type of trauma, type of TBI, and past medical history (stroke or use of anticoagulant/antiplatelet therapy) between the two groups. While traffic accidents (46.9% [305/650]) were the leading cause of injury followed by falls (42.0% [273/650]) in the vasopressor group, falls (50.4% [4,851/9,634]) caused the most injuries in the non-vasopressor group. Glasgow Coma Scale scores and systolic blood pressure on admission to the ED were significantly lower in the vasopressor group than in the non-vasopressor group (median [interquartile range]) (4 [3–6] vs. 6 [3–7] and 144 [107–172] vs. 151 [128–179] mmHg, respectively). The median (interquartile range) ISS was 25 (20–26) in the vasopressor group, with the ISS being significantly higher than that in the non-vasopressor group (25 [16–25]) (SMD, 0.336). The vasopressor group had more treatments than the non-vasopressor group, including prehospital intravenous infusion (8.5% [55/650] vs. 4.5% [429/9,634]) and blood transfusion in the first 24 hours (44.9% [292/650] vs. 24.3% [2,345/9,634]).

GCS, Glasgow Coma Scale; ISS, Injury Severity Score; IV, intravenous injection; PMH, past medical history; PS, propensity score; SBP, systolic blood pressure; SMD, standardized mean difference; TBI, traumatic brain injury.

Table 1 shows the baseline characteristics of the PS-matched patients. Following PS matching, 649 patients in each group were included. The area under the receiver operating characteristic curve for PS was 0.73. The characteristics of PS-matched patients were finely balanced in terms of absolute SMD.
Although two groups was slightly imbalanced about type of TBI, the characteristics between the groups were almost similar (sTable1).

The results of the multivariable logistic regression analysis and PS matching for the primary outcome are presented in Table 2. Mortality on hospital discharge was higher in the multivariable logistic regression vasopressor group than in the non-vasopressor group (adjusted OR, 5.71; 95% CI: 4.56–7.16). For PS-matched patients, mortality on hospital discharge was 81.66% (530/649) in the vasopressor group and 50.69% (329/649) in the non-vasopressor group. PS matching analysis illustrated that death on hospital discharge was higher in the vasopressor group than in the non-vasopressor group (OR, 4.33; 95% CI: 3.37–5.57).

Table 3 shows the results of the multivariable logistic regression analysis and PS matching for the secondary outcomes. For PS-matched patients, ED mortality in the vasopressor group was higher than that in the non-vasopressor group (8.01% [52/649] vs. 2.77% [18/649]) (OR, 3.05; 95% CI: 1.77–5.28). There was no significant difference in complications of cognitive disorders between the two groups (5.55% [36/649] vs. 5.39% [35/649]) (OR, 0.97; 95% CI: 0.60–1.57) (Table 3).

Subgroup analysis suggested that the use of vasopressors for severe TBI was associated with higher mortality on hospital discharge for each factor (Fig. 2). As for past medical history, stroke patients had significantly better outcomes than non-stroke patients (adjusted OR [95% CI]) (3.33 [95% CI: 0.84–13.21] vs. 5.87 [95% CI: 4.67–7.39]; p = 0.039).

Discussion

This was a retrospective cohort study conducted to evaluate the effect of vasopressor use on mortality among patients with severe TBI, using a nationwide trauma database in Japan. With robust analyses to adjust for severity, it was found that the use of vasopressors was significantly associated with higher mortality, not only on hospital discharge but also in the ED. Subgroup analysis also reiterated the same and suggested that the use of vasopressors for severe TBI was associated with higher mortality on hospital discharge in almost all cases.

Hypotension (defined as systolic blood pressure <90 mmHg) is a well-known risk factor associated with the occurrence of secondary cerebral damage and poor outcome, especially after TBI [7-9]. In contrast, the large IMPACT prospective database found that a systolic blood pressure <120 mmHg is also a strong predictor of unfavorable neurological recovery [22]. Based on these findings, pharmacological elevation of blood pressure is frequently incorporated in severe TBI management algorithms aimed at preventing or treating cerebral ischemia caused by reduced cerebral perfusion pressure [5]. Further, vasopressors may be used as volume-sparing resuscitating agents to prevent brain edema. However, our study showed that neither mortality nor the occurrence of cognitive dysfunction was improved by the use of vasopressors. Therefore, these results suggest the potential harms of this therapy. A similar recommendation that vasopressors should be avoided in any cases of severe TBI was provided by Lund [23].
Only a few studies have tried to assess whether the use of vasopressors is beneficial for severe TBI, although a number of studies have focused on which type of vasopressor should be used [24, 25]. Our findings could pave the way for future guidelines regarding the use of vasopressors for the treatment of severe TBI.

The penumbra, the brain tissue surrounding the impacted core of the TBI, becomes particularly vulnerable to cell death by hypoxic insults, and preservation of this area is one of the important objectives for using vasopressors. The overall effects of vasopressor use on the brain are probably a consequence of systemic arterial contraction but are also influenced by other factors such as brain injury, integrity of the blood–brain barrier, and the status of cerebral autoregulation [26]. In the abnormal state of autoregulation after severe TBI, excessive elevation of intracranial pressure favors edema formation by increased capillary hydrostatic pressure across the blood–brain barrier, causing brain herniation [27]. This may also result in unwanted hemodynamic effects, such as intracranial hemorrhage, leading to increased mortality [28, 29]. Additionally, vasopressors cause an array of adverse effects among other organs of the body [30, 31]. Catecholamine surge after TBI can lead to peripheral insults induced by the release of proinflammatory substances and result in increased vascular permeability [30]. In this situation, the accentuated proinflammatory response could trigger the development of acute respiratory distress syndrome [31]. Other potential negative systemic side effects of vasopressors include arrhythmia, diuresis, and increased left ventricular afterload.

In patients with severe TBI, low arterial blood pressure, major surgery with bleeding, and blood transfusion can contribute to secondary insults to the brain and aggravate the primary insults and cerebral edema, increasing the risk of developing severe lung injury, or even multiple organ failure. Traditionally, it has been suggested that the clinical effects of brain injuries vary depending on individual characteristics such as sex, age, past medical history, and type of TBI. With multiple confounding factors, planning treatment for TBI is especially challenging. In this study, we used the largest trauma dataset in Japan that allowed us to adjust for confounding factors. To the best of our knowledge, our study included the largest cohort of its kind, to date, that has focused on patients with severe TBI. This study also reveals the utility of vasopressors in some specific conditions (e.g., in patients with TBI with a history of stroke). In cases of stroke, structural cerebrovascular changes may increase resistance; thus, normal cerebral blood flow can be maintained using vasopressors. However, the use of vasopressors in such a subgroup did not reduce mortality at the time of hospital discharge. These results suggest that vasopressors should be avoided in most cases of severe TBI.

The authors acknowledge a few limitations of this observational study. First, PS matching analysis has the risk of residual selection bias. Some differences in the two groups may still exist, even after PS matching, particularly if data on important confounding factors are not included in the analysis. Second, although the use of vasopressors (mainly catecholamine) for resuscitation was recorded in the JTDB, important information such as the type and dose of vasopressors used and the timing of vasopressor administration was not included, leading to selection bias. Detailed information about the dose, type of vasopressor, mode of administration (bolus and/or continuous), and patients’ vital signs when the
vasopressor was initiated was not included in the JTDB. However, all TBI patients transported to JTDB-participating hospitals were treated based on the guidelines for the management of severe TBI, which state that vasopressors are recommended to maintain a systolic blood pressure >110 mmHg [32]. Third, the incidence of cognitive disorders could have been underestimated in our study as the JTDB includes clinical data until hospital discharge, which would largely cater to the acute phase of the injury. Cognitive symptoms after TBI would be accurately evaluated only after recovering from an altered state of consciousness in the acute phase. Additionally, the data included in this study primarily involved cases of blunt trauma, which cannot be extended to penetrating TBI. Lastly, this was an observational study, and there may be other unknown confounding factors. The results of this investigation could not establish causality and remain limited to associations. a randomized controlled trial regarding intracranial pressure ICP management or vasopressors use for maintaining CPP, and regarding blood transfusion or vasopressors use with hypotensive TBI patients.

Blood pressure augmentation to avoid hypotension has been considered to attenuate secondary cerebral damage. Therefore, the management of blood pressure using vasopressors during the emergent phase and intensive care unit phase is based on low-quality evidence and strong recommendations according to recent guidelines for the management of severe TBI [5, 9, 32, 33]. However, our results suggest that vasopressors should be avoided in most cases of severe TBI. Our study has the potential to result in policy and guideline changes for the treatment and management of severe TBI.

Conclusions

In a Japanese population with severe adult TBI associated with blunt trauma, the use of vasopressors was associated with increased mortality, both on hospital discharge and in the ED. Our results could not reach the conclusion that vasopressor therapy is beneficial in improving clinical outcomes in patients with sTBI.

Abbreviations

AIS, Abbreviated Injury Scale; CI, confidence interval; ED, emergency department; ISS, Injury Severity Score; JTDB, Japanese Trauma Data Bank; OR, odds ratio; PS, propensity score; SMD, standardized mean difference; TBI, traumatic brain injury.

Declarations

Ethics approval and consent to participate: This study was approved by the Ethics Committee of Osaka University Graduate School of Medicine (Osaka, Japan) (approval number: 16260). Personal identifiers were removed from the JTDB database beforehand; thus, the requirement for informed consent was waived.

Consent for publication: Not applicable.
Availability of data and materials: The data that support the findings of this study are available from the JTDB, but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of the JTDB.

Competing interests: The authors declare that they have no competing interests.

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Authors’ contributions: SH, T. Sobue, and TK designed the study and wrote the manuscript. SH, TK, and AH performed the statistical analysis. HO and T. Shimazu critically revised the manuscript for important intellectual content. All authors read and approved the final manuscript.

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References

1. Maas AIR, Menon DK, Adelson PD, Andelic N, Bell MJ, Belli A, et al. Traumatic brain injury: integrated approaches to improve prevention, clinical care, and research. Lancet Neurol. 2017;16:987-1048.

2. SAFE Study Investigators; Australian and New Zealand Intensive Care Society Clinical Trials Group; Australian Red Cross Blood Service; George Institute for International Health; Myburgh J, Cooper DJ, et al. Saline or albumin for fluid resuscitation in patients with traumatic brain injury. N Engl J Med. 2007;357:874-84.

3. Nichol A, French C, Little L, Haddad S, Presneill J, Arabi Y, et al. Erythropoietin in traumatic brain injury (EPO-TBI): a double-blind randomised controlled trial. Lancet. 2015;386:2499-506.

4. Farhad K, Khan HM, Ji AB, Yacoub HA, Qureshi AI, Souayah N. Trends in outcomes and hospitalization costs for traumatic brain injury in adult patients in the United States. J Neurotrauma. 2013;30:84-90.

5. Carney N, Totten AM, O’Reilly C, Ullman JS, Hawryluk GW, Bell MJ, et al. Guidelines for the Management of Severe Traumatic Brain Injury, Fourth Edition. Neurosurgery. 2017;80:6-15.

6. McGinn MJ, Povlishock JT. Pathophysiology of traumatic brain injury. Neurosurg Clin N Am. 2016;27:397-407.

7. Stocchetti N, Le Roux P, Vespa P, Oddo M, Citerio G, Andrews PJ, et al. Clinical review: neuromonitoring - an update. Crit Care. 2013;17:201.
8. Corps KN, Roth TL, McGavern DB. Inflammation and neuroprotection in traumatic brain injury. JAMA Neurol. 2015;72:355-62.

9. Carteron L, Taccone FS, Oddo M. How to manage blood pressure after brain injury? Minerva Anestesiol. 2017;83:412-21.

10. Aoki M, Abe T, Saitoh D, Hagiwara S, Oshima K. Use of vasopressor increases the risk of mortality in traumatic hemorrhagic shock: a nationwide cohort study in Japan. Crit Care Med. 2018;46:e1145-51.

11. Miotto EC, Cinalli FZ, Serrao VT, Benute GG, Lucia MC, Scaff M. Cognitive deficits in patients with mild to moderate traumatic brain injury. Arq Neuro Psiquiatr. 2010;68:862-8.

12. Olsen A, Brunner JF, Indredavik Evensen KA, Finnanger TG, Vik A, Skandsen T, et al. Altered cognitive control activations after moderate-to-severe traumatic brain injury and their relationship to injury severity and everyday-life function. Cereb Cortex. 2015;25:2170-80.

13. Butcher N, Balogh ZJ. AIS>2 in at least two body regions: a potential new anatomical definition of polytrauma. Injury. 2012;43:196-9.

14. Härtl R, Gerber LM, Iacono L, Ni Q, Lyons K, Ghajar J. Direct transport within an organized state trauma system reduces mortality in patients with severe traumatic brain injury. J Trauma. 2006;60:1250-6; discussion 1256.

15. Katayama Y, Kitamura T, Hirose T, Kiguchi T, Matsuyama T, Takahashi H, et al. Pelvic angiography is effective for emergency pediatric patients with pelvic fractures: a propensity-score-matching study with a nationwide trauma registry in Japan. Eur J Trauma Emerg Surg. 2019; doi:10.1007/s00068-019-01154-w.

16. Japan Trauma Care and Research. Japan Trauma Data Bank Annual Report 2014–2018. 2019. [Online]. Available: http://www.jast-hp.org/trauma/pdf/jtdb2019e.pdf. Accessed 10 Oct 2020.

17. Hussmann B, Schoeneberg C, Jungbluth P, Heuer M, Lefering R, Maek T, et al. Enhanced prehospital volume therapy does not lead to improved outcomes in severely injured patients with severe traumatic brain injury. BMC Emerg Med. 2019;19:

18. Beynon C, Hertle DN, Unterberg AW, Sakowitz OW. Clinical review: traumatic brain injury in patients receiving antiplatelet medication. Crit Care. 2012;16:

19. Marshall LF, Marshall SB, Klauber MR, van Berkum Clark M, Eisenberg HM, Jane JA, et al. A new classification of head injury based on computerized tomography. J Neurol Surg. 1991;75:S14-20.

20. Iaccarino C, Schiavi P, Picetti E, Goldoni M, Cerasti D, Caspani M, et al. Patients with brain contusions: predictors of outcome and relationship between radiological and clinical evolution. J Neurol Surg. 2014;120:908-18.

21. Gallo V, Egger M, McCormack V, Farmer PB, Ioannidis JP, Kirsch-Volders M, et al. Strengthening the Reporting of OBservational studies in Epidemiology–Molecular Epidemiology STROBE-ME: an extension of the STROBE statement. J Clin Epidemiol. 2011;64:1350-63.
22. Maas AI, Murray GD, Roozenbeek B, Lingsma HF, Butcher I, McHugh GS, et al. Advancing care for traumatic brain injury: findings from the IMPACT studies and perspectives on future research. Lancet Neurol. 2013;12:1200-10.

23. Grände PO. Critical evaluation of the Lund concept for treatment of severe traumatic head injury, 25 years after its introduction. Front Neurol. 2017;8:315.

24. Johnston AJ, Steiner LA, Chatfield DA, Coles JP, Hutchinson PJ, Al-Rawi PG, et al. Effect of cerebral perfusion pressure augmentation with dopamine and norepinephrine on global and focal brain oxygenation after traumatic brain injury. Intensive Care Med. 2004;30:791-7.

25. Sookplung P, Siriussawakul A, Malakouti A, Sharma D, Wang J, Souter MJ, et al. Vasopressor use and effect on blood pressure after severe adult traumatic brain injury. Neurocrit Care. 2011;15:46-54.

26. Rosner MJ, Rosner SD, Johnson AH. Cerebral perfusion pressure: management protocol and clinical results. J Neurol Surg. 1995;83:949-62.

27. Nordström CH. Physiological and biochemical principles underlying volume-targeted therapy - the “Lund concept”. Neurocrit Care. 2005;2:83-95.

28. Malhotra AK, Schweitzer JB, Fox JL, Fabian TC, Proctor KG. Cerebral perfusion pressure directed therapy following traumatic brain injury and hypotension in swine. J Neurotrauma. 2003;20:827-39.

29. Chico-Fernández M, Barea-Mendoza JA, Pérez-Bárcena J, García-Sáez I, Quintana-Díaz M, Marina L, et al. Concomitant traumatic brain injury and hemorrhagic shock: outcomes using the Spanish Trauma ICU Registry (RETRAUCI). Am Surg. 2020:0003134820949990.

30. Caplan HW, Cox CS. Resuscitation strategies for traumatic brain injury. Curr Surg Rep. 2019;7.

31. Robertson CS, Valadka AB, Hannay HJ, Contant CF, Gopinath SP, Cormio M, et al. Prevention of secondary ischemic insults after severe head injury. Crit Care Med. 1999;27:2086-95.

32. Guidelines Committee on the Management of Severe Head Injury, Japan Neurosurgical Society, Japan Society of Neurotraumatology, Japanese Association for the Surgery of Trauma. Guidelines for the Management of Severe Head Injury. 4th ed. Tokyo: Igaku-shoin; 2019.

33. Geeraerts T, Velly L, Abdennour L, Asehnoun K, Audibert G, Bouzat P, et al. Management of severe traumatic brain injury (first 24hours). Anaesth Crit Care Pain Med. 2018;37:171-86.

Tables
Table 1. Characteristics of patients with and without vasopressor use (all patients and PS-matched patients)

| Characteristic          | All patients |               |               | PS-matched patients |               |               |
|-------------------------|--------------|---------------|---------------|--------------------|---------------|---------------|
|                         | Non-vasopressor | Vasopressor   | Controls      | Non-vasopressor    | Vasopressor   | Controls      |
| Age (years)             | N = 9634     | N = 650       | SMD           | N = 649            | N = 649       | SMD           |
| Sex (male)              | 67 (51.7%)   | 60 (57.8%)    | 0.092         | 60 (54.7%)         | 60 (57.8%)    | 0.011         |
| Year of onset           |              |               |               |                    |               |               |
| 2004–2006               | 476 (4.9%)   | 45 (6.9%)     | 0.163         | 51 (7.9%)          | 44 (6.8%)     | 0.019         |
| 2007–2009               | 1248 (13.9%) | 118 (18.2%)   | 109 (16.8%)   | 118 (18.2%)        |               |               |
| 2010–2012               | 2265 (23.5%) | 154 (23.7%)   | 157 (24.2%)   | 154 (23.7%)        |               |               |
| 2013–2015               | 3059 (31.8%) | 177 (27.2%)   | 185 (28.5%)   | 177 (27.3%)        |               |               |
| 2016–2018               | 2586 (26.8%) | 156 (24.0%)   | 147 (22.7%)   | 156 (24.0%)        |               |               |
| Type of trauma (blunt)  | 9198 (95.5%) | 621 (95.5%)   | 0.003         | 617 (95.1%)        | 620 (95.5%)   | 0.022         |
| Cause of trauma         |              |               |               |                    |               |               |
| Motor vehicle accident  | 3417 (35.5%) | 305 (46.9%)   | 0.185         | 289 (44.5%)        | 304 (46.8%)   | 0.012         |
| Fall                    | 4851 (50.4%) | 273 (42.0%)   | 298 (45.9%)   | 273 (42.1%)        |               |               |
| Sports                  | 69 (0.7%)    | 2 (0.3%)      | 4 (0.6%)      | 2 (0.3%)           |               |               |
| Other                   | 1297 (13.5%) | 70 (10.8%)    | 58 (8.9%)     | 70 (10.8%)         |               |               |
| Type of TBI             |              |               |               |                    |               |               |
| Isolated TBI            | 5325 (55.3%) | 283 (43.5%)   | 0.091         | 327 (50.4%)        | 283 (43.0%)   | 0.039         |
| w/Skull fracture        | 2802 (29.1%) | 269 (41.4%)   | 193 (29.7%)   | 268 (41.3%)        |               |               |
| w/Cerebrovascular injury| 33 (0.3%)    | 11 (1.7%)     | 2 (0.3%)      | 11 (1.7%)          |               |               |
| w/Other head injury     | 1474 (15.3%) | 87 (13.4%)    | 127 (19.6%)   | 87 (13.4%)         |               |               |
| SBP on arrival          | 151 (126–179) | 144 (107–172) | 0.291         | 144 (120–170)      | 144 (107–172) | 0.051         |
| GCS score on arrival    | 6 (3–7)      | 4 (3–6)       | 0.515         | 4 (3–6)            | 4 (3–6)       | 0.005         |
| Prehospital IV          | 429 (4.5%)   | 55 (8.5%)     | 0.164         | 52 (8.0%)          | 55 (8.5%)     | 0.017         |
| Blood transfusion       | 2345 (24.3%) | 292 (44.9%)   | 0.443         | 297 (45.8%)        | 291 (44.8%)   | 0.019         |
| Surgery for TBI         | 3831 (39.8%) | 272 (41.8%)   | 0.042         | 302 (46.5%)        | 271 (41.8%)   | 0.096         |
| PMR of stroke           | 275 (2.9%)   | 17 (2.6%)     | 0.015         | 15 (2.3%)          | 17 (2.6%)     | 0.02          |
| Anticoagulant/antiplatelet therapy | 633 (6.6%) | 32 (4.9%) | 0.071 | 28 (4.3%) | 32 (4.9%) | 0.029 |
| ISS                     | 25 (16–25)   | 25 (20–26)    | 0.336         | 25 (21–26)         | 25 (20–26)    | 0.075         |

Table 2. Primary outcome comparisons between patients with and without vasopressor use before and after PS matching

|                      | Total | Non-vasopressor | Vasopressor | Crude OR (95% CI) | Adjusted OR (95% CI) |
|----------------------|-------|-----------------|-------------|------------------|---------------------|
| All patients         | 10,284 | 9,634          | 650         |                   |                     |
| Death at hospital discharge | 4,405 (42.83%) | 3,874 (40.21%) | 531 (81.69%) | 6.63 (5.42–8.13) | 5.71 (4.56–7.16)   |
| PS-matched patients  | 1,298  | 649            | 649         |                   |                     |
| Death at hospital discharge | 859 (66.18%) | 329 (50.69%) | 530 (81.66%) | 4.33 (3.37–5.57) |                     |

CI, confidence interval; OR, odds ratio; PS, propensity score.

Table 3. Secondary outcome comparisons between patients with and without vasopressor use before and after PS matching
|                          | Total    | Non-vasopressor | Vasopressor | Crude OR (95% CI) | Adjusted OR (95% CI) |
|--------------------------|----------|-----------------|-------------|-------------------|----------------------|
| All patients             | 10284    | 9634            | 650         |                   |                      |
| Death at ED              | 816 (7.93%) | 781 (8.11%)    | 35 (5.38%)  | 0.65 (0.46–0.91)  | 0.77 (0.54–1.11)     |
| Cognitive disorder       | 1298     | 649             | 649         |                   |                      |
| PS-matched patients      | 70 (5.39%) | 18 (2.77%)     | 52 (8.01%)  | 3.05 (1.77–5.28)  |                      |
| Death at ED              | 71 (5.47%) | 36 (5.55%)     | 35 (5.39%)  | 0.97 (0.60–1.57)  |                      |

CI, confidence interval; ED, emergency department; OR, odds ratio; PS, propensity score.