Out-of-Hospital Hypertonic Resuscitation Following Severe Traumatic Brain Injury
A Randomized Controlled Trial

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Context Hypertonic fluids restore cerebral perfusion with reduced cerebral edema and modulate inflammatory response to reduce subsequent neuronal injury and thus have potential benefit in resuscitation of patients with traumatic brain injury (TBI).

Objective To determine whether out-of-hospital administration of hypertonic fluids improves neurologic outcome following severe TBI.

Design, Setting, and Participants Multicenter, double-blind, randomized, placebo-controlled clinical trial involving 114 North American emergency medical services agencies within the Resuscitation Outcomes Consortium, conducted between May 2006 and May 2009 among patients 15 years or older with blunt trauma and a prehospital Glasgow Coma Scale score of 8 or less who did not meet criteria for hypovolemic shock. Planned enrollment was 2122 patients.

Intervention A single 250-mL bolus of 7.5% saline/6% dextran, 7.5% saline (hypertonic saline), or 0.9% saline (normal saline) initiated in the out-of-hospital setting.

Main Outcome Measure Six-month neurologic outcome based on the Extended Glasgow Outcome Scale (GOSE) (dichotomized as >4 or ≤4).

Results The study was terminated by the data and safety monitoring board after randomization of 1331 patients, having met prespecified futility criteria. Among the 1282 patients enrolled, 6-month outcomes data were available for 1087 (85%). Baseline characteristics of the groups were equivalent. There was no difference in 6-month neurologic outcome among groups with regard to proportions of patients with severe TBI (GOSE ≤4) (hypertonic saline/dextran vs normal saline: 53.7% vs 51.5%; difference, 2.2% [95% CI, −4.5% to 9.0%]; hypertonic saline vs normal saline: 54.3% vs 51.5%; difference, 2.9% [95% CI, −4.0% to 9.7%]; P = .67). There were no statistically significant differences in distribution of GOSE category or Disability Rating Score by treatment group. Survival at 28 days was 74.3% with hypertonic saline/dextran, 75.7% with hypertonic saline, and 75.1% with normal saline (P = .88).

Conclusion Among patients with severe TBI not in hypovolemic shock, initial resuscitation with either hypertonic saline or hypertonic saline/dextran, compared with normal saline, did not result in superior 6-month neurologic outcome or survival.

Trial Registration clinicaltrials.gov Identifier: NCT00316004

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Current therapy following severe TBI is focused on minimizing secondary injury by supporting systemic perfusion and reducing intracranial pressure.
intraventricular perfusion pressure in animal models and patients with severe TBI. Hypertonic saline has also been shown to have beneficial vasoregulatory, immunomodulatory, and neurochemical effects on the injured brain.

Previous trials have suggested that early administration of hypertonic fluids to patients with severe TBI may improve survival, but no large definitive trials have been reported and the effects on neurologic outcome are not known. Furthermore, all studies have focused on patients with severe TBI and hypovolemic shock; thus, the effect of early administration of hypertonic fluids for patients with severe TBI in the absence of hypovolemic shock is also not known.

We hypothesized that administration of hypertonic fluids as early as possible after severe TBI in patients without hemorrhagic shock would result in improved 6-month neurologic outcome.

METHODS

This study was conducted by the Resuscitation Outcomes Consortium, a multicenter clinical trial network including 11 regional clinical centers in the United States and Canada. The trial involved 114 emergency medical services agencies within the catchment area served by the consortium. Two trials with 2 distinct patient cohorts, one for hypovolemic shock and the other for TBI, were conducted simultaneously using the same intervention.

This report describes the outcome of the TBI cohort. This was a double-blind, 3-group, randomized controlled clinical trial comparing a 250-mL bolus of 7.5% saline (hypertonic saline) vs 7.5% saline/6% dextran 70 (hypertonic saline/dextran) vs 0.9% saline (normal saline) as the initial resuscitation fluid administered to injured patients with suspected severe TBI in the out-of-hospital setting. This dose of hypertonic saline and hypertonic saline/dextran was selected because it was the dose used in all previous prehospital trials and thus had a proven safety record. Previous studies suggested that the expected serum sodium level on admission would be 145 to 155 mEq/L. Details of the initial study design have been published.

Patient Population

Patients were included in the TBI cohort based on the following: blunt mechanism of injury, age 15 years or older, Glasgow Coma Scale (GCS) score of 8 or less, and ineligibility for enrollment in the hemorrhagic shock cohort. The hemorrhagic shock cohort included all patients with systolic blood pressure of 70 mm Hg or less or of 71 to 90 mm Hg with a concomitant heart rate of 108 per minute or greater. All patients meeting the shock criteria were included in the shock cohort, because a low score may be attributable to poor cerebral perfusion secondary to shock and may not be indicative of severe TBI, and GCS score thus becomes much less reliable as a predictor of TBI in this setting. Furthermore, a previous trial of hypertonic resuscitation for patients with out-of-hospital GCS score of 8 or less and systolic blood pressure less than 100 mm Hg had closed for futility, so the current study focused on the effect of these resuscitation strategies for patients with TBI but without evidence of hypovolemic shock.

Exclusion criteria included known or suspected pregnancy, age younger than 15 years, out-of-hospital cardiopulmonary resuscitation, administration of more than 2000 mL of crystalloid or any amount of colloid or blood products prior to enrollment, severe hypothermia (<28°C), drowning, asphyxia due to hanging, burns on more than 20% of total body surface area, isolated penetrating head injury, inability to obtain intravenous access, more than 4 hours between receipt of dispatch call to study intervention, prisoner status, and interfacility transfer.

Intervention

Out-of-hospital personnel were trained to administer unlabeled study fluid as the initial resuscitation fluid once intravenous access was established. In the event that a participating out-of-hospital crew arrived after crystalloid had already been initiated (eg, by ground service before aeromedical crew arrival), they were allowed to administer study fluid as long as the patient still met inclusion criteria and had received less than 2 liters of crystalloid and had not received any mannitol, colloids, or blood products. Once study fluid had been administered, additional fluids could be given as guided by local emergency medical services protocols. Subsequent in-hospital care was not prescribed, with the exception of protocol-specified monitoring of serum sodium levels during the first 24 hours. Investigators encouraged the implementation of established guidelines for care of critically ill patients with trauma.

Outcome Measures

The primary outcome was 6-month neurologic status based on the Extended Glasgow Outcome Score (GOSE). Additional assessment of neurologic outcome included the GOSE at discharge and 1 month following discharge, and the Disability Rating Score (DRS) at discharge, 1 month following discharge, and 6 months following injury. The GOSE and DRS following discharge were determined by the use of a structured telephone survey. If the patient was unable to respond to the survey, information was gathered from a family member or caregiver. Previous reports have validated the use of information obtained from caregivers to assess both the GOSE and DRS.

Other secondary outcomes included 28-day survival, survival to hospital discharge, ICP, interventions required to manage intracranial hypertension, fluid and blood requirements in the first 24 hours, physiologic parameters of organ dysfunction, 28-day acute respiratory distress...
syndrome–free survival, Multiple Organ Dysfunction Score,\textsuperscript{19,20} and nosocomial infections.\textsuperscript{19,20} Diagnosis of multiple organ dysfunction was subject to patients having the required physiologic measurements available during their intensive care unit (ICU) stay. Measures of resource utilization included ventilator-free days in the first 28 days, days alive outside the ICU, and days alive outside the hospital within 28 days. TBI severity was assessed using the Abbreviated Injury Score for the head (head AIS) and the Marshall Score to grade the head computed tomography findings.\textsuperscript{21,22}

Randomization and Blinding

All study fluids were purchased from Biophausia Inc (Stockholm, Sweden). Study fluids were provided in identical intravenous bags and shipped to a single distribution center, where they were labeled with a randomly generated numeric code. The randomization scheme was 1:1:1:4 for hypertonic saline, hypertonic saline/dextran, and normal saline, respectively. This ratio was chosen because it can be shown that this is technically the most efficient ratio for this setting.\textsuperscript{23} Patients were individually randomized by administration of a blinded bag of study fluid.

All out-of-hospital personnel, clinicians, investigators, and patients remained blinded to the treatment assignment until end of study. Because of a labeling issue at the onset of the study, the randomization scheme was initially biased toward enrolling more patients into the normal saline group. After the issue was resolved, randomization continued according to protocol, with final distribution among the groups approaching the planned ratios.

Sample Size and Power Calculations

To assess neurologic outcome, we dichotomized the GOSE into good outcome (moderate disability or good recovery [GOSE $\leq 4$]) vs poor outcome (severe disability, vegetative state, or death [GOSE $>4$]). A 15% relative reduction in the prevalence of poor outcome was considered clinically relevant. Review of the literature suggested that 40% to 57% of this population would have a poor outcome.\textsuperscript{24,25}

A 49% incidence of poor outcome was estimated, and hypertonic fluids were assumed to offer a 15% relative reduction (absolute reduction, 7.5%) in the risk of poor outcome. Based on a previous trial that used a GCS score of 8 or less as an out-of-hospital enrollment criterion, we anticipated that approximately 10% of the patients enrolled in the TBI cohort would have a less severe injury and have other reasons for altered mental status, such as alcohol or drug intoxication.\textsuperscript{26} These patients would be unlikely to meet criteria for poor outcome, regardless of treatment. Therefore, we estimated a sample size of 2122 patients to provide an overall power of 80% (1-sided study-wide $\alpha = .025$, 62.6% power for each of the 2 comparisons) for an attenuated absolute reduction of 6.75% (based on the 10% contamination with truly uninjured patients) for each individual agent vs control, accounting for the primary analysis and 2 interim analyses.

Data Analysis

The primary analysis was designed as modified intent-to-treat, with all patients who had fluid connected to the intravenous tubing included regardless of how much fluid was administered. Per the a priori trial design, patients for whom the fluid bag was opened but not connected to the intravenous line were not included in the primary analysis; thus, we were not permitted to collect hospital and outcome data on such patients because they were not considered enrolled in the trial. Tests for differences in proportions were used for the primary analysis.

Initial analyses of the data indicated the absence of 6-month neurologic outcome data for 15% of the study cohort. Therefore, in addition to the completer analysis, we performed an analysis using multiple hot deck imputations (20 imputations) to estimate the 6-month neurologic outcome. For these imputations we used data from patients who were discharged alive based on 1-month GOSE data or discharge GOSE (if 1-month data were not available), length of hospital stay, and treatment group.\textsuperscript{27} For 28-day survival, patients with missing 28-day vital status who were known to be discharged alive prior to 28 days were assumed to be alive at day 28.

Significance was defined as $P < .05$ based on 2-sided tests. Differences in means or proportions with 95% confidence intervals are also presented. Medians with interquartile ranges (IQRs) are reported for skewed variables. Analyses were performed using SAS version 9.2 (SAS Institute Inc, Cary, North Carolina) and S-plus version 7.0 (Tibco Spotfire, Somerville, Massachusetts). A priori secondary analyses included patients with an Abbreviated Injury Score for the head (head AIS) of 4 or greater and 2 or greater, those with documented intracranial hemorrhage, and those requiring emergent craniotomy. Secondary outcomes were assessed using t tests or $\chi^2$ analyses as appropriate.

Trial Monitoring

Trial monitoring was conducted using a group sequential stopping rule for each comparison of hypertonic saline/dextran vs normal saline and hypertonic saline vs normal saline, based on a level 0.0125 one-sided group sequential test with O’Brien-Fleming boundary relationships for efficacy and a nonbinding futility boundary that corresponds to a boundary in the Wang and Tsiatis\textsuperscript{28} power family of boundary shape functions, as implemented in the unified family of Kittelson and Emerson\textsuperscript{29} with boundary shape parameter $P = .80$ and $\beta = .9875$. In making a decision to terminate the clinical trial, the data and safety monitoring board was also presented with estimates of the 95% confidence intervals for treatment effects after adjustment for the sequential stopping rule, the
Bayesian predictive power of eventual statistical significance based on non-informative (flat) prior distributions, and conditional power estimates defined for a spectrum of hypothesized treatment effects.

**Regulatory Oversight**

The study was conducted under the US regulations for Exception From Informed Consent for Emergency Research (21 CFR 50.24) and the Canadian Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans. The protocol was reviewed and approved by the US Food and Drug Administration and Health Canada. The protocol was also approved by all institutional review boards (United States) and research ethics boards (Canada) in the communities in which the research was conducted. Consent was obtained for continuation in the trial after hospital arrival. Details of the community consultation and public disclosure processes have been published elsewhere.\(^\text{30,31}\)

**RESULTS**

Between May 2006 and May 2009, 1331 patients were randomized (FIGURE 1). For 49 of these, the study fluid package was opened but fluid was not administered. Reasons included that out-of-hospital personnel recognized that the patient did not meet inclusion criteria; intravenous access could not be obtained or was lost prior to fluid administration; the sterility of the bag had been broken; or out-of-hospital personnel were unsure of inclusion/exclusion criteria and elected not to administer the fluid.

Complete 6-month neurologic outcome data were available for 1087 of 1282 treated patients (85%). The primary reasons for loss of follow-up included refusal of the patient to consent to contact after discharge, inability to obtain consent because of rapid discharge of minimally injured patients or lack of legal next of kin for severely disabled patients, or inability to locate the patient after discharge. Neurologic outcome data were available at either discharge or 1 month for 1217 of 1282 patients (95%). Because of the nature of this trial we were unable to track patients screened but not enrolled, and adequate epidemiologic data to estimate the potentially eligible population were not available.

At a planned data and safety monitoring board review for the second formal interim analysis (April 27, 2009), data from 1073 participants were reviewed and the futility boundary was crossed, with a boundary crude difference of 0.016 in proportion of good outcome for each of the 2 comparisons of treatment to normal saline. The crude differences for the group comparisons ranged from \(-0.017\) to \(-0.041\), depending on the assumptions made regard-
The primary outcome was 6-month neurologic function based on the GOSE. As noted, this outcome measure was obtained for 85% of the study population. To address the possibility that data might not be missing at random, we performed multiple hot deck imputations to account for missing neurologic outcome data, based on patients alive at discharge. Thus, the results for 6-month GOSE are presented as both the complete and imputation analysis, with the multiple imputation analysis results considered the primary result. Figure 2 illustrates the distribution of GOSE by category. When dichotomized to assess GOSE of 4 or less (severe disability, vegetative state, or death), there were no significant differences between the groups based on either the complete cases or imputed analysis (Table 3). There was also no difference in the DRS between treatment groups 6 months after injury.

There was no difference in 28-day survival, development of organ failure, or duration of ICU and hospital stay. Interestingly, there appeared to be a higher rate of nosocomial infection in the hypertonic fluid groups, which was

### Table 1. Demographic, Injury Severity, and Out-of-Hospital Care Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Hypertonic Saline/Dextran (n = 359)</th>
<th>Hypertonic Saline (n = 341)</th>
<th>Normal Saline (n = 582)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD), y</td>
<td>38.5 (18.6)</td>
<td>38.6 (17.3)</td>
<td>39.5 (19.2)</td>
</tr>
<tr>
<td>Male sex, No. (%)</td>
<td>273 (76.3)</td>
<td>277 (81.2)</td>
<td>426 (73.3)</td>
</tr>
<tr>
<td>Blunt trauma, No. (%)</td>
<td>352 (98.6)</td>
<td>334 (97.9)</td>
<td>575 (99.0)</td>
</tr>
<tr>
<td>Out-of-Hospital fluids, mean (SD)</td>
<td>0.88 (0.71)</td>
<td>0.85 (0.65)</td>
<td>0.82 (0.63)</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>3.3 (1.9)</td>
<td>3.3 (1.8)</td>
<td>3.3 (1.8)</td>
</tr>
<tr>
<td>Head AIS, mean (SD)</td>
<td>3.3 (1.9)</td>
<td>3.3 (1.8)</td>
<td>3.3 (1.8)</td>
</tr>
<tr>
<td>Score, No. (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>3 (0.8)</td>
<td>2 (0.6)</td>
<td>1 (0.2)</td>
</tr>
<tr>
<td>2</td>
<td>30 (8.5)</td>
<td>31 (9.3)</td>
<td>55 (9.7)</td>
</tr>
<tr>
<td>3</td>
<td>44 (12.5)</td>
<td>45 (13.5)</td>
<td>74 (13.1)</td>
</tr>
<tr>
<td>4</td>
<td>81 (22.9)</td>
<td>67 (20.1)</td>
<td>129 (22.8)</td>
</tr>
<tr>
<td>5</td>
<td>125 (35.4)</td>
<td>126 (37.8)</td>
<td>202 (35.7)</td>
</tr>
<tr>
<td>6</td>
<td>3 (0.8)</td>
<td>1 (0.3)</td>
<td>1 (0.2)</td>
</tr>
<tr>
<td>9 (unknown)</td>
<td>6 (1.7)</td>
<td>6 (1.8)</td>
<td>11 (1.9)</td>
</tr>
<tr>
<td>Marshall Score, first head CT, No. (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diffuse injury I</td>
<td>102 (29.2)</td>
<td>98 (28.0)</td>
<td>175 (31.2)</td>
</tr>
<tr>
<td>Diffuse injury II</td>
<td>128 (36.7)</td>
<td>122 (37.5)</td>
<td>184 (32.9)</td>
</tr>
<tr>
<td>Diffuse injury III</td>
<td>44 (12.6)</td>
<td>40 (12.3)</td>
<td>67 (12.0)</td>
</tr>
<tr>
<td>Mass lesion</td>
<td>59 (16.9)</td>
<td>43 (13.2)</td>
<td>105 (18.8)</td>
</tr>
<tr>
<td>Out-of-hospital advanced airway, No. (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time from dispatch call to fluid administration, mean (SD), min</td>
<td>33.7 (18.5)</td>
<td>34.1 (25.2)</td>
<td>35.7 (24.1)</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>29.8 (20.9-42.0)</td>
<td>28.0 (20.0-40.0)</td>
<td>29.8 (20.3-45.0)</td>
</tr>
<tr>
<td>Total out-of-hospital time, mean (SD), min</td>
<td>54.9 (23.8)</td>
<td>56.0 (30.4)</td>
<td>57.3 (30.3)</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>51.6 (38.0-66.0)</td>
<td>47.9 (37.6-65.7)</td>
<td>51.1 (38.2-68.0)</td>
</tr>
<tr>
<td>Out-of-hospital fluids, mean (SD), L</td>
<td>0.88 (0.71)</td>
<td>0.85 (0.65)</td>
<td>0.82 (0.63)</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>0.70 (0.35-1.25)</td>
<td>0.65 (0.35-1.25)</td>
<td>0.65 (0.35-1.15)</td>
</tr>
<tr>
<td>Air transport, No. (%)</td>
<td>150 (41.9)</td>
<td>139 (40.8)</td>
<td>217 (37.4)</td>
</tr>
</tbody>
</table>

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(Reprinted) JAMA, October 6, 2010—Vol 304, No. 13 1459.
primarily related to an increased rate of bloodstream and urinary tract infection. There were no statistically significant differences in the rate of adverse events between the treatment groups. Importantly, we observed no increase in progression of intracranial hemorrhage in the hypertonic fluid groups.

As a predefined subgroup analysis we also evaluated the GOSE between the treatment groups for patients with evidence of severe anatomical TBI (head AIS ≥4) and for those with head AIS of 2 or greater to exclude those with minor or no TBI. As noted in Table 3, restricting the analysis to these subgroups revealed no statistically significant differences among the treatment groups with regard to the proportion of patients with a GOSE of 4 or less.

### Comment

To our knowledge, this is the largest randomized clinical trial of hypertonic resuscitation following TBI. We were unable to demonstrate any improvement in 6-month neurologic outcome or survival in this patient population. Previous clinical trials in this patient population have been limited and with mixed results. A meta-analysis of the subset of patients with severe TBI enrolled in out-of-hospital studies conducted prior to 1997 of hypertonic saline/ dextran for injured patients with hypovolemic shock (n = 233) suggested an odds ratio for improved survival of 2.12 (P = .048). In 2004, Cooper et al reported a randomized controlled trial of hypertonic saline

### Table 2. Admission Physiology, Neurologic Monitoring, and Interventions

<table>
<thead>
<tr>
<th>Table 2. Admission Physiology, Neurologic Monitoring, and Interventions</th>
<th>Mean (SD)</th>
<th>Difference, % (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertonic Saline/Dextran (n = 359)</td>
<td>Hypertonic Saline (n = 341)</td>
<td>Normal Saline (n = 582)</td>
</tr>
<tr>
<td><strong>Admission physiology</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systolic blood pressure, mm Hg</td>
<td>141.2 (33.1)</td>
<td>136.9 (33.5)</td>
</tr>
<tr>
<td>GCS score</td>
<td>5.2 (3.6)</td>
<td>5.5 (3.8)</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>3.0 (3.0-6.0)</td>
<td>3.0 (3.0-7.0)</td>
</tr>
<tr>
<td>Hemoglobin, g/dL</td>
<td>12.0 (2.2)</td>
<td>12.5 (2.2)</td>
</tr>
<tr>
<td>Serum sodium, mEq/L</td>
<td>146.1 (5.1)</td>
<td>147.1 (5.1)</td>
</tr>
<tr>
<td>Hemoglobin, g/dL</td>
<td>12.0 (2.2)</td>
<td>12.5 (2.2)</td>
</tr>
<tr>
<td>Serum sodium, mEq/L</td>
<td>146.1 (5.1)</td>
<td>147.1 (5.1)</td>
</tr>
<tr>
<td>Hyperventilation, No. (%)</td>
<td>31 (9.1)</td>
<td>31 (9.6)</td>
</tr>
<tr>
<td>MAnnitol in first 5 d, No. (%)</td>
<td>31 (9.1)</td>
<td>31 (9.6)</td>
</tr>
<tr>
<td>Total mannitol in first 12 h, g/kg</td>
<td>6.7 (26.1)</td>
<td>5.6 (21.6)</td>
</tr>
<tr>
<td>Hours with CPP &gt;25 mm Hg in the first 12 h</td>
<td>1.2 (2.8)</td>
<td>1 (2.9)</td>
</tr>
<tr>
<td>Hours with CPP &gt;25 mm Hg in the first 12 h</td>
<td>1.2 (2.8)</td>
<td>1 (2.9)</td>
</tr>
</tbody>
</table>

### Abbreviations:

- CI: confidence interval
- CPP: cerebral perfusion pressure
- GCS: Glasgow Coma Score
- IQR: interquartile range
- NA: not available

- aFor differences in means/proportions across all 3 treatment groups.
- bCells too small for valid interval estimation using normal approximation or for χ² P value.
vs normal saline for 229 patients with an out-of-hospital GCS score of 8 or less and systolic blood pressure less than 100 mm Hg. No difference in 6-month GOSE was observed between the treatment groups. Mortality in that study was greater than 50%, owing to the fact that those patients had both hypovolemic shock and evidence of severe TBI.

Despite these findings, hypertonic solutions have been increasingly used in the hospital for management of intracranial hypertension in the setting of TBI. Previous studies have used concentrations ranging from 1.6% to 23.4% saline with documented reduction in ICP and improved cerebral perfusion. No studies have demonstrated improvement in clinical outcome. Data from randomized controlled trials in support of improved neurologic outcome using mannitol, recommended as the primary osmotic agent for management of increased ICP, have likewise been limited.

Because cerebral edema can occur early after injury, we hypothesized that starting hyperosmolar therapy in the out-of-hospital setting could reduce the subsequent sequelae leading to secondary brain injury. It is possible the lack of effect observed is attributable to the need for a more prolonged course of hyperosmolarity continuing into the early hospital period or the dilutional effects of crystalloid therapy, which did not differ after initial treatment. We did not observe any difference in the initial ICP reading between the treatment groups, but this analysis is limited because only 27.5% of the cohort received an ICP monitor and the timing of ICP monitor placement was variable. In most cases these monitors were placed after admission to the ICU; thus, early effects on ICP may not have been detected, and some patients may have died prior to ICP monitor placement. Furthermore, the clinical decision to begin ICP monitoring was made after randomization. Thus, if the initial resuscitation fluid improved the clinical appearance of the patient, this could have resulted in a decrease in the use of ICP monitors in these groups. We did not observe any difference between the treatment groups in the rate of ICP monitoring (Table 2). The increase in serum sodium level observed in the hypertonic fluid groups was consistent with previous studies.

Despite the plethora of animal data demonstrating modulation of the inflammatory response by hypertonic fluids, we did not observe any difference in rates of organ failure and in fact saw a slight increase in the rate of nosocomial infection in the hypertonic fluid groups. While previous studies have suggested that hypertonicity preserves T-cell function and thus should protect the host from subsequent infection, recent work has also shown up-regulation of the A3 receptor on neutrophils following injury that could increase susceptibility to infection with hypertonic therapy. Further work in this area is necessary to define these mechanisms of action.

Given that patients were enrolled based on the out-of-hospital GCS score, we anticipated that approximately 10% of these patients would have an altered mental status due to intoxicating substances and not significant TBI. To account for this possibility we increased our proposed sample size and planned a subgroup analysis to evaluate outcome for those patients with anatomical evidence of TBI based on a head AIS score of 2 or greater. Restricting the outcome analysis to this subgroup did not show any difference in 6-month GOSE between the treatment groups. Likewise, restricting the analysis to the sub-

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**Figure 2. Six-Month Extended Glasgow Outcome Score by Treatment Group—Imputed and Completer Analyses**

The Extended Glasgow Outcome Scale (GOSE) is an 8-point scale (1=dead, 2=vegetative, 3=lower severe disability, 4=upper severe disability, 5=lower moderate disability, 6=upper moderate disability, 7=lower good recovery, 8=upper good recovery). Imputed analysis represents the results for the entire cohort who received the intervention, using multiple hot deck imputations (20 imputations) for missing 6-month GOSE data. Completer analysis represents data from all cases with complete 6-month outcome data, including those who discontinued intervention but received partial infusion of study fluid.
The strengths of this study include the randomized, blinded, placebo-controlled design; large sample size; early administration of hypertonic flu-
ids; and generalizability across several communities in the United States and Canada. The primary limitation is that TBI management in the hospital was not controlled. Therefore, not all patients with severe TBI received ICP monitoring, and the timing of monitor insertion was variable. Furthermore, some patients received additional hypertonic fluids after hospital arrival and others were treated with mannitol, based on neurosurgeon preference. Lastly, the challenge of obtaining complete 6-month follow-up in a trauma population resulted in a 15% rate of missing data for the primary outcome. These data were not likely missing at random, with less severely disabled patients being more difficult to locate along with the ability of surviving patients to refuse continued follow-up. We accounted for this using multiple hot deck imputations to handle missing variables and only imputed data based on patients discharged alive. Conclusions were unchanged when compared with the complete data analysis.

In summary, in this randomized controlled trial, we were unable to demonstrate any improvement in 6-month neurologic outcome or survival for trauma patients with presumed severe TBI (out-of-hospital GCS ≤8) without evidence of hypovolemic shock, who received a single bolus of hypertonic fluids compared with normal saline in the out-of-hospital setting. While this does not preclude a benefit from such treatment were it administered differently, at present there appears to be no compelling reason to adopt a practice of hypertonic fluid resuscitation for TBI in the out-of-hospital setting.

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Author Contributions: Dr May had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Bulger, Brasel, Schreiber, Kerby, Tisherman, Newgard, Slutsky, Coimbra, Emerson, Minei, Baker, Christenson, Davis, Aufrheide, Sopko, Hata, Hoyt.

Acquisition of data: Bulger, Brasel, Schreiber, Kerby, Tisherman, Newgard, Coimbra, Emerson, Minei, Bardarson, Kudenchuk, Baker, Christenson, Idris, Davis, Fabian, Aufrheide, Callaway, Williams, Banek, van Heest, Hata, Hoyt.


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Statistical analysis: May, Emerson.

Obtained funding: Bulger, Kerby, Newgard, Slutsky, Christenson, Idris, Davis, Hoyt, Hoyt.

Administrative, technical, or material support: Kerby, Slutsky, Coimbra, Bardarson, Kudenchuk, Baker, Christenson, Idris, Davis, Callaway, Williams, Banek, Vaillancourt, van Heest, Sopko, Hata, Hoyt.

Financial Disclosures: None reported.

Funding/Support: The Resuscitation Outcome Consortium (ROC) network.

Study concept and design: Bulger, Schreiber, Newgard, Slutsky, Coimbra, Emerson, Minei, Christenson, Callaway, Williams, Hata, Hoyt.

Study supervision: Bulger, Schreiber, Newgard, Emerson, Minei, Christenson, Callaway, Williams, Hoyt.

Financial/Conflict of Interest Disclosures: None reported.

Additional Contributions: We thank all of the emergency medical services agencies that participated in this project.

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Individuals are considered sincere when there is little or no discrepancy between the goals they seek and those they claim to be seeking.