A trial of intracranial pressure monitoring in traumatic brain injury

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Expanded abstract

Citations
Chesnut R, Temkin N, Carney N, Dikmen S, Rondina C, Videtta W, Petroni G, Lujan S, Pridgeon J, Barber J, Machamer J, Chaddock K, Celix J, Cherner M, Hendrix T: A trial of intracranial pressure monitoring in traumatic brain injury. N Engl J Med 2012, 367:2471–2481.

Background

Intracranial pressure (ICP) monitoring is considered the standard of care for severe traumatic brain injury (TBI) and is used frequently, but the efficacy of treatment based on monitoring in improving the outcome has not been rigorously assessed.

Methods

Objective: The objective was to compare efficacy of guideline-based management in which a protocol for monitoring intraparenchymal ICP was used (ICP group) or a protocol in which treatment was based on imaging and clinical examination (exam group).

Design: A multicenter randomized controlled trial was conducted.

Setting: The trial was set in ICUs in Bolivia or Ecuador.

Subjects: Patients had severe TBI (n = 324) and were 13 years of age or older.

Interventions: Patients were randomly allocated to ICP monitoring or clinical exam-based monitoring.

Outcomes: The primary outcome was a composite of survival time, impaired consciousness, functional status at 3 and 6 months, and neuropsychological status at 6 months; neuropsychological status was assessed by an examiner who was unaware of the protocol assignment.

This composite measure was based on performance across 21 measures of functional and cognitive status and was calculated as a percentile (with 0 indicating the worst performance, and 100 the best performance).

Results

There was no significant between-group difference in the primary outcome, a composite measure based on percentile performance across 21 measures of functional and cognitive status (score 56 in the pressure-monitoring group versus 53 in the imaging-clinical examination group; \(P = 0.49\)). Six-month mortality rates were 39% in the pressure-monitoring group and 41% in the imaging-clinical examination group (\(P = 0.60\)). The median lengths of stay in the ICU were similar in the two groups (12 days in the pressure-monitoring group and 9 days in the imaging-clinical examination group; \(P = 0.25\)), although the number of days of brain-specific treatments (for example, administration of hyperosmolar fluids and the use of hyperventilation) in the ICU was higher in the imaging-clinical examination group than in the pressure-monitoring group (4.8 versus 3.4, \(P = 0.002\)). The distributions of serious adverse events were similar in the two groups.

Conclusions

For patients with severe TBI, care focused on maintaining monitored ICP at 20 mmHg or less was not shown to be superior to care based on imaging and clinical examination.

Commentary

Traumatic brain injury (TBI) causes significant morbidity and mortality in the US, with 1.7 million TBIs accounting for 275,000 hospitalizations and 52,000 deaths annually [1]. At least 5.3 million Americans currently live with TBI-related disabilities, with an estimated $60.4 billion in direct and indirect costs [2,3].

Many interventions have been investigated to reduce TBI-associated sequelae. One such intervention is invasive intracranial pressure (ICP) monitoring, which is important...
as patients with ICP of greater than 20 mmHg have worse outcomes [4]. The most recent guidelines for TBI management recommend ICP monitoring in select patients [5].

Invasive ICP monitoring is not without risks, such as infection, hemorrhage, malfunction, and additional costs. As with any monitoring in the intensive care unit, the goal is to both obtain accurate data and initiate interventions that positively affect outcomes.

Several single-center and multicenter observational studies have shown that ICP-targeted management of TBI is associated with worse outcomes, including prolonged mechanical ventilation, worse functional status, and higher risk of pneumonia, acute kidney injury, and mortality [6-8]. However, the results of these studies may be biased because they were retrospective and sicker patients may be more likely to receive ICP monitoring.

Chesnut and colleagues conducted a prospective randomized clinical trial to determine whether ICP monitoring improves outcomes. Since invasive ICP monitoring is considered standard of care, this study would have lacked clinical equipoise if conducted in the United States. Therefore, this study had to be conducted in centers that deviated from this ‘standard’. The authors recruited patients from Bolivia and Ecuador, where ICP monitoring was not the standard of care. The authors concluded that ICP-guided care was not superior to care based on imaging and clinical exam alone, contrary to published guidelines.

The study results suggest that management of TBI based on invasive monitoring with a goal ICP of not more than 20 mmHg was not superior to management based on imaging and clinical exam alone, thus questioning the added risks of invasive ICP monitoring. Since this may significantly impact clinical practice patterns, a detailed assessment of the study and results must be undertaken.

Strengths of this study include its randomized, prospective nature as well as the comprehensive nature of the endpoint measured, which factored in measurements of in-hospital morbidity and quality of life after discharge.

This study has limitations. First, the authors did not report differences in pre-hospital care or care after hospital discharge. Although the study was randomized, there may have been important differences in care, including pre-hospital transport times and management and post-discharge care and rehabilitation. Unfortunately, this information is not provided in the article and may confound the results.

Second, the authors used a composite primary endpoint. They attempted to incorporate measurements of many quality-of-life issues, including mortality, cognitive impairment, and motor impairment. They produced an original, thoughtful, and complex, yet unvalidated, composite tool to measure TBI outcomes.

Third, they assessed therapeutic intensity as a secondary outcome and calculated it based on an original scoring system that bears some resemblance to the Therapeutic Intensity Level Scoring System [9] and Pediatric Intensity Level of Therapy Scoring System [10]. Based on this scale, the groups received different intensities of therapeutic interventions, with the control group receiving more treatment with mannitol, hypertonic saline, and hyperventilation ($P < 0.001$). However, the ICP group was noted to have increased barbiturate use ($P = 0.02$). What weights to assign these different interventions in a measure of treatment intensity is open to debate. Furthermore, detailed review of the protocols applied to the two groups shows small but key differences. For example, the first intervention for elevated ICP in the monitored group was to drain cerebrospinal fluid (CSF) if a ventriculostomy was in place, which was not an option in the exam group. The ICP group had 346 hours of CSF drainage, but according to a personal communication with the author, the majority of this was in a single patient who had intraventricular blood and was clearly an outlier. Other differences included the use of neuromuscular blocking agents in the ICP group and furosemide in the exam group. The results of the study would have been more robust had these differences been minimized. Finally, no information regarding the number of head computed tomography scans performed in each group was provided. Although the author stated there was not a significant difference (personal communication), this raises the question of additional radiation exposure and increased hospital costs.

An important question is whether the findings of this study can be generalized to other countries. There are differences in pre-hospital care, in-hospital management of patients with TBI, and rehabilitation services across different countries, and these differences may affect outcomes. Although invasive ICP monitoring did not affect outcomes in resource-poor settings, whether it improves outcomes in resource-rich countries is not known.

This trial is to be commended for its courage in questioning a therapy that has gained widespread acceptance as the ‘standard of care’ as well as its attempt to study this monitoring modality in a randomized controlled fashion. Though not providing a definitive answer as to whether ICP monitoring improves outcomes, it adds to the discussion in a thought-provoking way. ICP may represent a non-specific marker of dynamic brain pathology that draws attention to dynamic changes occurring in the brain, and both waveform analysis and cerebrovascular reactivity index can be derived from ICP monitoring, which may help guide therapy [11]. Moreover, the present study highlights the importance of clinical examinations in patient management, reminding us that reliance on monitors and measured parameters should not substitute for clinical acumen. Chesnut himself states in a commentary that
‘use of a safe and accurate quantitative index of ICP, the ICP course, and its response to treatment is much preferable to treating semi-empirically’ [12]. To this end, invasive ICP monitoring is better viewed as a tool providing another piece of data for the clinician to take into account and not as an endpoint in and of itself dictating treatment.

**Recommendation**

There is insufficient evidence at this time to abandon treatment based on ICP monitoring, and further studies need to be done before changing practice. ICP monitoring must be used as part of a multimodal approach to the patient and viewed as an additional tool available to the clinician to manage patients with TBI.

**Abbreviations**

CSF: cerebrospinal fluid; ICP: intracranial pressure; TBI: traumatic brain injury.

**Competing interests**

The authors declare that they have no competing interests.

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