Review

Year in review 2008: Critical Care - trauma

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Abstract

Eleven papers on trauma published in Critical Care during 2008 addressed traumatic brain injury (TBI), burns, diagnostic concerns and immunosuppression. In regard to TBI, preliminary results indicate the utility of either magnetic resonance imaging (MRI) or ultrasound in measuring optic nerve sheath diameter to identify elevated intracranial pressure (ICP) as well as the potential benefit of thiopental for refractory ICP. Another investigation demonstrated that early extubation of TBI patients whose Glasgow Coma Scale score was 8 or less did not result in additional incidence of nosocomial pneumonia. Another study indicated that strict glucose control resulted in worse outcomes during the first week after TBI, but improved outcomes after the second week. Another paper showed the prolonged neuroprotective advantages of progesterone administration in TBI patients. There was also guidance on improved classifications of renal complications in burn patients. Another study found that patients with inhalation injuries and increased interleukin-6 (IL-6) and IL-10 and decreased IL-7 had increased mortality rates. One literature review described the disadvantages of prolonged immobilization or additional use of MRI for ruling out cervical spine injuries in obtunded TBI patients already cleared by computerized tomography scans. Other investigators found that higher N-terminal pro B-type natriuretic peptide (NT-proBNP) levels may be useful markers for post-traumatic cardiac impairment. Finally, an experimental model showed that both splenic apoptosis and lymphocytopenia may occur shortly after severe hemorrhage, thus increasing the threat of immunosuppression in those with severe blood loss.

Introduction

During 2008, Critical Care published a number of papers in regard to injured patients. Several of these articles focused on identification and treatment strategies for traumatic brain injury (TBI) and increased intracranial pressure (ICP). Another examined acute renal injury in burn patients, and one described a novel method of assessing serum markers of burn severity. Finally, there were several articles applicable to the multiple-injury patients, including those articles that examined the potential adverse effects of prolonged cervical immobilization or the practice of adding magnetic resonance imaging (MRI) to computerized tomography (CT) scans in ruling out cervical spine injuries. Another article demonstrated the utility of obtaining N-terminal pro B-type natriuretic peptide (NT-proBNP) in identifying multiple organ dysfunction syndrome and decreased cardiac index in multiple-injury patients. The final paper was an experimental investigation that demonstrated two of the potential mechanisms for post-traumatic immunosuppression.

Traumatic brain injury

One of the challenges in treating patients with TBI is the identification of increased ICP. Increased ICP is known to affect outcomes [1] but often requires invasive catheters to determine its presence and monitor its course. One non-invasive modality that has been examined is the use of ultrasound to detect changes in the optic nerve sheath diameter (ONSD). Many of the research studies evaluating the use of ONSD in elevated ICP have used indirect methods to confirm elevated ICP, including CT findings and transcranial Doppler [2-6]. In their 2008 Critical Care article in regard to this subject, Soldatos and colleagues [7] used an intraparenchymal catheter to directly measure ICP and found that an ONSD of 5.7 mm was the optimal cutoff value for detecting an ICP of greater than 20 (sensitivity and specificity of 74.1% and 100%, respectively) and that ultrasound measurement of ONSD correlated well with invasive and noninvasive measurements of ICP. These results are similar to those of Geeraets and colleagues [8] and Kimberley and colleagues [9], suggesting that an ONSD of between 5 and 5.8 can reliably detect elevated ICP, depending on what sensitivity and specificity are desired. While ultrasound is a commonly available and easily performed test, many patients with head injury or other causes of increased ICP may also undergo an MRI. Using MRI, Geeraets and colleagues [10] performed a study in which they measured ONSD to determine how well it
correlates with ICP and found similar results; a diameter of 5.82 mm is the optimal diameter for predicting the presence of elevated ICP.

Patients with TBI and an elevated ICP that is refractory to all other medical and surgical treatment modalities may very well benefit from high-dose barbiturate therapy, although this practice remains controversial in some centers given its associated high morbidity [11]. However, few data comparing different barbiturate regimens exist. In another 2008 trauma paper in Critical Care, Pérez-Bárcena and colleagues [12] randomly assigned 44 patients with TBI and refractory intracranial hypertension to receive high-dose thiopental or high-dose pentobarbital drips. The authors showed better ICP control with thiopental (odds ratio = 5.1, confidence interval = 1.2 to 21.9; \( P = 0.027 \)). Though encouraging, this trial involved a small patient cohort with an unblinded study design. In addition, there were more patients with brain swelling on the initial CT scan in the pentobarbital group, so these preliminary results must be interpreted with caution.

Current practice management guidelines dictate the need for airway protection in patients with TBI and a Glasgow Coma Scale (GCS) score of 8 or less for respiratory protection and prevention of aspiration into the lungs [13]. It has been suggested, however, that prolonged endotracheal intubation leads to increased risk of pneumonia as well as longer intensive care unit (ICU) and overall hospital stays [14]. The 2008 study by Manno and colleagues [15] in Critical Care begins to address this issue. They randomly assigned 16 patients to an early extubation arm versus a delayed extubation arm. Although the study was designed to assess study feasibility and to determine the necessary sample size for a higher powered study, it did show that none of the patients in the early extubation arm developed nosocomial pneumonia compared with 37% in the delayed extubation arm. Although the study was underpowered for significant statistical analysis, this very impressive trend was compelling and the concept should be examined further [15].

Prevention of secondary injury is perhaps the most important goal in the critical care of those with TBI. The recent critical care literature has demonstrated significant debate concerning the optimal range and method of glycemic control in many critical illnesses [16-18]. It remains to be seen, however, how strict the glycemic control needs to be in severe TBI. Several recent studies have suggested that hypoglycemic episodes lead to metabolic derangement with lactate production and secondary brain injury [19-23]. A study from Zurich, Switzerland, by Meier and colleagues [24] published in 2008 in Critical Care showed that there appears to be a temporal relationship between glucose control and benefit. In their investigation, they compared a strict glucose control regimen maintaining a blood glucose of between 3.5 and 6.5 mmol/L (63 and 117 mg/dL) with a regimen maintaining a blood glucose of between 5 and 8 mmol/L (90 and 144 mg/dL). The lower glucose range was associated with a trend toward increased mortality during the first two weeks. The tight control was also associated with markedly increased insulin and norepinephrine requirements, increased frequencies of hyperglycemic and hypoglycemic episodes, and significantly elevated ICP during the first week after admission. In contrast, during the second week, maintaining patients in the lower glucose range resulted in lower ICP and reduced rates of pneumonia, bacteremia, and urinary tract infections when compared with the group in the higher glucose range. Based on these results, future research efforts should take into account these temporal factors. Lastly, with regard to TBI, progesterone has been shown to have neuroprotective benefits in several different animal models of injury [25-27]. More recently, it was shown to have improved mortality and, in patients with moderate brain injury, better outcomes than those receiving placebo for up to 30 days after injury in humans with TBI [28]. In the first issue of Critical Care for 2008, Xiao and colleagues [29] report the results of a related investigation, in which they followed outcomes for up to 6 months to determine whether the benefit was persistent. They found that, at both 3 and 6 months, patients who received progesterone had outcomes that were more favorable as measured by the GCS and the modified functional independence measure. More importantly, progesterone was associated with decreased mortality at 6 months. Along with existing experimental and preliminary clinical data, the results of this article indicate that hormonal intervention in the injured patient may now hold future therapeutic promise.

**Burns**

Two papers last year in Critical Care focused on burn care. Acute renal failure is a relatively uncommon occurrence in burn patients, but it carries a high mortality [30]. It also has been a difficult area to study, due to the variability in nomenclature and definitions used to describe kidney injury in the existing literature. In addition, many of the previous studies have been retrospective, using inconsistent data collection methods and reproducible definitions for variables, outcome measures, and reporting templates. In a 2008 Critical Care article, Steinwall and colleagues [31] used a classification system developed by the Acute Dialysis Quality Initiative Group to prospectively describe the incidence and mortality of acute renal injury. Their study involved a cohort of patients with at least 20% total body surface area burns who were admitted to a national burn center. Using the consensus-based approach and its prospective definitions, the authors found that 24% of their cohort developed some significant degree of acute kidney injury, with 5% developing a reproducible criterion for acute renal failure. Overall mortality for patients with the prescribed degree of kidney injury was 14% and, as expected, increased with severity of kidney injury: up to 83% in patients with kidney failure as defined by the classification system. This article adds to the knowledge base of the incidence and associated factors related to kidney disease in burn patients and also provides guidance for future studies.
As in the case of renal failure, inhalational injury is a harbinger of increased mortality in burn patients [32]. Despite the development of a number of potential scoring systems aimed at quantifying the potentiating effect of inhalation injury and dermal injury, none of those approaches has been altogether successful in terms of accurately predicting mortality [33]. However, Finnerty and colleagues [34] have shown that children with burn injury have a marked alteration in inflammatory cytokines when compared with healthy children. Therefore, in a follow-up study published last year in *Critical Care*, the same group, Gauglitz and colleagues [35], showed that patients with elevated interleukin-6 (IL-6) and IL-10 and decreased IL-7 serum levels upon admission had a significantly greater risk for mortality. This finding provides one potential tool for critical care providers to better predict those burn patients who would be at greater risk for mortality earlier in the course of care.

**Diagnostic issues**

A basic tenet of current trauma care practice is that the cervical spine remains vulnerable to further injury in patients who have received significant blunt force trauma to neck and who may have underlying unstable cervical spine injuries. In an analysis of over 34,000 patients with major trauma, Goldberg and colleagues [36] identified cervical spine injuries in 2.4% of cases. Approximately 70% of those cervical spine injuries were clinically significant insults [36]. This rate of cervical spine injury may be as high as 34% in the subpopulation of patients with severe blunt trauma who require ICU admission [37]. Although there are good data and resulting established criteria for ruling out a significant cervical spine injury in patients with normal mentation [38], there is still some debate with regard to the removal of cervical spine immobilization in those who remain comatose. Many sources recommend a CT scan of the cervical spine [39], but others have recommended that, in patients with a CT that indicates no cervical injury, an MRI may be necessary to detect injuries that may be missed by standard CT [40,41]. At the same time, others question some of the risks associated with getting an MRI or whether prolonged cervical immobilization outweighs the risks of missed injury. In another 2008 *Critical Care* trauma paper, Dunham and colleagues [42] conducted a review to begin the process of answering these questions. The researchers concluded that the risks of secondary brain injury from cervical immobilization or an MRI (largely due to consequences of protracted supine positioning) are much greater than the risk of cervical spine instability [42]. They specifically found that the risk for spinal instability in comatose or obtunded patients with unremarkable bony imaging was 2.5%. This was compared with a 26% increased rate of ICU complications with prolonged cervical collar use and a 27.8% incidence of increasing ICP from less than 15 mm Hg to greater than 20 mm Hg. This complication presumably occurs by virtue of the supine patient positioning (versus some degree of head elevation) that is used to maintain cervical spine precautions.

In addition, neurosurgical ICU patients transported to radiology had a greater number of secondary brain injury events compared with those undergoing studies within the unit. The MRI patients, for example, had significant ICP increases during out-of-ICU transportation. With regard to leaving the protective confines of the surgical ICU, 51% of brain-injured patients transported out of the unit developed complications such as hypoxia, hypotension, and increased ICP. With respect to the positioning required for MRI, Dunham and colleagues [42] noted not only that the flat position likely increases ICP but also that the head ‘down’ position significantly increases the risks of aspiration and ventilator-associated pneumonia. They suggest that, in some comatose or obtunded patients with a negative CT scan and no apparent spinal deficit, early collar removal is a safe and reasonable strategy.

Early post-traumatic mortality is determined by the initial traumatic impact and early resuscitation, whereas late mortality is usually associated with the development of septic inflammatory response syndrome and progresses to multiple organ dysfunction syndrome and death. This pathophysiological pathway is thought to be responsible for up to 80% of trauma deaths in the ICU [43]. Cardiac function has been described as having particular relevance to multiple organ failure, but the traditional methods of assessment, such as pulmonary artery catheterization or echocardiography, either are invasive or may not be readily available in an ICU. Therefore, Kirchhoff and colleagues [44] performed a cohort study in which they compared serum NT-proBNP levels in patients with minor signs of organ dysfunction, specifically those with a Marshall Multiple Organ Dysfunction Score (MODS) of 4 or less, with those with major signs of organ dysfunction, namely a MODS of greater than 4. The researchers found that, at admission, both groups had slightly elevated NT-proBNP but that, at 24, 48, and 72 hours after admission, the patients with the higher MODS had significantly higher levels than those with the lower MODS. They also showed that NT-proBNP levels correlated with decreased cardiac index, thus suggesting that NT-proBNP measurement may serve as a useful marker for post-traumatic cardiac impairment.

**Post-traumatic immunocompromise**

Hemorrhagic shock states are known to produce a variety of immunosuppressive effects that can lead to increased susceptibility to infections and post-traumatic complications like multiple organ dysfunction syndrome, multiple organ failure, or adult respiratory distress syndrome [45-48]. While the exact mechanisms of post-traumatic immunocompromise still are not well understood, several studies have suggested that functional and immunological alterations in the spleen may play a major role [49-51]. In an experimental design developed by Hostmann and colleagues [52], a murine model of resuscitated hemorrhagic shock was used to detect apoptosis in the spleen as a marker of cellular injury and...
reduced immune functions. The authors showed a distinct lymphocytopenia immediately after severe hemorrhage that did return to baseline within the subsequent 72 hours. They also showed a rapid activation of splenic apoptosis in a biphasic onset, both immediately after the hemorrhage and 72 hours after the bleed was induced. This article thus demonstrated two of the potential mechanisms for post-traumatic immunosuppression and provides us with important hypothesis-generating clues for future interventions.

Conclusions
The 2008 volume of Critical Care contained several articles related to the care of injured patients. Proper care of patients with TBI is necessary to prevent secondary brain injury. New therapeutic interventions may be on the horizon. Consistent approaches to the identification of renal injury and inhalational trauma in patients with thermal injury may improve future research and care. Reports from that volume also suggest that the CT scan is an adequate method of identifying un-}

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

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