Critical Illness Related Corticosteroid Insufficiency in Trauma – A Review

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Abstract

Critical Illness Related Corticosteroid Insufficiency is an intense inflammatory condition associated with steroid tissue resistance. Although traditionally felt to be uncommon, it is being increasingly recognized in severely stressed patients with prolonged intensive care unit stays. Over the last decade the spectrum of CIRCI in trauma has been better defined. Trauma patients with CNS (brain and or spinal cord) injury, burns or blunt multisystem injury are at particular risk. The diagnosis is best established using a random cortisol level combined with an Adrenocorticotropic Hormone (ACTH) stimulation test. A low cortisol level and or a low response to the ACTH stimulation test in the setting of refractory shock makes the diagnosis. Stress dose hydrocortisone therapy is essential and improves outcome. CIRCI should be suspected in any elderly trauma victim with a prolonged ICU stay that exhibits shock. Drugs known to inhibit cortisol synthesis (like etomidate) are probably best avoided in this trauma subset. CIRCI in trauma has a bimodal distribution. The first peak occurs early (within 48 hours) after injury and is associated with shock and the attendant inflammatory response. The second peak occurs a week or more into the hospital course. This peak is usually associated with sepsis. Inflammatory cytokines (particularly IL-6) are elevated during both peaks but their exact role in establishing the diagnosis remains unclear. Physicians continue to search for the Eucorticoid state that achieves a balance between the inflammation initiated by the injury and the anti-inflammatory response anchored by endogenous steroid production. The administration of exogenous steroids to achieve this balance is an approach that seems to hold promise.

Introduction

Acute injury activates the hypothalamic-pituitary-adrenal axis. The HPA axis in combination with the autonomic nervous system and immune system produces the hormonal and cytokine milieu that allows the patient to respond to the initial insult [1]. This neuroendocrine and immune response involves a complex orchestration of hormones, cytokines, prostaglandins and other substances. An insufficient response increases the likelihood of significant morbidity and mortality. This review highlights our current understanding of critical illness corticosteroid insufficiency in the setting of trauma.

Critical Illness Corticosteroid Insufficiency is a pro-inflammatory state characterized by corticosteroid resistance [2]. One interesting theoretical concept proposed is that the level of adrenal response is inadequate for the given level of stress [3]. Put another way the inflammation initiated by the initial insult must be balanced by the anti-inflammatory effect of endogenous steroids. This has been referred to as the ‘Eucorticoid’ state [4]. When this balance is achieved, survival is likely. When inflammation proceeds unabated, end-organ injury and death becomes a major risk. The term Critical Illness Corticosteroid Insufficiency or CIRCI has replaced the older terms relative adrenal insufficiency and functional hypoadrenalism. When describing adrenal insufficiency in trauma we will use CIRCI to encompass all of these patients.

The incidence of CIRCI varies with the underlying illness, its’ severity and duration [5]. Traditionally thought to be uncommon, CIRCI has been recognized in critically ill patients with sepsis, cirrhosis, coronary artery disease, trauma and many other conditions. The diagnostic criteria used to establish CIRCI will have an impact on the incidence [6-8].

Background and clinical context

Studies on the surgical stress response document a significant burden of inflammation related to open surgery. Increased cortisol levels are seen after major surgery [9]. Minimal access procedures produce a diminished inflammatory response and this observation supports why these procedures are often preferable in an elective setting [10]. Fast track adjuvants (including epidural anesthesia and steroids) can blunt the inflammatory response [11].

Burn injury studies in both children and adults highlight a large increase in circulating cortisol and catecholamines after significant injury [12,13]. In the setting of burn injury serum cortisol levels remain elevated for several months after injury. Past 100 days the levels begin to approach baseline (uninjured patients). Urinary catecholamines are also persistently elevated. These findings reflect the tremendous inflammatory and metabolic burden patients with large body surface area burns endure. HPA dysregulation has been noted in pediatric burn patients. Low baseline cortisol levels 72 hours after injury with an ACTH stimulation test that does not increase serum cortisol more than 9 micrograms per deciliter has been documented in a small cohort of pediatric burn patients [14].

In contrast, blunt trauma patients present a different pattern of cortisol response. Initially after injury the serum cortisol levels may be dramatically elevated only to return to a more normal level some 10 days out from injury [15]. Serum cortisol levels greater than 25 ug/dl are common after multi-system trauma [3]. The presence of hemorrhagic shock complicates this picture further as some of these patients may have low cortisol levels [16].

Keywords: Critical Illness Related Corticosteroid Insufficiency - CIRCI, Interleukin 6 - IL-6, Adrenocorticotropic Hormone - ACTH, Central Nervous System - CNS

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A trauma-induced catecholamine surge has been documented in a small cohort of bluntly injured patients. This increase in adrenaline has been linked to markers of tissue damage, coagulopathy and death [17]. The association between elevated adrenaline levels and poor outcome cannot be assumed to be causal. The findings from this small preliminary study raise the possibility that an over exuberant neuroendocrine response may itself produce injury. Further work must be done to duplicate and expand these findings before firm conclusions can be reached.

Defining critical illness corticosteroid insufficiency biochemically has been problematic. Random cortisol levels may not accurately reflect adrenal function. The serum cortisol level reflects bound and free cortisol. 90% of cortisol is bound primarily to cortisol binding globulin [18]. The free cortisol is metabolically active. The bound cortisol is an inaccurate surrogate for the active moiety. In addition diurnal variation in cortisol levels can be misleading, making some researchers question the reliability of the random cortisol level [13]. In critical illness a random cortisol level of < 15 ug/dl has been proposed as an indication of CIRCI [18]. In severely stressed patients, a random cortisol level less than 25 ug/dl should raise the issue of CIRCI [3]. The ACTH stimulation test has been suggested as an effective way to detect inadequate adrenal response [19]. Hoen S et al. [20] noted a significant increase in interleukin 6 in trauma patients who were unable to respond to ACTH stimulation with a change in cortisol of greater than 9 ug/dl. These early non-responders had greater volume requirements (colloid) and required more vasopressor support than responders. Fann SA et al. [21] have suggested that the need for vasopressor support and the need for intubation are predictive of the development of adrenal insufficiency in the setting of trauma. These are two indicators of profound shock and clinically should guide our thinking about this condition. Bernard F et al. [22] have shown that the incidence of CIRCI (for head injured patients) will vary based upon the criteria used to establish it. Strict biochemical criteria (a serum cortisol less than 10ug/ dl or a change in cortisol less than 9 ug/dl on ACTH testing) in the clinical context of refractory shock, establishes the diagnosis of CIRCI. 

Special trauma patient subsets at significant risk

Elderly trauma patients may have an impaired adrenal response due to aging [23]. These are patients with multiple co-morbidities and declining physiologic reserve. Prolonged critical illness itself has been linked to CIRCI in chronically ill medical patients [24]. Older patients with prolonged ICU stay are at risk [25,26]. Trauma among our seniors is increasing as many remain active and continue to drive in their later years. This is the population that we must carefully evaluate when the clinical situation raises the issue of CIRCI.

Patients with CNS injury; particularly closed head injury and or spinal cord injury may be at risk [22,27,28]. Direct trauma to hypothalamus or pituitary may occur. Increased Corticotrophin releasing hormone MRNA after brain fluid percussion injury in rats has been described [29]. In another animal study using a fluid percussion injury model, the administration of large doses of methylprednisolone was associated with pituitary apoptosis and increased mortality with a biochemical profile suggestive of CIRCI [30]. Chronic repeated head injury is also associated with pituitary injury [31]. Taken together this data suggests that blunt cerebral trauma activates the HPA axis, but in the setting of massive doses of methylprednisolone, hypothalamic and pituitary dysfunction (anatomic and physiologic) may occur.

Spinal cord injury may increase the risk of CIRCI. Lecamwasam published a case report of a patient with acute spinal cord injury who developed acute adrenal insufficiency after large dose glucocorticoid therapy [32]. Weant KA et al. [33] reported 2 patients with cervical cord injuries who developed acute adrenal insufficiency. Huang TS et al. [34] have documented impaired HPA axis responses to corticotrophin releasing hormone among 25 men with spinal cord injuries. All of these patients were more than 1 year out from the initial injury. 6 patients had no response to insulin hypoglycemia and minimal or no response to ACTH stimulation. Many cord injured patients are treated with pharmacologic dosages of methylprednisolone in an effort to minimize swelling. Chen’s work cited above suggests that in the setting of cerebral trauma methylprednisolone may cause harm [30]. Supraphysiologic dosages of steroids may suppress endogenous production and disrupt the potential balance of a eucorticoid state. Very high cortisol levels have been documented in a small cohort of spinal cord injured patients [28]. All of these patients received pharmacologic methylprednisolone therapy to reduce cord edema. More than half of these patients were non-responders on ACTH testing.

As noted previously, patients who have sustained burns are a unique group. The prolonged inflammatory response seen in the setting of burns likely produces corticosteroid resistance. Prolonged elevations in serum cortisol have been clearly described. Disrupted cells in burn wound tissue may activate the conversion of cortisol to cortisone by enhancing 11B Hydoroxysteroid dehydrogenase [35].

This deactivation of cortisol may be another mechanism of corticosteroid resistance in burns.

Direct adrenal injury related to blunt trauma is an unlikely cause of CIRCI. In contrast, patients with bilateral adrenal hemorrhage in the setting of hærin-induced thrombocytopenia may be at increased risk [36].

Drugs may play a role in the development of CIRCI. Propofol, ketoconazole, rifampin and etomidate are some of the drugs that can impact adrenal function [5]. Of those mentioned, etomidate has generated the most concern. It is extremely useful because it does not produce cardiovascular dysfunction in patients about to be paralyzed and intubated. It causes adrenal suppression (for 24–48 hours) by inhibiting the 11B hydroxylase essential for steroid production [37]. Using this drug in the trauma setting may place patients at increased risk [38]. Although serum cortisol levels are lower [24] after etomidate is given, the short-lived nature of this suppression may not cause major problems for young trauma victims. For those older patients with multi-system trauma and for those with severe sepsis or septic shock, etomidate is best avoided.

A recent prospective double blind study evaluated intravenous hydrocortisone for 48 hours versus placebo in patients (two-thirds were trauma victims) who received etomidate. No difference in 28 day mortality or ICU stay was noted [39]. In this study patients who received steroids were able to wean from vasopressors more quickly. Overall hydrocortisone did not have an impact on outcome regarding etomidate induced adrenal suppression. Steroids cannot be recommended for this indication.

CIRCI and the link to sepsis

The animal and human data is compelling regarding the link of sepsis to hypothalamic-pituitary-adrenal axis activation and suppression. Mitchie HR et al. [40] have administered endotoxin to human volunteers and have documented a spike in TNF(tumor necrosis factor), ACTH and epinephrine early after the dose. The systemic and biochemical response can be muted by pretreatment with
ibuprofen. This implies that the cyclooxygenase pathway is involved in mediating the effects of endotoxin. Endotoxin administered in rats, will produce the same ACTH spike as noted above. Administering an antibody to interleukin 6 abrogates this response almost completely [41]. High levels of inflammatory cytokines (particularly TNF) can directly inhibit adrenal cortisol synthesis [5]. Cationic peptides isolated from neutrophils called defensins also impact the HPA axis. These peptides can decrease cytokine production and reduce ACTH induced steroidogenesis [5]. The clinical impact of these observations in regard to the evolution and management of CIRCI remain uncertain.

The clinical presentation and diagnostic work-up

The major manifestation of CIRCI will be the presence of refractory shock. Adequate volume resuscitation and possibly vasopressor therapy has been started. Those patients who remain in shock in spite of these measures may have corticosteroid insufficiency. Rushing GD et al. [16] have documented low serum cortisol levels in 15 patients with hemorrhagic shock. There was a 13% mortality and prolonged hospital stay. If it is seen in the setting of sepsis, all of the systemic manifestations of infection may be present. Fever, tachycardia, tachypnea, altered mental state, with diminished urine output may be seen. If hemodynamic monitoring is available a hyperdynamic state is typical. Retrospective Corticus data suggests that mortality will be increased in those patients who do not respond to the ACTH stimulation test [42]. Although the major Corticus prospective study findings did not support a difference in outcome between responders and non-responders, time to shock reversal was shorter in those patients treated with steroids [43].

The biochemical work-up is the essential next step. A baseline cortisol level in the single digits in the setting of refractory shock is diagnostic. A positive ACTH stimulation test (with the change in cortisol of less than 9 micrograms per deciliter after the administration of 250 micrograms of ACTH) in the setting of refractory shock also makes the diagnosis. The Surviving Sepsis campaign (2008) addressed the issue of corticotrophin stimulation testing for patients in septic shock. These guidelines emphasize that clinicians should not wait for results of ACTH testing to administer steroids. Indeed these authors do not advise ACTH stimulation testing at all in the setting of septic shock [44]. Assessing glucocorticoid tissue resistance remains an open question. T cells from burn-injured mice lose sensitivity to glucocorticoids [45].

Elevated serum cytokines in non-responders to the ACTH stimulation test have been documented in a small cohort of CIRCI patients [46]. Most of the cytokine increase was accounted for by the group with sepsis.

There is little published guidance regarding the use of serum cytokines to support the diagnosis of CIRCI. Electrolyte imbalance (low sodium or high potassium) is usually not present. Hypoglycemia may be present in profound adrenal insufficiency. Eosinophilia may be seen and should raise the issue of CIRCI when noted [26].

Imaging is usually not helpful in the management of these patients. CT scanning early on may reveal trauma to the adrenals. The absence or presence of adrenal trauma is not predictive of the future development of CIRCI [28]. In the setting of septic shock a recent study suggests that adrenal size (reflecting adrenal hyperplasia) might be associated with prognosis. The larger glands may be protective in the setting of septic shock [47]. Although this is a preliminary finding, it raises the question of whether extremely small glands are associated with CIRCI. There is one instance where imaging may be helpful: the presence of Heparin-Induced Thrombocytopenia. Although uncommon, adrenal hemorrhage may be present and this can be documented potentially with CT [36]. MRI may be done to assess for spinal cord trauma or ligamentous injury but it has little role to play in the evaluation of trauma patients with CIRCI overall.

In summary, based on published literature, CIRCI has a bimodal distribution in the trauma setting. The first peak occurs early after trauma (within the first 24-48 hours). Persistent shock in the absence of occult hemorrhage and in spite of adequate volume resuscitation may be the initial clue. The second peak occurs later in the hospital course (usually by the second week). Sepsis is the initiating factor in this phase.

Treatment

Once the diagnosis of CIRCI has been made, the case for treatment should be based on improved outcome. Some authors have argued that non-response to the ACTH stimulation test need not be treated [14]. They also cite the prospective Corticus data showing no difference in outcome between responders and non-responders in the setting of septic shock [43]. Others who advocate no intervention, believe that CIRCI is similar to the Sick Euthyroid Syndrome.

In the setting of trauma, non-response to the ACTH stimulation test has been linked to increased vasopressor requirements, more volume needs and worse outcomes when compared to patients who respond (20,46). Guillemondegui et al., studied 82 ACTH stimulation test non-responders. 66 were treated with steroids and 16 were not. Treatment reduced mortality by 50% [19]. Rivers noted improved survival and shorter time to shock reversal in a surgical cohort of patients with CIRCI treated with hydrocortisone [26]. In the retrospective Corticus study, 477 patients with severe sepsis or septic shock from 20 European ICU’s were studied. An ACTH test was performed on the day of sepsis onset. Non-responders with any baseline cortisol level had a 1.38 odds ratio for mortality. Corticosteroid treatment improved outcome in both univariate and multivariate analysis [42]. Although the strength of some of this data can be challenged, taken together these studies and others suggest that clinical outcome may be improved if biochemically defined CIRCI is treated in the trauma setting. Randomized controlled trials regarding the management of CIRCI in trauma are currently not available. The recommendations regarding treatment below are based largely on observational studies that provides our best evidence at this time.

A stress dosage of hydrocortisone (200-300 mgs/day) for 10-14 day period is usually recommended. During this time, weaning from vasopressors should occur. For those patients with CIRCI and septic shock, mineralocorticoid therapy with fludrocortisone along with hydrocortisone may improve outcome [48]. Routine mineralocorticoid therapy as an adjunct does not seem to be warranted in most instances of CIRCI [3].

Steroid therapy is associated with significant adverse events. Steroid related GI tract ulcerations with bleeding, and superinfections are seen [43]. The ideal length of treatment is unknown. Several authors emphasize that abrupt cessation of steroids is associated with rebound inflammation that can seriously jeopardize recovery. For this reason they recommend a 3-5 day taper [3].

Outcome

Declining responsiveness to ACTH stimulation has been associated with multi-organ dysfunction and worse outcome in CIRCI [49]. This change in ACTH response reflects ongoing HPA dysfunction and not a
shortcoming of the test itself. In this study, renal replacement therapy and thrombocytopenia were risk factors for a decline in delta cortisol. In another small study evaluating cytokine levels in CIRCI, elevated IL-6, IL-10 and TNF levels were detected in non-responders with low (less than 9 micrograms per deciliter) delta cortisol [46]. C-reactive protein was higher in these patients and their overall outcome was worse when compared to patients with normal adrenal function or those with a low baseline cortisol levels. Even after head injured CIRCI patients have been weaned from ventilator support, elevated IL-6 levels have been documented [50]. Although causation is not proven by this data, it certainly suggests that HPA inhibition may occur in the setting of increased inflammatory cytokines and this may impact outcome.

Table 1 describes a summary of several clinical studies spanning a decade on CIRCI in blunt and penetrating trauma. 511 patients are included in this compilation. The average patient age was 41 years. The average ISS was 27 and the baseline cortisol was 17 micrograms per deciliter. Ventilator days were prolonged (average 17) as was length of stay (37 days). The mortality rate for this collected series was 17%.

Recent burn center data for CIRCI, reveals an older patient cohort with a mean age of 51 years and large body surface area of injury (mean 33%). When compared to controls, CIRCI patients had a longer length of stay (mean 66 days vs. 35 days), prolonged ventilator days and a higher mortality (17% vs. 2.5%). Patients with more co-morbidities and inhalation injury were more likely to develop CIRCI [51].

New Horizons

A recent multi-center trial of continuous intravenous hydrocortisone for patients with traumatic brain injury was conducted [52]. 150 multiple trauma patients with cerebral injury were studied. The incidence of pneumonia and ventilator free days were the primary outcome variables. The pneumonia rate was dramatically reduced in the steroid group and ventilator free days were increased. These results were documented for both the CIRCI and non-CIRCI cohorts. The thinking is that the early administration of hydrocortisone blunted the systemic inflammatory response and changed the environment in a way that the patients’ endogenous immune system could fight off potential infection. This takes us back to the concept of a Eucorticoid state. If the inflammatory response is balanced by endogenous or in this case exogenous steroids, survival and decreased morbidity will occur. Of course these results will have to be duplicated, but they remain provocative regarding what we may be able to accomplish to assist in recovery from severe injury.

Conclusion

Critical Illness Corticosteroid Insufficiency remains a significant risk to survival in the trauma setting. Although initially thought to be uncommon, CIRCI is being recognized more frequently with more rigorous testing. It has a bimodal presentation. Early on the severity of the insult and attendant inflammatory response may place the patient at risk. Later in the hospital course, sepsis is usually the initiating event. Serum cortisol monitoring and the ACTH stimulation test are critical studies that should be a part of the diagnostic work-up. Refractory shock should always raise this diagnosis. To avoid unnecessary morbidity and mortality this condition must be recognized quickly and rapidly treated. Stress dose hydrocortisone therapy is usually effective. The ideal duration of therapy remains unclear. CIRCI in trauma must be viewed as a condition of urgency.

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Table 1: Collected Series of CIRCI in Blunt & Penetrating Trauma.

| Year of study | Number of patients | Age | ISS | Mean Cortisol | Stimulation Test | Rx’d Hydrocortisone | LOS | Vent days | Mortality Rate |
|---------------|--------------------|-----|-----|---------------|----------------|-------------------|-----|-----------|----------------|
| Offner et al. [15] | 22 blunt | 30 | 34 | 30 mg/dl | no | no | 49 | 12 | 7% |
| Hoen S et al. [20] | 34 blunt | 34 | 29 | 19 mg/dl | yes | no | 28 | 8 | 12.5% |
| Molina PE [1] | 8 blunt | 47 | 28 | 7.5 mg/dl | yes | yes | 19 | 5 | 13% |
| Bernard F et al. [22] | 113 blunt | 35 | 23 | 10.2 mg/dl | yes | yes | 27 | 13% |
| Rushing GD et al. [16] | 15 blunt | 50 | 23 | 15.8 mg/dl | no | 22 | 19 |
| Fann SA et al. [21] | 63 | 35 | 22 | <20 mg/dl | yes | 66 | 11 | 25% |
| GuillaMondequi OD et al. [19] | 82 | 50 | 31 | <20 mg/dl | yes | 70 | 38 | 34% |
| Walker ML et al. [28] | 70 | 64 | 49 | 23 | 18 mg/dl | yes | 56 | 1 | 10.7% |
| Roquilly A et al. [52] | 113 blunt | 35 | 30 | 20 ug/dl | yes | 313 | 37 | 17 | 16.7% |
| Total | 511 | 41 | 27 | 17.2 mg/dl | no | 313 | 37 | 17 | 16.7% |

Abbreviations: ISS = Injury Severity Score LOS = Length of Stay Vent Days = Ventilator Days
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