Chapter 17
Novel Modalities to Diagnose and Prevent Compartment Syndrome

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Background

Acute compartment syndrome (ACS) of an extremity following trauma remains a diagnostic and therapeutic enigma. This phenomenon is because the clinical signs and symptoms of ACS are variable and nonspecific, and there are multiple causes of ACS after trauma (Table 17.1). Furthermore, the pathophysiology of ACS is poorly understood, which leads to confusion regarding the ideal diagnostic strategy. For decades now, ACS has been considered to be a pressure-related pathophysiologic

| Causes of acute compartment syndrome after trauma |
|--------------------------------------------------|
| 1. Accumulation of mass within the muscle compartment |
| Bleeding |
| Edema |
| Tissue injury caused by direct trauma |
| Inflammation and/or toxins |
| Postischemic reperfusion |
| Insertion of internal fixation hardware |
| 2. Reduction of compartment volume |
| External compression from bandages, splints, casts |
| Changes in the limb length related to fracture treatment |
| External fixation |
| Traction |
| Internal fixation |

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process, with diagnostic recommendations largely based on pressure thresholds – either the absolute intracompartment pressure (ICP) or some measure of the perfusion pressure (defined as the difference between ICP and the patient’s blood pressure). However, defining a single pressure threshold that accurately defines when fasciotomy is needed has not proven possible, and multiple studies suggest that clinical use of such thresholds leads to overtreatment [1–3].

Current treatment of ACS is surgical fasciotomy, which if not performed before the onset of cellular necrosis results in intractable pain, muscle fibrosis and contracture, and sensory deficits. Fasciotomy restores perfusion by increasing compartment volume so that the ICP falls, allowing tissue perfusion to be restored in viable vascular beds. This benefit is consistent with the traditional view of ACS as a “pressure problem.” However, it is more appropriate to consider ACS as a metabolic problem, and both diagnostic and therapeutic strategies aimed at assessing cellular viability and reversing metabolic abnormalities may not only make our diagnosis of ACS more precise but also provide opportunities for less invasive ways to prevent ACS and mitigate the sequelae of ACS when it does occur.

New Concepts in ACS Pathophysiology

Compartment syndrome has been considered to be the result of tissue ischemia caused by decreased perfusion of the limb from elevated ICP, which may occur from any one of the several mechanisms: arterial spasm, collapse of arterioles, or collapse of the venous system. However, a recent series of publications by researchers in London, Ontario, Canada, has shown that ACS is accompanied by both a local and a systemic inflammatory response that may play a significant role in the pathogenesis of ACS and the resultant tissue damage [4–9]. This knowledge “opens the door” to using inflammatory mediators as potential biomarkers for understanding the progression or resolution of ACS and anti-inflammatory therapies as either primary or adjunctive medical therapy for ACS.

In a series of animal studies [4–9], the relationship between elevated ICP and the inflammatory response on skeletal muscle microcirculation and cell viability was measured in rats using intravital videomicroscopy. In one study, the level of tissue injury in an experimental compartment syndrome was compared between normal rats and those with neutropenia induced by injection of high-dose cyclophosphamide [5]. Muscle cellular injury was reduced more than 50% in neutropenic compared to healthy rats. Furthermore, the control rats had higher tissue injury (23.0% ± 4.0% of cells) compared to the neutropenic group (7.0% ± 1.0% of cells) after 90 minutes of elevated ICP (p = 0.00005) [5].

In a second study using a similar experimental model, local and systemic cytokine activation was measured and showed that animals with ACS have a systemic inflammatory response in addition to the local injury [4]. Remote organ damage was measured in the liver using intravital videomicroscopy techniques. Leukocyte activation, increased serum TNF-α levels, and necrotic hepatocytes were noted after 2 hours of ACS, all indicating systemic effects from ACS induced in one limb.
These same researchers have also demonstrated that attenuation of the inflammatory response may be beneficial in cases of ACS. Manjoo et al. [9] studied the effect of indomethacin on the level of tissue injury measured in rats with induced elevations in ICP for 45 or 90 minutes. Tissue injury was reduced almost to baseline levels in rats that were pretreated with indomethacin; a smaller benefit was noted when indomethacin administration was delayed [9]. In another study using the same model, treatment with a different anti-inflammatory molecule (CO-releasing molecule 3) led to statistically significant improvement in the number of continuously perfused capillaries, resulted in decreased tissue injury, reversed the ACS-associated increase in TNF-α, and reduced leukocyte adherence [6].

This series of papers indicate that ACS is accompanied by a substantial inflammatory response in the involved limb that also affects other organ systems. This information suggests that mitigation of the tissue injury caused by ACS may be possible by decreasing inflammation. Further work to determine the benefits of inflammatory modulation in a larger animal model is necessary, followed by clinical research in humans. The ideal therapeutic agent, mode of delivery, and timing are all unknown questions at this time.

**New Diagnostic Modalities**

The traditional diagnosis of ACS has been based on the clinical examination, supplemented when needed by intracompartment pressure measurement. Newer diagnostic approaches to ACS include near-infrared spectroscopy (NIRS), biomarker analysis, and radiofrequency identification (RFID) implants. These new approaches are discussed in the following paragraphs.

**NIRS**

In the context of measuring tissue perfusion in possible cases of ACS, NIRS uses differential light reflection and absorption to estimate the proportion of hemoglobin saturated with oxygen approximately 2–3 cm below the surface of the skin [10–14]. The depth of tissue interrogation is determined by the distance between the light source and the receptor.

The potential use of NIRS to diagnose ACS is supported by preclinical studies. In 1999, Garr et al. induced compartment syndrome in anesthetized pigs and showed that NIRS was a better predictor of neuromuscular dysfunction than compartment perfusion pressure [10]. However, its clinical use has shown mixed success. A case report published in 2011 outlined the benefits of using NIRS in three cases of lower extremity ACS [13]. NIRS differentiated between adequately perfused lateral compartments and poorly perfused deep posterior compartments of one patient with ACS. In another unresponsive patient diagnosed with ACS on clinical means, NIRS was able to detect perfusion deficits. In a third patient, NIRS demonstrated changes
in muscle perfusion due to anesthetic induction within seconds [13]. However, also in 2011, Bariteau et al. described seven patients with a clinical diagnosis of ACS who had oxyhemoglobin saturation (rSO2) and ICP values measured in each compartment of the affected lower extremity before undergoing fasciotomy. No statistically significant association was observed between rSO2 and ICP or perfusion pressure [11].

Many questions remain about the clinical use of NIRS in traumatized limbs for diagnosing ACS, such as the ideal penetration depth, the effect of skin pigmentation and subcutaneous fat, the effect of skin abrasion or degloving, and subcutaneous or intramuscular bleeding. It has been demonstrated that injured limbs have a hyperemic response to injury, resulting in increased rSO2 values [14].

One large theoretic advantage of NIRS is its potential ability to demonstrate changes in tissue perfusion corresponding with the onset of ACS. However, continuous NIRS has not been as well studied. Two recent papers have reported on continuous NIRS in a clinical setting. Shuler et al. [15] reported on 109 patients who had NIRS recording of the tissue perfusion in all four leg compartments of both legs (anterior, lateral, superficial posterior, and deep posterior): 86 had unilateral leg injuries while 23 did not. Mean NIRS values were between 72% and 78% in injured legs, between 69% and 72% in uninjured legs, and between 71% and 73% in bilaterally uninjured legs. NIRS values were typically >3% higher in injured limbs without ACS than in uninjured compartments. In contrast, all seven limbs with clinically diagnosed ACS had at least one compartment where NIRS values were 3% or more below an uninjured control compartment [15].

One problem with NIRS that may be even more of an issue when using NIRS in a continuous manner is the reliability of data capture. Shuler et al. noted that “Missing data were encountered in many instances” [15]. Schmidt et al. [16] used blinded continuous NIRS and blinded continuous ICP monitoring in 191 patients with leg injuries [16]. Data capture was unreliable with NIRS, with simultaneous data available from both injured and control limbs just 9% of the expected time, compared to 88% of the expected time with continuous ICP [16]. At this time, it is not clear whether these data-capture issues with NIRS represent clinically immature technology that needs further development for this purpose or represent fundamental problems using NIRS to measure soft tissue oxygenation when that same tissue has had traumatic injury [16].

**Biomarkers**

Markers of muscle injury or metabolic disturbances such as acidosis may be measured systemically or locally.

Odland et al. have applied tissue ultrafiltration (TUF) catheters to the problem of diagnosing and treating acute compartment syndrome [17, 18]. TUF uses small diameter, flexible, hollow fiber catheters connected to suction in order to remove interstitial fluid, which may contain biomarkers of injury and contribute volume to the muscle compartment. The possible therapeutic benefits of TUF are discussed later in this chapter. In their study, serum and ultrafiltrate levels of creatine kinase
and lactate dehydrogenase were measured hourly. The study demonstrated that biomarker concentrations were up to 80-fold higher in ultrafiltrate than in serum.

A promising metabolic marker for ACS is tissue pH, as measured by intracompartment probes. Basic science studies done at the University of Aberdeen in Scotland have revealed a correlation between local pH and depletion of high-energy phosphate stores [personal communication, Alan Johnstone 2018]. These same researchers have demonstrated that in a small study continuous pH measurement may be better than continuous pressure measurement at diagnosing ACS [19]. Further research is ongoing that should better define the role of pH monitoring on this setting.

**RFID Chips in Diagnosing Compartment Syndrome**

Current methods of diagnosing ACS rely on catheters or sensors that must remain attached to the patient and are therefore subject to inadvertent removal or may interfere with aspects of patient care, such as transportation, imaging, splinting, or the performance of surgical procedures. An “always-on” device that can be applied in a minimally or noninvasive manner and that does not interfere with transportation or other care of the patient would be a significant advance. New sensor technology that utilizes radiofrequency electromagnetic fields (radiofrequency ID tags or RFIDs) to transfer data exists and is beginning to be used for monitoring physiological systems [20]. Such devices do require a battery and are powered externally by the very fields used to read their data. RFID pressure sensors have been developed and used for compartment monitoring [20].

**Potential Therapeutic Advances in Treating ACS**

In at-risk or early phases of compartment syndrome, before tissue necrosis has started to occur, other means of improving tissue perfusion and oxygenation may exist. One method would be to increase the tolerance of the muscle to ischemia. Other areas of theoretical interest include methods to decrease intramuscular pressure (tissue ultrafiltration, foot pumps, mannitol, and diuretics), improving tissue oxygenation, mitigating the effects of ischemia (free-radical scavengers and other pharmacologic measures), and small-volume resuscitation with hypertonic saline.

**Tissue Ultrafiltration**

Compartment syndrome is associated with accumulation of mass within a muscle compartment of relatively fixed volume, which is what results in the progressive increase in intramuscular pressure, which eventually sets in motion the vascular
embarrassment and associated metabolic changes that result in tissue death. By removing even small amounts of fluid, loss of the associated fluid mass within the compartment may decrease compartment pressures. In an animal study, the potential benefits of tissue ultrafiltration (TUF) were demonstrated [18]. A bilateral infusion model of compartment syndrome in the pig hindlimb was used; each animal had one control limb and one treatment limb. Muscle pressure was measured with an indwelling catheter. Three TUF catheters were inserted into the anterior compartment of each limb. The catheters in the treatment limb were connected to negative pressure suction. The ICP pressure was lower in the treated limbs, and the muscle histology was more normal in the treated limbs compared to the control legs [18]. This study was followed up with clinical research on TUF in a small human trial. A pilot study demonstrated the feasibility of TUF in ten patients with tibial shaft fractures treated with nailing [17]. A small-randomized clinical trial of TUF versus pressure monitoring alone was also performed [21]. Fourteen patients were randomized to two groups: one used TUF plus monitoring while the other was monitored alone. Anterior and deep posterior pressures were measured for 24 hours in all patients. Outcome measures between the two catheters were compared using T-tests. The mean intramuscular pressure was smaller than in the control group in both the anterior and posterior compartments. Because of insufficient power, these differences were not statistically significant. Interestingly, two patients in the control catheter group developed compartment syndrome – 8 and 16 hours after the initial surgery and catheter insertion. No compartment syndromes were diagnosed in the treatment group.

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**Hypothermia**

Tissue cooling has been used to improve tissue viability in transplant and replant surgery and is of known value in treating soft tissue injury. Sanders and colleagues, using a well-developed rat model (as described in the “New Concepts in ACS Pathophysiology” section of this chapter), showed that hypothermia to 25 °C reduced tissue damage score by 50% and also seemed to attenuate the inflammatory response [22]. Although this information is interesting, at this time the role of induced hypothermia for treating ACS in humans is not defined.

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**Conclusion**

New concepts and ideas about the pathophysiology of ACS show promise at developing more physiologic, targeted treatments. These developments include modulation of the inflammatory response and new diagnostic modalities. Moreover, these advancements all address the current shortcomings in the diagnostic armamentarium that trauma surgeons can use, and which should lead to more precise methods
to diagnose ACS and less invasive ways to prevent and treat it. Many of the approaches discussed in this chapter may be used simultaneously, and a multimodality approach to diagnosis and therapy, for example, using tissue ultrafiltration, intermittent plantar compression, small-volume resuscitation, and pharmacologic therapy, may someday be able to supplant fasciotomy for many patients.

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