Epinephrine auto-injection radically increases risk for clostridial infection and necrotizing fasciitis

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Abstract

Clostridial perfringens is a bacteria commonly found on skin flora. Due to the optimal growth environment intramuscular epinephrine injections predispose patients to the rapid development of clostridial myonecrosis. There have been only four cases, including this one, reported in the last 60 years of pediatric Clostridium perfringens infections post-epinephrine injection. We detail the successful management of a 16 year old, immunocompetent female who developed gas gangrene and necrotizing fasciitis on her thigh secondary to clostridial infection after utilization of an Epinephrine Auto-Injector and review the pediatric literature of patients with Clostridial perfringens secondary to epinephrine injection. We define common clinical signs and symptoms of clostridial infection from the review of the literature. The relevance of our findings is to raise awareness among emergency physicians when patients present following an injection in order to reduce diagnostic delay that could result in amputation or death.

Introduction

Necrotizing fasciitis (NF) is within the general category of necrotizing soft tissue infection (NSTI). There are three variants of NSTI: i) NF, ii) necrotizing cellulitis, and iii) myonecrosis. However, NF is often used in clinical settings as a broadly inclusive term for overlapping types of NSTI, thus for the purposes of this paper, the authors will use the broadly inclusive term NF. Clostridial myonecrosis is an infection primarily of muscle tissue, of which 90 percent are Clostridium perfringens.1 Clostridial myonecrosis or gas gangrene is a highly lethal necrotizing soft tissue infection of skeletal muscle caused by toxin- and gas-producing Clostridium species.2

Currently, 90% of contaminated wounds demonstrate clostridial organisms, but less than 2% develop clostridial myonecrosis.2 This emphasizes the significance of both host and local wound elements in the development of this process, rather than the mere incidence of the organisms in the wound. For this reason, there have been only four cases,1-4 including this one, reported in the last 60 years of pediatric Clostridium perfringens infections secondary to epinephrine injection. This review of the literature describes these cases, as well as, a recent case detailing the manifestation of gas gangrene and necrotizing fasciitis caused by Clostridium perfringens.

To the best of our knowledge, this is the first reported pediatric case in the last 35 years. We describe risk factors related to the epinephrine injection that contributed to the development of Clostridia infection, as well as antibiotic therapy, pain medications, surgical debridement, and wound vacuum assisted closure (VAC) therapy management.

Case Report

After a consult at a Pediatric Urgent Care (Day 0), the 16-year-old female was transferred to the Emergency Department (ED) with suspected Compartment Syndrome. Approximately 24 hours prior, she reported administering an [Epinephrine AutoInjector (EAI)]; Brand: EpiPen, dose: (0.3mg)] due to an allergic reaction with the symptoms of hives and difficulty breathing. Upon presentation to the ED, no clinical signs of anaphylaxis were evident.

Upon arrival (Day 0), patient presented with tenderness and mild swelling of the left thigh and the initial vital signs were: [Heart Rate (HR)] 120bpm; [Blood Pressure (BP)] 100/55mmHg; Respiratory rate 18bpm; Temperature 98.8°F. The emergency physician prescribed fluids, oral and intravenous (IV) pain medication. Within 5 hours she demonstrated tachycardia (HR 140’s) with an increase in temperature to 101°F (a further increase may have been masked by the administered analgesics/antipyretics). Despite IV pain medication, the patients’ pain continued to increase. The initial lab tests demonstrated the patient had leukocytosis with a white blood cell count of 20.2 thousand/ μL. The red blood cell count (4.73 million cells/ μL), the hemoglobin (13.8 g/dL) and hematocrit (42.6%) were all normal. The patient was followed for hematologic issues. Blood cultures taken upon arrival later demonstrated no apparent growth. An ultrasound verified a non-specific edema within the soft tissue of the affected area, with no drainable fluid. (Day 1) Patient was administered [non-steroidal anti-inflammatory drugs (NSAIDs)] for inflammation, IV broad spectrum antibiotics to contain infection and analgesics to control for the increasing pain. The patient was preliminarily diagnosed with bacterial myositis and sepsis (Day 1). Once admitted the pediatric physician became concerned due to the rapid progression of the symptoms and consulted a surgeon for evaluation. The surgeon ordered an immediate MRI which demonstrated extensive gas in the quadriceps muscles, (especially in the vastus lateralis and adductor muscles) along with extensive edema and enhancement. These findings were concerning for an infection with a gas-forming organism. The patient went to the operating room (Day 1) for leg incision, drainage, and skin debridement (Figure 1A). Post-operative pathology showed necrotic skeletal muscle with acute inflammation. Clostridium perfringens was identified in the three specimens (Table 1, for a procedure overview).

On day 3, she underwent a second surgery for leg irrigation, further debridement, and placement of Wound VAC. Broad-spec-
trum antibiotics were adjusted in accordance with lab susceptibility testing results. Blood cultures indicated no growth. The patient continued on fluids, targeted antibiotic therapy and [patient-controlled analgesia (PCA)] for pain control. Patient started having muscle spasms in the affected leg; consequently diazepam was added to medications.

On day 15, the patients’ main complaint was pain, PCA support was continued with antibiotic therapy. By day 20, blood and urine cultures obtained were negative. Patient underwent fourth surgical procedure on day 23 for [split thickness skin grafting (STSG)] with leg debridement and wound vac closure (Figure 1B). On day 26, patient moved from Pediatric Intensive Care Unit to the floor and on day 28 Wound VAC takedown with transition dressing changes that include topical antiseptic and antibiotic ointments. Antibiotic therapy included cefepime as an intravenous piggy back, urine was positive for gram negative rods (low level contaminated), blood cultures negative. Patient was discharged home after a LOS of 34 days. The patient continued physical therapy and follow up in the outpatient wound clinic (Figure 1C) for 5 months and successfully returned to physical activities. The patients’ final diagnosis was necrotizing fasciitis, gas gangrene caused by *Clostridium perfringens*, with infective myositis of the left thigh.

According to EAI guidelines, patients are to go to the ED immediately after injection. However, this patient waited 24 hours before being admitted into the ED for suspected compartment syndrome. Initially, an infected EAI was suspected to be the culprit for contamination of *Clostridial* spores, clarification testing of the EAI was not performed.

**Patients and methods**

A review of the literature using the PubMed database for gas gangrene, myonecrosis and adrenaline in which subjects were ≤18 years old, was performed. The references of qualifying articles were examined to identify other articles. Articles with patients older than 18 years old, gas gangrene or myonecrosis attributed to another bacterial species were excluded from this study. The following data was collected: sex, age, site of injection, time post-injection of hospital admittance, signs/symptoms shown upon admittance, bacterial species, medications given, surgical procedures used and outcome. The study was submitted to the Mercy Institutional Ethics Review Board and received expedited approval.

**Results**

In this review of the published literature, we found three pediatric patients (≤18 years old) with *C. perfringens* infections secondary to epinephrine injections, with only one patient surviving (Table 2). Commonly, all were healthy females with similar presenting symptoms most notably; persistent severe pain, fever, inflammation, elevated heart rate, low blood pressure, and feeling of malaise. Of those reported, the initial predominant symptom was pain at the injection site, starting between 6-12 hours post-injection.

**Discussion**

Classically, clostridial myonecrosis

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**Table 1. Patient treatment timeline.**

<table>
<thead>
<tr>
<th>Day</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Initial surgical debridement</td>
</tr>
<tr>
<td>3</td>
<td>Wound irrigation, secondary debridement and wound VAC</td>
</tr>
<tr>
<td>8</td>
<td>Further wound irrigation, application of dermal repair scaffold dressing and wound VAC</td>
</tr>
<tr>
<td>23</td>
<td>Final debridement, STSG and wound VAC take down</td>
</tr>
</tbody>
</table>

VAC, vacuum assisted closure; STSG, split thickness skin grafting. Patient underwent four surgical procedures within 23 days.

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Figure 1. Photographs representing course of treatment: day 1, intra-operative photo of left thigh post surgical debridement (A); day 23, fresh STSG of left thigh (B); day 54, healed left thigh (C).
Table 2. Pediatric clostridium infections following epinephrine injections.

<table>
<thead>
<tr>
<th>Author</th>
<th>Sex</th>
<th>Age (year)</th>
<th>Site of injection</th>
<th>Time post-injection symptoms of hospital admittance</th>
<th>Signs/</th>
<th>Species</th>
<th>Medications</th>
<th>Surgery</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present case</td>
<td>♀</td>
<td>16</td>
<td>Thigh</td>
<td>24 hours</td>
<td>P,F,H,G,B, I, muscle spasm</td>
<td>C. perfringens</td>
<td>Piperacillin/ taxobactam, cefepime</td>
<td>Yes</td>
<td>Alive</td>
</tr>
<tr>
<td>Gaylis (1968)</td>
<td>♀</td>
<td>18</td>
<td>Thigh</td>
<td>3 days</td>
<td>P,F,H,G,M, I,C,BP</td>
<td>C. welchii*</td>
<td>penicillin</td>
<td>Yes</td>
<td>Expired</td>
</tr>
</tbody>
</table>

P: pain; I, inflammation/swelling; F, fever; BP, low blood pressure; HR, elevated heart rate; M, malaise/bli; C, crepitus; G, gas gangrene. *C. perfringens was formerly known as C. welchii.

References


