Toxic shock syndrome toxin-I producing *Staphylococcus aureus* in fulminant necrotizing fasciitis

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**Abstract**

We describe a patient presenting with fulminant monomicrobial necrotizing fasciitis (NF) and toxic shock syndrome (TSS) caused by *Staphylococcus aureus* producing toxic shock syndrome toxin-I (TSST-I). To our knowledge, TSST-I positivity combined with NF has been described only once in previous literature. We discuss the clinical and pathophysiological aspects of both necrotizing fasciitis and toxic shock syndrome and provide a comprehensive overview of causative micro-organisms and the different toxins they produce. We conclude that intravenous immunoglobulin (IVIG) treatment might be considered not only in streptococcal but also in staphylococcal NF.

**Case**

A 67-year old man was admitted to our Intensive Care Unit (ICU) with septic shock. Five days earlier he presented to the Emergency Department with a mild fever (up to 38 degrees Celsius) and paravertebral back pain. An abdominal ultrasound revealed no pathologic findings (no abdominal aneurysm, gall stones or pancreatitis). His symptoms were attributed to muscular pain possibly caused by influenza (endemic at the time) and he was discharged.

Over the course of 5 days he deteriorated, with severe and progressive back pain with radiation to both legs (which caused an inability to walk) and high fever (up to 41 degrees Celsius). On examination he was tachycardic (heart rate 130 bpm), hypotensive (initial systolic BP 70 mmHg, 99 mmHg after 3 L fluid resuscitation). He was alert with a respiratory rate of 20/min and oxygen saturation results showed a slight leucopenia (3.8 × 10⁹/L), C-reactive protein 360 mg/L, creatinine 280 µmol/L and lactate 5.4 mmol/L. Blood cultures were drawn, the patient received antibiotics (ceftriaxone 2000 mg according to local protocol) and was admitted to the ICU for further treatment. On repeated examination he developed a discolouration of the skin on the dorsal side of his left thigh extending to just below the knee. It was blue/purple with surrounding erythema and mildly tender on palpation. A CT-scan was performed, showing induration of skin and subcutaneous tissue in the left thigh. No abdominal or retroperitoneal pathology was detected. Because necrotizing fasciitis was suspected, the patient was started on penicillin, clindamycin, gentamicin, intravenous immunoglobulins (IVIG) and underwent emergency surgical exploration. The subcutaneous tissue and fascia were swollen and excisional biopsies showed extensive presence of gram positive cocci. The diagnosis of necrotizing fasciitis was confirmed. Extensive surgical debridement of skin, subcutaneous tissue and muscle fascia of the dorsum of the left upper leg was performed. The patient developed severe septic shock with multi-organ failure and required high dose norepinephrine, vasopressin and steroid treatment. His condition slightly improved after surgery, antibiotics and IVIG administration. However, soon after the initial stabilisation he developed new skin lesions on his right leg for which he underwent a second surgical procedure. During surgery, similar findings to those in the left leg necessitated another debridement. Upon return to the ICU his condition had deteriorated further, with persistent anuria, severe metabolic acidosis with lactate levels of more than 15 mmol/L and cardiac failure. At that point, further treatment was considered futile and the patient died immediately after withdrawal of treatment. Methicillin sensitive *Staphylococcus aureus* was found in blood cultures, urine cultures and tissue cultures. Toxic shock syndrome toxin 1 gene was detected by polymerase chain reaction (PCR) in the *Staphylococcus aureus* strains isolated. Genes for other staphylococcal toxins (enterotoxin A-D, exfoliative toxin A/B and Panton-Valentine leucococidin) were not present. Influenza testing had not been performed.

**Background**

Necrotizing fasciitis caused by a single microorganism is reported in the minority of cases. More specifically, necrotizing fasciitis due to *Staphylococcus aureus* as the only pathogen is not often described in...
Necrotizing fasciitis

Necrotizing fasciitis (NF) is an infectious disease involving subcutaneous tissue and muscle fascia. It is characterised by fever, tenderness, swelling and erythema. Pain disproportionate to the examination findings is frequently described. Other findings that may develop later are numbness of the skin, crepitus and skin discoloration and/or necrosis. These late features are quite specific but absent in more than half the cases [1,4]. Diagnosis can be difficult as symptoms may be atypical or absent; a patient may present with profound shock without skin abnormalities. A high index of suspicion is essential. The disease typically has a rapidly progressive course with extension of infection and development of symptoms of systemic toxicity [2].

Necrotizing fasciitis can be divided into categories based on the microbial etiology (Table 1). Four types have been described [1,2,4-6]. Type 1 NF is polymicrobial, usually with at least one anaerobic bacterium (e.g. *Bacteroides*, *Peptostreptococcus*, *Fusobacterium*, or *Clostridium* species) in combination with a facultative anaerobic bacterium (usually non-group A streptococci or enterobacteria such as *E. coli*, *Klebsiella* or *Proteus* species). Usually 3 or more microorganisms are involved. When located in the scrotal/perineal area this type of necrotizing fasciitis is known as Fournier gangrene. Risk factors especially apply for type 1 NF and include age of more than 50 years, diabetes mellitus, peripheral vascular disease, HIV, alcohol abuse, and poor nutritional status [1,7].

Type 2 NF is usually monomicrobial and involves group A hemolytic streptococci. However combinations with *Staphylococcus* are described [1,2]. This type usually affects younger patients without comorbidities. More than in type 1 NF there is a history of trauma or surgery [1].

Type 3 NF is caused by *Vibrio vulnificans* (particularly associated with chronic liver disease) or *Aeromonas* species [8]. A fourth type has been described, caused by fungal organisms (*Candida, Zygomycetes*) [1,2,5,6,9]. In fact type 3 and 4 are also monomicrobial and could be considered a subcategory of type 2. Most cases of NF (55-80%) are polymicrobial with anaerobic presence in approximately 25% of cases [1,6,10].

The use of imaging studies in diagnosing NF is limited. CT might show fascial thickening, oedema, subcutaneous gas and abscess formation, but these findings are not helpful in differentiating between NF and other soft tissue infections. It could be useful if other causes for shock are considered but the clinician should be aware that surgery should not be delayed [2,4].

NF is a surgical emergency. Early surgical exploration is cornerstone in both confirming the diagnosis and in management. Macroscopic findings are generally sufficient to confirm the diagnosis and include swollen, non-vital, gray necrotic tissue, lack of bleeding due to thrombosed vessels and foul-smelling (‘dishwater’) discharge [1]. When macroscopic findings are inconclusive, frozen section procedure is necessary to confirm diagnosis of NF. This is based on histological findings which include necrosis of fascial tissue, polymorphonuclear leucocyte infiltration and thrombosis in arteries and veins of the fascia. Deep fascial biopsy for Gram-staining and cultures will usually reveal the pathogen involved (82% of cases) [10].

When NF is confirmed, aggressive surgical debridement and fasciectomy should be performed. Re-exploration or second-look surgery is recommended within 24 to 48 hours to assess progression of infection and the need for further debridement. On average, 3-4 operative procedures are needed [2,10].

Treatment further consists of supportive care and empirical broad-spectrum antimicrobial treatment, which should cover Gram-positive, Gram-negative and anaerobic pathogens. Clindamycin should be added for both synergism and the benefit of lowering toxin production. This has been proven only in group A streptococcal NF, and failure to add clindamycin to the antimicrobial treatment has shown to increase mortality from 14% to 60% in patients with group A streptococcal disease [6].

Coverage of methicillin resistant *S. aureus* (MRSA) should depend on local incidence [2]. Intravenous immunoglobulin (IVIG) administration is also recommended for neutralising or binding of superantigens and toxins in group A streptococcal disease [2].

Mortality has been shown to vary greatly. Between 1 death per 3 to 5 patients (21% [10], 25-35% [1], 29% [13]) to more varying percentages ranging from low to very high (6-76% [7]) are found in the literature. A recent review found average rates between 20 and 40% [2]. Patients either die in an early stage due to refractory shock and/or ARDS or succumb to multiple organ failure later on.

The time between admission and surgery is strongly associated with mortality, with cumulative survival decreasing with increasing time between admission and surgery [1,2,6,10]. Mortality risk is further affected by both localisation of the infection and underlying conditions of the patient. Perineal and cervical localisation are associated with a higher mortality [14], and immunocompromised patients have a 2-fold higher risk of dying in the course of NF [2].

The pathogens involved also influence mortality, although only clostridial and fungal pathogens are consequently described to lead to increased mortality; for group A streptococcal NF data are inconsistent although most describe a better prognosis in type I than in type II NF [1,6,7].

Table 1. Necrotizing fasciitis divided into categories based on the microbial etiology [1,4-6,8,10-12].

<table>
<thead>
<tr>
<th>Type</th>
<th>Polymicrobial</th>
<th>40%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type I</td>
<td><em>Aerobes (gram +)</em></td>
<td>40%</td>
</tr>
<tr>
<td></td>
<td><em>Streptococci</em></td>
<td>9-42%</td>
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<tr>
<td></td>
<td><em>S. aureus</em></td>
<td>5-36%</td>
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<tr>
<td></td>
<td><em>Enterococci</em></td>
<td>17-20%</td>
</tr>
<tr>
<td></td>
<td><em>Aerobes (gram -)</em></td>
<td>43%</td>
</tr>
<tr>
<td></td>
<td><em>E. coli</em></td>
<td>7-16%</td>
</tr>
<tr>
<td></td>
<td><em>Klebsiella</em></td>
<td>4-23%</td>
</tr>
<tr>
<td></td>
<td><em>Pseudomonas</em></td>
<td>10-14%</td>
</tr>
<tr>
<td></td>
<td><em>Anaerobes (Bacteroides, Peptostreptococcus, Clostridium)</em></td>
<td>7-18%</td>
</tr>
</tbody>
</table>

| Type II | Haemolytic streptococci (group A-G) | 22-40% |
|         | *Staphylocoeci* | 13-20% |
|         | *E. coli* | 2-22% |

| Type III | *Vibrio vulnificans* | 50% |
|          | *Aeromonas* | 4-17% |

| Type IV | *Candida* | 5% |
|         | *Zygomycetes* | ? |

* Population of 61 Taiwanese patients [8]
† Case reports
Toxic shock syndrome

Toxic shock syndrome (TSS) is an acute systemic illness caused by specific bacterial toxins. These toxins act as superantigens and interact with antigen-presenting cells by binding to class II Major Histocompatibility Complex (MHC) molecules, and with T-cells by binding to specific variable regions on the β-chain of the T-cell antigen receptor. This induces T-cell activation in up to 5-30% of the patients T-cells, which is much more than in normal T-cell response (when less than 0.01% of T-cells are activated [15,16]). Furthermore the superantigens cause production of cytokines such as TNF-α, TNF-β and several interleukins. This leads to the development of capillary leak and shock [16-19].

There are several bacteria capable of producing toxic shock causing toxins, with streptococci and staphylococci forming the largest groups. The superantigens causing TSS are known as pyrogenic toxin superantigens and are produced by either streptococci or staphylococci [15].

Streptococci are capable of producing exotoxins (e.g. streptococcal pyrogenic exotoxin (SPE) serotype A produced by group A beta-haemolytic streptococci). This results in a syndrome known as streptococcal toxic shock syndrome or toxic shock like syndrome. Other SPE serotypes include B, C, F, G, H and J. Due to massive T cell activation by the superantigen, shock can be rapidly progressive, and this syndrome is associated with mortality rates as high as 50 to 80% [16-18]. The incidence of TSS in group A streptococcal disease is unclear. Symptoms consistent with TSS in patients with group A streptococcal disease have been reported to occur in up to 30-50% [16,18] which seems to be an overestimation. For patients with NF, presence of streptococcal TSS increases mortality considerably from less than 40% to 67% [6].

Staphylococci also are capable of producing exotoxins. They include TSST-1, Staphylococcal Enterotoxins (SEs) serotype A, B, Cn, D, E, I [18], G and H [19], leucocidin and exfoliative toxins A and B. Most SEs and TSST-1 are the superantigens that can cause staphylococcal TSS.

Staphylococcal TSS can be divided into menstrual and non-menstrual TSS, with the menstrual type being associated with tampon use that facilitates TSST-1 production by introducing oxygen [18]. The incidence of menstrual TSS is declining. Nowadays, approximately one third to half of TSS cases are non-menstrual, with the clinical setting varying from postoperative wound infections, mastitis, sinusitis, osteomyelitis, arthritis, burns and post-influenza respiratory infections [18,20]. While the majority of S. aureus in menstrual TSS (93%) produces TSST-1, this toxin was present in only 40-63% of non-menstrual TSS strains [21,22].

Staphylococcal TSS is characterised by sudden onset of fever, rash, gastrointestinal complaints (nausea, vomiting, diarrhoea and abdominal pain) and capillary leak with hypotension due to vasodilatation, possibly leading to multiple organ failure [19]. Case fatality rates for both menstrual and non-menstrual staphylococcal TSS are by far not as high as for streptococcal TSS [17,18]. Menstrual TSS (i.e. predominantly TSST-1 mediated) disease has a slightly lower mortality rate than non-menstrual TSS. These findings suggest a relatively low T cell activation potency for TSST-1 compared to the other toxins involved in staphylococcal TSS (enterotoxin B and C) and the exotoxins involved in streptococcal TSS [19].

Discussion

Our patient developed profound septic shock and, despite aggressive surgical and antibiotic treatment and maximum supportive care, died shortly after the diagnosis of NF was made. Staphylococcus aureus was cultured from blood, tissue and urine. No other pathogens were found. Presence of TSST-1 was confirmed post-mortem. It is possible he developed staphylococcal TSS as a complication of influenza, but this remains unknown since an influenza PCR was not performed.

Little is known about the incidence of staphylococcal TSS in staphylococcal NF. To our knowledge, only one case report has been previously published that describes a patient with a (monomicrobial) NF and TSST-1 mediated staphylococcal TSS [3].

It is reasonable to assume that presence of staphylococcal superantigens and thus staphylococcal TSS worsens prognosis in NF similar to streptococcal NF.

Treatment with IVIG is usually recommended in streptococcal disease. IVIG neutralize the superantigens, decrease cytokine production and have been shown to reduce mortality in streptococcal TSS [18,19].

It has been shown however that IVIG also have activity against TSST-1 produced by staphylococci [22]. Patients with staphylococcal TSS might therefore also benefit from treatment with IVIG. Our patient showed a slight improvement after IVIG and might be an example of this.

Conclusion

We report the case of a 67-year-old male who died of profound shock and multi organ failure due to necrotizing fasciitis in combination with TSST-1 mediated staphylococcal toxic shock syndrome. This has only been described once previously in literature. Both NF and TSS are known for progressing rapidly; high clinical suspicion index and aggressive surgical management in combination with antibiotic treatment and supportive care are essential. Given the mechanisms of action it is reasonable to assume IVIG have a role in the treatment of patients with staphylococcal TSS. With the rising incidence of MRSA related infections [1,6,11,24] the role of IVIG might turn out to be even more important in controlling these infections.

References


22. Schlievert PM (1986) Staphylococcal enterotoxin B and toxic-shock syndrome toxin-1 are significantly associated with non-menstrual TSS. Lancet 1: 1149-1150. [Crossref]
