Lesson of the month 2: Toxic shock syndrome

Authors: Tamer Shalaby, Samantha Anandappa, Nicholas John Pocock, Alexander Keough and Angus Turner

Toxic shock syndrome (TSS) represents a fascinating example of immune activation caused by infection resulting in a dramatic and challenging clinical syndrome. TSS is commonly associated with tampon use and still causes significant morbidity and mortality in young healthy women. A misconception is that TSS presents with a skin rash and only occurs in women and children; however, it can occur in males and can present without skin changes. TSS presents initially as a febrile illness and within a few hours can progress to severe hypotension and multiple organ failure (MOF). Staphylococcus aureus and group A beta haemolytic streptococcus (GABHS) can secrete toxins from a small or hidden focus of infection and hence blood culture and sensitivity (C+S) tests can be negative, thereby making diagnosing this condition challenging. Clindamycin is superior to penicillin in the treatment of this condition and significantly decreases the mortality rate in TSS. However, there is also an important role for intravenous immunoglobulins (IVIG). Early intensive care unit (ICU) as well as surgical team involvement (in selected cases) is required to avoid mortality which may approach 70%.

KEYWORDS: Toxic shock syndrome (TSS), group A beta haemolytic streptococcus (GABHS), Staphylococcus aureus, necrotising fasciitis, multiple organ failure, intravenous immunoglobulins

Presentation
A 45-year-old female patient presented to the accident and emergency department (A&E) having been generally unwell for the past 12 hours. She had begun to develop rigors. Her past medical history was of birdshot chorioretinopathy, an autoimmune disease causing posterior uveitis, for which she was taking prednisolone and mycophenolate mofetil. She had never smoked and had no recent travel history. She worked as a teaching assistant and was a mother to two children who were well. She did not consume alcohol, had never smoked and had no recent travel history.

On examination she had a temperature of 39.7°C and tachycardia at 110 beats per minute. However, her blood pressure (BP), oxygen saturations and respiratory rate were within normal limits and she was fully alert and orientated. Electrocardiography (ECG) showed sinus tachycardia and chest X-ray was unremarkable. A urine dipstick test only revealed a trace of blood and was negative for nitrites and leucocytes. Bloods on admission showed venous lactate of 1 mmol/l (blood results are shown in Table 1). Initially this gave an impression of sepsis with unknown origin (in view of fever on a background of an immunosuppressed status). The treatment plan included iv antibiotics (Penicillin) after blood cultures were taken, doubling the dose of adrenaline and vasopressin were subsequently commenced with good effect. Intravenous immunoglobulin (IVIG) was also started. Despite vasopressors and aggressive fluid resuscitation the patient remained oliguric and became progressively more acidic. Continuous veno-venous haemofiltration (CVVHF) was initiated. After 3 days on...
intensive care she was weaned off the CVVHF and vasopressor support, and transferred to ward care. Four days later she was discharged home from the ward, with advice from the gynaecologist to not use tampons again. A high vaginal swab subsequently grew Staphylococcus aureus, while two sets of blood C+S tests were negative.

Discussion

Toxic shock syndrome (TSS) is an acute systemic illness associated with infection by strains of *S aureus* which produce toxic shock syndrome toxins (TSST-1) or streptococcus pyogenes M types I and III – mainly group A beta haemolytic streptococcus (GABHS)-producing streptococcal pyogenic exotoxin A (SPEA). TSST-1 and SPEA are ‘super-antigens’ that interact with antigen-presenting cells (APCs) and T Cells to induce T Cell proliferation and massive cytokine production including tumour necrosis factor (TNF)-α, TNF-β, interleukin (IL)-1, IL-2 and IL-6, causing severe shock.

*S aureus* infection can be divided into menstrual and non-menstrual cases (ie postmenopausal women, men and children). Menstrual cases are associated with vaginal colonisation with a toxigenic strain of *S aureus*-producing TSST-1 followed by penetration of a sufficient concentration of TSST-1 across the epithelium, aided by abrasion from barrier contraception or tampon use. The absence of, or depression in, titres of neutralising antibody to the toxin, as in an immunosuppressed host, can enhance the severity of TSS.

The focus of infection in non-menstrual cases of *S aureus* TSS can be pharyngitis, burns, cellulitis or even simple skin abrasion. GABHS TSS occurs more commonly in patients who are intravenous drug users, have had surgical implants and potentially in those with simple blunt trauma. It is enhanced in an immunosuppressed patient including those with diabetes, alcoholism and steroid use.

Clinical presentation of *S aureus* TSS is usually an abrupt onset of fever (ie a temperature of >38.9°C), vomiting, diarrhoea and abdominal pain, macular erythroderma involving palms and soles with desquamation 1–2 weeks later, mucous membrane hyperaemia (vaginal, oesophageal and conjunctival), sore throat, myalgia, headache and photophobia. It is worth mentioning that our patient had not complained of gastrointestinal tract (GIT) disturbance, nor had a skin rash. Hypotension with multiple organ failure (MOF) occurs usually within 72 hours with an estimated mortality rate of 5%.

GABHS TSS differs in that it is usually present with flu-like symptoms and febrile illness, but GIT disturbance is much less likely and rash occurs in only 10% of cases. Infective myositis and necrotising fasciitis occurs in 70% of patients as GABHS usually arises from a deep-seated area of the skin or after minor blunt trauma. Hypotension and MOF develops within 4–8 hours in 95% of patients with a mortality rate of up to 70%.

Investigations

Clinical suspicion is the most important step, as there may be no clues. Common laboratory findings include abnormal liver function tests with hypoalbuminemia, acute renal failure, low platelet count and raised creatinine kinase. Skin, vaginal and pharyngeal swabs (according to the focus of infection) can confirm the source of infection. Blood C+S may be positive and, in cases of GABHS TSS, antistreptolysin O titre (ASOT) may be detected.

Failure to treat the infection with penicillin is common. Clindamycin is superior to penicillin because of its potency.

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<tr>
<th>Table 1. Blood results in the first 24 hours.</th>
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AST = aspartate aminotransferase; CRP = C-reactive protein; K = potassium; Na = sodium; WBC = white blood cell.

*Low result. †High result.

 Normal reference levels: Na = 133–146 mmol/l; K = 3.5–5.3 mmol/l; Urea = 2.7–7.8 mmol/l; Creatinine = 46–80 μmol/l; CRP = <5 mg/l; Albumin = 35–50 g/l; AST = <32 U/l; Alkaline phosphatase = 30–130 U/l; Haemoglobin = 115–165 g/l; WBC = 3.4–11 × 10⁹/l; Neutrophils = 1.7–8 × 10⁹/l; Platelet count 140–450.
toxic shock syndrome (TSS) should be suspected in immunosuppressed patients as well as in menstruating ladies using tampons who present with febrile illness with no obvious source of infection. Hypotension and multiple organ failure occur within hours and can be resistant to aggressive fluid and antibiotic therapy. Infective myositis and necrotising fasciitis can occur in 70% of group A beta haemolytic streptococcus (GBHS) TSS cases. Blood culture and sensitivity can be negative in TSS. Clindamycin is superior to penicillin in upper case and there is a role for intravenous immunoglobulins.

in suppressing bacterial toxin synthesis as well as facilitating phagocytosis. Clindamycin also suppresses lipopolysaccharide-induced monocyte synthesis of TNF-α. Intravenous immunoglobulin (IVIG) therapy reduces T cell production of pro-inflammatory cytokines, and significantly increases the plasma neutralising activity against superantigens, thereby resulting in improved survival in many TSS studies. IVIG therapy has also demonstrated an anti-TSST-1 effect and significantly improved survival in cases of S aureus TSS.

Identification of potentially implicated foreign bodies – especially tampons, surgical drainage and irrigation or excision of infected sites such as implants or necrotizing fasciitis – is of high priority, and input from surgical teams can be essential.

Our case emphasises the importance of maintaining a high level of suspicion for TSS in patients who are at high risk of the condition (eg menstruating ladies using tampons, patients who are immunosuppressed, and patients who have trauma, burns, cellulitis, pharyngitis or implants) and who are presenting with a febrile illness and signs of shock or hypoperfusion. It is important to take a formal and accurate menstrual and sexual history in selected cases.

Fluid resuscitation and intravenous antibiotics after blood C+S is essential and early ICU involvement for possible vasopressors, haemofiltration and IVIG use is required. Surgical excision of any focus of infection may result in a significant decrease in morbidity and mortality.

**References**


**Address for correspondence:** Dr T Shalaby, Maidstone Hospital, Hermitage Lane, Maidstone, Kent ME16 9QQ. Email: t.shalaby@nhs.net