A case of streptococcal myositis (misdiagnosed as hamstring injury)

Norbert Kang, Dimitrios Antonopoulos, Atul Khanna

Abstract
Streptococcal myositis is a very rare bacterial infection of muscle with a high mortality. Diagnosis is difficult because of the paucity of clinical signs and symptoms at the onset. However, presentation of the disease appears to have changed over the last 50 years. A case of streptococcal myositis is presented (misdiagnosed as hamstring injury), which more closely reflects the current presentation of the disease. Some of the features that may help emergency clinicians to recognise the onset of the condition are highlighted.

Keywords: necrotising fasciitis; myonecrosis; streptococcal myositis

Streptococcal myositis is an acute infection of muscle by invasive group A β haemolytic streptococcus causing myonecrosis without abscess formation. It differs from the more frequent and relatively benign streptococcal pyomyositis, which is characterised by the formation of abscesses in muscle and which has a good prognosis.

The standard surgical texts are misleading: “Streptococcal myositis resembles acute clostridial gas gangrene and was not described until World War II. After an incubation period of 3 to 4 days there is swelling, edema, and purulent wound exudate. These signs are followed by pain which rapidly becomes severe. Gas is present and the infected muscle changes from pale and soft to bright red, striped with purple and finally purple and gangrenous. The seropurulent discharge has a sour odour.”

The principle source of this description is MacLennan who collected a series of eight cases of streptococcal myositis in soldiers with battle wounds during World War II. This description differs from more recent reports including our own experience of the condition.

Case report
A 23 year old labourer presented with 24 hours of increasing pain in the medial aspect of his right thigh. There was no history of recent trauma. Slight erythema, tenderness, and swelling were noted. The pain was worse with movement especially extension and abduction. Sensation and pulses were intact throughout the lower limb. He was afebrile. A diagnosis of

cally stable as they are at risk of subsequent deterioration. Despite these measures, however, mortality remains high.

hamstring strain was made. He was given naproxen, a support bandage, prescribed a course of physiotherapy, and discharged. He returned the next day with increasing pain in his right thigh and vomiting. The whole of his right lower limb, particularly the thigh, was grossly swollen, and "bruised". An ultrasound scan showed no evidence of deep vein thrombosis, muscle rupture, haematoma or abscess, but did show diffuse oedema of the muscle. Blood results were normal except for urea 13.6 mmol/l, creatinine 0.228 mmol/l, and creatinine kinase 2491 IU/l. Plain radiographs of the right thigh showed diffuse oedema of muscle but no foreign body and no fracture. Seven hours after admission the patient developed haemorrhagic bullae. He was afebrile, but hypotensive and tachycardic. Ten hours after admission his right thigh, groin, and scrotum became gangrenous (fig 1). At operation (72 hours after the onset of symptoms) the skin of the right lower limb, and right lower quadrant of the abdomen, were necrotic. Because the muscles of the thigh were necrotic, a hip disarticulation was performed. The patient remained in intensive care for four weeks and returned home eight weeks after admission.

Discussion
In 1994, the Public Health Laboratory Service (PHLS) in the UK initiated a system of enhanced surveillance for all invasive group A streptococcal infections. Between 1 July 1994 and 30 June 1996 there were 1092 reported cases of invasive group A streptococcal infection in England and Wales. Of these, only four were described as invasive group A streptococcal infections of muscle without abscess formation.

Our review of previously published cases of streptococcal myositis and those recently reported to the PHLS indicated a male to female ratio of 2:1. This compares with a male to female ratio of 0.9:1 for all invasive group A streptococcal infections. The typical patient was aged 30 to 40 years with no significant previous medical history. A portal of entry for infection was present in only five of previously published cases and none in the four cases recently reported to the PHLS. This compares with the open battle wounds described by MacLennan. The first symptom was usually increasing pain in the affected muscle. This is in contrast to simple hamstring injury or the description in the surgical texts which describe "swelling, edema, and purulent wound exudate" preceding the onset of pain. The most frequent misdiagnoses were flu/viral illness, deep vein thrombosis, or muscle strain/rupture/haematoma/abscess. The lower limbs were affected in approximately 50% of all cases. Skin involvement with haemorrhagic bullae, blue mottling, and ecchymosis were late features. In further contrast to MacLennan, intramuscular gas was not a feature of any recent cases. However, when plain radiographs or ultrasound were performed at initial presentation, oedema of the affected muscle was seen. In addition, serum creatinine kinase levels (where recorded) were always raised at initial presentation.

For the emergency room physician faced with a possible case of streptococcal myositis, the most significant symptom appears to be pain in the infected muscle that worsens rapidly over 24-48 hours. Once the index of suspicion has been raised, ultrasound and plain radiographs will help to distinguish streptococcal myositis from more common diagnoses while a raised serum creatinine kinase will clinch the diagnosis well before streptococci can be cultured. The recommended treatment remains aggressive surgical debridement of all infected tissue with high dose benzylpenicillin (2.4 g, every four hours) plus clindamycin (0.6–1.2 g, every six hours) in severe cases.

I gratefully acknowledge the help of Dr M Monninckendam and Dr R C George in providing data on cases of invasive group A streptococcal infection reported to the PHLS between 1 July 1994 and 30 June 1996.

References
2 MacLennan JD. Streptococcal infection of muscle. Lancet 1943;ii:582-4.