Lower limb bilateral pyomyositis in a diabetic patient: case report and literature review

ABSTRACT
Pyomyositis is a bacterial infection of skeletal muscle tissue occurring mostly in immunocompromised patients, including diabetics. The main agents are Gram-positive cocci. We report a case of a 42 years old, male, diabetic, who presented the emergency room referring pain in thighs and inability to flex his legs. We performed a magnetic resonance which revealed extensive purulent collections in both lower limbs, thus confirming clinical suspicion, and allowing proper antibiotic treatment. In this case, we showed that imaging tests facilitate early diagnosis and treatment through direct location of the lesions, and guides invasive procedures such as biopsy and abscess aspiration when needed. (Clin Diabetol 2020; 9; 2: 134–137)

Key words: pyomyositis, diabetes mellitus type 2, magnetic resonance imaging

Introduction
Scriba, in 1885, reported the first case of pyomyositis [1], as a primary acute bacterial infection of skeletal muscle’s tissue [1, 2]; it occurs mainly in immunocompromised patients [3], malnurished [4] or, exceptionally, in immunocompetent patients in tropical areas [3]. In fact, it was originally described as a tropical disease and is now being seen, with increasing frequency, in temperate regions; pyomyositis holds a mortality rate of about 10% and may also be responsible for complications such as abscesses, septicemia, and shock [2].

The pathogenesis of primary pyomyositis remains unclear, although trauma and immune impairment can be predisposing factors [2]. It affects individuals, preferably during the first and second decades of life, and is more prevalent in men [1]; patients over 30 years of age usually have associated comorbidities [5], and the most described in the literature are human immunodeficiency virus infection [6], diabetes mellitus, malignant tumors, liver cirrhosis, renal failure, transplantation and use of immunosuppressive drugs, and intravenous drug abuse [1, 3]. Its initial signs and symptoms are usually subtle and can be easily overlooked or misdiagnosed as a more benign condition [2], specially erysipelas.

Pyomyositis can affect any muscle group, infecting by contiguity or hematogenous spread [1, 5]. The main agents are Gram-positive cocci [3], and the most common etiologic agent is Staphylococcus aureus, in up to 85–95% of the cases [1, 2, 5] — with increasing frequency of resistance to methicillin (MRSA) [2].
Case report

The patient is a 42 years old, Brazilian male, security agent, referred to the emergency department complaining of a acute pain of severe intensity on both thighs, restraining flexion of both lower limbs for the last seven days. He reported that a less severe pain had been present for over two months, and, one month earlier, he was hospitalized for 15 days due to erysipelas in his right leg — received analgesics without improvement at that time. He denied smoking or trauma, reporting, just social drinking; as a family background his father was diabetic and his mother had arterial hypertension. Physical examination of the patient showed a hyperemic, swollen and painful right leg, especially in the calf; his body temperature was 38.2°C pulse of 98 ppm and blood pressure of 150 over 80 mm Hg. Laboratory workup showed increased leukocyte count (18,452 hg/mm³) and elevated c-reactive protein 15 mg/dL. Throughout this hospitalization, he was also diagnosed with type 2 diabetes mellitus and systemic arterial hypertension, having started treatment since then, and had good glucose control throughout hospitalization with metformin.

During the first hospitalization, a magnetic resonance image (MRI) of the right leg was performed (Figure 1 — description on the legend), but it might had been neglected. Given the progression of symptoms and severity of the clinical setting, a new MRI of both thighs was performed (Figure 2 — description on the legend), which showed purulent collection affecting both legs, rising the diagnosis of bilateral pyomyositis stage 2. Patient started to receive ceftriaxone plus teicoplanin empirically, and after 2 weeks was discharged with complete resolution.

Discussion

The diagnosis of pyomyositis can be challenging due to the nonspecific clinical signs at onset [2]. It may be confused with bone or joint infection processes, thrombophlebitis, panniculitis or systemic diseases [2]. The differential diagnosis includes muscle trauma, deep vein thrombosis, osteomyelitis, cellulitis, septic arthritis, and malignant tumors [2].
In the case of diabetic patients, it appears that muscular and circulatory disorders which, together with granulocyte dysfunction and decreased cellular immunity, increase the risk of developing pyomyositis [4]. Laboratory findings are nonspecific and the blood count may show leukopenia or leukocytosis with left deviation, depending on the etiologic agent and the patient immune response [1, 4]. There may be increased inflammatory markers and generally does not affect kidney function [3, 5]. Blood cultures are positive in less than 40% of patients and secretion cultures are positive in only 21–41% of cases [6].

Usually, it only affects one muscle group, but in 11% to 43% of patients may have widespread outbreaks in various muscles, being quadriceps muscle the most affected, followed by the gluteal and iliopsoas muscles [7]; iliopsoas pyomyositis occurs more commonly secondary to gastrointestinal or urinary tract infections [7].

Pyomyositis presents three evolutionary stages: invasive, purulent and late. The invasive stage is characterized by insidious onset, where only about 2% of affected patients seek medical support. The purulent stage occurs about 10–21 days after symptom onset and is characterized by the abscess location and, at this stage, about 90% of patients seek medical care due to the severity of the symptoms. If the disease is not identified and treated properly with antibiotics and/or surgical drainage, evolution occurs for the so-called late period associated with significant systemic involvement like septic embolization, septic shock, and even death [8]. Our patient had a bilateral phase 2 pyomyositis, due to an initial misdiagnosis of erysipelas, when he was still on initial phase.

Imaging tests such as ultrasound, computed tomography (CT) and magnetic resonance imaging (MRI) are useful to identify the number, size, extension, location of abscesses, and guide needle biopsy [9].

Ultrasound has high sensitivity for fluid collections detection; it may demonstrate initial muscle edema as a hypoechoic lesion with imprecise limits affecting one or more muscles that, with latter disease evolution, can lead to the formation of abscess [3].

CT scan shows enlargement and decreased attenuation of the affected muscle, with a blurring of surrounding fat. The involvement of a muscle group that is disproportionate to the involvement of the subcutaneous tissue helps distinguish myositis from primary cellulitis. Intramuscular collections can be observed and contrast material is used to help differentiate viable and necrotic muscles by demonstrating enhancement around the abscess [5, 7]. CT scan, as well as ultrasound, can help guide needle aspiration [3] and are also useful for surgical planning, with abscess drainage muscle followed by culture-guided antibiotics [5, 7].

MRI is the gold standard for the diagnosis by detecting early findings of muscles diffuse inflammation [5], and may also guide biopsy of the affected muscles [10]. MRI can show extension of infection and evaluates adjacent structures, such as joints, bone, and other surrounding soft tissues, differentiating pyomyositis from other differential diagnosis.

Treatment with antibiotics in conjunction with surgical drainage or suction, when applicable, is generally sufficient; prognosis is good with early diagnosis and appropriate treatment, depending on the underlying comorbidities [11, 12]. Usually, one antibiotic with S. aureus coverage such as oxacillin, ciprofloxacin or cephalosporins is enough, but when dealing with immunocompromised patients, association of a minoglicoside or even glycopeptides may be useful [5].

Untreated pyomyositis complications may include compartment syndrome, progression infection to septic arthritis, osteomyelitis and even death, which ranges between in up to 10% of the cases [5, 7]. In the long term, patients may experience weakness, and muscle dysfunction [5, 7].

In conclusion, pyomyositis may have subtle symptoms that are easily misinterpreted as signs of a more common and benign infection. Imaging tests facilitate early diagnosis and treatment through the direct location of the lesions and, when it’s necessary, assist invasive procedures such as biopsy and abscess aspiration. The early recognition of the disease allows an earlier treatment preventing the development of complications.

Conflict of interest

We state that the authors have no conflict of interest.

REFERENCES