REVIEW

Tropical pyomyositis (myositis tropicans): current perspective

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Tropical pyomyositis, a disease often seen in tropical countries, is characterised by suppurative involvement of skeletal muscles, usually as single or multiple abscesses. The most common organism implicated is *Staphylococcus aureus*. In 20%–50% of cases there is a history of trauma to the affected muscles. Commonly involved muscles are quadriceps, glutei, pectoralis major, serratus anterior, biceps, iliopsoas, gastrocnemius, abdominal and spinal muscles. Early diagnosis is often missed because of lack of specific signs, unfamiliarity with the disease, atypical manifestations, and a wide range of differential diagnosis. Diagnostic techniques like ultrasound and computed tomography/magnetic resonance imaging are very useful in diagnosis. The diagnosis is confirmed either by biopsy or aspiration of pus from the affected muscles. The initial antibiotic of choice is cloxacillin. Incision and drainage are important components of management. Treatment for Gram negative or anaerobic organisms should be instituted, whenever indicated. Physicians should become more familiar with this potentially life threatening but curable infective disease entity.

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Traqair credited Virchow for the earliest mention of tropical pyomyositis, though Scriba in 1885 described this entity for the first time. Levin et al in 1971 reported the first case from a temperate region. Since then many cases have been reported from various geographical regions of the world. This increase in incidence is attributed to heightened awareness of this disease, an increase in the number of immunocompromised patients, and improvement in diagnostic techniques.

Since this entity is characteristically found in tropical areas, various terms like tropical pyomyositis, myositis tropicans, tropical skeletal muscle abscess, and tropical myositis are used. With increasing recognition from temperate regions, it is also referred to as non-tropical myositis, infectious myositis, or spontaneous bacterial myositis. Although the classical presentation is with muscle abscess, the hallmark of the disease is not an abscess but finding myositis on a biopsy specimen of involved muscle. Therefore, some authors prefer the term myositis instead of pyomyositis. The term “tropical pyomyositis” should be restricted to primary muscle abscess arising within the skeletal muscle. It should not be used to describe (a) intermuscular abscesses, (b) abscesses extending into muscles from adjoining tissues such as bone or subcutaneous tissues, and (c) those secondary to previous septicemia.

PREVALENCE AND INCIDENCE

Tropical pyomyositis is common throughout the tropics and has been widely reported from Asia, tropical Africa, Oceania, and the Caribbean islands. It accounts for 1%–4% of all hospital admissions in some tropical countries, In Uganda about 4% of all hospital admissions and in Ecuador 2.2% of all surgical admissions have been reported as due to tropical pyomyositis. In India it has not been widely reported but there have been sporadic case reports. Tropical pyomyositis is also increasingly reported from temperate regions. Recent data indicate that up to 75% cases of tropical pyomyositis in temperate areas are immunocompromised—HIV infected, or with diabetes, leukaemia, chronic renal failure, asplenia, scleroderma, rheumatoid arthritis, or Felty’s syndrome—while some patients have been on cancer chemotherapy or immunosuppressive drugs after transplantation of solid organs. Tropical pyomyositis has also been described after bicycle accidents and vigorous exercise. The incidence of tropical pyomyositis in HIV infected patients with or without AIDS is around 31%. Although cases are seen throughout the year, maximum incidence is from July to September (the monsoon season).

AETIOLOGY

*Staphylococcus aureus* is the organism most commonly cultured from the abscess. It is seen in up to 90% of cases in tropical areas and 75% of cases in temperate countries. Group A streptococcus accounts for another 1%–5% of cases and other organisms uncommonly implicated are streptococcus (groups B, C, G), pneumococcus, neisseria, haemophilus, aeromonas, serratia, yeinia, pseudomonas, klebsiella, and escherichia. Rarely salmonella, citrobacter, fusobacterium, anaerobes, and mycobacterium are seen.

In tropical regions, pus cultures are sterile in 15%–30% cases and 90%–95% of patients also, have sterile blood cultures. Blood cultures are positive in 20%–30% of cases in temperate regions. Better microbiological culture techniques in the temperate regions may account for these variations.
PATHOGENESIS
Pyomyositis is an intriguing disease of unclear pathogenesis. Organisms reach skeletal muscles during transient bacteremia. Skeletal muscle tissue is intrinsically resistant to bacterial infections under normal circumstances. It is thought that there is sequestration of iron by myoglobin, which is an essential nutritional requirement of proliferating bacteria. This results in slower growth of bacteria, allowing celluar and humoral defences to enter infected zones and thereby preventing establishment of infection. In a series of autopsied cases of staphylococcus septicemia, abscesses in skeletal muscle were found in fewer than 1% of cases. Even thereby preventing establishment of infection. In a series of autopsied cases of staphylococcus septicemia, abscesses in skeletal muscle were found in fewer than 1% of cases. Even though not proven, include nutritional deficiencies and viral and parasitic infections. It is postulated that abundant iron is available after trauma, resulting in profuse growth of bacteria.

It has also been suggested that an abnormality of the immune system may be the underlying cause in many cases. Some researchers have proved that the lymphocytes, particularly T-cells, in patients with tropical pyomyositis are not primed adequately against staphylococcus during the course of infection. Other predisposing factors implicated include a preceding viral infection (especially arbovirus) or nematode infection (Dracunculus medinensis) but their role in pathogenesis is uncertain. Moreover, as these abscesses are intermuscular rather than intramuscular, these should not be included under tropical pyomyositis.

Intravenous drug abuse is another important risk factor for tropical pyomyositis. The factors responsible are impaired cellular and humoral immunity, defective bactericidal capacity of neutrophils, increased bacterial colonisation of skin, and direct injection of contaminated materials.

Tropical pyomyositis is also increasingly documented in persons infected with HIV. Mechanisms include muscle damage caused by HIV infection per se, zidovudine therapy, infections caused by parasites and mycobacteria, and impaired host defences including dysfunction of helper T-cells and granulocytes. In some studies, it has been demonstrated that the CD4 counts are less than 150/μl in such cases. Also, those with HIV infection have an increased incidence of staphylococcus carrier state compared with those not infected with HIV.

PATHOLOGY
In early stages of tropical pyomyositis, muscles show oedematous separation of fibres, followed by patchy mycotoxicosis progressing to complete disintegration. The fibres are surrounded by lymphocytes and plasma cells. Muscle fibres may heal without abscess formation or degenerate, progressing to suppuration with bacteria and polymorphonuclear leucocytes.

CLINICAL FEATURES
The disease is seen in all age groups, although young males are the most susceptible group. Maximum incidence is seen at 10–40 years of age with a male to female ratio of 1.5:1. Muscles frequently involved are quadriceps, glutei, pectoralis major, serratus anterior, biceps, iliopsoas, gastrocnemius, abdominal and spinal muscles. Usually, a single group of muscle is affected, but in 12–40% of cases multiple groups are involved either sequentially or simultaneously. The clinical picture is divided into three stages.

(1) Invasive stage
This first stage is characterised by a subacute onset of variable fever, painful firm swelling, and minimal systemic symptoms with or without erythema (as infection is deep seated). The area is tender with a wooden consistency. Aspiration, if attempted at this stage, yields no pus as the phlegmonous inflammatory process is diffuse. Firm swelling, absence of erythema, and mild pain may divert the attention of a physician away from an infectious aetiology. The invasive stage may resolve itself, mimicking fibromyalgia or may progress to next stage of suppuration.

(2) Suppurative stage
From the second week to third week, abscess forms in the muscle. High spiky fever, with more severe systemic symptoms marks the beginning of the supplicative stage. Most cases present at this time. The classical signs of abscess, fluctuation and erythema, may be lacking because the overlying muscle is tense. Needle aspiration at this stage yields pus. Regional lymph nodes are not involved.

(3) Late stage
If the abscess remains untreated, dissemination of infection occurs. Bacteremia, followed by septicemia, septic shock, acute renal failure, and metastatic abscesses are some of the complications described.

ATYPICAL PRESENTATIONS
The patient may present acutely with fever and chills or local symptoms. In a rare patient, the picture may be that of toxic shock syndrome. At times the invasive stage may be prolonged and the patient may present with pyrexia of unknown origin. Rarely, it may present as an acute abdomen or spinal cord compression or compartment syndrome. When localised to neck muscles, it can be mistaken for cervicobrachial neuralgia.

DIFFERENTIAL DIAGNOSIS
Tropical pyomyositis is a great masquerader. Its differential diagnosis is wide and includes pyrexia of unknown origin, muscle contusion, septic arthritis, osteomyelitis, cellulitis, muscle haematomata, deep vein thrombosis, muscle rupture or muscle strain, osteosarcoma of muscle, trichinosis, leptospirosis, and polymyositis. Occasionally, the stage of invasion by Cysticercus cellulosae is characterised by fever, muscle tenderness, and eosinophilia and needs to be differentiated from tropical pyomyositis. Rarely, focal or diffuse inflammatory myopathy due to trypanosomiasis or toxoplasmosis presents as severe myalgia mimicking pyomyositis. It must also be distinguished from spontaneous gangrenous myositis (caused by Streptococcus pyogenes) characterised by gangrenous necrosis. Therefore, a diagnosis of tropical pyomyositis should be considered in any patient presenting with muscle pain, fever, and/or leucocytosis.

DIAGNOSIS
Early diagnosis is critical for saving the tissue and also the life of patient but is often missed because of unfamiliarity with the disease, atypical presentations, a wide range of differential diagnoses, and lack of early specific signs. Aspiration of pus from the muscle or muscle biopsy with culture and tissue staining in cases of absent macroabscesses is the gold standard for diagnosis (pus may be sterile in 15%–30% of cases). Muscle biopsy also helps to exclude osteosarcoma, polymyositis, trypanosomiasis, toxoplasmosis, cysticercosis, and trichinosis.

Non-invasive radiological methods are helpful in establishing the diagnosis. These techniques are also used for subsequent follow up of the patient. Ultrasound is the initial
screening tool for reasons of economy and easy availability. Hypoechoic areas with an increase in muscle bulk are seen on ultrasound. Computed tomography/magnetic resonance imaging (MRI) are the best imaging techniques for early diagnosis. Computed tomography shows areas of low attenuation with loss of muscle planes and a surrounding rim of contrast enhancement as characteristic of pyomyositis. Computed tomography is also useful in differentiating tumours, haematoma, and thrombophlebitis from abscesses. However, at times a computed tomogram alone may be unreliable in distinguishing muscle abscess from swollen muscles. MRI shows hyperintense rim on T1 weighted images with peripheral enhancement on gadolinium DTPA scan. A case has been reported in which areas of signal attenuation were visible on MRI but a computed tomogram was normal. Also, MRI is the imaging modality of choice for pelvis. Therefore, MRI may be a better modality than computed tomography for early diagnosis. Gallium scintigraphy is an extremely sensitive tool, but its inability to provide anatomic details, lack of specificity, and high cost reduces its usefulness. It is best reserved for a group of patients in which clinical suspicion is high but computed tomography/MRI is inconclusive. It is also utilised for detecting unsuspected but possible metastatic abscesses.

Laboratory investigations reveal anaemia, leucocytosis (shift to the left), raised erythrocyte sedimentation rate, and acute phase reactants. In an asymptomatic patient with muscle mass and no other features of infection (fever, chills, erythema) but who has leucocytosis and raised erythrocyte sedimentation rate, pyomyositis should be suspected. In cysticercosis and trichinosis, eosinophilia is present. Blood cultures have low positivity, varying from 5%–10% in the tropics to 20%–30% in temperate regions but are essential in all suspected or proven cases. Serum levels of muscle enzymes—that is, aldolase, creatine phosphokinase, and lactic dehydrogenase—are normal or slightly raised despite evidence of muscle destruction. Raised creatine phosphokinase along with characteristic electromyography changes (short duration, low amplitude polyphasic potentials), favour the diagnosis of pyomyositis. Deranged liver and renal functions, along with fever and severe diffuse muscle pain, in leptospirosis that is confirmed by serological tests. Immunodeficient states must be ruled out and tests for HIV, diabetes, rheumatological disorders, and malignancies should be carried out, along with immunoglobulin levels.

TREATMENT

Once diagnosis is established, attention should turn to aggressive management. Surgical debridement and drainage, accompanied by parenteral antistaphylococcal β-lactamase resistant penicillin (cloxacillin 1–2 g every six hours), is the initial recommended treatment. Diffuse myositis without abscess may respond to an antimicrobial agent alone but abscess may develop eventually requiring drainage. In most clinical settings, treating S aureus infection with more than one drug, to which the organism is known to be susceptible, attains no significant benefit. Penicillin is the drug of choice for infections caused by penicillin susceptible staphylococcus. Drug combinations of penicillin with β-lactamase inhibitor are also effective but are best reserved for treatment of mixed infections. In case of penicillin allergic subjects, first generation cephalosporin (cefazolin) may be preferred for reasons related to cost, potency, and breadth of spectrum. For methicillin resistant staphylococcus, vancomycin in a dose of 15 mg/kg to a maximum of 1 g, given every 12 hours is a suitable alternative. Another glycopeptide, teicoplanin, in a dosage of 400 mg/day in a single dose is equally effective. For vancomycin intermediate sensitive staphylococcus, linzolid or dalfopristine-quinupristine are effective alternatives.

If group A streptococcus is isolated from the pus, treatment should be changed to penicillin, while awaiting culture and sensitivity reports. For Gram negative bacilli, addition of an aminoglycoside, that is, gentamicin in doses of 5–6 mg/kg/day intravenously, should be considered in addition to cephalosporins. The possibility of associated anaerobic infection is an indication for introducing metronidazole (20–30 mg/kg/day) intravenously or orally at intervals of eight hours.

For HIV infection and immunosuppressed patients, broad spectrum empirical antibiotics against Gram positive, Gram negative and anaerobic organisms should be administered. In addition to antistaphylococcal antibiotics, patients should also receive aminoglycosides and clindamycin.

Secondary spread of metastatic infection from involved muscles usually requires four to six weeks of parenteral high dose antimicrobial therapy. Otherwise, treatment should be continued till wound is clean, the leucocyte count becomes normal, and the patient is afebrile for 7–10 days. Continuation or recurrence of fever after surgical drainage while the patient is receiving appropriate antimicrobials suggests the presence of other foci, development of drug resistance or, less commonly, drug fever.

Although there are no recommendations to prevent pyomyositis, it is believed that nasal carriage should be eliminated in patients with a history of pyomyositis or bacteraemic staphylococcal infection. It should be treated with topical mupirocin nasal formulation. Alternatives include rifampicin (600 mg each day) or cloxacillin (500 mg four times a day) for 10 days.

PROGNOSIS

The natural history is progressive suppuration with either spontaneous drainage and gradual resolution or eventual bacteraemia, and secondary infection leading to fatal outcome. Despite advances in diagnosis and treatment, mortality varying from 0.5% to 2% still occurs.

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