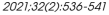
CASE REPORT





Neglected tuberculous trochanteric bursitis in an adolescent girl: A case report and literature review

¹Division of Paediatric Orthopaedic Surgery, Children's Hospital Zagreb, Zagreb, Croatia

²Department of Pulmonology, Allergology and Immunology, Children's Hospital Zagreb, Zagreb, Croatia

³Department of Paediatrics, School of Medicine, University of Split, Split, Croatia

⁴Department of Paediatric Radiology, Children's Hospital Zagreb, Zagreb, Croatia

⁵National Reference Laboratory for Tuberculosis, Croatian National Institute of Public Health, Zagreb, Croatia

⁶Department of Medical Microbiology and Parasitology, School of Medicine, University of Zagreb, Zagreb, Croatia

⁷Department of Pathology, Sestre Milosrdnice University Hospital Centre, Zagreb, Croatia

⁸Department of Pathology, School of Medicine, University of Zagreb, Zagreb, Croatia

Extrapulmonary tuberculosis (TB) occurred in 15% of all TB cases globally in 2018, while musculoskeletal TB accounts for approximately 10% of all extrapulmonary TB cases in developed countries. [1,2] Tuberculous trochanteric bursitis (TTB) is uncommon and accounts for less than 2% of musculoskeletal TB. [3] However, in the first half of the 20th century, TTB was not as uncommon as today. [4,5] Longstanding medical literature until 1950 described only 105 cases of TTB, not distinguishing between isolated infection of the bursa, trochanter, or combine infection of both the bursa and trochanter. [6]

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Correspondence: Josip Vlaic, MD, PhD. Division of Paediatric Orthopaedic Surgery, Children's Hospital Zagreb, Ulica Vjekoslava Klaica 16; 10000 Zagreb, Croatia.

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ABSTRACT

Tuberculous trochanteric bursitis (TTB) is an extremely rare form of extrapulmonary tuberculosis. Due to a low clinical suspicion and poor collaboration among medical professionals, the diagnosis of TTB can be often delayed. In this report, we describe a case of neglected TTB in an adolescent girl that initially presented with right thigh swelling and fluctuance. The patient underwent repeated unsuccessful surgical treatment; however, dull pain and periodic wound drainage remained for eight years. Complete excision of fistula and trochanteric bursa and one year of oral antituberculous drug therapy led to complete recovery. This case report highlights tuberculosis as a diagnostic challenge, when rare localizations are affected. In addition, this report addresses several diagnostic pitfalls and reviews the literature regarding TTB in adolescent patients. Orthopedic surgeons need to consider TTB, when swelling, fluctuance or repeated wound drainage are present on the thigh.

Keywords: Adolescent, anti-tuberculosis drugs, surgery, tuberculous trochanteric bursitis.

Recently, most of the published data on this subject are case series and case reports with only two adolescent patients reported.^[3,7,8]

Mainly due to a low clinical suspicion, lack of symptoms, and poor collaboration among medical professionals, the diagnosis of TTB can be delayed. Therefore, it is not surprising that TTB may be easily overlooked and neglected, delaying treatment and affect the patient's quality of life.

In this report, we present a case of TTB in an immunocompetent adolescent who initially presented

with recurrent thigh swelling. After several unsuccessful treatment regimens over eight years, she remained symptomatic, developing prolonged wound drainage of a cutaneous fistula. This report addresses several diagnostic pitfalls in TTB confirmation and reviews the literature regarding epidemiology of TTB in adolescent patients.

CASE REPORT

Patient's history

A 14-year-old girl presented to the tertiary referral Children's Hospital orthopedic outpatient clinic due to dull pain and periodic wound drainage on the lateral side of her right thigh. Her mother noted two previous surgeries on that area performed at a county hospital. The first one, eight years ago, was performed due to right thigh swelling and fluctuance. These symptoms were initially suspected to be caused by pyomyositis of the vastus lateralis. A small longitudinal incision on the lateral side of the thigh was done and a purulent cyst was evacuated. Samples were sent for microbiological, cytological, and pathological examination. The laboratory reports were not evaluated properly by the treating surgeon, and the girl was discharged from the hospital with recommendations for oral antibiotics to treat pyomyositis. A month after the operation, purulent wound drainage appeared. The child was continued to treated with oral antibiotics and underwent frequent dressing changes, until complete wound closure was achieved. However, 10 months later, swelling, pain and purulent drainage reappeared at the place of the initial incision. The patient underwent repeated surgical incision and evacuation of what was considered to be a recurrent subfascial abscess. After surgery, the patient was instructed to continue with antibiotics and to dressing changes while waiting for wound closure. Unfortunately, her wound was not completely closed. For the next seven years, she suffered from monthly purulent drainage accompanied by moderate swelling and dull thigh pain that was most intense a few days before the appearance of drainage. During that period, there were no complaints about fever or malaise.

Physical examination and initial considerations

Seven years after the second surgery, physical examination performed at our institution showed a chronic wound on the lateral right thigh with mild drainage and local erythema. Palpation was slightly painful; however, there was full, painless range of motion of the hip and knee. A swab was

taken for microbiological examination and an initial laboratory work-up was done. Microbiology results became positive for Pseudomonas stutzeri, while the laboratory findings were within the normal range. Initial radiographs of the femur were deemed normal. A thigh ultrasound scan revealed a sharply bordered hypoechoic collection between muscles and subcutaneous fat tissue that measured 3×1 cm in size, which was considered to be an abscess. Oral ciprofloxacin 1,000 mg was prescribed for 10 days; however, there was no change in her exam. Therefore, due to clinical presentation and according to previous diagnostic work-up findings, recurrent abscess was suspected. The patient was scheduled for surgical excision followed by continuous intravenous antibiotic therapy. However, while waiting for surgery, she developed intense pain in the right hip.

Radiological findings

Hip radiographs revealed a lytic lesion within the greater trochanter with partial reactive sclerosis (Figure 1). Unfortunately, due to the initial symptoms, as well as the site of the previous surgeries and ultrasound findings, surgery was performed without additional imaging studies.

Surgical technique

Surgery started with scar tissue excision around the previous wound. However, due to the



FIGURE 1. Hip anterposterior radiograph of a 14-year-old patient. In the right hip region, a lesion of the greater trochanter with surrounding reactive sclerosis of the lateral aspect, and without periosteal reaction is visible. Soft tissue edema is present around the right greater trochanter.

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FIGURE 2. Intraoperative photographs. (a) Right thigh from lateral side - a long subfascial cutaneous fistula going from the middle third of the thigh to the greater trochanter region is visible; (b) Completely excised fistula and affected bursa before sending to pathology.

intraoperative finding of a cutaneous fistula, the skin incision was extended to track the fistula proximally. The fistula was marked with a probe, and a detailed exposure showed a subfascial fistula connected to the enlarged trochanteric bursa (Figure 2a). While operating on the trochanteric bursa, it partially ruptured and whitish liquid tissue, resembling pus was observed. At this point, TB was suspected and the material was taken for acid fast laboratory diagnostics. The fistula and trochanteric bursa were excised completely and sent to pathology (Figure 2b).

Postoperative findings

Postoperative intravenous antibiotic therapy was administered for 10 days with 2 g of ceftazidime. The direct microscopic smear examination from the specimen was negative for acid-fast bacilli. The pathology report described tissue changes characteristic for fistula, without specifying the underlying disease. The wound healed uneventfully. There was no longer any local tenderness and the patient was discharged home with recommendations to continue with antibiotic therapy.

Prior to first postoperative visit, the microbiology culture returned positive for *Mycobacterium tuberculosis* (*M. tuberculosis*). The experienced pathologist who reviewed the recent histological findings was notified. The specimen was re-reviewed and corrected identified *M. tuberculosis* and amended the report to possible TB infection (Figure 3). While discussing these results with the patient's mother, it was found that a close relative that lived with them was TB positive at the time of the index surgery eight years ago. The genotyping results of both *M. tuberculosis* strains were identical, confirming the likelihood of transmission. In the retrospective analysis of the initial cytology report from eight years ago, it was

clearly reported as granulomatous inflammation with necrosis. Therefore, we concluded that the initial presentation with swelling and fluctuance on the right thigh was as a result of TTB. Hip and thigh magnetic resonance imaging (MRI) performed soon after surgery did not show any signs of TB.

Pulmonology work-up and oral TB therapy

Due to these findings, a complete pulmonology work-up was indicated. On pulmonary examination, the patient was immunocompetent, afebrile, had non-labored breathing, with normal breath sounds

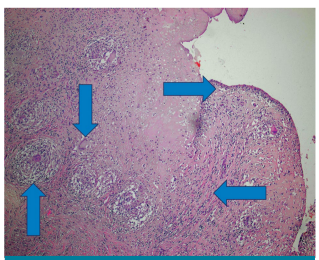


FIGURE 3. Microscopic analysis of trochanteric bursa biopsy showing ill-circumscribed granulomas composed of epithelioid and multinuclear cells and some mononuclear cells in the stroma. Some multinuclear cells are of Langhans type. Surface of the specimen is partly ulcerated with preserved synovial cells denoted by arrow. (H-E, ×100).

Down arrow-multinuclear giant cell of Langhans type, up arrow-poorly formed granuloma composed of epithelioid cells, left arrow-mononuclear inflammatory cells, right arrow-synovial cells on the surface.

on lung auscultation. Bacille Calmette-Guérin (BCG) vaccination after birth was confirmed by her visible BCG scar. There was no history of previous pulmonary TB, and her chest radiograph showed no signs of pathologic lesions. Culture of sputum samples were negative for *M. tuberculosis*. Oral TB treatment with four drugs was initiated (isoniazid 300 mg daily; rifampicin 450 mg daily; pyrazinamide 1 g daily; ethambutol 600 mg daily) for two months and continued with two drugs (isoniazid 300 mg daily; rifampicin 450 mg daily) for 10 months.

The patient completed one year of TB therapy and was followed closely by a pulmonologist and orthopedic surgeon. Twenty months after surgery, she complained about snapping and pain of the right hip, which was successfully managed with physical therapy. At the most recent four-year follow-up, there was no evidence of local recurrence, and the patient returned to all normal activities of daily living.

An informed consent was obtained from the parents of the patient.

DISCUSSION

The presented case of TTB took eight years to be correctly diagnosed and successfully treated. Unfortunately, during this delay, a great burden to the patient and family was created. Therefore, we believe that this report highlights all diagnostic pitfalls present in this case and attempts to explain why TTB remained neglected.

Based on the data published in the last Global Tuberculosis Report by the World Health Organization (WHO), the number of new TB cases has trended down over the last decades. Nevertheless, in developing countries, TB remains an important source of mortality. The ongoing migrant crisis may also affect the future TB incidence in Europe. In Croatia, the latest report on TB states the incidence of TB to be 9.1 per 100,000 population, while musculoskeletal TB was reported in only one patient in 2018.

Presentation of musculoskeletal TB may be insidious over a long period, as the diagnosis remains elusive and delayed. Vertebral involvement (*i.e.*, tuberculous spondylitis, or Pott's disease) is the most common type of skeletal TB, accounting for about half of all cases of musculoskeletal TB.^[1] However, when other sites are affected, such as the trochanteric bursa, TB is not initially considered in the differential diagnosis.

A standardized search of the published data on publicly available search platforms was performed to identify studies dealing with TTB. To the best of our knowledge, only a few cases of TTB in children were reported (Table I).^[5,7,8,10] The first description of, what is now considered to be TTB in a child, was given by Macnab, [10] a surgeon at Suffolk General Hospital, United Kingdom, in 1870. In his report, following minor trauma, the child's hip was slightly swollen over the greater trochanter, hot, sore on palpation, while walking was painful. Recommended therapy was rest, supportive ambulation, and immobilization. However, six months later, swelling over the hip had suddenly become painful, rapidly increased in size, and erupted, discharging a large quantity of pus. Macnab, then, instructed carbolic acid to be injected into the abscess three times daily, and series of dressing to be applied to the wound. Finally, after one month, the wound over the hip diminished in size, healthy granulation tissue was present, with little discharge. Hip range of motion was symmetric to the unaffected side. According to the physical exam and treatment described, one could notice that Macnab, indeed, witnessed the natural course of TTB. There is a remarkable resemblance between our recent case and Macnab's description 150 years ago what should now provoke disappointment to all physicians involved.

In the first half of the 20th century, TTB was not uncommon.^[5,6] Among 15 cases of TTB, McMurray^[5] reported two of them in children. Similar to our case, a 14-year-old girl complained to a thigh swelling and pain for more than a year. Affected bursa was aspirated, but still the patient developed bony erosion of the greater trochanter in a one-year follow-up. Therefore, the author strongly emphasizes that greater trochanteric bone lesions were often found secondary to TB infection of the trochanteric bursa. Seen backwards, the described erosions of the greater trochanter on radiographs in our patient should raise suspicion of TTB even before surgical intervention. Nevertheless, the diagnostic challenge in regards to TB of the bone, joint, and bursa in children was recognized long ago. In 1931, Milgram^[4] reported the diagnostic inaccuracy between Legg-Calvé-Perthes (LCP) disease and TB of the hip. However, currently, LCP disease is not a diagnostic challenge with TB due to imaging methods used. In the last three decades, two radiological studies reported on two adolescent patients with TTB.^[7,8] As in our case, X-ray findings indicate the slight lytic changes of the greater trochanter and soft tissue reaction.[7,8]

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		Long-term outcome	N A	₹	NA	Ϋ́	Four years after surgery, complete recovery
TABLE I	Literature review of pediatric and adolescent cases of tuberculous trochanteric bursitis ^[5,7,8,10]	Treatment	Carbolic acid injected into the abscess. Series of dressing	Aspiration of abscess Rest	Surgery, chemotherapy	Ϋ́	Two surgeries (abscess evacuation) and antibiotics Series of dressing* Surgery (excision of fistula and trochanteric bursa) and oral antituberculous medications
		Diagnostic method	Observation	Guinea pig inoculation	Y Z	Υ	diagnostics
		Plain radiographs	V V	Normal radiographs at time of diagnosis Trochanteric erosion 1 year later	Trochanteric erosion, STC	STS	Lytic lesion and partial reactive sclerosis of the greater trochanter
		Other TB sites	N A	Not noted	N A	ΑN	Not noted
		Clinical presentation	From pain and swelling to wound with discharge	Mild pain and swelling	Pain for 6 months	Ϋ́	Present study 6*-14/F Home exposure Thigh swelling and Not noted I fluctuation* Chronic wound on the lateral thigh with a minor secretion and local redness
		Predisposing factors	Trauma to hip	₹	Home exposure	ΝΑ	Home exposure
		Age/sex	11/M	14/F 7/NA [†]	13/M	17/F	6*-14/F
		Publication year	1870	1947	1981	1996	
		Author	Macnab ^[10]	McMurray ^[5]	Mabille et al.[7]	Jaovisidha et al. ^[8]	Present study

TB: Tuberculosis; NA: Not available; † Case not described; * At first presentation; STC: Soft tissue calcification; STS: Soft tissue swelling.

As seen in Table I, in the present time, the lack of familiarity with TTB in children comes from its' extremely low incidence and, therefore, data paucity in orthopedic literature. [3,5,7,8,10]

Tuberculous trochanteric bursitis can occur through hematogenous transmission from a remote site of infection or by direct spread from an adjacent infectious site. According to published data, the primary focus of TTB could be identified in nearly half of the cases.^[3] In our case, chest radiographs showed no abnormalities, and there were no MRI findings suggesting nearby structures as infectious foci. Moreover, no other present or past evidence of any other foci of TB were found. Thus, we assumed that hematogenous spread of *M. tuberculosis* bacilli after inhalation; however, without pulmonary involvement, it led to the development of primary TTB.

When analyzing our case, several clinical missteps were identified, starting with the initial presentation. At first, when a six-year-old child presents with swelling and fluctuance over the thigh, without history of trauma, the differential should include infection and tumor. Therefore, it is not surprising that the diagnosis of musculoskeletal TB is often mistaken as malignancy. [1] However, chronic wounds with secretion and minor pain should always raise suspicion on musculoskeletal TB. [1,5] Despite that, the treating orthopedic surgeon overlooked the lack of improvement on antibiotic therapy, misread pathology report, and approached this case as a recurrent pyomyositis.

In this case, the crucial event leading to proper diagnosis of TTB was, when the whitish liquid, resembling pus was noted intraoperatively, and TB smear was taken. A definitive diagnosis of TTB requires the obtaining of specimens from the trochanteric bursa for TB microscopy and culture, while the finding of granulomas and cultivation of *M. tuberculosis* are the gold standards for diagnosing TTB.^[11]

Optimal treatment for TTB remains debatable and involves mainly TB drugs, which are administered for 6 to 18 months, combined with surgical treatment.^[12] Although TB drugs can cure the disease at any stage, it is worth noting that, in some patients, the treatment of TTB with TB drugs alone may lead to TB recurrence.^[13]

In conclusion, we emphasize that TB still represents a diagnostic and treatment challenge, particularly in areas of low incidence. Therefore, it is a responsibility of orthopedic surgeons to consider TB as a possible source of disturbances in children, to avoid cases of neglected TTB in the future.

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