Case report

Miliary tuberculosis with empyema, a case report

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Abstract
We report a 39 year-old patient who presented with chest pain, malaise, lassitude, anorexia, weight loss, fever, chills, productive coughs, pleural pain in the left thorax area and septic empyema without hemoptysis. Laboratory investigations’ including tuberculosis (TB) skin test by PPD, HIV, HBV and HCV serologic tests were negative. The CBC showed anemia and leucocytosis. The ESR was elevated and he had hypoalbuminaemia. Both chest radiograph and high-resolution CT scan showed miliary infiltrates and diffused reticulonodular lung lesions. He was diagnosed with miliary tuberculosis (MTB) via direct staining, PCR and culture from open window region washing sample but not from broncho-alveolar lavage. He was treated with antituberculosis drugs.

Keywords: Miliary Tuberculosis, Empyema, Mycobacterium tuberculosis

Introduction
Miliary tuberculosis (MTB) is a type of disseminated and active tuberculosis (TB) that presents radiopathologic signs of TB micronodules as well as microbiologic findings [1]. MTB counts for 1 to 2% of all TB cases and about 8% of all forms of extrapulmonary tuberculosis in immunocompetent individuals [2]. Also, MTB is often under-diagnosed in the elderly, resulting in diagnosis of disseminated TB upon autopsy [3]. The chest radiograph plays an important role in the initial detection and final diagnosis of MTB. Positive sputum smear is found in about one-third of MTB patients [4]. Histological demonstration of granulomas in tissues (e.g. in lung, liver, and bone marrow) is usually required to make a rapid diagnosis. The molecular detection of Mycobacterium tuberculosis DNA by polymerase chain reaction is useful. IS6110-specific primer, which can amplify all M. tuberculosis complex strains, was used for amplification [5]. The main advantages of PCR are its rapidity, sensitivity and specificity [6].

Case history
A 39 year-old smoking man from a middle-class family with a rural background presented with chest pain, malaise,
lassitude, anorexia, weight loss, fever, chills, productive coughs, pleural pain in the left thorax area and septic empyema without hemoptysis. He used to live in Japan from 1990 to 2008. He came back to Iran in May 2008. His disease started in Japan and continued here in Iran. He had chest wall pain due to a left pleural effusion that was treated with antibiotics and a chest tube one year ago.

A bacteriologic examination of pleural fluid performed after chest tube showed the presence of methicillin-resistant *Staphylococcus aureus* and the patient received antibiotics according to the antibiogram. But he returned to our hospital with chest pain and the afore-mentioned signs and symptoms. TB skin test with PPD was negative. There were no laboratory findings indicating immunodeficiency or HIV. The HIV, HBV and HCV serologic tests were negative. The CBC showed anemia and leucocytosis. The ESR was elevated (82mm/hr Wintrobe’s method) and he had hypoalbuminaemia. Both chest radiograph and high-resolution CT scan showed miliary infiltrates and diffused reticulonodular lung lesions (Figs. 1, 2).

![Fig.1: Chest radiograph (X-ray) showing empyema in left lower lobe](image1)

![Fig. 2: High-resolution computed tomographic scan of the chest showing military infiltrates and diffuse reticulonodular lung lesions](image2)

On abdominal CT scans hepatosplenomegaly was confirmed. Pleural effusions necessitated surgical drainage. Broncoalveolar lavage (BAL) was carried out. BAL and washing samples from open window (Fig. 3) were processed for *M. tuberculosis* and polymerase chain reaction to rule out the possibility of mycobacterial infection. The direct smears were stained by standard Zeil Nelson staining technique. BAL samples staining in were negative for acid-fast bacilli. However, washing samples from open window were positive. The samples were subjected to the standard phenol chloroform DNA extraction process and were cultured on Lowen-Stein medium [6]. The extracted DNA was then amplified. PCR was carried out on open window region washing sample (OPWS) using IS 6110 insertion sequence-based primers yielding a 390 base pair (bp) product (Fig. 3) [6]. The PCR result from related washing samples from open window indicated the existence of *M. tuberculosis* infection. *M. tuberculosis* was grown after 40 days in Lowensten-Jensen medium. Immediately
after diagnosis of TB, oral antituberculous treatment with four drugs was started (isoniazide: 300mg/daily, rifampin 600mg/daily, pirazinamide 1.5g/daily, ethambutol 1g/daily) for two months and then continued with two drugs (isoniazide: 300 mg/daily, Rifampin 600mg/daily) for four months. Finally, in order to prevent bone tuberculosis, we continued treatment with two drugs (isoniazide: 300mg/daily, rifampin 600mg/daily) for two more months. The patient was treated completely.

Discussion
Extrapulmonary tuberculosis accounts for up to one third of all cases of tuberculosis [7]. The clinical manifestations and laboratory findings of MTB may be insidious and non-specific. [1,8,9]. In the current case, he had no specific clinical manifestations or laboratory findings. In miliary TB, hematologic abnormalities such as anaemia, leukopenia, leukocytosis, monocytesis, leukemoid reactions, thrombocytopenia, agranulocytosis and pancytopenia are well recognized but their significance is controversial. Erythrocyte sedimentation rate is usually elevated but albumin is frequently decreased in MTB [10].

In the present study, elevated ESR and hypoalbuminaemia were observed. These findings disappeared with treatment. Miliary infiltrates and radiologic features were present in our patient; these radiologic features have been reported in 40-100% of patients with miliary TB [8,10]. Preliminary diagnosis of TB is based on AFB in

![Open window region in chest of patient](image1)

![FLASH-PCR was carried out using using IS 6110 insertion sequence based primers giving 390 base pair(bp) product specific for Mycobacterium tuberculosis. Tub No1= Positive Control, Tub No 2= positive control, Tub No 3= Negative control, Tub No 6= patient’s sample](image2)

![Fig. 4: FLASH-PCR was carried out using using IS 6110 insertion sequence based primers giving 390 base pair(bp) product specific for Mycobacterium tuberculosis. Tub No1= Positive Control, Tub No 2= positive control, Tub No 3= Negative control, Tub No 6= patient’s sample](image3)
suspicious samples. Acid-fast bacilli have been seen in TB granulomas in 0-44% in previous studies [10]. Advances in molecular techniques have provided a new approach to the rapid diagnosis of TB by PCR. Depending on the clinical material, sample preparation method and target nucleic acid, the sensitivity of \textit{M. tuberculosis} detection by PCR varies between 42% and 100% [11,12]. TB for our patient in this study was diagnosed through OPWS which is an exceptional and rare specimen in the world. Our patient had AFB staining confirmed by PCR study and culture in Lowenstein Jensen medium.

**References**


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