**Abstract**

The *Mycobacterium terrae* complex, consisting of three saprophytic species, (*M. terrae*, *M. nonchromogenicum*, and *M. triviale*), they rarely cause diseases in immunocompetent people. There are few case reports of lung involvement by *Mycobacterium terrae* complex. We reported a 13 years old immunocompetent girl with a cavitary opacity seen on chest imaging and bronchoscopy revealed an endobronchial lesion, sputum and tissue specimens revealed growth of *Mycobacterium triviale*. Our patient is unique as it is the first case reported in Saudi Arabia with this type of *Mycobacteria*, in addition there is no case reported in the literature with *Mycobacterium triviale* with endobronchial lesion, the patient treated with a combination of rifampicin, ciprofloxacin and ethambutol for six months, she showed good response with resolution of radiological abnormalities and she did not have relapse after one year of follow up.

**Keywords:** Immunocompetent girl; *Mycobacterium triviale*, Bronchial asthma

**Case Report**

A 13 years old girl referred to King Khalid University hospital, Riyadh, from another hospital, she is a known case of bronchial asthma on PRN inhalers. She was admitted with 3 weeks history of productive cough, dyspnea, fever and chest discomfort. There was no history of hemoptysis, anorexia, weight loss or night sweats, no history of contact with sick person or exposure to pets or animals, she has no previous Hx of extra pulmonary infections. In the referring hospital she was managed as community acquired pneumonia with cefuroxim and moxifloxacin, the patient showed improvement of symptoms. In the referring hospital the sputum for AFB was also positive and microscopy showed long branching filaments suspicious of *Nocardia* and sputum PCR for *Mycobacterium tuberculosis* was negative and because of residual abnormality on chest X-ray and CT lungs which raised possibility of underlying lymphoma, the patient referred to our hospital for further intervention, the patient had history of Bell’s palsy, she left the school because of poor performance, she is living with her family in Riyadh with an average social class. Examination in our hospital revealed young girl, thin and short for her age, pale but looking well, fully conscious, not dyspnic, she is a febrile and other vital signs were normal, there was no clubbing or preferred lymphadenopathy, chest exam revealed decrease breath sounds on the right side otherwise normal, systemic examination was normal. Initial investigations revealed CBC: WBC 7.1 Hb 10 platelets normal ESR : 95, urea and electrolytes and LFT are normal, chest X-ray (Figure 1) showed right paratracheal and hilar lymphadenopathy which was confirmed on CT lung (Figure 2) showed right lower lobe consolidation with cavitation with bilateral hilar lymphadenopathy.

In our hospital, sputum for AFB was also positive and culture was positive for atypical *Mycobacteria* after one week bronchoscopy showed Shiny mass bleed to touch blocking most of the Rt main bronchus, BAL was done but Bx was not taken because of the risk of bleeding. BAL results revealed No AFB But culture was also +ve for atypical *Mycobacteria*, Cytology revealed several polymorphs consistent with inflammatory process because an underlying lymphoma couldn’t be ruled out the pt. underwent Rt side thoracotomy, which revealed multiple mediastinal casing lymph nodes which were eroding to the lumen of the Rt. Main and Rt. Upper lobe bronchi which were explored and lymph nodes specimens taken for investigation. Histopathology results of the specimens revealed: Reactive follicular lymphoid hyperplasia with histiocyte proliferation with mixed inflammation, no malignant cells, there are filamentous acid fast bacilli organisms suggestive of either nocardiosis or atypical mycobacteria (*Figure 3*). From TB lab Bx stain revealed:long filamentous AFB, Culture was positive for atypical mycobacteria which is sensitive to Rifampicin ,etambutol but resistant to isoniazid and Streptomycin and the specimens were send to another hospital for further processing. Pt was referred to infectious group who started the patient on Rifampicin, Clarithromycin, Ciprofloxacin and etambutol. Pt was discharged after 2 weeks in good condition.

After about one month, report of the molecular and biochemical processing of the specimen revealed *Mycobacterium triviale*. Patient was seen monthly in the thoracic surgery and ID clinic, clinically she was doing well and follow up chest X ray showed progressive resolution of the abnormal findings. In the first visit in the clinic, clarithromycin was discontinued and the patient was continued on rifampicin, ciprofloxacin and ethambutol for six months, she gained weight and ESR came back to normal, after completion of treatment she was followed for one year. She did not show any evidence of relapse either clinically or radiologically (*Figure 4*).

**Discussion**

The original description of the *Mycobacterium terrae* complex subspecies was by Richmond and Cummings in 1955 [1]. The *Mycobacterium terrae* complex includes *M. terrae*, *Mycobacterium nonchromogenicum*, and *Mycobacterium triviale* [2]. Members of the *M. terrae* complex are Runyon group III, non-pigmented, slow-growing organisms. Because of the presumed nonpathogenic nature of these organisms, there has been little effort to distinguish the species of this complex in the clinical setting [3]. *M. terrae* organisms are much...
less common mycobacterial pathogens than are other nontuberculous mycobacteria at any site of infection [4]. Up to 18% of isolates of nontuberculous mycobacteria that are recovered from humans are M. terrae complex organisms [5,6] and these isolates are rarely causing clinical disease, thus signifying nonpathogenic colonization [7]. But despite the common opinion that M. terrae complex isolates are nonpathogenic, these organisms are occasionally identified in the clinical laboratory in the setting of clinical disease of the joints, tendons (tenosynovitis), lungs, gastrointestinal tract, and genito-urinary tract.

There are few case reports of human disease that involve the lungs caused by M. terrae complex organisms [8-15], in a review by Scott Smith and colleagues of a total of 54 cases with M. terrae complex diseases, Lung disease was found in 26% of cases (14 patients out of 54), in many of the reports of the case series of this review members of M. terrae complex were not identified. In this review no predisposing factors are found in 44% of patients with disease caused by M. terrae complex and there was no correlation between the presence of a comorbid condition and an undesirable outcome in these cases [4]. Pulmonary infection with M. terrae can result in a cavitary process with noncaseating granulomas in tissue samples [16]. Following this review there is only one further case of Mycobacterium terrae involving the lung reported in Spain by Diaz ricoma [17] and another case reported by carbonara with disseminated M. terrae infection involving the lung in advanced HIV patient [7]. In the literature there is no case reported with lung disease by M. terrae complex; identified the member Mycobacterium triviale, though it may be part of the cases reported with M. terrae complex disease, in addition there is no case reported with M. terrae complex involving the lung with endobronchial lesion.

The optimum antimicrobial therapy for M. terrae complex has not been established [18]. In vitro antibiotic susceptibility testing is preferred for better selection of the best regimen of therapy, which may have to be continued for a prolonged period to increase the possibility of cure. Treatment with clarithromycin, ethambutol, and rifampicin was recommended and the duration of treatment needs to be for at least 12 months after clinical response [4].

Our patient had recurrent chest infections, which were probably due to post obstructive pneumonia due to obstruction of the right main bronchus by the endobronchial lesion, the patient acute symptoms responded initially to the pneumonia treatment; but her radiological abnormalities, her history of recurrent pneumonia and the presence of endobronchial lesion raised the possibility of chronic underlying disease and though the patient sputum grown atypical Mycobacteria which can alone explain the radiological findings, the possibility of underlying lymphoma still couldn’t be ruled out especially it could be a risk factor for this unusual infection there is reported cases of lymphoma with endobronchial lesion; we thought of this rare possibility in this patient and the growth of these organisms could be just a contaminant; but the possibility of a real infection with atypical Mycobacteria with underlying lymphoma could not be completely excluded.

Giving the patient medical treatment for atypical Mycobacteria and observing for the response was an option but because the treatment usually will takes long time it would not be a favorable option if the patient had an underlying lymphoma, so we preferred to perform right sided thoracotomy and exploration to take good biopsies mainly to rule out lymphoma in addition to relieve the bronchial obstruction.
In conclusion we add to the literature another case with lung disease involved by the member of M. terrae complex; Mycobacterium trivialae and it is the first reported case with endobronchial lesion with this organism.

References