Early Disseminated Mycobacterium Abscessus Complex Infection in an Infant with Coexisting Cystic Fibrosis and Progressive Familial Intrahepatic Cholestasis

Case report and literature review

Sanjeeewani A. Weerakoon,1 Maya Al Salti,2 Jahlila Mohsin,1 Hilal Al Hashami,3 Tawfiq Al Lawati,3 *Hussain Mohsin3

Abstract: Mycobacterium abscessus complex (MABSC) is a rapidly growing mycobacterium and may rarely cause disseminated infections in immunocompromised patients. In patients with cystic fibrosis (CF), it peaks between the ages of 11 and 15 years. We present a five-month-old infant with coexisting CF and progressive familial intrahepatic cholestasis (PFIC) who had pulmonary and cutaneous dissemination of MABSC infection. The management of this disseminated infection in an infant with two coexisting chronic diseases was challenging and resulted in the rapid deterioration of lung function and progression of PFIC to liver cirrhosis with a fatal outcome.

Keywords: Cystic Fibrosis; Atypical Mycobacterium; Mycobacterium Abscessus Complex; Progressive Familial Intrahepatic Cholestasis; Case Report; Oman.

Mycobacterium abscessus complex (MABSC) belongs to a group of rapidly growing mycobacteria (RGM) that can be clinically significant. Diseases caused by RGM have been increasingly reported during the past few decades due to various factors including improvement of molecular diagnostic methods and the increase in population at risk for non-tuberculous mycobacterial (NTM) diseases.1 MABSC is implicated in causing a wide range of clinical diseases including localised skin and soft tissue infections and disseminated infections such as osteomyelitis, lymphadenitis, cutaneous, lung and bloodstream infections, particularly in an immunocompromised host.1–3

The presence of underlying chronic lung diseases predisposes an individual to pulmonary infections.2 NTM has been increasingly reported as a significant pathogen among cystic fibrosis (CF) patients and has been linked to poor prognosis among this group.4–7

Progressive familial intrahepatic cholestasis (PFIC) is an autosomal recessive liver disease that can present with early-onset progressive liver failure.8

Herein, we report a case of disseminated infection due to M. abscessus in an infant with coexisting CF and PFIC.

Case Report

A full-term male infant who has a 10-year-old sibling with CF, was referred to a tertiary care hospital at the age of two months with a chronic cough for one month and failure to thrive. He was diagnosed with CF based on elevated sweat chloride level (>60 mmol/L) and genetic testing that confirmed the presence of a homozygous 3120 + 1G >A cystic fibrosis transmembrane conductance regulator (CFTR) gene mutation. The infant received intravenous antibiotics therapy and was started on pancreatic enzymes and fat-soluble vitamins. His throat swab culture was negative and he was discharged with an outpatient follow-up.

At the age of five months, the infant was admitted through the paediatric emergency department with a two-day history of fever, cough and difficulty in breathing. At the time of presentation, the infant was in respiratory failure with a respiratory rate of 70 breaths per minute, low oxygen saturation of 76% in room air, bilateral diffused crackles and wheezing. He required immediate tracheal intubation and mechanical ventilation. The infant looked icteric with a palpable liver 4–5 cm below the costal margin with no splenomegaly. Other systemic examinations were unremarkable.

The initial laboratory tests revealed a haemoglobin level of 12.1 g/dL with leukocytosis (predominantly neutrophils) at 26.2 × 10⁹ cells/L. He had elevated liver enzymes including alanine transaminase of 460 IU/L, aspartate aminotransferase of 768 IU/L, aspartate aminotransferase of 768 IU/L and alkaline phosphatase of 316 IU/L. The total bilirubin was 159 μmol/L (conjugated bilirubin at 121 μmol/L)
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Reaction (PCR) were all negative. The infant continued to experience low-grade fever over the second week and had progressive worsening of his respiratory parameters which prompted performing a flexible bronchoscopy and obtaining bronchoalveolar lavage (BAL) fluid. The BAL sample was sent for bacterial, mycobacterium and fungal cultures and extended viral PCRs. The BAL sample microscopy showed a positive result for acid fast bacilli (AFB) through Ziehl-Neelson staining, while the tuberculosis PCR test done using GeneXpert (Cepheid, Sunnyvale, California, USA) was negative, raising the suspicion of NTM. A computed tomography (CT) scan of the chest showed bilateral cystic bronchiectasis and nodules [Figure 1].

Given the lack of improvement on broad-spectrum antibiotics, the isolation of NTM from BAL and the CT changes, the child met the American Thoracic Society and the Infectious Diseases Society of America (ATS/IDSA) diagnostic criteria for non-tuberculous mycobacterial pulmonary disease. On day 20 of admission, anti-tuberculous drugs—rifampicin (15 mg/kg/dose once daily), ethambutol (20 mg/kg/dose once daily), clarithromycin (7.5 mg/kg/dose twice daily)—were commenced while meropenem was continued.

The child developed a confluent macular-papular skin rash involving the face, trunk, limbs, palms and soles but sparing the mucus membranes. There was neither evidence of desquamation, erythoderma or angioedema clinically nor any blood eosinophilia. Disseminated cutaneous infection due to atypical mycobacteria was highly suspected after excluding other causes including drug-related hypersensitivity reaction. Around the same time, the AFB positive peripheral blood culture was also positive for Gram-positive bacilli [Figure 2]. These were subsequently...
identified as *M. abscessus* by matrix-assisted laser desorption ionisation-time of flight mass spectrometry (MALDI-TOF MS). The central line blood culture remained negative for bacterial growth. The BAL culture was identified as *M. abscessus* through the line probe assay method using INNO-LiPA (LiPA Innogenetics, Ghent, Belgium). Azithromycin-based combination therapy with parenteral amikacin (18 mg/kg/dose once daily) and meropenem (imipenem was not available) was started as an intensive phase regimen based on the US Cystic Fibrosis Foundation (CFF) and European Cystic Fibrosis Society (ECFS) recommendations for eight weeks.\(^8\) The isolate was not tested for standard drug sensitivity due to technical limitations. Subsequently, the treatment was switched to the continuation phase that included oral azithromycin (10 mg/kg/dose once daily), levofloxacin (10 mg/kg/dose twice daily) and nebulised amikacin (250 mg/dose twice daily).

Furthermore, the infant had progressive cholestasis with elevated conjugated bilirubin and transaminitis. Based on a family history of PFIC in his first degree cousins, he underwent a liver biopsy which showed cholestatic hepatitis with fibrosis. A whole-exome sequence sample was sent to Centogene labs (Centogene N.V., Rostock, Germany), which found a homozygous mutation in the *ATP binding cassette subfamily B member 11* (ABCB11) gene, confirming the diagnosis of autosomal recessive PFIC type 2 in addition to CF. Given the early onset of NTM lung disease and the disseminated infection, the patient was extensively investigated for primary immunodeficiency disorders (PIDs) including Mendelian susceptibility to *M. abscessus* infection. No PIDs were found.

During his five-month stay in the hospital, the infant remained on respiratory support; mostly non-invasive ventilation with recurrent ICU admissions requiring intubation and mechanical ventilation. Multiple respiratory cultures obtained through the endotracheal tube and BAL remained negative for *M. abscessus* over the initial three months of treatment. However, a repeat BAL sample once again showed a positive result for *M. abscessus* in the fourth month of treatment. Two samples were positive for *Aspergillus* species with no other laboratory findings suggestive of allergic bronchopulmonary aspergillosis. The infant received four weeks of voriconazole (10 mg/kg/dose twice daily) that was switched to amphotericin B due to transaminitis. Despite being on ursodeoxycholic acid, his liver cirrhosis progressed and he developed portal hypertension.

Despite receiving anti-bacterial and anti-tuberculous therapy and the supportive PICU care, the infant developed an acute-on-chronic deterioration of respiratory status. This led to rapid respiratory failure and death after five months of inpatient hospital stay.

**Discussion**

Disseminated infections due to *M. abscessus* are rare and occur mostly in immunocompromised patients.\(^9,10\) The current patient was investigated thoroughly for PIDs but none were found. To the best of the authors’ knowledge, cases of coexisting conditions of CF and PFIC have never been reported before. Although CF is a known risk factor for NTM lung disease, it is not usually associated with disseminated NTM infection. Additionally, the onset of NTM lung disease in the infant was much earlier than what has been reported in the CF literature.\(^12,13\) PFIC type 2 typically results from a failure in the bile salt exporter pump (BSEP) protein which is responsible for exporting bile acids from the hepatocyte into canaliculi. This results in early-onset and progressive liver failure.\(^14\) Reviewing the literature, in general, PFIC or liver cirrhosis has not been shown to increase the susceptibility to NTM infection. It is possible that the coexistence of these two conditions resulted in a secondary immunodeficiency status which led to this disseminated infection. However, this conclusion cannot be made based on this single case.

The presence of at least one of the following characteristics would define the disseminated disease: cutaneous manifestations, involvement of more than one organ with or without cutaneous involvement, involvement of more than two groups of lymph nodes or positive blood culture.\(^12\) Cutaneous presentations can manifest as skin infections with discrete nodules, ulceration, multiple abscesses, diffuse maculopapular eruptions or, in contrast, as reactive skin lesions such as erythema nodosum, pustular psoriasis, generalised pustulosis and acute febrile neutrophilic dermatosis (Sweet syndrome).\(^14\) The current patient did have maculopapular skin eruptions or, however, though the lesions were not biopsied, the patient showed dramatic improvement after the commencement of anti-NTM agents.

Lymphadenitis is the most common presentation of NTM in immunocompetent children.\(^15\) NTM infects patients with underlying chronic lung diseases such as bronchiectasis, emphysema and healed tuberculosis. It has emerged as a frequently isolated pathogen among CF patients over the past few decades with prevalence rates of up to 20% in USA and 13% in Europe.\(^4,5\) According to the US CFF registry data (2016), *M. abscessus* was found to be the second most common isolate after *M. avium complex* accounting for 16% and 72% of positive cultures respectively.\(^15\) The isolation of *M. abscessus* from
sputum culture does not necessarily indicate infection. Accurate differentiation of lung infection from lung colonisation is challenging but important since the infection contributes to a more fulminant disease outcome in CF patients. Furthermore, the recovery of *M. abscessus* from routine bacterial cultures is often masked and easily missed due to overgrowth of other common respiratory tract pathogens and colonising bacteria. In order to diagnose NTM lung disease, the following criteria should be fulfilled: clinical (persistent respiratory symptoms), radiological (nodular and/or cavitary opacities on radiograph or bronchiectasis with nodules on CT scan) and microbiological evidence. Microbiological diagnostic criteria for NTM lung disease requires persistent growth of NTM in respiratory specimens: one positive AFB culture results in either two separate expectorated sputum samples or in one bronchial wash or lavage. The presence of a central venous catheter is a common predisposing factor for *M. abscessus* bloodstream infections. Identification of the NTM from blood culture may be difficult and often misdiagnosed with *Corynebacterium spp* which are usually considered to be skin contaminants. This case highlights the importance of careful interpretation of Gram-positive bacilli which helps in the timely management of NTM disseminated infections.

Accurate and urgent identification of NTM down to the species level is important in deciding the correct antibiotic choice. Different laboratory methods are used for the identification of NTM including biochemical tests, high-performance liquid chromatography and molecular tests. A reverse hybridisation assay such as ‘INNO-LIA Mycobacteria v2' (LiPA Innogenetics) is designed to amplify the mycobacterial 16S–23S rRNA internal transcribed spacer region. However, it cannot differentiate *M. abscessus* on a sub-species level. A recent study shows that MALDI-TOF MS can be used with promising results to identify NTM species including *M. abscessus* (concordance results with a reverse hybridisation-based assay showing 96.9% agreement).

Management of pulmonary and extra-pulmonary MABSC infection is faced with multiple challenges including intrinsic resistance to the common, standard anti-tuberculous agents, discordant *in vitro* susceptibility and clinical response, limited proven-to-be-effective drugs, serious side-effects and the long duration required for clinical and microbiological cure. To overcome these difficulties, a multi-disciplinary team which includes a chest physician, infectious disease expertise, clinical microbiologist, clinical pharmacist, physiotherapist, community nurse, the patient and his/her family is needed. The ATS/IDSA 2007 guideline recommends a multi-drug, macrolide-based regimen along with surgical resection, if possible, for both pulmonary and extra-pulmonary infections. As per the US CFF and ECFS consensus recommendations, the treatment is divided into an initial intensive phase (which can range from three to 12 weeks) followed by a continuation phase. The intensive phase includes an oral macrolide, intravenous amikacin and one or more of the following: intravenous tigecycline, imipenem or ceftolozin. The decision to switch to the continuation phase should depend on the response to treatment, the severity of infection and if the regimen is tolerated by the patient. The continuation phase includes an oral macrolide, inhaled amikacin and two to three out of the following antibiotics: minocycline, clofazimine, moxifloxacin and linezolid. Antibiotic choices should be guided but not dictated by drug susceptibility testing. Inducible resistance to clarithromycin is another challenge and it is due to the presence of the *erythromycin ribosome methyltransferase (erm)* gene in two of the *M. abscessus* sub-species: *M. abscessus subsp. abscessus* and *M. abscessus subsp. bolletii*. In comparison, *M. abscessus subsp. massiliense* does not have inducible resistance to clarithromycin due to the non-functioning *erm* gene, making therapeutic response to macrolide-based therapy more effective and efficient. Unavailability of recently validated techniques such as multi-locus sequence typing of *hsp65*, *rpoB* and *secA* genes for differentiation into sub-species levels, was an additional limitation in optimising antibiotic treatment in our case.

**Conclusion**

The current case report describes a fatal case of disseminated *M. abscessus* infection in an infant who had two chronic comorbid conditions of CF and PFIC along with the diagnostic and therapeutic challenges associated with case management. This report highlights the importance of having a high index of suspicion for NTM infection in children with CF presenting with early deterioration of lung function. More research with regard to the optimal management of NTM infection in this young population is required.

**AUTHORS’ CONTRIBUTION**

SW and MAS drafted the initial manuscript focusing on the microbiology aspect. HM added the clinical presentation and discussion sections. HAH reviewed the manuscript and updated the infectious disease aspect. JM reviewed and updated the microbiology aspect. TAL wrote and reviewed the liver disease aspects. All authors approved the final version of the manuscript.
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