CASE REPORT

A FIRST CASE REPORT OF TALAROMYCES MARNEFFEI INFECTION PRESENTING AS A NON-RESOLVING PNEUMONIA IN A NON-HIV DIABETIC PATIENT FROM BANGLADESH

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Abstract:
Talaromycosis is a systemic fungal infection caused by dimorphic fungus Talaromyces marneffei. It is endemic in Southeast Asia and generally complicates immunocompromised individuals, especially those infected by human immunodeficiency virus (HIV). Here, we report an autocthonous case of talaromycosis occurring in a non-HIV-infected Bangladeshi type 2 diabetic patient who presented with non-resolving pneumonia. To the best of our knowledge, this is the first talaromycosis case being reported from Bangladesh.

Key words: Talaromyces marneffei, nonresolving pneumonia.

Introduction:
Talaromyces marneffei, previously known as Penicillium marneffei, is a dimorphic fungus and can cause disseminated mycosis mainly in immuno-compromised individuals.1 Geographically it is limited in Southeast Asia and it commonly occurs in patients with human immunodeficiency virus (HIV) infection and rarely occurs in patients with normal immunity.2,3 However, the incidence of talaromycosis is gradually increasing in non-HIV patients. Here we report a case of talaromycosis who presented with non-resolving pneumonia.

Case Report:
A 58-year-old farmer, diagnosed case of diabetes with poor glycaemic control, presented with 1-month history of fever and cough. He did not have any history of traveling outside Bangladesh. Before presenting to our center, he required repeated admissions at different hospitals and was treated with broad-spectrum antibiotics without much benefit. Later on, empiric anti-tuberculosis drugs were added. He was febrile with features of left sided consolidation. He had neutrophilic leukocytosis (17400/cmm) and chest imaging revealed consolidation (Figure: la) with multiple cavity lesions (Figure 1b). Bronchoscopy showed a tongue like projection in left upper bronchus (Figure: 1c) and bronhoalveolar lavage was sent for bacterial and fungal identification.

Received: 03.12.2022 Accepted: 12.12.2022
DOI: https://doi.org/10.3329/bjm.v34i1.63430

Citation: Haque HF, Ahmed AKMS, Hoque T, Saha RC, Afroz F. A first case report of Talaromyces marneffei infection presenting as a non-resolving pneumonia in a non-HIV diabetic patient from Bangladesh. Bangladesh J Medicine 2023; 34: 62-64.

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culture, which revealed growth of *Pseudomonas* and *Talaromyces* respectively. Histopathological findings of bronchoscopic lung specimen (Figure: 1d) were consistent with the diagnosis of talaromycosis. Antibody against HIV appeared negative.

He was treated with injection colistin (according to sensitivity) and oral itraconazole. Other medications included insulin and paracetamol. Patient was discharged after two weeks with markedly improved clinical and radiological status. Antifungal treatment was continued up to 12 weeks.

**Discussion:**

The first reported human case of talaromycosis was in an American missionary with Hodgkin disease who lived in Southeast Asia. Almost 80% of the patients are immunocompromised like HIV infection and others are acquired cellular immune deficits such as transplant recipients, individuals with hematologic malignancies, those treated with steroids or cytotoxic agents, diabetes mellitus. *T. marneffei* can infect various organs like lung, liver and skin.

The clinical manifestations of talaromycosis vary from isolated skin lesions to respiratory failure and
circulatory collapse. Diagnosis can be made by demonstrating the characteristic morphologic findings of this fungus in biopsy material or in blood smears of patients with fungemia. The antifungal therapy causes clinical and microbiologic resolution of infection in up to 95 percent of patients. Our case was non-resolving pneumonia. Fungal culture, microscopic examination and histopathological study were done to establish the diagnosis and a long-term antifungal therapy was required for successful treatment outcome.

Conclusion:
We emphasize, careful history taking, physical examination and relevant investigations are required to reveal double pathology and to treat simultaneously for alleviating complications and mortality.

Consent: Informed consent was taken from the patient regarding publication of this case and accompanying images.

Acknowledgments:
We express our acknowledgements to the Department of Immunology and Histopathology, BIRDEM General Hospital, Dhaka, Bangladesh for support in laboratory evaluation of the patients.

Conflict of interest: Nothing to declare.

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