
**CASE REPORT**

**PULMONARY COCCIDIOIDOMYCOSIS PRESENTING AS A MASS, AN UNCOMMON DISEASE ENTITY IN ETHIOPIA**

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**ABSTRACT**

Coccidiodomycosis is a disease caused by the spores of the fungi coccidiodes immitis and pulmonary coccidioidomycosis comes after inhalation of the spores which are mainly found in desert areas of the United States, central and South America. Reported cases from outside the endemic areas have always history of travel to these areas. There are no reports so far from Ethiopia or the whole Africa. We report here a case of pulmonary coccidiodomycosis with no history of travel to such areas.

A 24 years old female patient from Samre, South-Eastern Tigray, presented with right side chest pain and productive cough of yellowish sputum which sometimes is blood streaked. She had completed anti-tuberculosis treatment without any improvement. With a preliminary diagnosis of pulmonary mass, surgical exploration was made and histology of the excised tissue showed appearances consistent with pulmonary coccidioidomycosis. There was marked clinical and radiological improvement after three weeks of treatment with ketoconazole. Though there are no reported cases from Ethiopia and Africa as a whole, Coccidiodomycosis should be considered as differential diagnosis especially for patients from arid areas like that of our patient before any empirical treatment.

**INTRODUCTION**

Coccidiodomycosis is a disease caused by the soil-inhabiting spores of the fungi coccidiodes immitis, which is often found in desert regions with incubation period of 7-21 days. About 60 % of infections cause no symptoms and the rest 40% symptoms range from mild to severe. Individuals with compromised immune status are at high risk for pulmonary coccidioidomycosis as well as for disseminated and cutaneous forms of the disease.

The disease can have an acute, chronic or disseminated form. Acute pulmonary coccidiomycosis is almost mild, with few or no symptoms and may resolves without treatment, while as chronic pulmonary coccidiomycosis can develop 20 or more years after initial infection and may present as lung abscess, empyema, bronchopleural fistula or scarring(fibrosis) including as mediastinal mass. (1,2,4)

Rates of infection are highest during dry months and are accentuated when soil is disturbed by windstorms. Exposure to contaminated balls of cotton or other fomites can result in infection beyond the endemic region, although rarely. Person to person transmission of pulmonary infection has not been reported. (5,6)

Coccidiodomycosis is usually asymptomatic. Those who become ill usually experience self-limited pulmonary syndromes. However, few patients develop complications or progressive forms of infection that display a broad variety of manifestations and pose difficult problems in management for the clinician (2, 3, 4)

Diagnostic workup for pulmonary coccidiomycosis includes proper history taking and physical examination, imaging studies with X-Ray finding of primary pulmonary disease including variable nonspecific infiltrates, hilar adenopathy and pleural effusions. Findings such as cavities and nodules demonstrate progression in to the complicated or residual stage of pulmonary coccidiomycosis. Although radiographic fea-
tures can be diagnostic, proof of coccidiodomycosis relies on histopathological, mycological and serological evaluation. Pathological diagnosis requires the demonstration of endosporulating spherules. (4,5,6).

Treatment options for coccidiodomycosis depend on the severity of the disease and the treatments generally consist of amphotericine B deoxycholate or azoles. Amphotericine is for the severe form of coccidiodomycosis while Ketoconazole, Fluconazole and Itraconazole are for mild form of the disease. (6-8)

Case presentation:

A 24 year old female patient from Samre, South-Eastern Tigray presented with productive cough of yellowish sputum which sometimes is blood stained, right sided chest pain and unquantified weight loss of 6 months duration. Complete blood count and organ function tests were normal. She was screened for HIV and was negative. Sputum was also negative for AFB. Culture and serology was not done due to limited set up. She completed anti-tuberculosis treatment without any improvement. Chest x-ray and CT scan showed right parahilar mass (Fig 1 and 2).

Fig.1: Pulmonary coccidiomycosis presenting as Right parahilar mass on CT-scan

Right thoracotomy was done with preliminary diagnosis of pulmonary mass. The patient made an uncomplicated recovery from her operation and was discharged after 5 days of hospital stay.

Histopathology of the excised tissue reported spherules with retractile walls and internal organizations that it are consistent with coccidiodomycosis. The patient had completely improved after treatment with ketoconazole 200mg twice daily for three weeks and was followed for six months which is the standard treatment.

On subsequent outpatient follow up the surgical wound had healed without any complications and had marked clinical and radiological improvement with no further symptoms (Fig 2).

Fig.2: Chest X-ray Pulmonary Coccidiomycosis before (a) and after (b) treatment with ketoconazole for three weeks.
Conclusion.: This particular case showed that coccidiodiomycosis could also be found in cases without any history of travel to the western hemisphere and that this disease, although rare, could be one cause of pulmonary disease in Africa, particularly in dry parts of northern Ethiopia. So the available diagnostic modalities should be carried out including thoracoscopic or open air biopsy if the facility permits before initiation of empirical treatment.

Consent: Written informed consent was found from the patient for possible publication of this case report and for the pictures posted.

REFERENCES