Idiopathic Pulmonary Fibrosis with Complications at Nnamdi Azikiwe University Teaching Hospital, Nnewi, Nigeria: A Case Report.

Anyabolu AE1, Enemuoh EH1, Ele PU1, Ugoeze FC1, Ufoaro CU1, Nwagbara CT1, Ukah CO1, Onwukamuche ME2, Ndukwe CO2.

ABSTRACT

Background: Idiopathic Pulmonary Fibrosis is a progressive diffuse lung disease, characterised by interstitial fibrosis, minimal inflammation, and architectural distortion interpersed with areas of normal lung tissue. It is rarely reported in Nigeria.

Aim: The aim is to present a case of idiopathic pulmonary fibrosis in our centre.

Methods: A review of a case of a 60 year old woman who presented to our centre with two year history of progressively worsening breathlessness and unproductive cough.

Results: A 60 year old foodstuff trader was managed in our center with two year history of progressive breathlessness and unproductive cough. She eventually developed complications of right sided heart failure and hypoxemia. The investigations she had included spirometry, chest X-ray, pulse oximetry, high resolution chest CT scan and lung necropsy. The results are all in keeping with idiopathic pulmonary fibrosis.She died after about 8 months steroid therapy as well as antifailure regimen.

Conclusion: Idiopathic pulmonary fibrosis is a chronic progressive fibrosing interstitial pneumonia of unknown cause. Lung transplant is the only effective treatment as response to medical therapy is generally poor.

Key Words: Idiopathic Pulmonary Fibrosis, Right sided heart failure, Hypoxemia

INTRODUCTION

Idiopathic Pulmonary Fibrosis (IPF) is a specific form of chronic progressive fibrosing interstitial pneumonia of unknown cause, occurring primarily in older adults, limited to the lungs and associated with the histopathologic and/or radiologic pattern of Usual Interstitial Pneumonia (UIP)1. It is the most common form of idiopathic interstitial pneumonia1 and the type with worst prognosis2. The prevalence of idiopathic pulmonary fibrosis in the USA is 14 per 100,000 and incidence of 6.8 per 100,000 but data is scarce on the epidemiology of IPF in Nigeria.

Development of IPF is influenced by both genetic and environmental factors. Family history of pulmonary fibrosis is strongly associated with increased risk of IPF3. Also, studies have shown increasing association with cigarette smoking4,5, environmental exposure to birds and cats6,7, gastro-esophageal reflux disease8,9, viral infection10 and occupational exposure to metal and wood dust11.

There is currently no evidence that any drug treatment improves survival of patients with IPF12. The prognosis of IPF is poor, with an estimated median survival of 2-5 years from diagnosis14,15.

CASE REPORT

A 60 year old female Nigerian of Ibo origin, who was a foodstuff trader, presented with a 2-year history of insidious onset progressive breathlessness and non productive cough. There was no history of fever, weight loss, night sweat, or contact with person having chronic cough. She had no chronic exposure to organic or inorganic dust. Though dyspnoea progressed to occur at rest, there was no orthopnea, paroxysmal nocturnal dyspnoea or leg swelling. She had no history of tobacco use.

Her physical examination findings included respiratory distress with respiratory rate of 28 cycles per minute; widespread fine end inspiratory crackles, pulse of 100 b/min and blood pressure of 140/80mmHg. Other systems were essentially normal.

Significant investigation findings included: Negative sputum smear examination for AFB; ESR of 4mm/hr; initial SPO2 of 94% and negative HIV screening test. Her spirometric result, as shown in table 1, depicted a restrictive pattern. Chest X-ray result showed loss of volume with crowding of ribs on the right, compensatory hypertrophy of the left lung, patchy opacification of most lung fields with reticulonodular background (figure 1).

Figures 2 to 5 are serial axial cuts of the patient's high resolution CT of the chest from the apex to the base. It shows diffuse involvement of both lungs with more apical sparing and basal pathology. There are reticulonodular opacities, marked peripheral interstitial

1Respiratory unit, Department of Internal Medicine, Nnamdi Azikiwe University Teaching Hospital, Nnewi, Nigeria.
2Department of Histopathology, Nnamdi Azikiwe Teaching Hospital, Nnewi, Nigeria.
Corresponding Author: Dr. Ugoeze Francis C. Respiratory Unit, Dept. of Medicine, Nnamdi Azikiwe University Teaching Hospital, Nnewi.
E-mail: ociso@yahoo.com.
thickening, cystic airway disease (mainly in the posterior segment) volume loss (more on the right) and regions of ground glass opacification characteristic of idiopathic pulmonary fibrosis.

She was commenced on tabs prednisolone 40mg daily (after meal), caps omeprazole 20mg daily, oral antibiotics when there was evidence of chest infection and supplemental oxygen when SPO₂ is < 92%. She was followed up forthnightly to monthly. She however progressively deteriorated despite treatment as she ultimately developed grade 2 digital clubbing as well as bilateral leg edema, raised jugular venous pulsation of 7cm, tender hepatomegaly and ascites suggestive of predominant right sided heart failure. She was managed in ICU as SPO₂ dropped to 63% but later died after 3days in ICU.

Lung necropsy result (figure 6 and figure 7) showed patchy interstitial fibrosis of varying intensity (ie some fibroblastic foci and other areas of dense collagenization) with interspersed areas of relatively normal lung parenchyma. The fibrotic areas show moderate inflammation consisting mostly of lymphocytes and a few plasma cells, neutrophils and eosinophils. It also showed alveolar destruction with cystic changes (honeycomb fibrosis) with smooth muscle hyperplasia and pulmonary oedema. All these are in keeping with usual interstitial pneumonia.

Table 1: Spirometry result.

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<th>PRED</th>
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<td>FVC</td>
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DISCUSSION

Idiopathic pulmonary fibrosis is commoner in older adults with the highest incidence in persons more than 50 years. In the United States of America, the age-adjusted incidence rate by both narrow-case and broad case finding criteria in a community based study was higher in men than in women with highest incidence among those aged 70-79 years.

Data are scarce on prevalence of idiopathic pulmonary fibrosis in Nigeria. This report represents one of the few cases of IPF reported in Nigeria and perhaps the first case confirmed by tissue diagnosis from South-Eastern Nigeria.

A surgical lung biopsy is required for definitive diagnosis of IPF. Limitation to extensive investigations in Nigerian patients with IPF (including lung biopsy) could partly account for the paucity of work on IPF in Nigeria. Also, the frequency of autopsy in tertiary centers is low due to many factors including sociocultural beliefs.

However, a diagnosis of IPF can be considered likely in the absence of a surgical lung biopsy specimen when certain major (4) and minor criteria (3) are met. This patient was extensively investigated with all findings in keeping with the diagnosis of IPF. Tissue diagnosis in this patient was possible probably because of her demise. The diagnosis of IPF in few published work done in Africa are based on high resolution CT scan.

There is no effective therapy for IPF. Corticosteroids alone were the mainstay for the treatment of IPF but the response to treatment is generally poor. Combined corticosteroid and cyclophosphamide therapy has no impact on survival in patients with IPF. Lung transplant is the only treatment with an impact on survival, but it concerns only a minority of patients and must be performed early in the disease process. Pifemidine, a novel antifibrotic and anti-inflammatory drug, reduces deterioration in the lung function and therefore represents an appropriate treatment option for patient with IPF.

Our patient was treated with steroid for about a year before she died. Her deterioration despite steroid therapy appears to corroborate the reported poor treatment response to steroids and poor prognosis of idiopathic pulmonary fibrosis.

CONCLUSION

Idiopathic pulmonary fibrosis is a chronic progressive fibrosing interstitial pneumonia of unknown cause commoner in older adults. It is rarely reported in Nigeria partly limitations due to extensive investigations required to establish the diagnosis. Lung transplant is
the only effective treatment as response to medical therapy is generally poor.

REFERENCES


