Case Report

Pulmonary Alveolar Microlithiasis (PAM)

Sarkar DN¹, Hossain M², Haque MM³, Zahin AKM⁴, Miah MS⁵, Ohab MA⁶, Hossain A⁷, Beauty S⁸

Abstract

PAM is a rare parenchymal Lung disease. Very few case report are available about PAM in Bangladesh. It is diagnosed incidentally during chest radiograph. It is a autosomal recessive disease and is associated with sporadic or familial mutation of SLC 34A2 gene. Many patients are asymptomatic and have either normal or restrictive pulmonary function. Some patients remain static and others progress into pulmonary dysfunction, respiratory failure and cor pulmonale. The disease is usually discovered up to 40 years and there is no definitive treatment of this disease. Chest radiograph, HRCT used lung biopsy (transbronchial or open) are the main investigations. This patient come with chest pain and dyspnoea on exertion and nonproductive cough and diagnosed as PAM incidentally during chest radiograph. On the basis of clinical features and laboratory finding, we diagnosed him a case of PAM a very rare condition. As there is no definitive treatment, we treat him symptomatically.

Case Report

A 25 years old businessman hailing from Haricharan, Pirgacha, Rangpur was admitted on Rangpur medical college Hospital on 17th September 2014 presented with chest pain for 5 to 3 years which is mild and not associated with exertion and is not radiated to another side and respiratory distress on exertion for 2 to 3 years which is gradually increasing. He has no recent history of respiratory tract infection and although he had a slight cough, sputum production was minimal. He has smoking history for 10 years and smoking 5 to 7 stick per day 3 to 4 years back but had stopped smoking because of increasing dyspnoea. He has past history of pulmonary tuberculosis 12 to 13 years back. Then he Completed CAT-1 anti-TB drugs. There is no history of consanguinity of marriage between his parents. General examination revealed patient was well looking having average body build according to his age, pulse-88 beat per minute, BP-120/80 mm of Hg, respiratory rate-18/min. He has no anaemia, jaundice, cyanosis or lymphadenopathy. All other Systemic examinations were revealed normal. Investigations reveals total count 8500/cmm, differential count- neutrophils 61%, lymphocytes 35%, monocytes 02%, eosinophils 02%, basophils 00%, haemoglobin 12.2g/dl, E.S.R 10mm at the end of 1st hour. (wintergreen method), random serum glucose 92mg/dl ,sputum for AFB- negative (2samples), mantoux test- negative, X-ray chest P/A view-shows suggestive of alveolar microlithiasis(figure I).

Figure I: Multiple milliary motling like shadow all over lung field
Echocardiogram shows good LV systolic function. CT Scan of chest suggestive of alveolar microlithiasis (figure II). Forced expiratory volume in one second of 74% or 2.93L. FEV1/FVC94%, PEFR=6.33 (68%)

Discussion

PAM is a rare disease that presents chronic evaluation, poorly defined etiology and pathogenesis and is basically characterized by numerous calcui(denominated calciferites, calcospherites or microlites) within air spaces1-9. Patients may remain asymptomatic for many years and do usually become symptomatic between the third and fourth decades3,6,10. At clinical presentation Patients usually demonstrate a lung disorder with restrictive pattern3,5-7,12. Adult patients commonly show progressive deterioration of the pulmonary function and dead usually occur in mid-life because of respiratory failure associated with corpulmonale7,11,12 and there is no important predominance of gender2,4,14. The disease presents a high incidence of familiar occurrence (approximately one third of the cases) suggesting an autosomal recessive inheritance pattern2,5,8,10. Most reported cases of pulmonary alveolar microlithiasis have occurred in turkey, Japan and Italy, However the disease is not region specific, it may be seen anywhere in the world21,22. Although the aetiology of PAM remain unclear, mutations of SLC 34A2 gene, which encodes a type to sodium dependent phosphate co-transporter (Napi-lb) are considered to be the cause of the disease14 SLC34A2 is primarily expressed in the lung and here only in alveolar type 2 cells14,15. These cells are responsible for production of surfactant. Loss of function of the gene due to mutations may lead to a decreased cell uptake of phosphate, which in turn may lead to formation of intra-alveolar microlithiasis as a result of phosphate chelating calcium in the extracellular fluid14,16. Although the SLC 34A2 gene is involved in phosphate homeostasis in several organs including the lung16,17, blood levels of calcium and phosphate are usually normal in PAM13,18,19,20. PAM is often diagnosed early in the clinical course of disease, although the result of the initial laboratory work up are often unremarkable, with no identifiable underlying disorder of calcium metabolism23. However, even with early recognition of the disease, the lack of treatment option results in a poor long term prognosis22,23. Although patients with PAM often experience a protracted disease course, they may be asymptomatic for years before respiratory insufficiency become manifest21,22, typically as seen in our case, patients present with progressive dyspnea, a restrictive pattern of lung disease and decreased diffusion capacity21,22,23. Respiratory insufficiency eventually progresses to pulmonary fibrosis, end stage lung disease and chronic pulmonary heart disease. Extra pulmonary involvement is uncommon22. Death is often due to a combination of pulmonary dysfunction and subsequent corpulmonale22,23. On radiographs, PAM is characterized by diffuse fine calcific micro nodules that involve both lung in a pattern that is classically described as sandstorm-like24,25,26,38.

Increased calcific densities are often move pronounced in the lower lung zones, a fact that has been attributed to the larger surface area and greater thickness of the lower part of the lungs27. High-resolution CT with thin section acquisitions and high spatial frequency, reconstruction algorithms is preferred for the evaluation of pulmonary alveolar microlithiasis because it allows detection of minimal structural changes of the lung parenchyma that are not optimally evaluated with radiography or with other CT techniques. The characteristics findings of extensive innumerable microlithiasis involving both lungs is noted with a predisposition for the posterior segments of the lower lobes and anterior segments of the upper lobes26-31. Additionally, the medial aspect of the lung appears to be more heavily involved than the lateral aspects. Microcalcifiths can also be seen along the bronchovascular and sub pleural interstitium, resulting in a thickened, micro nodular appearance of these structures26. Calcifications of the pleura also have been reported27. No definite treatment is available. Home oxygen therapy is necessary for the patients with respiratory insufficiency. Systemic corticosteroid and bronchoalveolar lavage have been shown to be ineffective. In general, No therapy has proved beneficial including whole lung lavage32,33. Lung transplantation has been performed in a few patients. Some patients have under gone bilateral sequential lung transplantation or unilateral lung transplantation34,35,36. Disodium etidronate, which is known to inhibit the microcrystal growth of hydroxyapatite has been used in the dose of 10 mg/kg per day orally for as long as on year with considerable regression of the calcific densities27.

In conclusion, PAM is diagnosed incidentally during chest radiograph. Most cases are asymptomatic for many years. But it may progress to corpulmonale of any other deteriorating evaluation. For this, patient need follow up examination for long time.
References


