Pulmonary alveolar microlithiasis with involvement of the sympathetic nervous system and gonads

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Pulmonary alveolar microlithiasis is a relatively rare disease, of obscure aetiology, manifested by the presence of intra-alveolar laminated concretions distributed diffusely throughout the lungs. In all the recorded cases disease has been limited to the lungs and similar lesions have not been observed in other organs or tissues.

The findings in a patient in whom lesions identical to those in the lungs were confirmed in the lumbar sympathetic chain and suspected to be present in the testes are recorded.

CASE REPORT

The patient, an African man aged 24 years from the Griqualand East area of South Africa, came under medical care in October 1963 when he presented with an attack of acute abdominal pain. No diagnosis was made. He had a similar attack three months later. Examination was largely negative. Radiographs of the pelvis showed scattered lines of calcification, not characteristic of that occurring in bilharzial involvement of the bladder. Laparotomy revealed no gross abnormalities. Because the possibility of a recurring pancreatitis could not be excluded, a sphincterotomy was performed.

He was seen by the author four months later, during an attack of abdominal pain associated with diarrhoea, vomiting, and urinary difficulty. He responded slowly to conservative treatment and it was decided to undertake further special examinations in an attempt to establish a diagnosis.

Special interrogation revealed that the patient had three brothers and three sisters. One sister had died at 23 years of age of a lung disease, but no details were available. Two brothers were available for medical examination: they were both in good health and chest radiographs were normal. There was no history of taking snuff or chewing tobacco. He smoked and drank beer in moderation. There was no opportunity for close contact with bats.

The patient appeared thin but was otherwise in good physical condition. There were no significant findings on general examination apart from slight hyperextendibility of the finger joints. Clinical examination of the respiratory, cardiovascular, and nervous systems revealed no abnormalities. The abdomen showed normal findings on examination.

The chest radiograph showed miliary opacities scattered throughout both lungs (Fig. 1). Repeated sputum examinations were negative for tuberculosis and showed no malignant cells. An intravenous pyelo-
FIG. 1. Chest radiograph showing diffuse miliary mottling.

FIG. 2. Linear opacities in paravertebral position with thickening at intervertebral level.

FIG. 3. Diffuse miliary mottling outlining both testes.
FIG. 4. Section of the lung showing intra-alveolar laminated concretions.

FIG. 5. Section of sympathetic ganglion (high power) showing laminated concretions. Sympathetic ganglion cells are seen.
inspection but was distinctly gritty on palpation. Skin and muscle biopsies were taken.

The patient made an uneventful recovery after the operation and when he was discharged from hospital in October 1964 had gained some weight. He was not seen again but his brother reported that he had died at home during March 1965, his death having been preceded by further attacks of abdominal pain. There was no indication of cardiac disease or heart failure prior to death.

Lung biopsy showed the histological appearance characteristic of pulmonary alveolar microlithiasis: many alveoli were filled with concentrically laminated microliths within the alveolar lumina. No inflammatory reaction was present (Fig. 4). The sympathetic chain and ganglia showed a heavy calcium deposition in the form of calciospherites similar to those in the lung (Fig. 5). The skin and muscle biopsies were normal on histological examination.

**DISCUSSION**

Pulmonary alveolar microlithiasis was first described more than a hundred years ago (Friedreich, 1856a, b, c). It was first so called by Pühr in 1933. Cases have been reported from most parts of the world. The present case is probably the first one from southern Africa. The disease affects males and females in roughly equal numbers. The ages of reported cases have ranged between 6 and 72 years. More recently, several asymptomatic cases in children have been described, often discovered on routine chest radiographs of the siblings of known cases.

The disease is a chronic one and probably always has a long asymptomatic phase. The condition is known to have been present for 25 years (Manz, 1954). Most cases are asymptomatic when discovered, usually during medical examination for employment or military service.

The aetiology and the nature of the condition have been much discussed. Many examples of familial incidence have been reported. Amongst the 26 cases described by Sosman, Dodd, Jones, and Pillmore (1957) there were five instances in which several members of one family were affected. This has suggested the possibility of heredity, environment, diet or contagion as aetiological factors. No consistent bacterial or other (e.g., fungal) infection has been demonstrated. The calcium metabolism has been specifically screened in many cases. Mikhailov (1954) found an elevated serum calcium in three cases. Renal stones have been reported in several cases (Badger, Gottlieb, and Gaensler, 1955; Sosman et al., 1957; Portnoy, Amadeo, and Henninger, 1964). O’Neill, Cohn, and Pellegrino (1967) did a particularly careful study of the calcium metabolism in three members of a family and found no derangement. No instance of parathyroid disease has been reported in a patient with microlithiasis. Chinachoti and Tangchai (1957) related the disease to the taking of snuff containing calcium, in Thailand. There has been no constant or consistent association with any type of occupational exposure, antecedent disease or involvement of other organs.

Involvement of other organs by a similar disease process has not been recorded. Sosman et al. (1957) described the finding of calcified nodules in the spleen in one of their cases. They considered these to be probably incidental, possibly due to previous infection with *Histoplasma capsulatum*. Two of Mikhailov’s (1954) cases showed a chronic nodular inflammatory process in the liver.

The disease has been regarded as an exudative process, the exudate providing a suitable intra-alveolar medium for subsequent calcification. A continuing process is suggested; new microliths continue to form and those present increase in size as the disease progresses. There is no reaction in the lung to the presence of the microliths.

The disease is usually asymptomatic, usually an incidental radiographic finding. Symptoms, when they arise, are those of pulmonary insufficiency with, finally, symptoms of heart failure due to cor pulmonale. There are no characteristic physical findings and the results of special laboratory examinations are essentially non-contributory.

The diagnosis is suggested by the radiological appearance of the lung fields. There are diffuse, fine, sand-like or granular opacities spread uniformly throughout both lungs. The granules are usually less than 1 mm. in diameter. They appear denser at the lung bases because of the greater depth of lung tissue. The density of the nodules is that of calcium and it is this calcific density, even more strikingly demonstrated under magnification of an over-exposed film, in the total absence of clinical signs and symptoms, that differentiates microlithiasis from other miliary conditions which may simulate it. Miliary tuberculosis, miliary metastases, sarcoidosis, pulmonary haemosiderosis, pulmonary eosinophilia and other rarer miliary diseases can usually be recognized by their appearance and distribution and the associated physical and laboratory findings.

Dust diseases, especially stannosis and berylliosis, have a different appearance and distribution and there will be a history of exposure.

The diagnosis of microlithiasis is confirmed by lung biopsy. The lung feels distinctly gritty on palpation. The microscopical appearance of
Pulmonary alveolar microlithiasis


Okuno, K., Chihaya, M., Kumamoto, S. et al. (1966). Two familial cases of microlithiasis alveolaris pulmonum. Runsho Hosha 2, 44.


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