Pulmonary sarcoidosis with an alveolar radiographic pattern

JP BATTESTI, G SAUMON, D VALEYRE, J AMOUROUX, B PECHNICK, D SANDRON, R GEORGES

From the Service de Pneumologie, Hôpital Avicenne, Bobigny, France, and the Groupe de Recherches U 82, INSERM, CHU, Xavier-Bichat, Paris

ABSTRACT  Thirty-three cases of sarcoidosis (4.4% of 746 patients) showed an alveolar radiological pattern. A study of pulmonary function was carried out in 25 patients and compared with that of 46 patients with the interstitial radiological type of sarcoidosis. Twenty-two cases have been followed up from one to six years after the initial examination. The radiographic lesions were most often bilateral and included nodules greater than 15 mm with ill-defined margins or diffuse, infiltrative, non-retractile opacities with fluffy margins. Bilateral mediastinal lymph nodes were present in 27 patients. In 20 patients an associated reticulation was found on radiography. In four patients an open lung biopsy was done. The granulomatous nodules were identical to those found in other forms of sarcoidosis, although they were more confluent in the affected areas. Clinical and functional findings did not differ from those in the more common forms of sarcoidosis. Alveolar sarcoidosis has a sudden course. The alveolar radiological patterns always disappeared, with or without steroid treatment, while reticular patterns persisted in four patients. Rapid radiological changes were observed. Some functional abnormalities persisted in cases that were followed. It is concluded that alveolar sarcoidosis is a distinct acute form of sarcoidosis.

Sarcoid granulomas are found in the alveolar septa and less frequently in the walls of bronchi, pulmonary arteries, and veins.¹ The radiological findings in pulmonary sarcoidosis may typically be of one of three types of diffuse interstitial opacities²: reticular, reticulomicronodular, or nodular (nodules less than 5 mm in diameter). Less frequently the radiological pattern is called “alveolar.”² The aim of the work reported here was to determine whether sarcoidosis with the alveolar radiological pattern is a distinct type of sarcoidosis.

Methods

Seven hundred and forty-six patients with pulmonary sarcoidosis were studied. The radiological criteria for alveolar manifestations were those of Felson²: opacities of the infiltrative type, non-retractile, with ill-defined margins and sometimes an air bronchogram, or nodules with ill-defined limits and diameters greater than 15 mm. Thirty-three patients (4.4%) satisfied those criteria. A study of pulmonary function was carried out in 25 of the 33 patients and the results were compared with those of 46 patients with the more usual interstitial radiological pattern.

The pulmonary volumes and the forced expiratory volume during the first second (FEV₁) were measured by spirometry. The functional residual capacity was measured by multiple-breath helium dilution. The standard values used were those of the European Communities.³ The transfer factor was evaluated by the single-breath carbon monoxide test (TLCO).⁴ The standards used have been published.⁵ The TLCO was scaled according to age and height (indicated as TLCO Ht).

The static ventilatory mechanics were estimated through the relation between the volume measured at the mouth and the transpulmonary pressure during slow expiration. The following indices were calculated: static expiratory compliance (CL) measured at tidal volume, elastic recoil at total lung capacity (PI max), and coefficient of retraction (CR).⁶ The following standards were used: Yernault et al⁷ for CL/TLC and values obtained by Turner et al⁸ for CR. The measurements of arterial oxygen tension (PaO₂) were made on blood obtained from the brachial artery.
Lung function was measured at rest in the sitting position. All data are presented as means and standard errors of the mean unless otherwise noted.

Results

CLINICAL FINDINGS
Of the 33 patients studied, 23 were men and 10 women. The mean age was 28.4 ± 1.85 years. One patient was under 20 years; 30 were 20–40 years; and 2 were over 40. There were 26 white and seven black patients (including two Africans and one North African).

In 21 patients (63.5%) the discovery of the radiological patterns had been made after systematic examination. Respiratory symptoms were not present in these patients. In 12 patients (36.5%) radiography was prompted by respiratory symptoms (cough, breathlessness during exertion, chest pain). In nine patients an x-ray examination had been done a year before; the appearances were normal in seven cases and showed enlarged hilar nodes in two cases. The clinical examination of the thorax gave negative results in all but one patient, in whom crackles were noted. Clinically, extrathoracic sarcoidosis was found in 13 patients; the lymph nodes were affected in nine, the skin in one, the parotid glands in one, the liver in two.

RADIOLOGICAL FINDINGS
The opacities of the alveolar type were usually bilateral and present in several zones. In 17 patients there were large nodules with ill-defined margins (fig 1), while 16 patients had diffuse infiltrative opacities, sometimes with an air bronchogram (fig 2). Spread to the middle and peripheral zones of the lung was the most frequent finding. The lesions extended to the costophrenic angle in only two patients, of whom one showed a pleural reaction. The presence of a clear zone inside the nodules was observed in six of the 10 patients in whom tomograms were performed. Bilateral mediastinal lymph nodes were present in 27 patients.

In 20 patients in addition to the alveolar patterns a reticular or reticulomicronodular pattern was observed. The patients showing only alveolar patterns constitute group A, while those showing alveolar and reticular patterns form group B.

PULMONARY FUNCTION
In 25 of the 33 patients respiratory function was studied when the diagnosis of sarcoidosis was made. The pulmonary function of 46 patients with an isolated interstitial radiological pattern was compared (group C) with that of the patients of group A and B. The age and sex of the patients studied, their

Fig 1 Chest tomogram showing large nodules with ill-defined margins and bilateral mediastinal lymph nodes.

Fig 2 Chest tomogram showing diffuse infiltrative opacities with air bronchograms and bilateral mediastinal lymph nodes.
smoking habits, and the main functional measurements are presented in the table. Vital capacity (VC), total lung capacity (TLC), and TLCO Ht were significantly diminished in the three groups. The other values, in particular the FEV1/VC, were not significantly changed. Nevertheless, when each case was studied an obstructive syndrome (FEV1/VC < 90% predicted value) was observed in five of 11 cases in group A, two of 14 cases in group B, and five of 46 cases in group C. Transfer factor impairment was frequent: TLCO Ht = predicted value −2 SD in five out of 11 cases in group A, nine out of 14 cases in group B, and fifteen out of 46 cases in group C. There were significant differences between groups A and B in the coefficient of retraction (eight from group A: mean = 3.8 ± 0.3; eight from group B: mean = 7.5 ± 0.7, p < 0.001; group C: 5.8 ± 0.3).

PATHOLOGICAL FINDINGS

Four patients required an open lung biopsy for a diagnosis to be made. One of the samples, which was done outside the infiltrated zone, showed lesions localised exclusively in the bronchial mucosa and peribronchial area. The three other samples contained large numbers of confluent masses of granulomas and between them rare bronchial or alveolar spaces bordered by a cuboidal epithelium. These follicles appeared to arise in the interstitium but protrude into the alveolar spaces, thus occluding them to a variable extent. In two cases, between large confluent granulomas small zones of normal pulmonary tissue were seen without any specific damage or sclerosis of the alveolar walls. Follicular lesions were observed in the wall of a small pulmonary artery and in the wall of a vein (in two cases).

NATURAL HISTORY

The course of the disease was followed radiologically in 22 patients. In 15 of them impaired respiratory function required steroid treatment for two years. The alveolar type patterns disappeared in all 15 and did not reappear for one year after the end of the treatment. On the other hand, the reticular patterns observed in 10 of the 15 are still present in three. Six of seven untreated patients (five in group A, one in group B) showed clearing of the radiological abnormality after three to nine months. In one the alveolar and reticular patterns are still present after 17 months, treatment being contraindicated. A rapid change in radiological appearance was seen often in a matter of weeks. Changes of respiratory function could be followed in only 10 patients, for periods varying from one to six years. A significant improvement of TLC and VC was observed (TLC % of predicted: mean increase = 13.5%, t = 2.34, p < 0.05; VC % of predicted: mean increase = 11%, t = 2.54, p < 0.05). The other values did not change significantly.

Discussion

There is a difference, according to most recent publications, in the frequency of the radiological alveolar pattern found in sarcoidosis. Considering only the nodular pseudotumoral forms, Romer estimated it to be 1%, while Kirkps found it in 2% and Sharma in 4% of their subjects with sarcoidosis. Shigematsu, considering all the alveolar patterns of sarcoidosis, estimated the frequency of this form to be 20%. Felson thought that sarcoidosis was the most important cause of disseminated chronic alveolar opacities. In our study the alveolar forms occurred in 4-4% of patients with the disease.

Whether this radiological pattern, as Shigematsu suggested, is related to a particular form of sarcoidosis requires correlation studies between the radiological and the pathological data. Unfortu-
nately such studies are rare and are interpreted differently. According to Sahn,13 the alveoli are invaded by mononuclear cells, which represent a non-specific reaction to the sarcoid granuloma present in the pulmonary interstitium. According to Shigematsu12 the alveoli contain epithelioid cells whose origin—endoalveolar or interstitial—is uncertain. According to Felson2 and Reed,14 the alveoli are compressed or filled with coalescent interstitial nodules.

The pathological observations made on lung biopsy specimens in four of our patients indicate that the granulomas are identical to those found in the common forms of the disease, but less diffuse and more confluent. Thus the radiological aspect of alveolar sarcoidosis appears to be due not to histological disorders of the alveoli but rather to the collapse of the alveolar walls by the confluence of interstitial granulomas. The cause of cavitation observed in some cases of alveolar sarcoidosis is unknown.15 Ischaemic or eosinophilic necrosis of conglomerate granulomas has been suggested.16 Nodules with cavitation were not observed in our biopsy specimens. The fact that in two cases we observed small zones of normal pulmonary tissue between large confluent granulomas suggests that the clear zones observed in the nodules' centres might correspond to undamaged parenchyma, surrounded by coalescent granulomas.15 The air bronchogram can occur in interstitial diseases such as sarcoidosis in this alveolar variety.14

Clinically, alveolar sarcoidosis seems to be an acute form, to judge by the young age of the patients, the presence of mediastinal nodes, and the frequent occurrence of a normal radiograph during the previous year. But the clinical findings are similar to those of the other acute forms of sarcoidosis. Likewise, the abnormalities of pulmonary function observed in the present group of patients did not differ from those observed in other patients with different radiographic patterns.17,18 The combination of reticulation with the alveolar-type patterns should be taken into consideration. This radiological feature is most probably due to the extension of the granulomatous process into the interstitial tissue. In these cases we have observed more severe impairment of the coefficient of retraction. This can be explained by the association of diffuse interstitial lesions and lung shrinkage.19

The course of the disease was characterised by the improvement of the alveolar pattern either spontaneously or with corticosteroid treatment. Interstitial patterns can, however, persist even with treatment. Spontaneous radiographic clearing (six out of seven patients) was more frequently observed with the alveolar pattern than with other forms of acute sarcoidosis.20

Rapid changes of radiological patterns are characteristic of alveolar sarcoidosis.21,22 The pathological findings cannot explain this. In fact, it is hard to conceive that confluent granulomas seen in the biopsy specimens, which seem to cause the radiological pattern, can improve spontaneously so rapidly. Although the radiological patterns disappear, more often the functional measurements do not change when they are already abnormal at the first examination. Indeed, even though the pulmonary volumes did in some cases improve, compliance and the result of the single-breath carbon monoxide test did not change, at least during the period of observation. Similar observations concerning pulmonary volumes have been made by Onal et al23 in cases of nodular pulmonary sarcoidosis with radiographic resolution.

References


Pulmonary sarcoidosis with an alveolar radiographic pattern.

J P Battesti, G Saumon, D Valeyre, J Amouroux, B Pechnick, D Sandron and R Georges

Thorax 1982 37: 448-452
doi: 10.1136/thx.37.6.448