Chronic pulmonary paracoccidioidomycosis masquerading as lymphangitis carcinomatosa

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Many pulmonary disorders may produce a non-specific granulomatous reaction. We describe a case of unsuspected chronic pulmonary paracoccidioidomycosis that emphasises the importance of performing fungal stains on lung biopsy specimens in all cases where an interstitial granulomatous reaction is present without an obvious cause.

Case report

A 57 year old caucasian female clerk was admitted for investigation of possible lymphangitis carcinomatosa. She complained of cough and of breathlessness and wheeze on exertion, which had developed gradually over a period of three years. Four years earlier she had had a left mastectomy for an intraduct carcinoma of the breast; this was grade T1 N0 M0, although at subsequent follow up she had not developed clinical metastatic disease. She had smoked cigarettes (total 60 pack years). She had worked as a clerk in Buenos Aires (1961–4) and Caracas (1965–7) but had not revisited South America in the last 17 years.

There were no abnormalities on examination. Investigations showed: haemoglobin 15.7 g/dl; white cell count $8.3 \times 10^9$/l (differential count neutrophils 65%, eosinophils 4%, lymphocytes 28%, monocytes 3%); erythrocyte sedimentation rate 17 mm in one hour; blood urea and electrolyte concentrations normal; liver function normal; serum calcium normal; aspergillus and avian precipitins negative; serum angiotensin converting enzyme activity 83 (reference range 16–52) nmol ml$^{-1}$ min$^{-1}$.

A chest radiograph showed diffuse bronchocentric interstitial shadowing in both mid and lower zones (fig 1). Retr-
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Paracoccidioidomycosis (or South American blastomycosis) is a systemic mycosis caused by Paracoccidioides brasiliensis and endemic in many parts of South America. It affects mainly agricultural workers with a male:female ratio of 50:1. Cases have been divided into four groups:

(a) acute pulmonary—with multiple cavitating lesions mimicking tuberculosis;
(b) chronic pulmonary—with a progressive fibrosis affecting both lower zones;
(c) acute disseminated—with skin papules, gastrointestinal infiltrations, rectal bleeding, lymphadenopathy, and hepatosplenomegaly;
(d) chronic disseminated—with oral ulceration and lymphadenopathy.

Types (a) and (c) are rapidly fatal if untreated but types (b) and (d) run a course over many years. The diagnosis is made by smear examination and culture of the morphologically distinct budding yeast forms of P. brasiliensis from sputum or oral ulcers, and is confirmed serologically by complement fixation tests. Treatment, which was relatively unsuccessful with amphotericin and sulphonamides, has been revolutionised by the introduction of ketoconazole, which has improved the response rate from 50% to 95%.2

The patient described here is atypical, being a female urban dweller. She falls into clinical category (b) (chronic pulmonary) with a latent period of at least 17 years. Nine cases of chronic pulmonary or disseminated paracoccidioidomycosis have been described in the United States.3 All the patients were men, had lived in South America or Mexico, and had latent periods of from three to 20 years after leaving the endemic area. Our patient had raised activity of serum angiotensin converting enzyme. Although this has been documented in a range of granulomatous conditions, including some fungal diseases,4 we believe this is the first report of increased serum angiotensin converting enzyme activity in paracoccidioidomycosis.

We believe that this case illustrates two important points. Firstly, it is essential to carry out silver stains for fungi on biopsy material in any patient with an unknown granulomatous condition, especially when the patient has visited an area where a systemic mycosis is endemic. Secondly, in a patient with a resected breast carcinoma causes of reticular basal shadowing other than lymphangitis carcinomatosa should be considered, especially when there is no additional evidence of metastatic disease.

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References

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