Acute fulminating intrapulmonary haemorrhage in Wegener’s granulomatosis

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Diffuse pulmonary capillaritis is a rarely described feature of Wegener’s granulomatosis. We report a case in which this appears to have been the cause of the acute fulminating course of the disease.

Case report

A 63-year-old retired school caretaker was admitted to hospital with a three-week history of painful mouth ulceration and hoarseness. He had a cough productive of blood-streaked sputum and had become increasingly breathless on exercise over the same period. He had mild generalised joint pains and had lost 7 kg in weight.

A routine chest radiograph in 1967 was normal. A further radiograph in 1974 showed some ill-defined shadowing at the right apex. Tomography showed scarring consistent with old tuberculosis, and the lesion did not progress. In 1978 he was treated for bilateral uveitis. In 1979, one year before his admission, a miniature radiograph, requested because of “recurrent chest infections” and haemoptysis, showed no change since 1974, and his symptoms settled.

There was no significant family history or occupational exposure to noxious or allergic agents. He smoked 30 cigarettes a day. On questioning he admitted that genital ulcers had developed four weeks previously.

Physical examination revealed a bright, cheerful man who was thin and anaemic. He had painful ulcers on the soft palate, gums, penis, and scrotum. There were inspiratory crackles at the right lung base. The liver edge was just palpable. The chest radiograph showed bilateral reticulo-nodular shadowing, more marked on the right side. A cavitating lesion was noted at the right apex.

Further investigations showed a haemoglobin of 7·3 g dl⁻¹ with a microcytic film; white cell count 11·3 × 10⁹ l⁻¹ with normal differential; ESR 143 mm hr⁻¹; blood urea 14·3 mmol l⁻¹; electrolytes and liver function tests normal; urine granular casts only; blood, sputum, and urine cultures negative; sputum smears for acid-fast bacilli negative; rheumatoid factor, ANF, anti-DNA antibody, and Australia antigen negative; serum immunoglobulins normal; viral titres negative; total haemolytic complement low but C3 normal; serological tests for syphilis negative. A biopsy of a buccal ulcer was taken.

At this stage our differential diagnosis included tuberculosis, Behçet’s disease, polyarteritis nodosa, Wegener’s granulomatosis, Goodpasture’s syndrome, and disseminated carcinoma.

On the second hospital day he became very breathless and his haemoglobin had fallen to 5·9 g dl⁻¹. Five units of blood were transfused and his haemoglobin rose to 11·0 g dl⁻¹. Hydrocortisone, 100 mg six-hourly, and antituberculous chemotherapy was begun. The next day he was still more breathless, tachypnoeic, and cyanosed. Breathing air his Po₂ was 3·7 kPa, Pco₂ 4·1 kPa, and pH 7·36. The Po₂ did not rise on 60% oxygen. The chest radiograph showed extensive consolidation of both lungs (fig 1); his heart size remained normal. The haemoglobin had fallen to 5·0 g dl⁻¹.

A diagnosis of intrapulmonary haemorrhage was made and continuous positive pressure ventilation was begun. Copious amounts of blood were aspirated via the endotracheal tube and repeated blood transfusions were necessary. At this stage the results of the mucosal biopsy showed a necrotising vasculitis. Intravenous cyclophosphamide 4 mg Kg⁻¹ daily was begun as additional treatment of his fulminating systemic vasculitis. However,

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Fig 1 Chest radiograph on third hospital day showing widespread consolidation in both lungs.
spread necrotising alveolar capillaritis (fig 2). Sections of
the cavitating lesion showed massive necrosis of the lung,
accompanied by arteritis and venulitis. A granulomatous
reaction featuring histiocytes and giant cells encompassed
the necrotic focus (fig 2). Necrotising arteritis was also
found in the spleen. The kidneys showed focal necrotising
glomerulonephritis. Granular deposition of IgG and
complement on glomerular capillary basement membranes
(but not in the alveolar capillaries) was demonstrated
using immunofluorescent techniques on frozen sections.
An unrelated finding was a recent myocardial infarct, the
result of coronary thrombosis and atherosclerosis. The
combination of the pulmonary lesion, systemic vasculitis
and focal glomerulonephritis indicated Wegener’s
granulomatosis.

Discussion

Acute massive pulmonary haemorrhage is a rarely
described feature of Wegener’s granulomatosis. Fauci and
Wolf1 reported a patient who died of acute pulmonary
haemorrhage, and Kjellstrand et al8 described a patient
with sudden onset of copious haemoptyses and almost
total opacification of the lungs on chest radiograph, who
responded to anti-human lymphoblast globulin, methyl-
prednisolone and azathioprine. In neither case is there a
report of the pathological changes in the lung. Alveolar
capillaritis is not a well-documented feature of Wegener’s
granulomatosis. Liebow3 does not refer to it in his
detailed review of pulmonary angiitis and granulomatosis
and it is only briefly mentioned by Spencer.4 In the case
reported here, alveolar capillaritis was the dominant
pathological lesion and we believe it was responsible for
the massive pulmonary haemorrhage and fulminating
course of the disease.

References

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