Successful treatment of post-influenza pseudomembranous necrotising bronchial aspergillosis with liposomal amphotericin, inhaled amphotericin B, gamma interferon and GM-CSF

R J Boots, D L Paterson, A M Allworth, J L Faoagali

Abstract
A case of aspergillus tracheobronchitis following influenza A infection in an immunocompetent 35 year old woman is described that required prolonged mechanical ventilation for airways obstruction. Treatment included liposomal amphotericin, inhaled amphotericin, gamma interferon and GM-CSF. Liposomal amphotericin therapy was associated with reversible hepatosplenomegaly. Inhaled corticosteroids with continued antifungal therapy were used for the management of severe recurrent airway obstruction. After a prolonged course of treatment she survived with fixed airways obstruction unresponsive to corticosteroids.

(Thorax 1999;54:1047–1049)

Keywords: Aspergillus; tracheobronchitis; influenza

A 35 year old registered nurse presented to hospital following a flu-like illness with increasing shortness of breath, non-productive cough, central chest pain, and wheeze. She was afebrile and in respiratory distress with widespread inspiratory wheezes and patchy inspiratory crackles.

She had been previously healthy. She lived on a rural property but kept no animals. There was no history of previous respiratory disease nor overseas travel. She took no regular medication and weighed 55 kg.

The chest radiograph at presentation was hyperinflated. The FEV/FVC was 1.01/1.72 (2.53/2.93) and the Pao2 was 48 mmHg (normal range 2–7.5).

The presumptive diagnosis was asthma and intravenous antibiotics, bronchodilators, and oral prednisone in a dose of 50 mg/day were commenced. Her condition deteriorated with bilateral perihilar alveolar infiltrates on the chest radiograph. She was intubated and mechanical ventilation commenced. She was difficult to hand ventilate and remained hypoxic despite 100% oxygen.

Bronchoscopic examination revealed thick mucoid inspissated secretions in the trachea and main bronchi. After the airways were cleared a white thick adherent membrane was present in the trachea and the main and segmental bronchi, extending beyond the vision of the bronchoscope. Attempts to remove the membrane resulted in bleeding. Amphotericin B 40 mg/day, intravenous flucytosine 2500 mg six hourly, and nasogastric itraconazole 200 mg eight hourly were commenced. Progressive improvement in the gas exchange was seen over the next eight hours with pressure control ventilation, continuous nebulised ipratropium bromide. Corticosteroid treatment was discontinued.

Histological examination of the endobronchial biopsy specimen showed necrotic debris, fibrinopurulent exudate, acutely inflamed bronchial mucosa, squamous metaplasia with marked reactive atypia and small numbers of fungal hyphae. The endobronchial biopsy specimens and bronchoalveolar lavage fluid grew Aspergillus niger. Viral cultures were negative. Paired serum samples for influenza A were positive >1:512. Serological examination for other respiratory viruses was negative.

Ten days after admission to hospital the membrane persisted only in the segmental bronchi of all lobes. Aspergillus niger was again isolated from the bronchial washings. Liposomal amphotericin (Ambisome Faulding and Co Ltd) was given at a dose of 250 mg/day for deteriorating renal function and nebulised amphotericin B, 10 mg twice daily, was added.

Hepatosplenomegaly was noted on the 15th hospital day which progressed to a liver span of 25 cm and spleen span of 20 cm by day 24. Liver function tests were mildly abnormal. Hepatitis B and C serology were negative and cultures from an open liver biopsy specimen were also negative. Histological examination of the biopsy specimen showed a non-specific chronic hepatitis of moderate severity with focal aggregates of mononuclear cells within the acinar tissue and normal liver architecture. No fungi were seen. Flucytosine and itraconazole treatment were discontinued.
Severe airways obstruction recurred on day 37. There was persisting membrane in the segmental bronchi of all lobes. Nebulised amphotericin was ceased due to concerns that it was contributing to the bronchospasm. Pressure control ventilation required inflation pressures up to 90 cm H₂O. The patient was hypercapnic to a peak PaCO₂ of 90 mm Hg (12 kPa). Continuous nebulised adrenaline, aminophylline, ketamine by infusion, and isofluorane by vaporiser had no significant effect on her condition. There was little improvement to day 40 when itraconazole and nebulised amphotericin were recommenced. Sedation and paralysis were stopped by day 52. Bronchoscopic examination showed membrane in the subsegmental bronchi only. Liposomal amphotericin was discontinued after 42 days and treatment with nebulised amphotericin and itraconazole were continued. There was a slow general improvement to day 58 with a reduction in FiO₂ to 0.35. The hepatomegaly decreased to 12 cm and the spleen to 6 cm.

From day 65 the course was complicated by episodic severe bronchospasm requiring reintroduction of sedation, paralysis, and pressure control ventilation. Bronchoscopic examination revealed a thick craggy membrane in the distal tracheal and lower airways with cobbly stoning of the mucosa. There was persisting lymphopenia of 0.68 × 10⁹/l. Further biopsy specimens were consistent with an inflammatory membrane but no fungus was seen nor grown. GM-CSF 400 µg daily was given subcutaneously for 10 days (Leukomax; Sandoz Australia Pty Ltd) and gamma interferon 200 µg subcutaneously (Imukin; Boehringer Ingelheim Pty Ltd) was given three times a week for two weeks. Liposomal amphotericin and continuous nebulised adrenaline were reinstituted and nebulised budesonide commenced.

The patient’s condition gradually improved. The hepatosplenomegaly recurred to a liver span of 24 cm and a spleen span of 22 cm. Liposomal amphotericin was discontinued on day 81. The lymphocyte count returned to normal by day 122. By day 131 the liver span had decreased to 18 cm and the spleen to 6 cm.

Lymphocyte function tests on the fifth hospital day were consistent with generalised T cell deficiency with poor mitogen responsiveness: CD4 0.16 × 10⁹/l (normal range 0.57–1.67 × 10⁹/l), CD8 0.09 × 10⁹/l (normal range 0.07–0.53 × 10⁹/l), CD19 0.04 × 10⁹/l (normal range 0.02–0.43 × 10⁹/l). HIV serological tests by ELISA were negative. The T cell deficiency persisted until day 107 after presentation. Lymphocyte reactivity to phytohaemagglutinin, concanavalin A, OKT3, pokeweed mitogen, Candida and Aspergillus antigens was absent on day 12 and returned to normal by day 35. Mantoux testing using avian and human tuberculin were negative despite previous BCG vaccination. The Mantoux test became reactive at 6 mm by day 131. She remained anergic to diphtheria, tetanus, Streptococcus, Trychophyton, and Proteus despite previous tetanus and diphtheria vaccination. During the first week of admission and again after four and six months immediate, and delayed skin prick tests to Aspergillus niger, Aspergillus fumigatus, and Aspergillus terreus remained non-reactive. A rise in titre to Aspergillus IgM antibody from 2560 to 5120 occurred. There was no IgG response detectable up to day 190. The appearance of a bone marrow biopsy specimen was consistent with increased myelopoiesis. Antinuclear and rheumatoid factors were not detected and serum immunoglobulins including IgG subclasses were normal.

Due to persisting airways obstruction 60 mg prednisone per day was commenced on day 124 with nebulised amphotericin as a prophylactic. The patient was weaned off the prednisone after one month with no measurable improvement in lung function. A high resolution CT scan of the chest revealed patchy linear shadowing throughout both lung fields and peribronchial thickening with mild central bronchiectasis. She was transferred to a general ward on day 95 and discharged home with independence on day 138. At discharge spirometric values were FEV₁, 1.04 l and FVC 1.89 l. She continues well three years later with spirometric values unchanged from discharge. She has returned to employment as a critical care nurse. The clinical course of antifungal and immunotherapy are set out in figure 1.

**Discussion**

This case demonstrates a prolonged persistence of the endobronchial membrane of aspergillus tracheobronchitis for more than 64 days despite antifungal and immunomodulation therapy. The expected duration of an airway membrane in this disease is not described. Host immune status determines the manifestations of aspergillus lung disease and the response to treatment. Patients with aspergillus tracheobronchitis may have less neutropenia and less exposure to corticosteroids and cytotoxic drugs than patients with parenchymal involvement.¹ The T cell abnormality may have contributed to this, although aspergillus pulmonary infection is more typical of neutrophil dysfunction or neutropenia. Two forms of fungal tracheobronchitis due to *Aspergillus* have been described.² Growth can
be intraluminal with superficial mucosal invasion; a fibrinopurulent membrane is prominent with fungal plugs occluding the bronchial tree. In the absence of parenchymal disease the chest radiograph will be normal. The second manifestation comprises discrete plaques over a small portion of the tracheobronchial tree. Focal pneumonia and abscess formation may result from direct extension.

Our patient had cutaneous anergy and lymphopenia involving T cells and NK cells but not B cells. This has been previously described with influenza A infection. Although there was no clear seroconversion or diagnostic rise in antibodies, the high antibody titre in the absence of vaccination suggested recent influenza infection. Persisting lymphopenia may have been a result of the treatment.

All previously reported cases of post-influenza aspergillus lung disease have died despite amphotericin treatment. The airways obstruction in this case was not clinically responsive to high dose inhaled and intravenous bronchodilators. After prolonged treatment with antifungal agents, recurrent airways obstruction resolved with inhaled corticosteroids while antifungal therapy was maintained. GM-CSF can enhance macrophage antifungal activity and stimulate neutrophil function. Gamma interferon promotes macrophage and NK cell fungal killing. The relative contributions of antifungal and cytokine therapy in this patient are uncertain.

Nebulised amphotericin has been used both prophylactically and therapeutically for aspergillus lung disease, including aspergillus laryngotracheobronchitis. It is generally well tolerated but may be associated with significant bronchospasm. Its contribution to airway obstruction is unclear as withdrawal and subsequent reintroduction did not result in recurrence of the bronchospasm. Liposomal amphotericin has the advantage of being able to be used at a higher dosage with a decreased incidence of renal failure. Hepatosplenomegaly occurred in our patient which resolved after withdrawal of liposomal amphotericin and recurred with its reintroduction. This was associated with a mild hepatitis only. This has not been previously reported. Itraconazole is well described in the primary and adjunctive treatment of fungal infection. In critically ill patients the gastrointestinal absorption of medication is uncertain. Enteral feeding and the common use of stress ulcer prophylaxis may also interfere with the bioavailability of azole based antifungal therapy in this situation.

Antagonism may occur between concurrent amphotericin B and itraconazole therapy.

Reactive airways disease during the acute illness has not been described in tracheobronchitis due to aspergillus. Fixed airway obstruction has persisted despite inhaled and systemic steroid trials followed for over three years. This case resulted in survival with an excellent functional outcome in a patient with aspergillus tracheobronchitis following influenza infection. Intensive support along with multimodal treatment may have a place in the management of these patients.

Successful treatment of post-influenza pseudomembranous necrotising bronchial aspergillosis with liposomal amphotericin, inhaled amphotericin B, gamma interferon and GM-CSF

R J Boots, D L Paterson, A M Allworth and J L Faoagali

Thorax 1999 54: 1047-1049
doi: 10.1136/thx.54.11.1047