Use of a Heimlich flutter valve for pneumothorax in cystic fibrosis

F P Edenborough, I Hussain, D E Stableforth

Abstract

The use of a Heimlich flutter valve in an adult patient with cystic fibrosis with hypercapnic respiratory failure which allowed resolution of a persisting pneumothorax after failure of conventional tube drainage is reported. The patient was managed at home and avoided surgical pleurodesis which could have jeopardised transplantation at a later date.

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Case report

The 24 year old woman with cystic fibrosis was an insulin dependent diabetic with a past history of nasal polypectomy, cholecystectomy, duodenal ulceration, and paroxysmal supraventricular tachycardia. She had been chronically colonised with Pseudomonas cepacia for approximately 12 months and her lung function was FEV1/FVC 0.55/1.01 (20%/32% predicted) with a PaCO2 of 6.9 kPa and PaO2 of 6.0 kPa breathing air. She had already had bilateral pneumothoraces, the right side requiring tube drainage in 1989, the left resolving spontaneously.

She presented in January 1992 with a small right apical pneumothorax, initially thought not to require tube drainage. She was commenced on antibiotics but by the tenth day repeat radiography showed the pneumothorax to have enlarged significantly. A drain was placed in the fifth intercostal space in the mid-axillary line. The pneumothorax resolved and the drain was removed seven days later.

During the next month a Portacath was inserted on the left side to facilitate venous access and a percutaneous gastrostomy was sited, both without complications. Chest radiographs performed after each procedure showed no evidence of pneumothorax.

Four months later she developed a left pneumothorax requiring five days of drainage via a tube in the left fifth intercostal space in the mid-axillary line. Ten days after discharge she had a further large left pneumothorax and another drain was placed in the fourth intercostal space in the mid-axillary line. Six days later suction was commenced but after five days this drain continued to bubble and, in an attempt to avoid surgery, it was removed. A further drain was inserted in the sixth intercostal space in the mid-axillary line. Despite this there was no resolution and a Heimlich flutter valve was attached to the drain 19 days after admission to facilitate mobility and she was discharged. The drain was removed after 63 days when the lung was fully inflated. She was discharged with an FEV1/FVC of 0.49/1.03. Except for minor surgical emphysema there were no complications. There has been no recurrence of pneumothorax 14 months later.

Discussion

Experience of 205 patients with cystic fibrosis with 395 episodes of pneumothorax has been reviewed.1-3 Pneumothoraces occur a little more frequently in men and may occur at any age. Patients are often ill with FEV1<50% predicted, low body weight, and have chronic colonisation of the lungs of Pseudomonas aeruginosa. Pneumothorax recurs in 50–83% of cases. Management has included observation, chemical sclerosants (tetracycline, silver nitrate, quinacrine), thoracentesis, thoracoscopy and talc poudrage or pleural abrasion, open thoracotomy and partial pleurectomy, segmentectomy or bullectomy. Penketh et al6 suggested that a persisting air leak at seven days is usually an indication for surgical intervention, but more recent opinion4 suggests that this should be avoided if possible in patients with cystic fibrosis. An exception to this may be thoracoscopic stapling, with or without pleurodesis, which is safer and less painful than thoracotomy7 and may find favour in the treatment of pneumothorax in those patients who could tolerate anaesthesia.

Our patient was unfit for anaesthesia. Although prolonged intercostal drainage (>7 days) has been associated with a high morbidity and immediate and delayed mortality,4 no study has reported the use of the Heimlich flutter valve in patients with cystic fibrosis. The valve was first described in 19655 and has become increasingly popular in outpatient management with both large2 and small3 calibre tubes. No major complications have been reported except accidental dislodgement and inadvertent re-attachment back to front, when they may induce tension pneumothorax.8 This can be prevented by secure taping of the union between drain and valve. Blockage and infection are theoretical problems but have not been reported.

We believe this to be the first reported use of a Heimlich flutter valve to manage a persisting pneumothorax in a patient with cystic fibrosis. The technique allowed resolution of the leak at home in a patient in hypercapnic respiratory failure with poor lung function in whom surgery might have been life threatening and which could have jeopardised transplantation at a later date.


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Empyema and mediastinitis complicating retropharyngeal abscess

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Abstract
A 21-year-old man with a retropharyngeal abscess complained of right sided chest pain, and chest radiography and thoracentesis revealed an empyema. A computed tomographic scan of the chest showed a posterior mediastinal abscess communicating with the right pleural cavity. Emergency thoracotomy was performed and the mediastinal abscess and empyema were drained.

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Acute empyema following a retropharyngeal abscess and mediastinitis in an adult is very rare and often fatal. This report presents a case that was cured with surgical intervention.

Case report
A 21 year old man was admitted with a high fever to our hospital complaining of dysphagia and a four day history of a sore throat for which he was given oral antibiotics. On admission physical examination disclosed diffuse erythema and swelling of the pharynx and swelling and tenderness of the right side of the neck, but cervical lymph adenopathy was not present. There were no abnormal dental findings and the tonsils appeared normal. A chest radiograph showed widening of the upper mediastinum, and radiography of the soft tissues of the neck showed retropharyngeal gas and widening of the retropharyngeal space. A transoral retropharyngeal tap was performed and pus was obtained consistent with a retropharyngeal abscess. Cultures grew α-streptococcus, β-streptococcus and staphylococcus. His white cell count was 13 100/mm³ and the serum level of C-reactive protein (CRP), a non-specific marker for acute inflammation, was 37.8 mg/dl.

Treatment was started with intravenous cefmetazole to which these organisms proved sensitive. The following day he developed right sided chest pain and radiographs showed a right pleural effusion. Pus was obtained during thoracentesis and a chest tube was inserted immediately. The pleural effusion grew the same microorganisms as the retropharyngeal pus and the patient's mediastinitis appeared consequent to the retropharyngeal abscess which had perforated into the pleural cavity. An oesophagogram was normal. The patient's condition rapidly improved, temperature became normal, and both the pharyngeal and cervical swelling improved, but a computed tomographic scan of the chest on the fifth hospital day showed an increasing encapsulated mediastinal abscess between the oesophagus and vertebral bodies (fig 1). Since the mediastinal abscess had not been drained adequately a right thoracotomy was performed on the seventh hospital day, and 200 ml of turbid, yellow-white fluid obtained. The upper mediastinal pleura appeared swollen between the superior vena cava and vertebral
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