Massive haemoptysis caused by congenital absence of a segment of inferior vena cava

M H Ashour, S K Jain, K M Kattan, A Karim El-Bakry, M Khoshim, F M Mesahel

Abstract
A patient with congenital absence of a part of the inferior vena cava is described. This resulted in spontaneous rupture of a bronchial vein leading to massive haemoptysis.

(Thorax 1993;48:1044-1045)

Anomalies of the inferior vena cava are often discovered incidentally during imaging studies performed for other reasons, since patients with these abnormalities are usually asymptomatic. Haemoptysis as a presenting feature of interrupted inferior vena cava has not been previously reported. We describe a patient presenting with massive haemoptysis.

Case report
A 28 year old Saudi woman, 36 weeks into her second pregnancy, was admitted as an emergency with massive haemoptysis. She coughed up approximately 800 ml of fresh blood over four hours. There was no history of chest pain, pneumonia, asthma, tuberculosis, bleeding disorders, cigarette smoking, stigma suggestive of hereditary telangiectasia, or previous haemoptysis. Her pregnancy had been uncomplicated to date. Physical examination was unremarkable except for a respiratory rate of 30/minute and poor air entry over the right lung. Fetal heart rate was 140/minute.

Chest radiography showed a hyperinflated right lung with mediastinal shift to the left. Arterial blood tensions breathing room air showed a pH of 7.41, Po2 of 58 mm Hg, and Pco2 of 28-7 mm Hg. Blood biochemical tests were normal.

At bronchoscopy there was blood in the right main bronchus. The right bronchus intermedius was occluded by a clot-like material which could not be sucked out. A caesarian section was performed and a healthy baby was delivered. During this procedure another major haemoptysis occurred. At thoracotomy the aygos vein was found to be massively dilated. The bronchial as well as the intercostal veins were also dilated to 2-3 times their normal size. In view of the uncertainty about the pathological condition of the right bronchus intermedius, bronchotomy was performed rather than proceeding to lobectomy. A clot was seen to fill the bronchus. While removing the clot fresh bleeding from a small spot in the bronchial mucosa located immediately below the middle lobe orifice was seen. The pattern of bleeding was not arterial and the mucosa surrounding the area of bleeding looked normal. The bleeding site was transfixed with interrupted Dexon sutures. Postoperative recovery was uneventful.

Ten days after the operation, in search of a cause of the haemoptysis, the patient underwent further investigations. Pulmonary angiography showed no evidence of an arteriovenous fistula or pulmonary embolism. The contrast medium filled a dilated aygos vein that directed the flow to the superior vena cava (fig 1). A cavogram attempted from the femoral vein showed absence of the prerenal segment of the inferior vena cava (fig 2), with

Figure 1 Early phase of pulmonary angiogram showing massively dilated aygos vein (arrows).

Figure 2 Cavogram showing venous return through the ascending lumbar veins (arrows) as an alternative to the absent prerenal segment of the inferior vena cava.
venous return through the ascending lumbar veins. The bronchial and intercostal arteries were normal on a thoracic aortogram. Echocardiography was normal. Computed tomography of the abdomen and chest showed no evidence of thrombosis in the inferior vena cava. The patient was followed up for 18 months and had no more episodes of haemoptysis.

Discussion
Although anomalies of the inferior vena cava are commonly discovered during imaging for other diseases,1,5 it seems that the anomaly in this patient directly contributed to the pathogenesis of the haemoptysis.

The bronchial venous system communicates freely with the pulmonary veins and alveolar capillaries on one side, and with the azygous vein on the other.6 A direct communication between the systemic venous circulation and the pulmonary circulation is therefore present. This makes the bronchial veins vulnerable to pressure changes in either the systemic venous or pulmonary circulations. These bronchial venous channels are thin walled and non-dissectible.5 Physiologically about one third of the blood accumulated in the bronchial venous plexuses is thought to return to the azygous vein, while the remainder blood flow returns to the pulmonary veins.7

Although we did not measure pressure in the azygous vein, we believe that the congenital interruption of the inferior vena cava caused azygous venous hypertension due to the massive increase in the azygous venous flow. As a result of this haemodynamic change the bronchial venous drainage to the azygous vein could become impaired and reversal of bronchial venous flow might occur. Engagement of the bronchial veins would therefore take place, similar to oesophageal varices in portal hypertension. These engorged, thin walled, relatively non-dissectible, submucosal bronchial veins may undergo rupture with manoeuvres associated with increased intrathoracic pressure and this could lead to haemoptysis.

On the basis of our experience with this case we suggest that an inferior vena cavaogram should be considered as one of the investigations when dealing with massive or recurrent haemoptysis of unknown origin.

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Comparison of nebulised aerosol deposition in the lungs of healthy adults following oral and nasal inhalation

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Abstract
A standard jet nebuliser was used to generate a radiolabelled aerosol and the pattern of deposition within the airways of eight healthy adults was studied with a gamma camera. Penetration of aerosol to the lung was greatly reduced when breathing through the nose compared with mouth breathing.

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Although the nose acts as a filtration system it has been argued that patient preference should determine whether a facemask or a mouthpiece is used when inhaling aerosols from jet nebulisers.1,3 This argument is based on clinical studies which failed to show any significant difference in clinical response when salbutamol was inhaled through these two routes.1,3 However, bronchodilators are generally used in supramaximal doses1,3 and hence the observed clinical response need not directly reflect the total dose reaching the lungs.

The dose of aerosol deposited in the lungs is determined by the total dose of drug inhaled and the pattern of deposition of that dose within the airways. The factors influencing the total dose of drug inhaled when using a jet nebuliser are complex and have previously been discussed.4 The purpose of this study was to determine what effect inhaling a wet, heterodispersed aerosol by the nasal rather than the oral route might have on the pattern of deposition within the airways. A standard technique12 was used in which the deposition of a radiolabelled aerosol was assessed with a gamma camera.

Methods
Eight men aged 21–32 years were studied. All were in good health, were non-smokers, and gave no history of lower respiratory tract dis-
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