A familial case of pulmonary arterial sequestration

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Abstract

The cases of a mother and infant son are reported, both with a rare type of pulmonary sequestration where the arterial supply to the lung arises from the systemic circulation. This is a familial case of arterial sequestration. In both patients, the lung parenchyma was radiologically normal.

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A previously well woman had her first haemoptysis at 19 years old. Chest x ray and ventilation perfusion lung scan at presentation were normal. A provisional diagnosis of tracheitis was made. She re-presented with haemoptysis five years later in the second trimester of her second pregnancy. Bronchoscopy showed a large clot in the bronchus intermedius and right lower lobe bronchus. Throughout the third trimester recurrent haemoptysis became increasingly frequent and continued after delivery. A computed tomography (CT) scan of her thorax showed an anomalous artery from below the diaphragm supplying the right lung base and pulmonary angiography showed absence of a pulmonary arterial supply to the postero-basal segment of the right lower lobe. An aortogram was not performed.

Twelve months later she had further haemoptyses. Aortography revealed an anomalous artery arising from the coeliac axis supplying the postero-basal segment of the right lower lobe (fig 1). Venous drainage from this area entered the left atrium. An aortogram was not performed.

Twelve months later she had further haemoptyses. Aortography revealed an anomalous artery arising from the coeliac axis supplying the postero-basal segment of the right lower lobe (fig 1). Venous drainage from this area entered the left atrium. The diagnosis of pulmonary artery sequestration was established with sequestration of the right lower lobe from its normal pulmonary artery supply.

At thoracotomy, an enlarged anomalous artery was found along the right side of the oesophagus and the free edge of the right pulmonary ligament. Histology of the excised lung showed changes of local severe prolonged pulmonary hypertension.

Her second son was born at 37 weeks’ gestation. A patent ductus arteriosus (PDA) was diagnosed at his first neonatal examination when he was tachypnoeic with an enlarged liver. The PDA was confirmed by echocardiography. He remained tachypnoeic during feeding, suffered frequent respiratory tract infections, and was failing to thrive, with a body weight on the third centile. His chest x ray showed cardiomegaly with clear lungs.

Angiography was performed with a view to closing the PDA. During this study, an anomalous artery arising from the coeliac axis was seen supplying the right lower lobe, which drained into the left atrium (fig 2). The anomalous vessel was coil embolised resulting in a partial occlusion. The PDA was closed completely.

On follow up, he continued to have frequent respiratory tract infections and his body weight remained on the third centile. Repeat angiography at 16 months of age showed complete occlusion of the anomalous systemic feeding artery. Presently, aged 3 years, he remains well.

Discussion

Embryologically, pulmonary sequestration results from anomalies of foregut budding in which pulmonary tissue can be sequestered from the bronchial tree (bronchial sequestration) and/or the pulmonary arterial circulation (pulmonary arterial sequestration).1 2 In this

Figure 1Mother’s aortogram showing the anomalous systemic arterial supply to the right lung base from the coeliac axis (arrow). Its location is identical to her son’s vessel.

Figure 2Infant aortogram showing the anomalous systemic arterial supply to the right lung base from the coeliac axis (curved arrow) with venous drainage to the left atrium (arrow head).
familial case with isolated pulmonary arterial sequestration, the lung receives an anomalous arterial supply from the systemic circulation and there are no parenchymal lung changes evident on chest X-ray or CT scan. It is known that most isolated pulmonary artery sequestrations in infancy are found at angiographic investigation for congenital heart disease or anomalous pulmonary venous drainage anomalies.1

Familial cases of pulmonary sequestration are rare. With only three reports to date, two in association with the Scimitar syndrome (right lung hypoplasia, cardiac dextraposition, and a systemic arterial supply and venous drainage) in a father and daughter and a father and son,2,3 and one in siblings from consecutive pregnancies.4 In the affected sibling case a combined bronchial and pulmonary arterial sequestration was present, with the anomalous systemic feeding artery arising from the descending thoracic aorta. The lung parenchyma was abnormal on ultrasound and chest X-ray. A PDA was present in one of the siblings.

Our report adds to the exceptionally rare description of familial pulmonary sequestration. Familial isolated pulmonary arterial sequestration, as we present here, has not been described previously.

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