IMAGING IN BRONCHOPULMONARY SEQUESTRATION

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ABSTRACT

Bronchopulmonary Sequestration is a rare congenital malformation of lower respiratory tract that lacks normal communication with the tracheobronchial tree. In 1977, Huber stated that aberrant arterial supply of sequestered lung could be encountered. The term pulmonary sequestration was first used in 1946 by Pryce. It was also termed as bronchoarterial malinosculation by clements in 1987. It is a non functional mass composed of dysplastic lung parenchyma, embryologically detached from the tracheobronchial tree and receiving its own blood supply from a systemic artery, usually 75% from thoracic or abdominal aorta and 25% of sequestrations receive their blood flow from the subclavian, intercostals, pulmonary, pericardiophrenic, innominate, internal mammary, celiac, splenic or renal artery. The venous drainage is variable (1) computed Tomography, magnetic resonance angiography are able to demonstrate the aberrant arterial vessel feeding the sequestration. The blood supply of 75% of pulmonary sequestrations is derived from the thoracicabdominal aorta. The remaining 25% of sequestrations receive their blood flow from the subclavian, intercostal, pulmonary, pericardiophrenic, innominate, internal mammary, celiac, splenic, or renal arteries.

INTRODUCTION

Bronchopulmonary sequestration is a condition where in a piece of tissue that ultimately develops into lung tissue is not attached to the pulmonary arterial blood supply. This sequestered tissue is not connected to the normal bronchial airway. So it fails to contribute to respiration. The condition is thought to be congenital in nature and is usually diagnosed by ultrasound by prenatal ultrasound. Pulmonary sequestration is a rare congenital abnormality, with an overall incidence of 0.15 to 6.4% of all congenital pulmonary malformations. It is characterized by a mass of nonfunctioning, embryonic, cystic pulmonary tissue that receives its blood supply from the systemic circulation (Clements et al., 1987) and has no connection with the bronchial tree. It preferentially affects the lower lobes, predominantly the left lower lobe. Both intralobar and extralobar sequestrations arise through the same pathoembryologic mechanism as a remnant of a diverticular outgrowth of the foregut. Gastric or pancreatic tissue may be found within the sequestration (Zach et al., 2002). The presentation is variable, ranging from no symptoms to hemoptysis. Clinically, pulmonary sequestration is latent until infection leads to symptoms. Recurrent pneumonitis of the sequestrated segment, purulent sputum and haemoptysis are the prevailing symptoms (Franco et al., 1998). Pulmonary sequestration can be present clinically at all ages, but most lesions tend to develop these infective complications at school age and adolescence. However, symptoms may also occur in infancy and preschool age group. A symptomatic adult has also been described (Saygi et al., 2001). Many a times, it is discovered incidentally on a chest radiograph taken for another reason, as in this case. About two-thirds of all pulmonary sequestrations are found in the posterior basal segment of the left lower lobe (Halkic et al., 1999). On radiograph, the initial impression is usually one of pneumonia, though the lesion may appear as air- or fluid-filled cysts, single or multiple. Till recently, aortography, with selective angiography, was usually necessary to diagnose sequestration and demonstrate its blood
supply. CT scanning and recently spiral CT angiography offer less invasive means of demonstrating the anomalous vascular supply (Kang et al., 2006). Surgical excision is usually curative; it should be conservative, sparing the normal lung parenchyma (Bonnard et al., 2004). Some authors advocate embolisation of the aberrant systemic artery at the time of initial catheterisation, which may result in complete radiological resolution of mass (Walsh, 2005 and Saida et al., 2006).

**Anatomical Classification**

Intralobar sequestration (ILS) in which the lesion is located within a normal lobe and lacks its own visceral pleura. Extralobar sequestration (ELS) in which the mass is located outside the normal lung and has its own visceral pleura.

**Intralobar Sequestration**

- The intralobar variety accounts for 75 percent of all sequestrations.
- Usually presents in adolescence or adulthood as recurrent pneumonias.
- The lung tissue lies within the same visceral pleura as the lobe in which it occurs.
- Males and females are equally affected with ILS.
- In ILS, the arterial supply usually is derived from the lower thoracic or upper abdominal aorta.
- Venous drainage is usually to the left atrium via pulmonary veins establishing a right to left shunt.
- Abnormal connections to the vena cava, azygous vein, or right atrium may occur.
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- Two thirds of the time, the sequestration is located in the paravertebral gutter in the posterior segment of the left lower lobe.
- Unlike extralobar sequestration, it is rarely associated with other developmental abnormalities.
- Patients present with signs and symptoms of pulmonary infection of a lower lobe mass.
- It is believed that sequestrations become infected when bacteria migrate through the Pores of Kohn or if the sequestration is incomplete.

**Extralobar Sequestration**

- The extralobar variety accounts for 25 percent of all sequestrations.
- ELS usually presents in infancy with respiratory compromise.
- Develops as an accessory lung contained within its own pleura.
- ELS has a male predominance (80%).
- Related to the left hemidiaphragm in 90% of cases.
- ELS may present as a subdiaphragmatic or retroperitoneal mass.
- In general, the arterial supply of ELS comes from an aberrant vessel cic aorta.
- It usually drains via the systemic venous system to the right atrium, vena cava, or azygous systems.
- Congenital anomalies occur more frequently in patients with ELS than ILS.

2 months baby came with the history of cystic lesion in the left lung during routine antenatal ultra Sonography (USG)
- Associated anomalies include Congenital cystic adenomatoid malformation (CCAM), congenital diaphragmatic hernia, vertebral anomalies, congenital heart disease, pulmonary hypoplasia, and colonic duplication.
- Since it is enveloped in its own pleural sac, it rarely gets infected so almost always presents as a homogeneous soft tissue mass.
- The mass may be closely associated with the esophagus, and fistulae may develop.

**Case Report**

**Radiograph Chest and Abdomen AP View Showing**

- Nasogastric and tracheal tube seen in situ.
- Bronchietatic change like appearance in left middle and lower zone.
- Cardiac shadow is not seen on the left side.
- Mediastinal shift to right.
- Tracheal shift to right.
- Thymus appears enlarged and displaced to right side and merging with right cardiac border.
- Left main bronchus apers dilated.
- Minimal pneumothorax noted on left side.
- Left dome of diaphragm appears flattened.

**Non Contrast Enhanced CT**

Heterogeneous partially aerated mass in the posterior aspect of the left hemi thorax, consisting of cyst like air spaces of varying sizes interspersed between solid tissue components with no communication with the bronchial tree.

**Sonography**

Ultrasound evaluation is ideally suited for evaluation in the antenatal period or in neonates. (Thind et al., 1985, May et al., 1993) Sequestration should be considered in the differential diagnosis in any fetus with a lung mass. The lesion usually has a homogenously echogenic appearance on ultrasound, but may show cystic changes (Felker et al., 1990).
Venous drainage of heterogenous lung tissue via draining vein crossing anterior to the descending thoracic aorta into the azygous vein.

Arterial supply for heterogenous lung tissue derived from single branch arising from the descending aorta.

Doppler ultrasound may be useful for evaluation of the supplying systemic artery. (Deeg et al., 1992) However, a small aberrant systemic artery may not be identified and the acoustic window may be impaired by surrounding air-filled lung or the bony thorax.

**DISCUSSION**

The anomalous blood supply of a pulmonary sequestration is the clue to its diagnosis. Identification was traditionally made, as in case report 1, by intra-arterial DSA. Currently, minimally invasive angiographic techniques such as CT and MR
angiography are also able to demonstrate the aberrant arterial vessel feeding the pulmonary sequestration, as described in the second case. Three-dimensional reconstruction techniques such as maximum-intensity-projection (MIP) or more recently volume rendering (VR) are useful, with the last method having the advantage of displaying simultaneously several ranges of tissue densities. It is now accepted that intra-arterial angiography is no longer mandatory to achieve the diagnosis of these malformations (Pryce et al., 1946) but instead should be reserved for the few cases in which embolization is contemplated (Clements et al., 1987). The dierential diagnosis of pulmonary sequestration is extensive, and is modiﬁed by the age at presentation. In the new born other retinences should be ruled out, such as acongenital diaphragmatic hernia, intrapulmonary bronchogenic cyst, cystic adenomatoid malformation or bronchial atresia. Later on life, several tumours and tumour-like conditions must be excluded including broncho pulmonary neoplasia, bronchiectasis or arteriovenous fistula (Rosado et al., 1993)

Treatment

Usually the sequestration is removed after birth via surgery. In most cases this surgery is safe and effective; the child will grow up to have normal lung function. In a few instances, fetuses with sequestrations develop problematic ﬂuid collections in the chest cavity. In these situations a Harrison catheter shunt can be used to drain the chest ﬂuid into the amniotic ﬂuid. In rare instances where the fetus has a very large lesion, resuscitation after delivery can be dangerous. In these situations a specialized delivery for management of the airway compression can be planned called the EXIT procedure, or a fetal laser ablation procedure can be performed. During this minimally invasive fetal intervention, a small needle is inserted into the sequestration, and a laser ﬁber is targeted at the abnormal blood vessel going to the sequestration. The goal of the operation is to use laser energy to stop the blood ﬂow to the sequestration, causing it to stop growing. Ideally, after the surgery, the sequestration steals less blood ﬂow from the fetus, and the heart and lungs start growing more normally as the sequestration shrinks in size and the pleural efﬁssion goes away.

Conclusion

Pulmonary sequestration is a heterogeneous group of lesions with anomalous connections between the various components of the lung. The primary role of imaging in an individual patient is to depict the aberrant vascular anatomy for diagnosis and accurate surgical planning. A high index of suspicion is required, both by the clinician and the radiologist, in making a presumptive diagnosis. Any persistent intrapulmonary opacity in the lower lobes of a child or young adult, on a chest radiograph, should prompt further evaluation with cross-sectional imaging. Non-invasive CT and MR angiographic techniques are useful for depiction of vascular anatomy. Multi detector CT angiography, which allows simultaneous imaging of the aberrant vascular anatomy as well as the lung parenchyma, has now become the ﬁrst-line examination in the preoperative assessment of pulmonary sequestration. Diagnostic catheter angiography is only required in select problematic cases.

REFERENCES


