A Patient with Intralobar Pulmonary Sequestration: 
A Rare Congenital Anomaly

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ABSTRACT

A 51-year-old male patient had a history of recurrent pneumonia since the second decade of life. Interestingly, chest X-rays indicated the same radiological pattern with left lower lobe involvement. A multislice spiral CT scan of the chest was performed and depicted an anomalous arterial vessel emerging from the descending thoracic aorta towards the left lower pulmonary lobe, characterising pulmonary sequestration. The findings were confirmed by thoracic aortography, and the patient was referred for surgical treatment.


Pulmonary sequestration is an anomaly characterised by a non-functioning pulmonary mass without communication with the normal bronchial tree, whose blood supply derives from an anomalous systemic artery.1,2 This is a rare anomaly with an incidence of 0.15% to 6.45% of congenital pulmonary malformations.3 Two forms have been described: extralobar sequestration, which has its own pleura, and intralobar sequestration, which is covered by the visceral pleura of a normal pulmonary lobe.4 A case of this rare congenital anomaly is reported, highlighting the main findings of the imaging analyses.

CASE REPORT

A 51-year-old male from the state of Goiás, Brazil, who lives in a rural area, presented with recurrent episodes of pulmonary infection, which were usually difficult to treat, for approximately 28 years. Some episodes were associated with haemoptysis, which is occasionally abundant. The subject smoked from the age of 17 to 22 years. His past medical history was positive for childhood asthma, Chagas disease (arrhythmogenic type), and pulmonary tuberculosis treated at the age of 24.

Chest X-rays always indicated a pattern of homogeneous hypotransparency in the posterior left lower hemithorax. A chest spiral angiotomography was performed, which indicated (Figure 1):

1) an anomalous blood vessel emerging from the descending aorta towards the left lower lobe without evidence of irrigation from the ipsilateral pulmonary artery for this segment;

2) prominence of arterial and venous vascularisation in the left lower pulmonary fields due to hypertension, without characterisation of arteriovenous communications;

3) mild left pulmonary artery hypoplasia; and

4) mild left lung hypoplasia.

Intralobar pulmonary sequestration was diagnosed, based on these findings. Aortography and heart catheterisation were performed to complement the diagnosis and for surgical planning (Figures 2 and 3). Heart catheterisation indicated coronary arteries without significant obstructions and left ventriculography with
no changes. Aortography demonstrated the presence of a large arterial branch originating from the descending aorta into the lower left lung.

Currently, the patient is asymptomatic and awaiting surgery.

DISCUSSION

The term ‘sequestration’ refers to a lung segment with an anomalous arterial vessel. In 80% of cases, the blood supply of this segment derives from the descending aorta, although the abdominal aorta, the subclavian artery, and the renal artery can also be sources of this arterial vessel. In the present case, the anomalous arterial supply originated from the thoracic descending aorta.

The classification of extralobar or intralobar pulmonary sequestration depends on the presence or absence of its own pleura. Extralobar pulmonary sequestration, which represents 25% of the cases, has its own pleura. This type is more prevalent in males (4:1), and it is located in the left hemithorax in 80% of the cases. Conversely, intralobar pulmonary sequestration is surrounded by normal lung tissue and is equally distributed between genders. It usually affects the left lower lobe.

The symptoms of intralobar pulmonary sequestration usually manifest around the second decade of life as recurrent respiratory infections. This late diagnosis, associated with histopathological findings, indicates that some forms of intralobar pulmonary sequestration may originate from a chronic inflammatory process of an infectious aetiology. Conversely, extralobar pulmonary sequestration is usually diagnosed during childhood, and it is associated with other congenital anomalies in over 60% of the cases. For this reason, extralobar pulmonary sequestration is considered to be a congenital disease.

The diagnosis of pulmonary sequestration can be suggested by imaging exams, such as ultrasound with colour Doppler, chest spiral angiotomography, magnetic resonance imaging, and aortography. However, a definitive diagnosis is achieved through an anatomopathological study.

Chest X-ray usually detects opacity in the posterior basal segment of the left lower lobe contiguous with the hemidiaphragm; however, it does not rule out other lesions of a similar aspect, and it does not distinguish between extralobar and intralobar types.

Ultrasound is more useful during the prenatal period and in newborns when it usually identifies an echogenic and homogenous intrathoracic mass with clear edges. Colour Doppler highlights the anomalous artery. Identifying an arterial branch from the thoracic or abdominal aorta using colour Doppler is pathognomonic of sequestration.

Arteriography is considered to be the gold standard for diagnosing pulmonary sequestration. It permits the identification of the anomalous artery as well as its origin, number, course, size, and venous drainage, which are invaluable information for preoperative planning. However, spiral angiotomography and magnetic resonance angiography (angioMRI), due to their accuracy and lower invasiveness, are good diagnostic options.

Tomographic findings include the presence of a homogenous mass or cystic formations. Spiral tomography can indicate the origin and course of anomalous systemic vessels.

Angio MRI allows for excellent visualisation of anomalous vessels, and arteriography can be omitted.
Resection of the involved lung parenchyma is the treatment for pulmonary sequestration. This type of surgery is easier in extralobar pulmonary sequestration because the anomalous mass has its own pleura. Conversely, in the intralobar type, which has the same pleura as the remaining lung and has infectious changes, the surgical procedure can be more difficult. However, after the surgery, postoperative evolution is usually very favourable.\textsuperscript{13}

Some authors have proposed a conservative treatment for cases without comorbidities; however, the presence of diseases such as haemoptysis, benign tumours, and pneumonias caused by more aggressive etiologic agents, among others, requires surgical treatment.

It is concluded that, despite the fact that pulmonary sequestration is a rare entity, this possibility should be investigated in patients with recurring pneumonias. Moreover, imaging exams demonstrated great diagnostic value and, for this reason, should be used before any surgical procedure.
CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

REFERENCES