Introduction

Pulmonary sequestration (PS) is a rare congenital malformation characterized by a mass of nonfunctioning lung tissue that does not communicate with the tracheobronchial tree and is vascularized by an anomalous systemic artery. It is composed of cystic embryonic tissue and contains disorganized non-aerated alveoli, as well as bronchi, cartilage, and respiratory epithelium. (1)

In 75% of cases of PS, the blood supply is derived from the thoracic or abdominal aorta, and venous drainage is via the systemic veins or via the pulmonary vein. Accidental transection of an anomalous systemic artery can cause massive hemorrhage with fatal consequences, and it is of paramount importance that anomalous vessels be identified in the preoperative period. (2)

The objective of the present study was to report an intraoperative finding of PS following hemorrhage resulting from the transection of an anomalous pulmonary vessel during resection of a carcinoid tumor.

Case report

A 39-year-old female patient presented with a history of recurrent pneumonia, together with sporadic episodes of productive cough and fever. In the two preceding years, the patient had had four episodes of moderate volume...
hemoptysis. In the preoperative investigation, a CT scan of the chest revealed a lesion in the intermediate bronchus (Figure 1), as well as cystic bronchiectasis in the right lower lobe (Figure 2). Fiberoptic bronchoscopy confirmed the tomographic finding of a wine-colored, exophytic endobronchial lesion at the root of the intermediate bronchus, and biopsy of the lesion revealed a typical carcinoid tumor.

Clinical reasoning based on those findings suggested bronchiectasis resulting from chronic bronchial obstruction. The patient underwent right posterolateral thoracotomy with bilobectomy (middle and lower lobes). During the intraoperative period, sectioning of the pulmonary ligament resulted in a hemorrhagic process originating from an anomalous arterial vessel that had irrigated the parenchyma of the PS (Figure 3). Hemostasis was achieved by ligation of the vessel. Histopathology results were consistent with the finding of PS. The postoperative course was satisfactory, without any complications, and the patient was discharged on postoperative day 5.

Discussion

Despite being defined as distinct entities, lung malformations constitute a spectrum of abnormalities, with quite similar clinical presentations, arising from flaws in the development of the primitive intestine and its differentiation into a respiratory system during the embryonic period. Chief among the most common malformations are cystic adenomatoid malformation, PS, congenital lobar emphysema, bronchogenic cysts, and pulmonary arteriovenous malformations. The reported annual incidence of these malformations ranges from 30 to 42 cases per 100,000 population, PS accounting for 0.15–6.45% of all cases.

Depending on its pleural covering, PS is classically divided into intralobar and extralobar. An intralobar PS (ILPS) share the pleural covering with the rest of the lung, whereas an extralobar PS (ELPS) is completely covered by its own visceral pleura. Although the extralobar form is well defined as a congenital abnormality, the intralobar form has a controversial pathogenesis, with some evidence indicating that, in many cases, it is an acquired disease.

Accounting for approximately 75% of all cases of PS, ILPS is more common in the lower lobes and on the left, involving the posterior basal segment. Its arterial supply is nearly always derived from the aorta or from one of its branches, typically one of a large diameter. Venous drainage is via the pulmonary veins into the left atrium, creating a left-to-left shunt. In a minority of cases, drainage is via the inferior vena cava or via the azygos system. In the case presented here, the PS was located in the right lower lobe and the blood supply came from an anomalous branch derived from the thoracic aorta.

Cases of ILPS typically occur in adolescents and young adults with a history of recurrent infections of the respiratory tract, hemoptysis, and dyspnea. Some ILPS patients develop cardiac symptoms, which are a consequence of the left-
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Cyanosis, and feeding difficulties are common. More rarely, recurrent episodes of respiratory tract infection or gastrointestinal symptoms can occur. Cases of ELPS can be accompanied by pulmonary, cardiac, and vertebral malformations, as well as by malformations of the chest wall and gastrointestinal tract, being most commonly accompanied by diaphragmatic hernia.

In general, an ELPS is located in the posterior costodiaphragmatic recess, between the lower lobe and the left hemidiaphragm. More rarely, it can occur in the mediastinum or in the abdominal region. An ELPS is typically irrigated by an artery originating directly from the aorta, and, in 80% of cases, venous drainage is systemic, occurring via the azygos-hemiazygos system or via the superior vena cava, which creates a left-to-right shunt. Unlike in cases of ILPS, clinical manifestations appear in the first six months of life in cases of ELPS. Neonatal asphyxia, dyspnea, cyanosis, and feeding difficulties are common. More rarely, recurrent episodes of respiratory tract infection or gastrointestinal symptoms can occur. Cases of ELPS can be accompanied by pulmonary, cardiac, and vertebral malformations, as well as by malformations of the chest wall and gastrointestinal tract, being most commonly accompanied by diaphragmatic hernia.

On chest X-rays, PS can be initially identified as a mass of homogeneous opacity. Diagnostic confirmation is made through chest CT, magnetic resonance imaging, or arteriography. Chest CT is more useful in detecting abnormalities in the lung parenchyma and only in some cases can it identify PS anomalous irrigation. Arteriography is the best test for diagnostic confirmation of PS, because it detects the anomalous artery irrigating the PS with precision. More recently, reconstruction with multichannel CT scanners

Figure 2 - In A, CT scan of the chest revealing cystic areas permeated by fibrotic tissue in the right lower lobe. In B, cystic areas in the posterior segment of the right lower lobe.

Figure 3 - In A, systemic artery, which branched from the thoracic aorta, irrigating the pulmonary sequestration. In B, intermediate bronchus containing an exophytic lesion.
and venous contrast has allowed definitive diagnosis because of its enhanced capacity to depict communication between the anomalous artery and the PS.\(^\text{10,13}\)

The treatment for PS is surgical resection of the sequestered lobe or segment by thoracotomy or even by video-assisted thoracoscopic surgery. In both cases, the success of the procedure depends on adequate knowledge of the vascular anatomy of the PS and on early ligation of the artery irrigating the PS, because, as previously stated, accidental transection of that artery can lead to massive hemorrhage, with fatal consequences.\(^\text{4,14}\) In the case reported here, there was significant hemorrhage, but it was contained by mass clamping and subsequent ligation with continuous sutures.

There have been few reports of PS accompanied by lung cancer, and there has been only one case in which the histological subtype was carcinoid tumor.\(^\text{16}\) In a report published in 1985, the PS and the carcinoid tumor were located in the left lower lobe.\(^\text{16}\) In the case presented here, the PS and the carcinoid tumor were located in different lobes, but both were located in the right lung.

A carcinoid tumor is a rare type of neuroendocrine tumor, derived from enterochromaffin cells, and occurs primarily in the gastrointestinal tract. When a carcinoid tumor occurs in the bronchopulmonary system, it can manifest as recurrent episodes of pneumonia, cough, hemoptysis, and chest pain.\(^\text{16}\) In the case reported here, the major symptom experienced by the patient was hemoptysis, and there was a history of recurrent pneumonia. These two characteristics are common to PS and to carcinoid tumors, a fact that made the preoperative diagnosis of PS difficult, because the symptoms of the patient were attributed to the neoplasm and to chronic obstruction.

Despite being rare, pulmonary malformations can present similarly to more common infectious pulmonary diseases, such as bronchiectasis. Therefore, we emphasize the importance of ancillary investigation in patients with a history of recurrent pulmonary infections and cystic changes in imaging studies, because these problems cannot always be attributed to a single, more evident cause. Since PS is a rare pulmonary malformation and has subtle clinical manifestations, it requires tests that are more complex, and, if unsuspected, it might appear only as an intraoperative finding following potentially fatal complications.

References

About the authors

Fernando Luiz Westphal
Coordinator of the Teaching and Research Center. Getúlio Vargas University Hospital, Federal University of Amazonas School of Medicine, Manaus, Brazil.

Luís Carlos de Lima
Physician-in-Chief. Department of Thoracic Surgery, Getúlio Vargas University Hospital, Federal University of Amazonas School of Medicine, Manaus, Brazil.

José Corrêa Lima Netto
Attending Physician. Department of Thoracic Surgery, Getúlio Vargas University Hospital, Federal University of Amazonas School of Medicine, Manaus, Brazil.

Maria do Socorro Lucena Cardoso
Professor of Pulmonology. Federal University of Amazonas School of Medicine, Manaus, Brazil.

Márcia dos Santos da Silva
Physician. Getúlio Vargas University Hospital, Federal University of Amazonas School of Medicine, Manaus, Brazil.

Danielle Cristine Westphal
Medical Student. Federal University of Amazonas School of Medicine, Manaus, Brazil.