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Sclerosing Hemangioma of the Lung: Report of a Case

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Abstract

Sclerosing hemangioma of the lung is extremely rare. Herein, a case of sclerosing hemangioma of the lung in a 44-year-old woman is reported. On admission, the patient complained of pain in the left lower neck and upper chest. Chest x-ray revealed a mass in lingular lobe of the left lung. The patient underwent surgery and a mass was removed through left posterolateral thoracotomy. Pathologic findings confirmed sclerosing hemangioma. Sclerosing hemangioma of the lung should be considered in asymptomatic patients or those with symptoms in chest or lower neck.


Keywords • Hemangioma • lung • surgery

Introduction

Sclerosing hemangioma of the lung (SHL) is generally solitary and considered a benign lesion. It grows slowly and lymph node metastasis is rarely reported. In 1956 Liebow and Hubbel defined pulmonary fibrosing hemangioma as a markedly vascular proliferation with a tendency to fibrosis, and associated with papillary vegetation, extensive histiocytic infiltration and hemorrhages at various stages of organization. Currently, it is considered as a tumor proliferation which, on the basis of data from electron microscopy and histochemistry, is felt by some authors to be of vascular while others underscore an epithelial origin which develops apparently from immature type 2 pneumocytes. However, the designation “Benign Fibrosing Pneumocytoma” suggested by Chan seem to be more appropriate. This lesion occurs most often in the middle-aged women. Herein, we describe a case of benign sclerosing hemangioma of the lung presenting with pain in the left lower neck and upper chest and good surgical outcome.

Case Presentation

A 44-year-old female was admitted to thoracic surgical ward with a history of pain in the left lower neck and left upper chest of two months duration. There was no history of cough, fever, or weight loss. On general physical examination, there were decreased respiratory sounds in the left upper chest. She was otherwise completely well. Chest x-ray showed a well-defined opacification in the lingula and left upper lobe (Fig 1). On CT scan, a stricture of the lingular bronchus with a well-defined consolidation was noticed. In bronchoscopic examination compression of the left upper lobe bronchus was detected. Pathologic examination of bronchoscopic biopsies was negative for malignancy. The patient underwent a left posterolateral
thoracotomy and a LUL with lingula lobectomy. Pathologic examination of the mass revealed benign sclerosing hemangioma of the lung. (Fig 2)

Discussion

Sclerosing hemangioma proposed by Liebow and Hubbel defines a subset of benign tumor and tumor-like lesions of the lung. This subset differs from plasma cell granuloma and pseudolymphoma and may include cases of xanthomatous pseudotumor, post-inflammatory pseudotumor, fibrous histiocytoma, alveolar angiofibroma, and pneumocytoma. The cells of origin were initially considered to be endothelial akin to a tumor of the dermis of the skin.³ Macroscopic appearance of the tumor was white to yellowish-brown in color and fleshy to fibrous in consistency. A thin fibrous pseudocapsule separated the tumor from adjacent pulmonary parenchyma and the surface of the tumor was hypervascular, tending to bulge out on sectioning. Subsequent electron microscopic studies suggested either alveolar pneumocytes or mesothelium as the origin. Multiple authors utilizing immunologic studies have supported the hypothesis that both bronchioles and alveolar type 2 pneumocytes constitute simultaneously the cells of origin.³,⁵,⁶ However other investigators now consider this tumor to be primarily a proliferation of the epithelial cells. The results of staining were positive for epithelial membrane antigen and subsequent apo-protein, which are the markers of epithelial cells, and negative for factor 8 related antigens of endothelial cells suggested that the origin of the SHL might be type 2 pneumocytes.⁴ Anti-lung surfactant apoprotein monoclonal antibody was detected in the rough endoplasmic reticulum in some of the cells in the solid areas. In spite of these studies the exact histogenesis of SHL remains uncertain as was also the case in some previous studies.²,⁷ Microscopically the histologic features of SHL varied and four major histologic patterns were found in the same histologic section: solid, papillary, hemangiomatosus and sclerotic patterns.⁴,⁵,⁶,⁶

Sclerosing hemangioma of the lung is recognized as a distinct clinicopathological entity. It is a benign neoplasm, probably of epithelial origin.
Sclerosing hemangioma of the lung

Clinically the tumor is asymptomatic and is mostly seen in middle aged women.\(^9\)

Chest pain, hemoptysis, fever, and productive cough are rarely seen.\(^1\) SHL is often detected incidentally as a round, well-defined, homogenous mass on routine chest radiography.\(^1,9\) Sometimes an air trapping zone surrounding SHL or air meniscus sign is observed. A paranchymal mass without calcification on chest X-ray is suggestive of benign SHL.\(^1\) Bilaterality is a rare finding.\(^1\) Metastasis to multiple lymph nodes and contralateral lung is also rarely seen.\(^1,11\) The lesion usually presents as a solitary mass or nodules. The greatest diameter of the tumor on chest roentgenogram has ranged from 13 to 82 mm.

The diagnosis is based on histo-pathohistologic examination of the biopsy material. Therapy is surgical and prognosis is excellent.\(^9\) In our case, pathologic findings were of solid and papillary type in accordance with the findings reported elsewhere.\(^4\)

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References

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