Intrapulmonary Solitary Fibrous Tumor of the Lung: A Rare Case Presentation

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ABSTRACT
Localized fibrous tumors of the lung arise from the visceral pleura and are pedunculated. They also project into the pleural cavity. The tumor with an entirely pulmonary location is extremely rare. We present here a rare case of intrapulmonary localized fibrous tumor with review of the literature.

Keywords: Immunohistochemistry, Sphindle cell neoplasim, Solitary fibrous tumor.

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INTRODUCTION
Solitary fibrous tumor (SFT) is a rare neoplasm (2.8 cases/1 lakh persons/year), which frequently arises from the pleura, especially the visceral pleura, but intrapulmonary SFT is extremely rare. They generally have an indolent course and are often an incidental finding on chest images. Here, we report a rare case of intrapulmonary localized fibrous tumor.

CASE REPORT
A 50-year-old female, housewife, presented with generalized dull aching pain in the left chest, associated with breathlessness [Modified Medical Research Council (MMRC) – grade III] of 3 months duration. Patient had no significant past history. Routine blood investigations were within the normal limits. Chest X-ray showed a homogenous opacity occupying the left mid and lower zones (Fig. 1). A computed tomography (CT) scan of the chest confirmed the presence of a large nonenhancing heterogeneous solid mass lesion measuring (11.2 × 7.8 × 12 cm) in the left lower lobe with no evidence of lymphadenopathy (Figs 2 and 3). Ultrasonography (USG)-guided
Localised fibrous tumors of the pleura are exophytic pedunculated masses that extend from the visceral pleura into the thoracic cavity, and inward growth into the lung parenchyma is rare. Such tumors are then termed “inverted.” In our case, the tumor did not show histologic continuity with the visceral pleura. In addition, alveolar pneumocytes and small bronchioles that retained their histologically benign appearances were entrapped within fibrous cells of the tumor. Hence, we presume this tumor to have arisen from the parenchyma of the lung. Early reports of “fibro-adenoma” of the lung may represent this spindle-cell tumor with entrapped alveolar epithelium. Yousem and Flynn suggest that this localized fibrous tumor arose not from the pleura, but rather from the lung parenchyma itself. Apart from many theories, two main hypotheses for their entirely parenchymal location have been proposed. First, the subpleural mesenchyma is in direct continuity with the connective tissue of the interlobular septa, and intrapulmonary fibromas may arise from the septal mesenchyma or invagination of the visceral pleura. Second, these tumors may originate from facultative fibroblastic elements, which can be seen in the submesothelial area of normal pulmonary parenchyma. These elements have ultrastructural and immunohistochemical features similar to those of subpleural connective tissue elements. Our case may be in agreement with the latter hypothesis because of entrapped small bronchioles within the tumor. Hence, with this parenchymal location of the lung, the tumor should not be pathologically mistaken for carcinosarcoma or adenocarcinoma with a metaplastic spindle-cell component.

CONCLUSION

A SFT arising from the lung parenchyma is extremely rare. Complete surgical resection is the treatment of choice. Due to the rarity of these tumors, a long-term follow-up of cases is needed to clarify their clinicopathologic behavior and rule out malignant transformation.

REFERENCES