Large cystic-solid pulmonary hamartoma: A case report

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Abstract

BACKGROUND
It now seems that all pulmonary hamartomas (PHs) are large cystic-solid lesions that are difficult to diagnose. However, few cases of large cystic-solid PHs have been reported. The present case report presents a large cystic-solid PH and provides a literature review of the imaging features, formation mechanism and histopathological basis of PHs.

CASE SUMMARY
A 53-year-old woman with no clinical symptoms underwent a chest computed tomography (CT) examination at our hospital. Nonenhanced CT images revealed a large, flat tumor with multiple air-containing cysts in the left thoracic cavity and a cystic part confined to the medial side of the tumor; the solid part of the tumor showed abundant fat and lamellar soft tissue components. Multiple small blood vessels were detected in the solid part of the tumor on contrast-enhanced CT images. Given the large size of the lesion, the patient elected to undergo surgery. Histological examination revealed PH. A detailed review of the patient’s CT imaging showed that the lesion had a small vascular pedicle to the left lower lobe, which was a clue to its lung tissue histological origin. According to immunohistochemical staining, the confined multiple air-containing cysts were caused by the entrapment of respiratory/alveolar epithelium.

CONCLUSION
This case shows the imaging manifestations of a large PH. Heightened awareness of its formation mechanism and histopathological basis may alert radiologists to
consider this diagnosis in their daily workflow.

**Key Words:** Lung; Lung benign lesion; Hamartoma; Computed tomography; Case report

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**Core Tip:** We describe a large pulmonary hamartoma (PH) and its preoperative computed tomography (CT) imaging features, including multiple air-containing cysts, a rich blood supply and a vascular pedicle. The CT imaging features, formation mechanism, and histopathological basis of a large PH are summarized in this case report.

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**INTRODUCTION**

Pulmonary hamartoma (PH) has been defined as a mesenchymal tumor consisting of varying combinations of cartilage, fibrous tissue, fat, smooth muscle, and respiratory epithelium derived from entrapped adjacent lung tissue[1]. It is the most common benign neoplasm and usually presents as solitary nodules in the lung. However, PH can show unusual characteristics and can be clinically and radiologically challenging to diagnose preoperatively. In addition, PHs larger than 10 cm and containing multiple air-containing cysts are rare. In this case report, we present a rare case of a large PH with multiple air-containing cysts. We aim to increase the awareness of its formation mechanism, histopathological basis, and computed tomography (CT) imaging features through a literature review. This diagnosis should be considered in the daily workflow to improve the accuracy of the preoperative diagnosis of this disease.

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**CASE PRESENTATION**

**Chief complaints**
A 58-year-old woman who had never undergone a chest CT examination had a CT scan as part of a routine physical examination. Her medical history was negative for any symptoms of discomfort.

**History of present illness**
The patient’s history was unremarkable.

**History of past illness**
The patient underwent a hysterectomy for myoma 4 years prior. The patient had hypertension for 10 years, but her blood pressure was stable under drug control.

**Personal and family history**
No personal and family history.

**Physical examination**
No abnormal positive indications were found in physical examination.

**Laboratory examinations**
The blood biochemistry results were normal. Pulmonary function testing, arterial blood gas evaluation and electrocardiogram results were normal.

**Imaging examinations**
Initial nonenhanced chest CT images revealed a well-defined tumor with multiple air-containing cysts confined to the medial side of the tumor, and the solid part of the tumor showed abundant fat and lamellar soft tissue components. The tumor was well defined except for a locally unclear boundary with the left lower lung lobe (Figure 1A and B). Further contrast-enhanced chest CT examination showed
multiple small blood vessels in the solid part of the tumor, and several blood supplies to the tumor were detected coming from the left lower lobe (Figure 1C and D).

**FINAL DIAGNOSIS**

The final diagnosis after histological confirmation was a large PH (Figure 2).

**TREATMENT**

Single-hole exploratory video-assisted thoracoscope surgery was performed. There was no adhesion between the tumor and the lung tissue, except for a thin vascular pedicle connecting the tumor to the left lower lobe. The pedicle was dissected, and the tumor was completely removed. Gross examination showed a soft and flat-shaped tumor measuring 14.5 cm × 11.0 cm × 2.5 cm in size (Figure 2A). The multiple cystic components within the tumor were confined to one side, and the diameter of the cysts ranged from 1 cm to 3.5 cm.

**OUTCOME AND FOLLOW-UP**

Electron microscopy suggested that the well-developed epithelium lacked significant cytological atypia in the cystic part. Other parts had mesenchymal components, including fat, connective tissue and smooth muscle (Figure 2B-E). Immunohistochemical staining of the tumor was consistent with the components of normal lung tissue. Smooth muscle cells were observed in the tumor (SMA +) and were positive for desmin. Ciliated respiratory epithelium that lined clefts tested positive for thyroid transcription factor-1, napsin A and cytokeratin 7, and basal cells located within these epithelia tested positive for S-100, which indicated that these epithelia represented entrapped bronchioles and alveolar walls. Immunostaining with HMB45 was negative. The proliferation index Ki67 was low (< 5%). The patient recovered well after surgery, and no obvious abnormality has been found by chest CT examination at annual follow-ups thus far.
PH is the most common benign tumor of the lung. It is relatively easy to make a preoperative diagnosis of PH with typical CT imaging findings, such as a well-defined nodule with a size of less than 2 cm, popcorn-like calcification and a fat density component. Large PHs over 10 cm are unusual, and large cystic-solid PHs are even rarer. The final diagnosis of a large cystic-solid PH depends on postoperative pathology. The most common cause of these cysts is entrapped pulmonary epithelium. Although entrapment of the pulmonary epithelium by PH is well known, in our experience, the CT imaging features of this phenomenon have not received sufficient attention. We decided to review the literature on cystic-solid PHs, analyze their CT imaging features, formation mechanism and histopathological basis, and then discuss the sources of the challenges during preoperative diagnosis.

To our knowledge, only eleven cases of cystic-solid PHs have been reported thus far, of which 6 PHs were larger than 10 cm. The reason for the cyst formation is still unclear. Nevertheless, the literature focusing on this issue is sparse. According to the study of Erber et al [2], the entrapment of respiratory epithelium in primary and metastatic intrapulmonary nonepithelial neoplasms is a frequent morphological pattern but to variable extents. Their study involved 38 patients with pulmonary metastases (81%) and 8 patients with primary pulmonary nonepithelial lesions. There are two types of histological distribution of the entrapped pulmonary epithelium. In type one, the entrapped pulmonary epithelium is distributed mainly in the peripheral portion of the tumor, and in type two, the entrapped pulmonary epithelium is found throughout the tumor, albeit to a varying extent. Although the number of patients
was limited, we thought this conclusion could be extrapolated to more primary and metastatic intrapulmonary nonepithelial neoplasms in the lungs. Because PH is the most common form of primary pulmonary nonepithelial lesions, the same applies to our case. Different types of histological distributions of entrapped pulmonary epithelium produce different CT images. Type one represents the histopathological basis of the cysts in the present case. The entrapped pulmonary epithelium was located at the margin of the tumor and connected to the adjacent lung tissue by a vascular pedicle. In this type, the cysts are dilated bronchioles lined by flattened epithelial cells and containing mucoid secretion to branching leaflet-like papillary spaces. All of the factors mentioned above result in differences in epithelial secretion and immunohistochemistry. Notably, varying degrees of fluid are observed in the cysts of cystic-solid PHs. According to a previous study,[2] glands in the entrapped pulmonary epithelium frequently show a reactive/regenerative appearance. Furthermore, gland size and type vary greatly from small acinar-type glands or microcystic spaces lined by flattened epithelial cells and containing mucoid secretion to branching leaflet-like papillary spaces. All of the factors mentioned above result in differences in epithelial secretion function. Therefore, in previous case reports, various degrees of fluid were observed in the cysts of cystic-solid PHs: The cysts may be well inflated[3-9] or partially[3] or even completely filled with fluid[10-12].

In addition, through a literature review, we found that the CT image density of cystic-solid PHs can vary from ground glass density to solid density depending on the proportion of the solid part. In some cases, the proportion of solid components in cystic-solid PHs is very low, and cystic-solid PHs show extreme CT imaging, that is, a ground glass nodule appearance[12]. It is difficult to distinguish cystic-solid PHs from adenocarcinomas, which often present as ground glass nodules, and the final diagnosis depends on postoperative pathology. The other extreme case is that if the cystic-solid PH is dominated by the solid part, the cystic part may be too small to be observed on CT imaging[13].

Previous studies have demonstrated a high frequency of rearrangements involving 6p21 or 12q14-15 in PH[14] and HMGI-C and HMGI(Y) protein expression as a consequence of rearrangements involving 6p21 and 12q15[15]. These findings support the view that mesenchymal components of PHs represent neoplastic mesenchymal proliferation rather than neoplasms. Today, even with advancements in medical therapy, pulmonary resection remains the most important treatment measure for patients with PH[16,17]. However, controversy exists about the indication for surgery. For large cysts dominated by cystic-solid PHs, although malignant transformation of PHs is exceptional, prompt surgical resection is the recommended treatment. The main reasons are as follows. First, larger cystic-solid PHs are often located under the visceral pleura, similar to the present case, and separated from the thoracic cavity by only a thin layer of pleura (Figure 2F), so the cystic part is more vulnerable to rupture and can lead to secondary pneumothorax[3,18]. In addition, Secretions into the cysts of cystic-solid PHs are difficult to expel from the lungs and may lead to secondary infection. The patients involved in the present case and in the large cystic-solid PH cases discussed above had very good prognoses with uneventful outcomes after surgery.

**CONCLUSION**

Due to its epithelial involvement, clinicians and radiologists should be aware that cystic-solid PH is a diagnostic possibility in adults with large intrathoracic cystic-solid tumors. Cysts in PHs can show different features on CT images depending on the type of histological distribution of the entrapped pulmonary epithelium. If large cysts dominating cystic-solid PHs are treated in a timely manner after discovery, the patient will have a good prognosis.

**FOOTNOTES**

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