Mediastinal hemangioma mimicking an invasive tumor growth: A case report
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1. Introduction

Mediastinal hemangiomas are rare neoplasms accounting for 0.5% of mediastinal tumors without any specific imaging findings [1,2]. They may exhibit an infiltrating appearance into the surrounding tissue, and extensive surgery is often required despite its benign nature [3,4]. We here report a case of a mediastinal hemangioma mimicking an invasive tumor growth that required a combined resection of the lung and diaphragm. This work has been reported in line with the SCARE criteria [5].

2. Case presentation

An asymptomatic 73 year-old man with a remote history of surgically treated gastric cancer at the age of 40 presented with a 50 mm-sized mass on a chest radiography (Fig. 1). He had no significant familial history without any drug use nor allergic history. Chest contrast-enhanced Computed Tomography (CT) revealed an irregular homogenous mass in the anterior mediastinum, which extended along the left diaphragm without any specific enhancing effects (19–27 Hounsfield Unit) (Fig. 2ab). Magnetic Resonance Imaging (MRI) revealed a septic cystic lesion with a high intensity on the T2-weighted image (Fig. 3a). Fat-suppressed T1-weighted images revealed that the mass contained less fatty tissue (Fig. 3b). An invasive thymic epithelial tumor or soft tissue tumor were suspected mainly because of its location and a radiological extent, and a radical excision was planned via video assisted thoracoscopic surgery by an attending thoracic surgeon at our institute.

With the patient in the right semi lateral decubitus position, a total of 3 ports were placed on the 6th, 7th, and 8th left intercostal anterior axillary lines, respectively. Under Carbon dioxide insufflation at a pressure of 8 mmHg, the lesion was visualized adjacent to the pericardium. The lesion was also observed to have grossly invaded into the diaphragm and lower lobe of the left lung. He underwent a total tumorectomy with a combined resection of the pericardial fat, left lung, and diaphragm. Since a complete resection was achieved, no intraoperative frozen section examination was performed. The postoperative course was uneventful and the patient was discharged on the third postoperative day.

Abbreviations: CT, computed tomography; MRI, magnet resonance imaging.
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The macroscopic findings of the specimen revealed a circumscribed mass embedded in the fat tissue (Fig. 4). The histopathological findings revealed dilated medium sized blood vessel proliferation in the adipose tissue. Those vessels were mainly muscular veins with a small number of arteries. Those findings were compatible with the diagnosis of a mediastinal hemangiomma (Fig. 5a). Only fibrous adhesions were observed between the tumor and resected lung and diaphragm without any histological invasion (Fig. 5b).

3. Discussion

Mediastinal hemangiomas are rare neoplasms accounting for 0.5% of mediastinal tumors and are frequently found in the anterior mediastinum [1,2]. They have no specific symptoms and may present with a variety of clinical manifestations such as a cough, respiratory distress, and chest pain [2,4]. They often require surgical intervention to avoid disease related morbidities such as massive hemoptysis or respiratory failure [6,7].
While most cases are generally visualized with well circumscribed masses [4,8,9], some can exhibit a wide range of radiological findings mimicking dumbbell tumors [3,10] or an infiltrative-like progression as shown in the present case [11]. Unlike other mediastinal tumors, even an incomplete resection would be feasible for hemangiomas without a local recurrence, malignant degeneration, hemorrhagic morbidity, or becoming symptomatic [1,9]. Therefore, a preoperative diagnosis is essential to avoid extensive surgery. While imaging examinations play a key role, mediastinal hemangiomas may pose a diagnostic challenge mainly due to its rarity. Positron emission tomography CT was not available in the present case and its diagnostic significance in hemangiomas is unclear [12].

On MRI, a fat component and high intensity on the T2-weighted images have been reported as characteristic of a mediastinal hemangioma and likewise of hepatic hemangiomas [12,13]. Part of those findings were detected in the present case but were not diagnostic because a hemangioma had not been raised as a differential diagnosis due to its rare frequency. Peripheral nodular enhancement on the dynamic MRI has been also reported to be diagnostic but was not performed in this case [14–16].

Calcified phleboliths derived from organized thrombi are the most diagnostic sign, and are detected more sensitively by CT than conventional radiography [12,17]. However, they are not so frequent and account for only 10–52.9% at most [2,13,18,19] and were not identified in this case.

On the other hand, it has been reported that peripheral puddle images showing a punctate contrast effect on the margin of the tumor in the aortic phase on the dynamic CT is highly specific for hepatic hemangiomas [20], and its usefulness has also been reported for mediastinal hemangiomas with a sensitivity of about 70% [14–16]. With a clinical suspicion of a hemangioma during the diagnostic work up, a dynamic CT might have been helpful and we could have chosen a more limited surgery and avoided the extensive resections in this case.

4. Conclusions

Mediastinal hemangiomas are a rare entity frequently developing in the anterior mediastinum. They could exhibit an infiltrating appearance and often are amenable to extensive surgery despite being totally benign in nature. Therefore, a preoperative diagno-
sis is essential to avoid extensive surgery. A hemangioma should be raised as a differential diagnosis of anterior mediastinal tumors, especially in cases with an infiltrative appearance suggesting the necessity for a combined resection of the surrounding organs.

**Declaration of Competing Interest**

The authors report no declarations of interest.

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**Ethical approval**

Not applicable.

**Consent**

Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

**Author contribution**

Shogo Yobita wrote this paper. All authors read and approved the final manuscript.

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**Fig. 5.** a) The histopathological findings showing dilated medium sized blood vessel proliferation in the adipose tissue (arrows, the scale indicates 200 μm). Those vessels were mainly muscular veins with a small number of arteries. b) Only fibrous adhesions were observed between the tumor and resected diaphragm (arrowheads, the scale indicates 1000 μm).

**Availability of data and material**

Not applicable.

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