A Mediastinal Liposarcoma Resected Using a Double Approach with a Thoracoscope

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Abstract

We report a case of a 45-year-old woman who underwent a complete resection of a liposarcoma using thoracoscopic and cervical approaches. General checkup and computed tomography revealed a large mediastinal tumor occupying the thoracic outlet, which had reached the posterior thyroid region, and another small tumor at a subcarina. A cervical method for evaluating a neck lesion and a thoracoscopic (video-assisted thoracic surgery) approach for assessing a mediastinal lesion were performed. This double approach provided excellent visualization and enabled us to perform fine manipulation even within the narrow thoracic outlet region. The patient was disease free at 11 months after surgery.

Keywords
► mediastinal tumor
► sarcoma
► thoracoscopy/VATS

Introduction

Liposarcomas rarely arise in the mediastinum and comprise less than 1% of all mediastinal tumors.1,2 Complete surgical resection is thought to be the best treatment for liposarcoma because the role of adjuvant therapy has not been established for these tumors. Due to the tendency for liposarcomas to be large in size, a long incision is usually required for complete resection of these tumors. We here present a case of a mediastinal liposarcoma, which was located from the neck to the mid mediastinum, and which was resected completely using a double thoracoscopic (video-assisted thoracic surgery [VATS]) and cervical approach.

A 45-year-old woman presented at a community hospital with a mediastinal mass detected during a general checkup. She had no prior symptoms. A chest X-ray revealed a large mass shadow in her right chest. Chest computed tomography (CT) imaging showed a heterogeneous, regular-shaped mediastinal tumor (lesion A), occupying the thoracic outlet, and which had attained a posterior thyroid region location. An additional tumor was also detected by CT at a subcarinal region (lesion B) (►Fig. 1A), considered to be a lymph node. Magnetic resonance imaging (MRI) showed lesion A as iso-intense on T1W1, heterogeneous high intensity on T2W1, and lesion B as high intensity on T2W1, which revealed no notable invasion to surrounding structures (►Fig. 1B). Bronchoscopic and transcervical fine needle aspiration cytology and biopsy did not detect malignancy. These findings indicated that the lesion was a soft tissue tumor, either a solitary fibrous tumor or a sarcoma of unknown origin.

The cervical approach via collar incision was initially performed before VATS approach. Because the mediastinal part of lesion A was larger than its cervical part, we concluded that the tumor should be pulled out from neck into the thorax. The tumor showed no adhesion or invasion to the tissue surrounding the cervical region and hence could be divided easily (►Fig. 1C). After dividing the tumor, the collar incision was closed and a lateral position was adopted for the VATS procedure. Three VATS ports were placed in the middle portion of the right chest (►Fig. 1D). In the thoracic cavity, the thoracic outlet was occupied by the tumor, which was almost free from the surrounding organs, except for fibrous adhesion.
in places (►Fig. 1E). The dissection was preceded along the parietal pleura until the apical thorax. At the apical thorax, we carefully tunneled through the thoracic outlet to the neck (►Fig. 1F). Lesion A was then pulled out gently into the thorax. Lesion B which was next to, but independent of, lesion A was also extirpated. Both tumors were completely resected with negative margin thoracoscopically.

Macroscopically, lesion A, measuring 12.7 x 10 cm, was encapsulated and the cut surface was solid and uniformly white (►Fig. 2A). Lesion B, measuring 3.5 x 2.7 cm, appeared pale, yellow, and gelatinous with a spotty hemorrhage (►Fig. 2E). Histology revealed, lesion A was a fibrous tumor which consisted of spindle cells with mild nuclear atypia (►Fig. 2B), and lesion B was an adipocytic tumor having abundant myxoid stroma and scattered hyperchromatic stromal cells and atypical lipoblasts (►Fig. 2F). Immunohistochemistry (IHC) with CDK4 and MDM2 antibodies showed weak nuclear positivity in lesion A (►Fig. 2C, D) and B (►Fig. 2G, H). Additionally, fluorescence in situ hybridization (FISH) revealed co-amplification of MDM2 and CDK4 in both lesions (►Fig. 2C, D, G, H). Finally, she was pathologically diagnosed with dedifferentiated liposarcoma forming two separate lesions.

The postoperative course for this patient was uneventful with no serious wound pain. She was discharged from the hospital on postoperative day 11, and was disease free at 11 months postsurgery.

**Discussion**

A mediastinal liposarcoma accounts for only approximately 9% of primary sarcomas of the mediastinum.1,2 Recent advances in the molecular genetic characterization have led to the classification of liposarcoma into three main histogenetic subtypes: well-differentiated/dedifferentiated liposarcoma, myxoid liposarcoma, and pleomorphic liposarcoma.3 The prognosis is highly dependent on histological subtype—Well-differentiated/dedifferentiated liposarcoma has a particularly protracted clinical course, and myxoid and pleomorphic subtypes tend to progress rapidly.2,4 Therefore, adequate pathological classification is essential for determining prognosis and therapy. In our current case, with their histology, it is considered that lesion B was well-differentiated liposarcoma, and lesion A was a low-grade dedifferentiated counterpart of lesion B. Demonstration of MDM2-CDK4 by IHC and FISH confirmed the diagnosis.

As with most soft tissue tumors, since radio and/or chemotherapy are believed to be ineffective, complete surgical excision is the appropriate treatment for liposarcoma. Although adjuvant therapy was not undertaken in our patient...
Because the tumor was resected completely with negative margin, a careful follow-up plan was definitely warranted.

Liposarcomas tend to be large in size, and a large thoracotomy is, therefore, often required for a complete excision with a negative margin. For these reasons, anterior or posterior thoracotomy, sternotomy, or clamshell procedures are usually considered. However, these approaches can result in postoperative pain, and morbidity and a prolonged hospital stay because of rib spreading and a long incision. Decker et al reported a successful resection for a large liposarcoma located in an inferior thorax by thoracoscopic intervention. These authors reported that a minimally invasive thoracoscopic procedure resulted in less pain and a shorter hospital stay. In our present patient, the classical neck collar incision and lateral thoracotomy would have been considered in most settings due to the large size, location, and suspicious adhesion of the lesion. Even with these approaches, however, visibility would be expected to be poor and involve a blind spot, especially at the thoracic apex, because of the large size and immovability of the tumor. However, the VATS approach generated excellent visualization without a blind spot and enabled us to perform a safe and more precise operation, even in a narrow space.

In conclusion, the VATS approach should be considered in cases with a complete and less invasive resection of a large liposarcoma located at a thoracic apical region.

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