Atypical uniportal thoracoscopic resection of a giant thymolipoma

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ABSTRACT

Thymolipoma is a rare, slow-growing, benign anterior mediastinal tumor consisting of mature adipose tissue and thymic tissue. Herein we describe the case of a completely excised giant thymolipoma with uniportal video-assisted thoracoscopic surgery, which was detected incidentally on chest X-ray, with the review of the literature. A 30-year-old male patient was hospitalized to the department of general surgery with the diagnosis of appendicitis. Preoperative chest x-ray revealed mediastinal enlargement extending to the diaphragm at right paracardiac area where cardiac contours were erased and cardiomegaly-like appearance was observed. In the contrast-enhanced computed tomography of the chest, a massive mass of soft tissue was detected in fat density. The mediastinal lesion was completely excised with uniportal VATS. Histopathologically, the tumor was reported to be thymolipoma. Thymolipoma it should be totally removed with VATS surgery for both diagnosis and treatment.

Keywords: thymolipoma, uniportal VATS, mediastinal mass, giant
**Introduction**

Thymolipoma is an uncommon benign tumor of the thymus consisting of thymic tissue and mature fatty tissue. It accounts for 2-9% of all tumors originating from thymus [1,2]. It was first described by Lange in 1916 and the term was used for the first time in the literature by Hall in 1949 [1,3].

Thymolipoma may be identified incidentally during a diagnostic screening for other medical problems and may grow large dimensions and manifest themselves clinically by signs of compression on adjacent structures such as shortness of breath, cough, palpitations, hypotension, tachypnea, the fullness of the neck veins and edema in the lower extremities [4].

It can be seen at any age, but it is most common in children and in young ages. It is difficult to distinguish radiologically from other mediastinal masses, however, hyperplastic thymus and adipose tissue-like density may be determinative in contrast-enhanced computed tomography of the chest. We describe the case of a completely excised giant thymolipoma via uniportal video-assisted thoracoscopic surgery (VATS) which was detected incidentally on chest X-ray, with the review of the literature.

**Case Report**

A 30-year-old male patient was hospitalized to the department of general surgery with the diagnosis of appendicitis. On preoperative chest X-ray, cardiomegaly-like appearance and mediastinal enlargement extending to the diaphragm were observed in the right paracardiac region and the contours of the right heart were erased (Figure 1).

After being healed from the emergency situation, a contrast-enhanced computed tomography of chest (CCT) was taken under elective conditions. A massive soft tissue mass of fat density and giant size was observed, which completely filled the anterior mediastinum, extending from the right paracardiac area to the diaphragm, with clear borders with the mediastinal structures, not disrupting the contour of the mediastinal tissues (lesion dimension; 213x65 mm) (Figure 2).

Although there was no additional disease in his medical history, when his anamnesis was deepened, it was found that there was shortness of breath and fatigue those had existed for a while and came with effort, were observed. Results of other clinical laboratory tests were normal. Biopsy and additional radiological examination were not performed in this case, because the patient was considered to have benign anterior mediastinal mass lesion and surgery was planned for complete excision. After the preoperative preparations were completed, the patient underwent left selective intubation, with a double-lumen endotracheal tube and the hemithorax was explored via uniportal VATS (3 cm incision) approach incision through the right fourth intercostal space on anterior axillary line. The mediastinal pleura was dissected and an encapsulated lesion of benign character was observed. After the thymic vein was clipped from the superior mediastinum, the left side mediastinal pleura was also opened (Figure 3).
The capsule was preserved and the lesion was dissected from the pericardial and diaphragmatic surfaces with the help of electrocautery. At this point, the best effort was taken to preserve the phrenic nerve from an unintentional injury. The lesion could not be removed in one piece because of its giant size, was extracted with two separate endobags in two pieces, by dividing it into two in the endobag while maintaining the safe oncological principles (Figure 4).

The incision was lengthened by 1 cm for lesion removal. Assistance was received from endoscopic and conventional surgical instruments that we constantly use in uniportal VATS application.

Tube thoracostomy was performed from the same incision and no postoperative complication was observed in this case. On the second postoperative day, the chest tube was removed and on the third day, he was discharged after an uneventful course (Figure 5).

On macroscopic examination of the specimen, encapsulated two masses weighing totally 666 g, with the size of 16.5x11x3.5 cm and 13x10.5x3.5 cm were detected. The cut surface of the masses was solid and yellow-colored. Histopathologically, the tumor was reported to be thymolipoma. It was characterized by mature adipose tissue surrounded by a thin fibrous capsule and admixed with strands of atrophic thymic tissue containing Hassall corpuscles and calcification in some areas (Figure 6). The patient has been followed up for 10 months without any problem.

Written informed consent was obtained from the patient for publication of his data.

Discussion

Thymolipomas are characterized by mature adipose tissue. It is a slow-growing, encapsulated tumor of both
mesodermal (fat) and endodermal (thymus) origin. Different from thymoma, it doesn’t show mitosis and atypia, mature adipose tissue can be clearly separated from thymus tissue, the surrounding tissues usually don’t show invasion and malignant behaviour, such as distant metastasis. These tumors appear especially in the 3rd and 4th decades and are mostly asymptomatic. However, they can rarely grow to huge dimensions and be symptomatic. In our case tumor was detected incidentally in the 3rd decade and although the lesion enlarged to giant dimensions, it didn’t cause serious symptoms. In some studies, cases of thymolipoma with progressive huge dimensions; it has been reported that there were symptoms of dyspnea, chest pain, cough and respiratory infections [4,5]. In the differential diagnosis of anterior mediastinal lesions, the most effective method in radiological evaluation is contrast-enhanced CCT [6,7]. There are several publications in which CCT is inadequate and MRI has advantages in cases where the borders of mediastinal structures cannot be clearly separated and benign-malignant distinction cannot be made clearly [8,9]. In our case, CCT was observed sufficient in differential diagnosis and additional radiological examination was not required.

The surgical approach preferred in mediastinal lesions growing huge dimensions is sternotomy and throracotomy [2,10]. Thoracoscopic surgery has not been recommended due to the inability to remove these giant size lesions completely, the spread of tumor cells to the pleura and the risk of recurrence and VATS applicability has been reported only in small size lesions [10,11]. Uniportal VATS is a commonly used surgical approach in our clinic for mediastinal lesions. This surgery was completed with right uniportal VATS because it was considered as a benign and encapsulated lesion. Since the lesion was too large to be extracted with a single endobag, the giant lesion dissected from all mediastinal structures was divided into two parts in the endobag and completely extracted with two separate endobags.

Radiologically, lipoma, liposarcoma, teratoma, cardiomegaly, pericardial fat pad, pericardial effusion, pericardial tumors, congenital diaphragm hernias should be considered in the differential diagnosis. Making the correct diagnosis for thymolipoma by fine-needle aspiration cytology is not always possible, because the cytological material should contain both thymic and lipomatous components for differential diagnosis with other lipomatous tumors. In our opinion, it is not very satisfactory to perform needle biopsy to such type of lesions for diagnosis. In our case the differential diagnosis and the type of surgical approach were considered via CCT so that a preoperative biopsy was not indicated. Both definitive diagnosis and treatment were achieved with complete surgical resection.

Although the thymolipoma grows to huge dimensions, it may not show any symptoms and progress without symptoms. Mostly heavier than 500 g when diagnosed. In our case was asymptomatic and weighed 666 g.

As a conclusion, although thymolipoma is a rare benign lesion with no malignant features and no recurrence, it can grow to giant sizes and cause compression signs and symptoms. Therefore, complete excision should be performed with surgery for diagnosis and treatment. It should be kept in mind by experienced clinics that VATS is a good choice of approach, particularly in cases considered to be benign, even in the tumors grown to huge sizes.

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