Giant Mediastinal Thymolipoma in 35-Year-Old Women

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ARTICLE INFO

Article Type:
Case Report

Article History:
Received: 6 January 2011
Accepted: 2 Feb 2011
ePublished: 28 May 2011

Keywords:
Mediastinum
Myoid
Thymolipoma
Thymus

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Introduction

Thymolipomas are rare anterior mediastinal tumors composed of mature adipose tissue and benign thymic tissue arising from thymus gland. This tumor accounts for only a small percentage of mediastinal masses.1,2 The majority of these tumors are clinically quiescent; however, symptomatic patients may present with myasthenia gravis, upper respiratory tract infections, dyspnea, tachypnea, and chest pain.3,4 Thymolipomas are benign neoplasms for which complete surgical excision is curative.

Case Report

The patient was a 35-year-old woman with a six-month history of difficulty breathing and chest pain. An anterior-posterior chest radiograph revealed a widened mediastinum with small lung volumes. During his diagnostic evaluation, a computed tomographic scan was performed and with huge mass in the anterior mediastinum with extension to the left and right side of pleural space. With postero-lateral thoracotomy the huge mass was resected. The patient discharged with good condition.
interpreted as being consistent with mixed obstructive and restrictive lung disease. During his diagnostic evaluation, a computed tomographic scan was performed. There was a huge mass in the anterior mediastinum overlying the heart with extension into the left and right hemithorax, predominantly consisting of fat with scattered linear soft tissue (Figure 1 and 2).

In left side postero-lateral thoracotomy, a large, encapsulated, vaguely lobulated mass weighting 1460g was found within the anterior mediastinum, and resected completely. In cut section, it was primarily consisted of mature-appearing adipose tissue with no area of hemorrhage or necrosis (Figure 3,4 and 5). The mass occupied approximately 80% of left and 50% of right pleural cavity, resulted in marked compression of posterior and inferior of right and left lungs. Histological examination revealed that the tumor was predominantly composed of lobules of mature adipose tissue intermixed with septa of thymic tissue containing abundant Hassall’s corpuscles. In addition, the mass contained numerous polygonal myoid cells that were distributed and localized predominantly in the interface of thymic tissue and adipose lobules. The patient discharged in a good condition on 6th day after surgery. She had no problem until 18 month follow up.

**Discussion**

Thymolipoma is a very rare mediastinal tumor composing mature adipose and thymic tissue arising from thymus gland. It is a rare and benign mesenchymal tumor of mediastinum that is often asymptomatic. This tumor accounts for only a small percentage of mediastinal. Clinically, most thymolipomas are identified incidentally during a diagnostic workup for other medical problems, although some nonspecific signs
like upper respiratory infection, chest pain, dyspnea, tachypnea, and chronic nonspecific chest symptoms have been identified at the time of presentation.\textsuperscript{3} Thymolipomas have been reported to be associated with certain autoimmune medical conditions, such as graves' disease and myasthenia gravis.\textsuperscript{1,6} On computed tomographic scan, the most common patterns seen in thymolipomas are linear whorls of soft tissue intermixed with fat or predominantly fat with scattered linear soft tissue attenuation. Radiographically, thymolipomas may be confused with more common lesions, such as mediastinal teratomas, thymic hyperplasia, lipomas, and cardiomegaly.\textsuperscript{7}

This broad radiologic differential diagnosis stresses the need for histologic evaluation in the diagnosis of anterior mediastinal masses. Anterior mediastinal liposarcomas have been described and liposarcomas of thymic origin do exist.\textsuperscript{8} Mediastinal liposarcomas may grossly resemble benign thymolipomas and may be associated with thymic tissue.\textsuperscript{8} Grossly, most thymolipomas are lobulated and encapsulated, range in size from 4 to 36 cm, and consist of yellow adipose tissue with no areas of hemorrhage and necrosis.\textsuperscript{9} Histologically, thymolipomas have a varying proportion of mature adipose and thymic components. In this case, the thymolipoma consisted of predominantly mature adipose tissue with scattered elongated aggregates and small round nodules of atrophic thymic epithelium embedded within fat. No lipoblast was identified. In one section of the tumor, reminiscent of myoid cells, was identified within the mature fat. Such clusters of myoid cells have been reported previously in thymolipomas\textsuperscript{9} such as present case, some thymolipomas show predominantly mature adipose tissue with only occasional remnants of thymic tissue. However, other thymolipomas have been reported to have a greater proportion of thymic tissue.\textsuperscript{3,5,7}

The histologic differential diagnosis for thymolipomas includes lipoma, well-differentiated liposarcoma, and thymic hyperplasia.\textsuperscript{3} The distinction between a lipoma and a predominantly fatty thymolipoma may be difficult. In the latter, extensive sectioning and immunohistochemical, staining for cytokeratin may highlight thymic epithelial elements in a thymolipoma. Liposarcomas typically have scattered nuclear atypia, lipoblasts, and no thymic epithelium. Thymic hyperplasia classically has unremarkable thymic architecture without presence of abundant adipose tissue. Thymolipomas may adhere to the adjacent structures and displace organs within the chest cavity, but invasion into adjacent structures has not been documented in the literature. The encapsulated and lobular nature of thymolipomas and lack of invasion into adjacent structures usually allow for a relatively uncomplicated surgical excision of the tumor. Our case represents an atypical clinical presentation of a giant thymolipoma. Eighty percent of thymolipomas present within the first 4 decades of life. Thymolipomas are often asymptomatic and are identified incidentally after diagnostic evaluation for nonspecific respiratory symptoms.\textsuperscript{9,10}

In our case, the patient had long-standing respiratory complaints, which appears to have been directly caused by thymolipoma. The compression of the right and left lung by the thymolipoma in our case resulted in increased pulmonary parechima resistance and reduced lung volume. The reduced lung volume resulted in chronic respiratory symptoms and a primarily restrictive lung pattern. The chronically increased pulmonary vascular resistance led to development of pulmonary hypertension, cor pulmonale, heart failure, and ultimately death.\textsuperscript{4,10}

This case highlights an atypical presentation of a thymolipoma, a rare, benign thymic tumor which can achieve a massive size, occasionally resulting in respiratory symptoms and death. Thymolipomas are benign neoplasms for which complete surgical excision is curative as our case.\textsuperscript{3,10}

Ethical issues: The study was approved by the Ethical Committee of the University.

Conflict of interests: No conflict of interest to be declared.

References