A Giant Right Anterior Mediastinal Thymolipoma with Extension to Left Hemithorax and Neck and Misdiagnosed as Liposarcoma on Needle Biopsy

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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 dyspnea, tachypnea, and chest pain, upper respiratory tract infections and rarely myasthenia gravis [3-6]. Diagnosis should be confirmed radiologically during the preoperative work-up, with pathognomonic features demonstrated on both computed tomography (CT) scan and magnetic resonance imaging (MRI) and with fine needle aspiration biopsy [4,7-9].

CASE PRESENTATION

The patient is 30 year old man who has been struggling with progressive chest pain and productive cough, dyspnea and right side neck swelling for the last two years. The symptom has been increased by the last two month. In physical examination in right side of neck a soft tissue mass was palpable. The history of past medical, he had a sever car accident in two year ago and chest-tube insertion in right hemithorax because of hemothorax and laparotomy. Breath sound in right hemithorax was absent. The heart biting was palpable in in five intercostal space in middle axilla line, others was normal. CXR, show complete opassification of right hemithrax. Pulmonary function tests showed a decrease in forced vital capacity (47% predicted) and forced expiratory volume (42% predicted) with a total lung capacity of only 72%. These findings were interpreted as being consistent with mixed obstructive and restrictive lung disease. During his diagnostic evaluation, a computed tomographic scan was performed, CT scan show a huge mass of anterior mediastinum and right hemithorax with extension to left hemithorax and neck, the mass was collapsed totally the right lung and shift mediastinum to the left side and compressed the heart. The Radiologist’s report was liposarcoma or teratoma (Figure 1-3). Haematological and serological investigations, including tumor markers (alpha-fetoprotein, lactate dehydrogenase, beta human chorionic gonadotrophin, and alkaline phosphatase) were normal and not contributory to the diagnosis. With these radiologist report, the patient underwent to US-needle biopsy identified atypical cell and suspicious to liposarcoma. On completion of the preoperative evaluation, a right extensive posterolateral thoracotomy was performed via the fifth intercostal space. A large, encapsulated, vaguely lobulated mass was found within the
anterior mediastinum. The mass arose in the anterior mediastinal fat, increasing in size as it extended above to the right side of neck and to the superior vena cava to the right inferior pulmonary vein, and extended to left hemithorax, compress the heart, diaphragm and shifted the mediastinum to the left. The mass occupied approximately 90% of right and 30% of left pleural cavity, resulted in marked compression and totally collapsed of right lung and partially collapse of left upper lobe (Figure 1-4). After tumor dissection and sectioning of the blood vessels, First right hemithorax and mediastinal masses was resected completely. For prevention of re-expansion pulmonary edema, we did not re-expanded the right lung. Neck portion was resected via the mediastinum. Left side mediastinal and pleural portion was resected complexly. Weight of mass was 5000 g (Figure 5). In cut section of resected mass, it was primarily consisted of mature-appearing adipose tissue with no area of hemorrhage or necrosis. The post-operative period was incident-free, and the patient was discharged on days 5 after surgery. Pathology examination results showed a tumor measuring 31×21×8cm, consisting of fatty tissue and thymic parenchyma (Figure 6,7). There were also foci of collagenous fibrosis and accumulation of cholesterol crystals. No signs of malignant disease were observed. Final pathologist report was thymolipoma. The patient remains asymptomatic during six month follow-up.
Thymolipomas (TL) are very rare, slow-growing mediastinal tumors, accounting for only 2%-9% of all thymus tumors [5]. Thymolipomas are characterized by mesodermic (fatty) and endodermic (thymic epithelium) elements [6]. Thymolipomas 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 [4]. This tumor accounts for only a small percentage of mediastinal masses [6,8]. They are lobulated and well encapsulated, with septal divisions [8]. They consist of large lobules of mature adipose tissue interspersed with small areas of thymic tissue [6]. Thymolipomas usually present as asymptomatic tumors [6]. When the patient does have symptoms, these are usually due to compression of adjacent structures [6]. In our patient presented with dyspnea and chest wall pain and was due to pulmonary and mediastinal compression. Occasionally, this benign tumor may be associated with certain autoimmune disorders, such as myasthenia gravis, hypogammaglobulinemia or red cell aplasia [6]. None of these disorders were observed in our patient.

Although the finding of soft fatty tissue within the tumor with no invasion of adjacent structures on imaging studies clearly suggests a diagnosis of lipoma of mediastinum, it is impossible to make a definitive diagnosis or to even distinguish benign disease from malignancy [6]. Differential diagnosis includes other adipose tumors, such as prominent epicardial fat pad, lipomas, liposarcomas or thymoliposarcomas [2]. Nevertheless, although radiological signs may be non-specific, CT and MRI can still offer useful data [2]. The characteristic signs of (TL) on CT-scan consist of a fatty tissue with strands of white tissue, probably corresponding to islets of normal thymic components [6]. On T1-weighted MRI, fatty tumor tissue is isointense, and in T2-weighted sequences it is suppressed, while the thymic tissue remains are enhanced [6]. Definitive diagnosis is based on histopathological findings. A preoperative fine needle biopsy aspirate (FNAB) is really necessary in all patients with radiological suspicion of thymolipoma [7-9].

CONCLUSION

Thymolipoma is a very rare benign mediastinal tumor, consisting of thymic and fatty tissue. Preoperative diagnosis is frequently based on CT and MRI findings. In review of literature, the role of FNAB in these patients is controversial [6,7]. It is not always easy to differentiate between a thymolipoma and other fatty mediastinal lesions, such as well-differentiated liposarcoma [6]. Romero Guadarrama [8] reported a false positive result on FNAB in a patient with an erroneous diagnosis of well-differentiated liposarcoma [7-9]. On the other hand, Gupta [9] recently published the case of a child diagnosed from a cytology specimen obtained by endoscopic ultrasound-guided fine needle aspiration (EUS-FNA). We decided to perform FNA-needle biopsy in our patient, but cannot have ruled out the need for surgery and after complete surgery of neck and mediastinal mass, pathologist report was Thymolipomas (TL).

The only curative treatment of (TL) is surgical excision. This treatment is helpful to reducing the compression to adjacent structures and relief of symptoms or autoimmune diseases [6,8,10]. Various surgical approaches have been described, including thoracotomy [2,4] sternotomy [6,11,7] or video-assisted thoracoscopy [10]. The decision must be tailored to tumor size and site. We used posterolateral thoracotomy and removed all of right side and left side of mediastinum and cervical portion parts of mass without any complications.

REFERENCES


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