An Unusual Pleural Location of Thymoma Mimicking Malignant Mesothelioma: A Case Report and Literature Review

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Abstract

Primary pleural thymomas are rare tumors often mistaken for malignant mesothelioma clinically and radiologically. This paper reports the case of a 39-year-old woman presented with a pleural tumor in the right thorax. The patient was surgically resected and histopathologically diagnosed as ectopic primary pleural thymoma lacking any evidence of a mediastinal tumor. The clinicopathological futures of this unusual thymoma are also discussed.

Keywords: Tymoma; Pleura; Pleural thymoma

Introduction

The usual location of thymoma is anterosuperior mediastinum, but its occurrence can be seen in extramediastinal regions such as the neck [1], within the thyroid [2], pulmonary parenchyma [3,4], and in the pleura [5-8].

Primary pleural thymomas are rare tumors with less than 30 cases reported to date [9-14]. It is often mistaken as malignant mesothelioma clinically and radiologically.

The present case of ectopic pleural thymoma lacks any evidence of a mediastinal tumor.

Case Report

A 39-year-old woman was hospitalized with the complaint of continuous chest pain for 2 months. She had a history of right pulmonary tuberculosis. There was no known history of smoking or occupational exposure to asbestos or other chemicals. There were no signs associated with myasthenia gravis.

Chest X-ray showed homogenous opacity on the right side, obscuring the upper and middle zones of the right lung. Chest CT (Figure 1) showed a pulmonary mass of medium density was centered on the small scissure of the upper lobe and right middle lobe, with regular contours, 46 mm in diameter. There was no pleural effusion, pulmonary hilar or mediastinal lymph node swelling, or mediastinal tumor, particularly in the anterosuperior regions. PET scan (Figure 1): Presence of hypermetabolic sides making suspect malignancy. Diagnosis of NSCLC to exclude.

The patient underwent thoracotomy.

Macroscopically

The tumor, which was housed in the parietal pleura mainly at the fourth and fifth costal levels, although encapsulated, measured 5 × 3.5 cm. A cut, there is a whitish homogeneous appearance, with mild hemorrhagic and necrotic changes were observed on the surface of the tumor.

Microscopically

Well encapsulated tumor proliferation arranged in layers diffuse, made of spindle cells with oval, mildly pleomorphic nuclei intimately admixed with a mild infiltrate of lymphoid cells scattered throughout the tumor. Perivascular spaces characteristic of thymoma were found, although they were scarce. A few scattered mitoses of the epithelial cells were observed (Figures 2-6).

Immunohistochemical analyses showed that almost all of the epithelial cells stained positive for CK AE1/AE3 (Figure 5), and vimentine (Figure 3), and were negative for Calretinin, WT1 (Figure 6) (eliminating mesothelioma), CD34 (eliminating solitary fibrous tumor) and TTF1 (eliminating NSCLC). The lymphoid cells were positive for CD20 (Figure 4) and were negative for CD3, TDT, CD1a, and CD99.

These histopathological observations were almost completely compatible with those of thymoma rather than mesothelioma, and thus the tumor was diagnosed as being a medullary thymoma type A CK+, CD20+ (Figures 7 and 8).

The postoperative course was uneventful. The patient is free from the disease for the 10 months after surgery. Since no lesions in the mediastinum or other sites in the pleura were seen clinically in the CT scan, the tumor was conclusively judged to be ectopic primary pleural thymoma.

Discussion

Primary pleural thymomas are rare tumors thought to originate from ectopic thymic rests. There has been a case report on primary pleural thymoma in which ectopic rests of involuted thymic tissue were noted within the parietal pleural tissue [10]. Giving credence to this theory, several cases of pleural thymoma have also been reported so far [5-8] but some have shown more advanced tumors with mediastinal involvement, [5-6,7,15]. It remains questionable as to whether or not the origin of such tumors was actually the pleura itself.

Thymoma, particularly invasive thymoma spreads frequently to the intrathoracic pleura, as the disseminated lesions are generally regarded as metastases [5-7,15]. In contrast, our case showed that the lesion is developed in the thoracic pleura. Since no other tumor was evident either at the mediastinum or at the surrounding regions.

Keywords:

Tymoma; Pleura; Pleural thymoma

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Figure 1: Chest CT showed a pulmonary mass centered on the small scissure of the upper lobe and right middle lobe, of medium density, with regular contours, 46 mm in diameter. PET scan: Presence of hypermetabolic sides making suspect malignancy.

Figure 2: HES x 5. Tumor proliferation arranged in layers diffuse made of spindle cells with oval, mildly pleomorphic nuclei intimately admixed with a mild infiltrate of lymphoid cells scattered throughout the tumor. Perivascular spaces characteristic of thymoma were found, although they were scarce. A few scattered mitoses of the epithelial cells were observed.

Figure 3: HES x 10. Tumor proliferation arranged in layers diffuse made of spindle cells with oval, mildly pleomorphic nuclei intimately admixed with a mild infiltrate of lymphoid cells scattered throughout the tumor. Perivascular spaces characteristic of thymoma were found, although they were scarce. A few scattered mitoses of the epithelial cells were observed.

Figure 4: HES x 20. Tumor proliferation arranged in layers diffuse made of spindle cells with oval, mildly pleomorphic nuclei intimately admixed with a mild infiltrate of lymphoid cells scattered throughout the tumor. Perivascular spaces characteristic of thymoma were found, although they were scarce. A few scattered mitoses of the epithelial cells were observed.

Figure 5: HES x 40. Tumor proliferation arranged in layers diffuse made of spindle cells with oval, mildly pleomorphic nuclei intimately admixed with a mild infiltrate of lymphoid cells scattered throughout the tumor. Perivascular spaces characteristic of thymoma were found, although they were scarce. A few scattered mitoses of the epithelial cells were observed.

Pleural thymomas can be presented as multiple pleural nodules or as a diffusion of thickening of the pleura encasing the lung, mimicking a mesothelioma. All the cases reported so far presented either a multiple pleural nodules or localized pleural masses with diffusion of pleural thickening encasing the lung as in our case.

Most of the patients had dyspnea, chest pain, weight loss, and pleural effusion. Primary pleural thymoma has been reported both in men and in women, aged 19 to 75 [9], with a slight male predominance.

WHO reports A, B1, and B2 Type with the characteristic histological features such as lobulations, minimal nuclear pleomorphism, staghorn vasculature, and perivascular spaces with lymphocytes for the biopsy samples. The closest differential diagnosis is malignant mesothelioma [9,12].
Immunohistochemical markers are useful aid in differentiating malignant mesothelioma. However, it should be interpreted with caution.

CK, clearly mark thymomas but do not allow us to reliably differentiate cortical and medullary epithelial cells. The cortical cells are immature, they are CD1 A+ TDT+, CD99+ CD3+, CD4+, CD8+. Medullary cells are mature CD1A, TDT negative, CD3+, CD4+, CD8+. The CD 20 has been described positive in the medullary forms (A).

In the present case, the epithelial cells were negative for calretinin (Figure 9) but were found stains of stromale, nerve and mast cells. Other mesothelial markers were negative (WT1). The diagnosis of solitary fibrous tumor was also suspected of CD34 and immunostaining was negative, eliminating this hypothesis. The lymphoid cells were positive for CD20, consistent with thymic origin. The tumor was assigned to the WHO medullary thymoma type A based on the morphology.

Moran et al. [4] describes Intrapulmonary thymoma as slow-growing neoplasm with a good prognosis when surgically resected. However, in cases with cytologic atypia or an invasive growth pattern, adjuvant therapy should be considered to prevent the possibility of recurrence or metastases.

Moran et al. [8] described only two cases of ectopic pleural thymoma treated surgically and there was no recurrence in these cases during the follow-up periods. However, several authors [5-8] described the poor outcome of the cases with local advancement. Therefore, a complete surgical resection for the lesion appears to be the treatment of choice. The study additionally proposes postsurgical radiation for local control is desirable. In the present case, even though adjuvant irradiation was not performed due to localized lesion, though a careful follow-up.

**Conclusion**

Primary pleural thymomas are rare tumors often mistaken as malignant mesothelioma. Thoracoscopy with biopsy is the best diagnosing means. Their classification and treatment is similar to conventional thymoma.

**References**

thyroid thymoma: a distinct clinicopathologic entity. Hum Pathol 19: 1463-1467.


