Recurrent thymoma in the retroperitoneal space: a rare case report

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Abstract

Thymoma is an epithelial neoplasm of the thymus, which commonly lies in the anterior mediastinum and recurrences of thymoma generally are locally, and retroperitoneal recurrence is considered to be rare. A 46-year-old Asian female was referred to our hospital after finding left retroperitoneal mass during a routine health examination in July 2012. Abdominal ultrasound examination showed left retroperitoneal mass and ultrasound doctors recommended further examination. Fluorodeoxyglucose positron emission tomography/computed tomography (CT): left retroperitoneal tumors with increased metabolism, the mean standardized uptake values was 6.3, and made diagnosis of neurogenic tumors. But no investigations were done at that time. One year later, she was admitted for further examination and treatment at our hospital under a diagnosis of a left retroperitoneal tumor. CA-125 60.12kU/L (0-35), CA-724 12.61kU/L (0-6.9) and CYFRA21-1 4.7 ng/mL (<3.3) mildly elevated. Other laboratory tests were normal. No abnormalities were found on physical examination. Her medical history consisted of a primarily resected invasive thymoma (Masaoka stage III; WHO B1) 10 years ago at the local hospital. Post surgery completely resection, she received once adjuvant chemotherapy and radiotherapy (drug and radiation dose were not clear). The patient without preoperative myasthenia gravis (MG), but after thymectomy the patient developed MG from 2002-2008 year. She was symptom free on pyridostigmine bromide tablets and prednisone therapy on last follow up. From 2008 to 2012 year, she had not any clinical symptoms.

CT demonstrated a well-circumscribed, homogenous solid mass (10.1×11.3 cm) without calcification and fat in the left retroperitoneal space, the CT value is about 32–37 Hounsfield units (HU). Contrast enhanced CT abdomen-thorax showed a heterogeneously medium enhancing, CT value is about 56–77 HU (Figure 1). The mass compressed adjacent organs such as stomach, adrenal and kidney, leading to ascend left diaphragm (Figure 2). There were not masses in the anterior mediastinum.

After multidisciplinary consultation, the patient agreed to accept surgery. According to the operative report, surgical resection was performed with the retroperitoneal approach, which revealed the tumor adhering neighboring diaphragm. The tumor was histologically diagnosed to be type B1 thymoma according to the World Health Organization classification. The retroperitoneal mass was an unusual local recurrence after thymectomy. The patients whose had under invasive thymectomy should be evaluated carefully when finding retroperitoneal mass during follow-up.

Introduction

Thymomas are one of the most common neoplasms arising in the anterior mediastinum, but it accounts for <1% of all adult malignancies.1 All thymomas should be resected due to their malignant potential, providing patients are otherwise healthy. But some patients postoperative have recurrence. Local recurrence is more common. It usually presents itself as a pleural or pericardial nodule. Distant metastasis is usually found in lung, liver and bone.2 In this paper, we present a patient with a subphrenic retroperitoneal mass. Finally, this appeared to be a recurrence of the primary thymoma. This location of a distant recurrence is unusual.

Case Report

A 46-year-old Asian female was referred to our hospital after finding left retroperitoneal mass during a routine health examination in July 2012. Abdominal ultrasound examination showed left retroperitoneal mass and ultrasound doctors recommended further examination. Fluorodeoxyglucose positron emission tomography/computed tomography (CT): left retroperitoneal tumors with increased metabolism, the mean standardized uptake values was 6.3, and made diagnosis of neurogenic tumors. But no investigations were done at that time. One year later, she was admitted for further examination and treatment at our hospital under a diagnosis of a left retroperitoneal tumor. CA-125 60.12kU/L (0-35), CA-724 12.61kU/L (0-6.9) and CYFRA21-1 4.7 ng/mL (<3.3) mildly elevated. Other laboratory tests were normal. No abnormalities were found on physical examination. Her medical history consisted of a primarily resected invasive thymoma (Masaoka stage III; WHO B1) 10 years ago at the local hospital. Post surgery completely resection, she received once adjuvant chemotherapy and radiotherapy (drug and radiation dose were not clear). The patient without preoperative myasthenia gravis (MG), but after thymectomy the patient developed MG from 2002-2008 year. She was symptom free on pyridostigmine bromide tablets and prednisone therapy on last follow up. From 2008 to 2012 year, she had not any clinical symptoms.

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Discussion

The thymus is a lymphoepithelial organ that is derived embryologically from the third and fourth pharyngeal pouches, which descend to the anterior mediastinum in the sixth week of human gestation.3 Thymomas are an uncommon heterogeneous group of anterior mediastinal tumors that are generally considered indolent. In this paper, we presented the case of a distant recurrence of an invasive thymoma after resection ten years later at an unusual location.

Thymomas occur in all ages, but there is a broad peak between 35 to 70 years of age. The gender distribution of thymoma is approxi-
mately equal, although it is slightly more common in women in older age groups. Patients with mediastinal thymomas are often clinically asymptomatic (50%-60%) or present as local symptoms (30% to 40%) or associated systemic parathyroid disease syndromes (30% to 50%). Local symptoms, vague chest pain, dyspnea, and cough are the common complaints of thymomas. MG is the most common systemic paraneoplastic syndromes. According to the latest classification of histologic criteria for thymic epithelial tumors by the World Health Organization (WHO) Consensus Committee, published in 2004, thymic epithelial tumors are classified into two major categories: five types of thymomas (types A, AB, B1, B2, and B3) and thymic carcinomas.

Recurrences of thymoma occur often locally. Lymphogenous and hematogenous metastases are uncommon. Previous literature reported very few cases of retroperitoneal invasive thymoma recurrence and lesions were transdiaphragmatic, but in our case the mass did not cross diaphragm, only invaded diaphragm. Three potential routes for the transdiaphragmatic spread of invasive thymoma have been described, namely: i) through the retrocrural space; ii) through the openings in the anteromedial diaphragmatic origin; and iii) direct invasion of muscular diaphragm and spread into the peritoneal/extraperitoneal space.

Conclusions

In the patient, the retroperitoneal mass was suspected not to be a recurrence of the prior thymoma because of its supposed localization and without a local recurrence in the anterior mediastinum post surgery ten years later. But, the surgery pathology showed B1 thymoma.

We concluded that the retroperitoneal mass was an unusual local recurrence after thymectomy. The patients whose had under invasive thymectomy should be evaluated carefully when finding retroperitoneal mass during follow-up.

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