Clinical analysis of 47 cases of solitary fibrous tumor

WEI GE, DE-CAI YU, GANG CHEN and YI-TAO DING

Department of General Surgery, Nanjing Drum Tower Hospital,
The Affiliated Hospital of Nanjing University Medical School, Nanjing, Jiangsu 210008, P.R. China

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Abstract. The aim of the present study was to summarize the clinical manifestations, diagnosis, treatment, and prognosis of solitary fibrous tumor (SFT). In total, 47 cases of SFTs diagnosed by postoperative pathology between January 2002 and September 2014 were retrospectively reviewed, and the general information, clinical manifestations, imaging techniques, treatment, pathology and follow-up findings were analyzed. Of the 47 patients, clinical characteristics were collected in 37 cases (18 men and 19 women; mean age, 44.1 years; age range, 13-72 years). The maximum diameters of the tumors were 1.5-25 cm, with a mean diameter of 8.8 cm. The symptoms were various and non-specific. Imaging examinations following iodinated contrast administration showed the SFTs to be well-defined, cystic or solid mass and enhanced. On color Doppler ultrasound, SFTs were described as hypoechoic, clear, irregular masses. All patients underwent surgical resection, and SFT was diagnosed by postoperative pathological and immunohistochemical examination. Of the 47 patients, 25 received complete follow-up of 5-130 months, with a median follow-up period of 35.2 months, that included a color Doppler ultrasound or computed tomography (CT) scan every 6-12 months. At the end of the follow-up period all patients were alive and healthy, with the exception of one patient, who presented with recurrence 15 months after surgery. The findings of the present study showed SFT to be a rare systemic disease with no particular clinical manifestations. In the cases reviewed in the present study, CT, magnetic resonance imaging scans and color Doppler ultrasound were important for the diagnosis of SFT, while the definitive diagnosis relied on pathological and immunohistochemical examinations. Surgery, the primary treatment for SFT, was performed, and, following complete removal of the tumor, the prognosis was favorable.

Introduction

Solitary fibrous tumor (SFT), also known as hemangiopericytoma, is a fibroblastic mesenchymal tumor that was first described by Klemperer and Rabin in 1992 (1). SFTs are rare entities accounting for <2% of all soft tissue sarcomas (2). Although SFTs are commonly found in the pleura, according to the literature, 50-70% of these tumors are extrapleural and can manifest in the head and neck, abdomen and pelvis (3). This condition is rare and only limited cases of SFT have been reported. Due to the rarity of this tumor, preoperative diagnosis is challenging and no consensus has been reached on standard treatment. The present study summarizes the diagnosis, treatment and prognosis of 47 cases of SFTs from the literature, in order to determine the clinicopathological profile of SFT. To the best of our knowledge, the present retrospective study on SFT is the largest reported to date.

Patients and methods

Patients. Patient files from the Department of Pathology of the Nanjing Drum Tower Hospital, The Affiliated Hospital of Nanjing University Medical School, (Nanjing, China) from the period between January 2002 and September 2014 were retrospectively reviewed, and 47 cases of SFT were identified, including 39 inpatients and 8 outpatients. The present study was approved by the Institutional Review Board of Nanjing Drum Tower Hospital. Informed consent for participation in the study was obtained from all patients. All patients had been diagnosed with SFT by a pathological examination. Patient information, including age and gender, details of the lesion, clinical symptoms at the time of diagnosis, auxiliary and pathological examination findings, treatment and follow-up, was recorded.

Auxiliary examination types and findings. The findings of different types of auxiliary examination, including computed tomography (CT), magnetic resonance imaging (MRI) scans and color Doppler ultrasound, were reviewed. A histopathological examination of paraffin-embedded surgical specimens was performed to confirm the diagnosis using standard hematoxylin and eosin staining and immunohistochemical techniques. Immunohistochemical staining was performed using a Dako EnVision System (Dako, Glostrup, Denmark), according to the manufacturer’s instructions. The primary antibodies included rabbit polyclonal antibodies against vimentin...
Results

Baseline characteristics. Out of the 47 screened patients, full clinical characteristics were collected from 37 (18 men and 19 women; age range, 13–72 years; mean age, 44.1 years) and are summarized in Table I. The maximum diameter of the tumors was 1.5–25 cm, with a mean diameter of 8.8 cm. The patients’ symptoms were various and non-specific, and included pain, cough, inhibited defecation and frequent micturition. In total, 23 of the 37 patients (62.2%) had no symptoms at all, and the tumor was discovered incidentally during physical examination. Common lesion locations included the lungs, pleura, chest wall and pelvic cavity, while certain rare sites, including the thyroid, groin, eyebrow bow, bladder, kidney, eye, mediastinum and sigmoid mesocolon, were discovered.

CT scan was the most frequent diagnostic imaging technique used. CT images captured following iodinated contrast administration from 30 of the 37 cases showed SFTs as well-defined, cystic or solid mass, and enhanced (Fig. 1). In total, 11 out of 37 patients underwent an MRI scan, in which SFTs were shown as lobulated, heterogeneous soft tissue with short T1-weighted (T1WI) and a flake long T2-weighted imaging (T2WI) signal. Contrast-enhanced MRI scans demonstrated heterogeneous contrast enhancement (Fig. 2). Color Doppler ultrasound was performed in 14 out of 37 patients and found the tumors to be hypoechoic, clear, irregular masses. All patients underwent successful surgical resection with no serious complications, and no postoperative mortality was recorded. No patient underwent other adjuvant therapy.

Pathological findings. All patients were definitively diagnosed with SFT by postoperative pathological and immunohistochemical examination. Macroscopically, most tumors appeared as solid, well-encapsulated, smooth to firm soft tissue masses, with a gray-white to red-brown color on the cut surface. Microscopically, the tumors were shown to be comprised of spindle or short spindle cells and various quantities of vascular tissue. In certain regions, the cells were arranged in short, ill-defined fascicles, whereas in others they were arranged irregularly.

Immunohistochemical findings. Immunohistochemistry showed the following: The positive rates of vimentin, CD34, CD99, bcl-2, actin, desmin, CK, S100, CD117, EMA and SMA were 13/13 (100.0%), 40/46 (87.0%), 25/31 (80.6%), 30/38 (79.0%), 6/21 (28.6%), 2/35 (5.7%), 0/18 (0%), 6/40 (15%), 2/18 (11.1%), 3/13 (23.1%) and 7/15 (46.7%), respectively. The immunohistochemical indexes were selected according to the location of primary lesion, as different tissues have different specific markers. The representative pathological and immunohistochemical images of a patient that presented with SFT in the diaphragm are shown in Fig. 3. The immunohistochemical results for this patient were as follows: CD34+, CD99+, neurofilament, bcl-2+, desmin+, actin, S100+, Ki67+ 1% and β-catenin*.

Follow-up information. A total of 25 patients received complete follow-up lasting 5–130 months (median follow-up period, 35.2 months), during which they underwent a color Doppler ultrasound or CT scan every 6–12 months. At the time of writing, all patients were alive and healthy. Only 1 patient (4%) presented with recurrent SFT 15 months after the first surgery, and had a disease-free survival following the second operation. The status of each patient at the last follow-up is summarized in Table I. The follow-up data from 22 patients of the total 47 (12 of the 37) were either lost or could not be obtained.
SFT was first described as a rare spindle cell tumor that arises from the visceral pleura (4); however, over time, SFT was identified in various other locations outside of the thoracic cavity, such as the meninges (5,6), orbit (7), nasal cavity (8),
Figure 2. A 48-year-old female presented with a headache that had lasted for 1 month. Magnetic resonance imaging scan showing a well-defined and round soft tissue mass measuring 5x5 cm. (A) T1-weighted image; (B), T2-weighted image; (C) arterial phase; (D) sagittal view.

Figure 3. Microscopic examination of a solitary fibrous tumor located in the diaphragm of a patient, showing proliferating spindle cells with a bundle or flow pattern, determined by hematoxylin and eosin staining. Clusters of spindle cells were readily identifiable, and no nuclear atypia, mitotic activity or necrosis was evident; magnification, (A) x100 and (B) x400. Immunohistochemical examination showing that the tissue was (C) β-catenin and (D) B-cell lymphoma-2 positive.
salivary gland (9), parapharyngeal space (10) and paranasal sinuses (11). The present study reviewed numerous cases of SFT manifesting in locations outside the thoracic cavity, such as the thyroid, groin, bladder, sigmoid mesocolon and posterior superior iliac spine, locations that had not been previously reported. According to the literature (12), SFT is usually encountered in middle-aged people with an equal distribution between genders; however, it has also been reported in young patients (13). Consistent with these findings, the gender ratio in the present study was 0.95 (18 men and 19 women) and the mean age was 44.1 years. The symptoms varied depending on the lesions; the majority of patients (23/37) presented with physical symptoms. The diameter of SFTs is usually large (commonly >8 cm) (14). In the present study, 50% patients had a tumor measuring >8 cm.

SFTs do not express any tumor markers, and diagnostic techniques include CT, MRI scans and color Doppler ultrasound, despite the fact that their results may not be specific (9). These imaging tests often provide the first clue to the identification of the tumors, depiction of local extent and invasion of adjacent structures, which is useful in guiding surgery. On CT scan, SFT appears as a well-defined, heterogeneous or homogeneous isodense mass, and shows moderate to marked enhancement following contrast administration (15,16). On color Doppler ultrasound, it has the appearance of a hypoechogenic, clear, and irregular mass (17). On MRI, it has been shown to exhibit intermediate signal intensity on T1WI images, and enhancement on T2WI images, which was in accordance with the findings of the present study (18,19). None of these methods, however, could achieve accurate diagnosis; the final diagnosis has been reported to rely on surgical resection and pathological examination (20,21).

In the present study, all patients underwent surgical resection. Macroscopically, the majority of SFTs appear as rounded, encapsulated masses of homogenous density, with a yellow/brown-to-white whorled appearance of the cut surface (22). Microscopically, SFTs are shown to be comprised of spindle or short spindle cells and varying quantities of vascular tissue (9). The micro and macroscopic results of the present study were in accordance with the findings of previous studies. Immunohistochemically, SFT shows immunoreactivity with vimentin and CD34, with the largest part of tumor also displaying positive results for bcl-2 and CD99; however, the specimens do not express EMA or S-100 proteins (23). The immunohistochemical analysis results of the current study validated the previous findings.

The primary treatment of SFT is surgical resection with negative margins (24). All patients from the present study underwent complete resection without adjuvant treatments, such as radiotherapy or chemotherapy. Tumors that cannot be completely excised or that show malignant histological features may respond to radiation and/or chemotherapy. Xue et al (25) described a case of non-recurrent malignant SFT of the nasal and paranasal areas, and found that the combination of cytoreductive surgery with intensity-modulated radiation and stereotactic body radiation therapies have a good result. Studies have also demonstrated that preoperative embolization may be employed prior to surgical resection for highly vascular tumors (26,27). Immunotherapy, for example with interferon, may also be effective (28).

A previous study showed that the prognosis of patients with SFT is favorable (29). Patients that underwent complete tumor resection showed 100% survival at a mean follow-up period of 1.9 years (29). In the present study, all 25 patients that received a complete follow-up were alive at the time of writing. The longest follow-up period among these patients was 130 months. The recurrence rate of SFT was 4%.

In conclusion, SFT is a rare systemic disease with no particular clinical manifestations. In the cases reviewed in the present study, CT and MRI scans and color Doppler ultrasound were important for the diagnosis of SFT; however, the final diagnosis relied on pathological and immunohistochemical examinations. Surgery was the primary treatment for SFT, and the prognosis following complete tumor resection was favorable.

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