Inflammatory Myofibroblastic Tumor 12 Years After Treatment for Synovial Sarcoma: A Case Report

Aadit Shah, Eduard Pey, Justice U Achonu, Ji Dong K Bai, Fazel Khan

1Department of Orthopaedics, Stony Brook University Hospital, Stony Brook, NY, USA; 2Stony Brook University Renaissance School of Medicine, Stony Brook, NY, USA; 3Department of Pathology, Stony Brook University Hospital, Stony Brook, NY, USA

Abstract: Inflammatory myofibroblastic tumors (IMTs) are mesenchymal neoplasms most seen in the abdominopelvic region, lung, and retroperitoneum; and less commonly seen in virtually any other site. We report a case of two lower limb masses consistent with diagnosis of IMTs. This is a 39-year-old woman with a history of right lower extremity popliteal fossa synovial sarcoma diagnosed 12 years prior and treated with chemotherapy, surgery, and radiation. She presented with two new – one anterior and one posterior – right thigh masses. Biopsies of the lesions demonstrated low-grade inflammatory spindle cell lesions at both sites. Wide resection was performed for both masses and further characterization of the surgical specimens was most consistent with IMT. At follow-up, the patient is well with no signs of recurrence 19 and 7 months postoperative to the resection of the anterior and posterior thigh masses, respectively. This case represents the first reported IMTs occurring as late as 12 years after primary cancer treatment, and the first occurring after synovial sarcoma.

Keywords: inflammatory myofibroblastic tumor, radiation, resection, synovial sarcoma

Introduction

Inflammatory myofibroblast tumor (IMT) is a neoplasm of mesenchymal origin, most typically affecting children and young adults, with a predilection for visceral soft tissues.1,2 The term IMT emerged as a discrete entity from the benign morphological group termed “inflammatory pseudotumors (IPT)” roughly two decades ago. Its evolving, and at times, inconsistently applied nosology makes it difficult or impossible to tabulate the total number of reported cases. However, data from more recent case reports and two notable case series with 38 and 84 cases, respectively, underlie the present academic understanding.3,4 IMT is most commonly seen in the lung, abdominopelvic region, and retroperitoneum, but may also be seen in virtually any site.2 Clinical presentation is typically due to the mass itself, or nonspecific symptoms resulting from the tumor’s mass effect such as abdominal pain, gastrointestinal complaints, cough, or chest pain.4–6 A constitutional syndrome has also been described in 15–30% of the patients, consisting of fever, weight loss, and malaise, with laboratory evaluation revealing microcytic anemia, increased erythrocyte sedimentation rate, thrombocytosis, and/or polyclonal hypergammaglobulinemia.3,4 Prognosis is influenced by a site-dependent 2–25% recurrence rate.4,7,8 Additionally, rare distant metastasis is documented in 21 cases, most commonly to lung and brain, followed by liver and bone.2,3,9–20

Correspondence: Aadit Shah
Stony Brook University Hospital,
Department of Orthopaedics, 101 Nicolls Road, HSC T-18 – 089, Stony Brook, NY, 11794, USA
Tel +1-631-786-9484
Email aadit.shah@stonybrookmedicine.edu
Histologic characterization of IMT is classically characterized by spindle cell proliferation in a myxoid to collagenous stroma, with a prominent inflammatory infiltrate of mostly plasma cells and lymphocytes. Three basic histological patterns have been defined and are often seen in combination within the same tumor: a myxoid/vascular pattern, a compact spindle cell pattern, and a hypocellular fibrous pattern. Rearrangements of the ALK (anaplastic lymphoma kinase) gene involving constitutive expression of its product have been documented, with ALK rearrangements and expression shown in approximately 50% of IMTs by both fluorescence in situ hybridization (FISH) and immunohistochemistry (IHC), respectively. 21 ROS1 gene rearrangements have been identified most often in IMTs affecting children, many of these IMTs also demonstrate kinase fusions, which are often detected with immunohistochemistry or FISH studies. There have also been cases of IMTs that lack FISH abnormalities, and potential FISH and IHC assay issues may arise yielding false negatives. Kinase fusion negative IMTs lack these FISH abnormalities, making it more difficult to detect gene rearrangements such as ROS1, RET, NTRK, and PDGFβR. 22 23 This case has been demonstrated previously in an 18-year-old woman who had an ALK-negative lung IMT. 24 Another case study demonstrated detection of CDK4 and MDM2 double amplification in IMT tumor immunohistochemistry. 25 Histological diagnosis of IMT carries a differential diagnosis including spindle cell sarcoma, spindle cell melanoma, sarcomatous carcinomas, dedifferentiated liposarcoma, gastrointestinal stromal tumors, dendritic cell neoplasms, desmoid fibromatosis, nodular fasciitis, and IPT. Of note, IMT is particularly a diagnosis of exclusion middle-aged or older adults, and when seen in skin or somatic soft tissue. 2

Our case, accordingly, is one of these hard-fought diagnoses of exclusion: IMT near the knee in a 39-year-old woman, interestingly occurring 12 years after chemotherapy, surgical resection and radiation for a synovial sarcoma at the knee.

Case Presentation

Our patient is a 39-year-old woman with a past medical history of right lower extremity popliteal fossa synovial sarcoma that was diagnosed in 2007 and treated with chemotherapy: 3 cycles pre- and 3 cycles post-surgery with Adriamycin, Ifosfamide, and Mesna (AIM); radiation; and surgery; as well as monoclonal gammopathy of undetermined significance (MGUS) followed with routine positron emission tomography/computed tomography (PET/CT) scans, Sjogren’s syndrome, and rheumatoid arthritis, who presented to the orthopaedic oncology clinic at our institution for evaluation of two newly discovered right thigh masses.

She was well-appearing and ambulated without any assistive devices. She had two palpable masses—one in the anterior thigh and the other in the posterior thigh. The skin overlying the anterior mass was intact, while the skin over the posterior one had a surgical scar with extensive fibrous and scar tissue likely from her previous radiation. She had a reduced range of motion of the knee, from 4 to 100 degrees, and no palpable lymphadenopathy. The remainder of the examination was normal.

Her diagnostic assessment revealed anterior and posterior subcutaneous soft tissue masses of the thigh. PET/CT studies revealed both tumors with standard uptake values (Figure 1). Several small, mildly hypermetabolic lymph nodes within the right external iliac and inguinal region were also identified, with the largest measuring 9 mm consistent with reactive lymph nodes.

Ultrasound-guided core biopsies were taken of the anterior and posterior thigh masses by interventional radiology. Both resulted in a similar histological picture—spindle cell, myxoid lesion with a mixed inflammatory infiltrate. The cellular lesions consisted of low-grade myofibroblastic cells with associated myxoid matrix and a mixed inflammatory infiltrate, with neutrophils, plasma cells, and occasional eosinophils. There also was some free hemorrhage. The spindle cells stained positively for smooth muscle actin and negatively for ALK1, S100, CD99, cytokeratin 7, BCL-2, EMA, CD56, CD34. The plasma cells had no significant expression of IgG4. FISH analysis for SS18-SSX, a gene rearrangement for synovial sarcoma, was negative. Essentially, no definitive diagnosis was made.

Since the imaging-guided biopsies did not result in definitive diagnosis, we decided to perform an open biopsy with excision of the masses. Because of the presence of extensive dense and fibrous tissue posteriorly from previous surgery as well radiation, it was decided to operate in a staged fashion starting with the anterior mass. The anterior mass was excised and sent to pathology for evaluation. On initial evaluation, pathologists identified the features of the mass most closely resembling IMT and nodular fasciitis. FISH analysis later performed was negative for USP6 (17p13), a gene rearrangement found in 85–90% of the nodular fasciitis cases. The open biopsy and excision of the anterior mass was complicated by delayed wound healing, and
resection of the second, posterior mass was postponed until complete healing of this anterior wound.

During this period, a surveillance MRI performed showed an interval enlargement of the posterior thigh mass (Figure 2). There was increased infiltration of the subcutaneous tissue surrounding the lesion. Following this MRI and once the incision from the anterior mass had healed, it was decided to proceed with wide resection of the posterior thigh mass (Figure 3). The definitive wound closure procedure for this resection was performed by plastic surgery. Pathologic evaluation of the posterior mass like the anterior mass resulted in IMT and nodular fasciitis as being the top of the differential with FISH analysis showing negative USP6 rearrangement. In a multicenter discussion about the pathology, the highest diagnosis on the differential was IMT.

The patient has been seen up to 7 months post-op after resection of the posterior thigh mass (19 months after resection of the anterior mass). She has been doing well without any signs of recurrence of the disease.
After resection, chemotherapy, and radiation of a soft tissue sarcoma, 80% of the recurrences are detected within the first 2 years post-resection. Later recurrence may still occur, but this is rare. In this patient, 12 years had passed since diagnosis and treatment of her synovial sarcoma. Local recurrence rates following wide resection are 8–30% (60–90% for simple excision). Postoperative surveillance recommendations include periodic physical examination, imaging of the primary site, and chest imaging to rule out metastatic disease. Routine evaluation of this patient did not reveal any recurrence or metastasis. She was also followed with PET/CT scans for her history of MGUS, which did not reveal any disease. Therefore, when she was evaluated 12 years after her synovial sarcoma resection, there was a low probability that the lesions were recurrence/metastasis. Synovial sarcoma recurrence/metastasis was thus excluded as a likely diagnosis, even considering its location right at the site of prior radiation. After initial biopsy of the two masses, negativity FISH for SS18-SSX of biopsy specimens further supported the diagnostic exclusion of recurrent synovial sarcoma, as up to 97% of the synovial sarcomas demonstrate this gene fusion.

Histopathologic interpretation of the anterior thigh mass was characterized by its storiform cellular myofibroblastic pattern (Figure 4), with mixed inflammatory infiltrate of plasma cells, macrophages, neutrophils and some multinucleated giant cells. Collections of lymphocytes were also noted in the periphery of the lesion. Consistent with nodular fasciitis’ higher incidence, as well as IMT being a diagnosis of exclusion, initial interpretation of this lesion was as a nodular fasciitis. However, once FISH came back negative for USP6 this changed the scenario as this test has a positive predictive value of 100% and a negative predictive value of...
90%, with 10–15% of the cases of nodular fasciitis being negative for the gene rearrangement. IgG4-related disease was also suspected due to her history of MGUS, but on immunohistochemistry, the plasma cells that were present did not significantly express IgG4.

The posterior mass was histologically similar to the former, and with similar FISH negatives for SS18/SSX and USP6, was evaluated as most consistent with inflammatory myofibroblastic tumor. The lesion tested negative for ALK1, a known marker for IMT. Only 50% of IMTs are said to carry this marker, and moreover, most of those are in younger patients. It is important to note, IMTs can be negative for a kinase fusion on immunohistochemistry/FISH and still be the diagnosis. Our patient being 39 years old places her at the upper end of the bell curve; therefore, the probability of identifying ALK1 in an IMT lesion is likely <50%.

It has been demonstrated that IMT can result from surgical treatment, post-chemotherapy, and post-radiation treatment. A multicenter study has reported on IMTs that have developed locally, and multifocally as metastatic disease; IMT has developed as the second tumor in two prior cases as reported on by the European pediatric Soft Tissue Sarcoma Study Group (EpSSG) in a multicenter study. In both cases, however, the primary sarcomas were treated with chemotherapy and radiotherapy, and the IMT developed outside of the treatment area within 1 and 5 years. The tumor has also been shown to develop as much as 12–19 years later. This is a unique case as the IMT developed locally within the radiation therapy field 12 years post-surgery, post-chemotherapy, and post-radiation. This tumor has been demonstrated to have both benign and aggressive features. It is described as a tumor with indeterminate biological potential: the recurrence rate is 25% and some can become malignant. In the case of this IMT, its features were benign and described as low-grade.

IMT is a rare neoplasm overall, with rudimentary epidemiology supporting 150–200 new cases per year in the United States. Its post-surgical/traumatic and post-radiation development is rare; especially locally and multifocally. With evidence of these tumors expressing benign and potentially aggressive behavior, 5-year event-free survival (EFS) and overall survival (OS) have been reported to be 82.9% and 98.1%, respectively. IMT has also responded to systemic therapy including chemotherapy and radiation treatment in select cases. The rare ROS1-rearranged IMTs have been found to respond to crizotinib, an antibody that has demonstrated significant reduction in tumor size. (Mai) Trabectedin, a marine-derived anticancer agent, is a microenvironment-targeting drug that is currently being investigated. It exerts a cytotoxic effect against the promoters of tumor-related inflammation. A similar case report involving an IMT in the posterior thigh found no recurrence at 2 years, but reported the presence of a potential red herring: a postoperative cyst. Regardless, early diagnosis and prompt surgical treatment is warranted, and should be pursued when possible.

Conclusion
As introduced earlier, IMT is a difficult diagnosis with a history of mischaracterization and misrepresentation in the literature and is a diagnosis of exclusion—particularly in a patient such as ours where a mass is seen in skin or soft tissue. The lack of positive and supportive molecular stains makes this patient’s lesions difficult to classify; however, its closest differential diagnosis given this histological presentation, nodular fasciitis, has been quite strongly ruled out with the absence of the USP6 gene rearrangement, and histological appearance is consistent with IMT. If both lesions are IMT, this would be the first published case to have occurred as far out as 12 years after primary cancer treatment with chemotherapy and radiation, and the first to have occurred secondary to synovial sarcoma. We endorse the necessity of early diagnosis and prompt surgical treatment of tumors presenting as in our patient.

Consent for Publication
The patient was aware of their participation in the case report and informed consent was obtained from the patient for publication. Institutional approval was not required to publish the case details.

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Disclosure
FK reports patent pending for computer navigation type use in bone sarcomas, outside of this work. The authors declare no other potential conflicts of interest.

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