Intraarticular fibroma of tendon sheath

Michael J Griesser, Paul E Wakely, Joel Mayerson

**ABSTRACT**

A 17-year-old male presented to us following a hyperflexion injury to his right knee sustained while playing soccer. Immediately after the traumatic event, he developed a large, tense knee effusion. Physical examination revealed limited range of motion. MRI revealed a lobulated mass in the posteromedial aspect of the knee joint. The mass was excised and sections submitted to pathology. A pathologic, microscopic, and immunohistochemical characteristics revealed the final diagnosis of fibroma of tendon sheath in the knee. At 12 months followup, the patient reported no subjective symptoms, such as pain or limitation of athletic activities and has full range of motion. Additionally, he has demonstrated no signs of recurrence. We report a case of fibroma of the tendon sheath originating from the synovial membrane of the joint capsule of the knee.

**Key words:** Femoral condyle, intraarticular fibroma, tendon sheath, fibroma

**INTRODUCTION**

Fibroma of tendon sheath (FTS), first described by Geschickter and Copeland in 1949, is a slow-growing fibrous nodule that frequently adjoins a tendon sheath, and has a predilection for occurring on the fingers and hands. Often affecting middle-aged patients, the onset is usually marked by noticing a small mass or swelling. FTS rarely originates from the synovial membrane of a joint capsule, similar to nodular fasciitis.

To our knowledge, only 11 cases of FTS originating from the synovial membrane of a joint capsule have been reported in the English literature, including 6 from the knee, 1 from the radioulnar joint, 1 from the scapholunate joint, 1 in the shoulder joint, 1 in the ankle joint, and 1 in the temporomandibular joint. We report here such a case of fibroma of the tendon sheath originating from the synovial membrane of the joint capsule of the knee, and only the second case that demonstrated erosion into bone.

**CASE REPORT**

A 17-year-old male presented to us following a hyperflexion injury to his right knee sustained while playing soccer. Immediately after the trauma he developed a large, tense knee effusion and was seen at another institution. The knee was aspirated, and a significant amount of blood was removed. Based on the mechanism of injury as well as the bloody effusion, ligamentous tears were suspected, and a magnetic resonance imaging (MRI) was done. He was referred to our institution 2–3 weeks later. The patient reported having significant pain and limitations in his right knee associated with squatting motion or hyperflexion activity. Examination revealed limited range of motion of the knee joint from 0° to 110° of flexion of the right knee (contralateral knee = 0°–140°). There was mild tenderness to palpation posterior to the right knee, with no evidence of effusion or a palpable mass after 3–4 weeks postinjury. The MRI revealed a lobulated mass in the posteromedial aspect of the knee joint, measuring 3.3 × 1.7 × 1.9 cm at its greatest dimension. The lesion was adjacent to the posterior cruciate ligament (PCL) and appeared densely adhered to both the PCL and the posterior joint capsule. The posterior medial femoral condyle had cortical involvement measuring 1.1 × 0.9 cm and was associated with a moderate knee joint effusion and associated prominent synovial enhancement and thickening. There was some T1 hyperintense signal with prominent enhancement of the more superior portion of the lesion [Figure 1]. The lesion was isointense on T1-weighted imaging and heterogeneously hyperintense on T2-weighted imaging.
At this point, the preoperative differential diagnosis included a nodular form of pigmented villonodular synovitis (PVNS), ganglion, and synovial chondromatosis. The patient subsequently underwent uneventful excision of the mass using a posterior approach to the knee [Figure 2]. Intraoperatively, the entire lesion was excised and was 3.5 × 2 cm in its greatest dimension. The mass was totally intrarticular and was densely adhered to the synovial membrane of the posterior portion of the joint capsule, the PCL, and the posterior medial femoral condyle. The lesion had some bony penetration as well. As the lesion was very difficult to visualize in its entirety, it was removed sharply from the bone so as not to prohibit a return for greater margins if the frozen section revealed a need to do so. The frozen section was diagnosed as benign intraarticular nodular fasciitis. There were no postoperative difficulties or need for marginal revision.

The pathologic specimen grossly consisted of a 3.3 × 1.6 × 1.6 cm tan–white, rubbery mass that weighed 4.5 g. The cut surface was tan–white and rubbery with approximately 15% comprising foci of hemorrhage. The foci of hemorrhage were sampled and showed focal recent hemorrhage and cystic degeneration. Microscopically, a proliferation of uniformly bland, but plump, fibroblastic/myofibroblastic cells were arranged in short intersecting bundles. Markedly elongated nuclei had smooth contours and an open chromatin pattern with occasional distinct nucleoli. The stroma was variably collagenous, harboring microscopic foci of myxoid change and an absence of necrosis, giant cells, hemosiderin deposits, foamy histiocytes, and extravasated red cells. Slit-like vascular channels were particularly noticeable at the periphery of the nodule on one side. Mitoses were rare. Immunohistology demonstrated diffuse positive staining with vimentin and smooth muscle actin with weaker staining for muscle-specific actin. Staining was negative with S-100, CD34, cytokeratin cocktail, and beta-catenin. These pathologic, microscopic, and immunohistochemical characteristics were suggestive of FTS in the knee [Figure 3].

At 12 months followup, the patient reported no subjective symptoms, such as pain or limitation of athletic activities. Extension of the knee joint was 0° and flexion was 140°, showing no limitation of range of motion. There was no swelling, tenderness to palpation, or signs of recurrence.

**DISCUSSION**

FTS is a slow-growing fibrous nodule typically attached to tendon sheath, of which approximately 80% occur in the fingers and hand. Rarely this lesion has been reported to arise from the synovial membrane of a joint capsule as in our case.
In 1979, Chung and Enzinger published a report on 138 cases of FTS, which to this day remains the largest series on FTS, and the basis of much of our understanding of these lesions. FTS typically develops in young males in the third to fourth decade, with male to female ratio of 3:1. However, most of these cases occur around the fingers, hand, and wrist, and very rarely are reported to be elsewhere anatomically. Additionally, only scattered case reports exist demonstrating FTS of the knee, particularly in an intraarticular location. In all, 26 cases of FTS around the knee have been reported. However, a detailed report regarding location is lacking in most of these cases. Additionally, only 6 reports have adequately described intraarticular locations for these lesions, with 2 occurring adjacent to the PCL, 1 in the suprapatellar pouch, and 3 in the posterior joint capsule. The lesion in our case was densely adhered to the posterior joint capsule, the PCL, and showed bony erosion into the medial femoral condyle. This case is the third originating from the PCL and the fourth emanating from the posterior joint capsule of the knee.

The typical MRI findings in FTS have been described in a case series by Fox et al. This included low intensity on T1-weighted images in 5 cases, low intensity and isointensity on T2-weighted images in 3 cases, and a slightly high intensity on T2-weighted images in 2 cases. In our patient, MRI revealed some T1 hyperintense signal in the superior portion of the lesion, with T1 isointensity elsewhere and heterogenous hyperintensity on T2-weighted imaging. However, the MRI findings of FTS are very similar to giant cell tumor of tendon sheath (GCTTS) and PVNS and cannot be relied upon for definitive diagnosis.

The only accurate way to diagnose intraarticular nodular FTS is by microscopic examination. The principal differential diagnoses include GCTTS, nodular fasciitis, and PVNS. This case was originally mistaken for nodular fasciitis. Several pathologic features typical of that entity were missing, including a lack of extravasated red cells, only minor foci of myxoid change, and an absence of a disordered “tissue culture” appearance to the proliferating spindle cells. Instead, the nodular pattern and the obvious presence of many slit-like small vascular channels is typical of FTS. Immunohistochemical staining does not distinguish between FTS and nodular fasciitis as both will demonstrate positive staining for smooth muscle actin, muscle-specific actin, and vimentin. GCTTS and PVNS are readily excluded pathologically, based on the absence of giant cells, xanthomatous histiocytes, and hemosiderin deposition. Finally, as more reports of FTS appear in the literature, it would seem prudent to add this condition to the list of differential diagnosis that have historically been considered for a painful, range of motion-limiting, intraarticular mass lesions. Bony penetration or cortical involvement can be seen in GCTTS, PVNS, synovial chondromatosis, calcific tendinitis, and perioseal chondroma, but is rarely reported in FTS or nodular fasciitis.

Entities such as FTS and GCTTS can be treated with marginal excision, alerting the patient and the family that recurrence is possible. Nodular fasciitis, however, can be diagnosed and treated by excisional biopsy, counseling the patient that there may be spontaneous regression of the lesion, even if it is incompletely excised. Symptomatic PVNS is commonly treated with either arthroscopic or open total synovectomy or possibly total joint replacement. Thus, differentiation helps determine treatment as well as how to counsel patients and their families appropriately.

In summary, this case illustrates the importance of an inclusive differential when examining intraarticular knee lesions, one which must include FTS.

**References**

1. Geschickter CF, Copeland MM. Tumors of Bone. 3rd ed. Philadelphia: J. B. Lippincott; 1949. p. 693-5.
2. Takakubo Y, Fukushima S, Asano T, Yamakawa M. Case Reports: intraarticular fibroma of the tendon sheath in the knee. Clin Orthop Relat Res 2005; 439:280-5.
3. Ogata K, Ushijima M. Tenosynovial fibroma arising from the posterior cruciate ligament. Clin Orthop Relat Res 1987; 215:153-5.
4. Pinar H, Ozkan M, Ozaksoy D, Pabuccuoglu U, Akseki D, Karaoglan O. Intraarticular fibroma of the tendon sheath of the knee. Arthroscopy 1995;11:608-11.
5. Hitotra T, Yamamoto T, Akisue T, Marui T, Nagira K, Ohota R, et al. Fibroma of tendon sheath originating from the knee joint capsule. Clin Imaging 2002; 26:280-3.
6. Hermann G, Hoch BL, Springfield D, Abdelwahab IF, Klein MJ. Intra-articular fibroma of tendon sheath of the shoulder joint: synovial fibroma. Skeletal Radiol 2006; 35:603-7.
7. Ahn JH, Lee YS, Lee DH, Ha HC. Intraarticular fibroma of the posterior compartment in the knee: A case report. Knee 2008; 15:155-8.
8. Moretti VM, de la Cruz M, Lackman RD, Fox EJ. Fibroma of tendon sheath in the knee A report of three cases and literature review. Knee 2010; 25:25.
9. Li TJ, Kitano M, Tsuneyoshi M, Sonoda S, Mimura T. Intra-articular fibroma of tendon sheath in the temporo mandibular joint. Oral Surg Oral Med Oral Pathol Oral Radiol Endod 1997; 84:407-10.
10. Misawa A, Okada K, Hirano Y, Sageshima M. Fibroma of tendon sheath arising from the radio-ulnar joint. Pathol Int 1999;49:1089-92.
11. Athwal GS, Bueno RA, Bansal M, Mintz DN, Athanasian EA. Intra-articular fibroma of tendon sheath involving the scapholunate and radiocarpal joints. Skeletal Radiol 2006; 35:599-602.
12. Ciatti R, Mariani PP. Fibroma of tendon sheath located within the ankle joint capsule. J Orthop Traumatol 2009; 10:147-50.
13. Bertolotto M, Rosenberg I, Parodi RC, Perrone R, Gentile S,
Griesser, et al. Case report: Fibroma of tendon sheath in the distal forearm with associated median nerve neuropathy: US, CT and MR appearances. Clin Radiol 1996;51:370-2.

14. Chung EB, Enzinger FM. Fibroma of tendon sheath. Cancer 1979;44:1945-54.

15. Hashimoto H, Tsumeyoshi M, Daimaru Y, Ushijima M, Enjoji M. Fibroma of tendon sheath: a tumor of myofibroblasts. A clinicopathologic study of 18 cases. Acta Pathol Jpn 1985;35:1099-107.

16. Smith PS, Pieterse AS, McClure J. Fibroma of tendon sheath. J Clin Pathol 1982;35:842-8.

17. McGrory JE, Rock MG. Fibroma of tendon sheath involving the patellar tendon. Am J Orthop 2000;29:465-7.

18. Hur J, Damron TA, Vermont Al, Mathur SC. Fibroma of tendon sheath of the infrapatellar fat pad. Skeletal Radiol 1999;28:407-10.

19. Okada J, Shinozaki T, Hirato J, Yanagawa T, Takagishi K. Fibroma of tendon sheath of the infrapatellar fat pad in the knee. Clin Imaging 2009;33:406-8.

20. Aynaci O, Kerimoglu S, Ozturk C, Saracoglu M, Yildiz K. Intraarticular fibroma of the tendon sheath arising from the infrapatellar fat pad in the knee joint. Arch Orthop Trauma Surg 2009;129:291-4.

21. Le Corroller T, Bouvier-Labit C, Sbihi A, Champsaur P. Mineralized fibroma of the tendon sheath presenting as a bursitis. Skeletal Radiol 2008;37:1141-5.

22. Fox MG, Kransdorf MJ, Bancroft LW, Peterson JJ, Flemming DJ. MR imaging of fibroma of the tendon sheath. AJR Am J Roentgenol 2003;180:1449-53.

23. Enzinger FM, Weiss SW. Soft Tissue Tumors. 5th ed. St. Louis: Mosby;2008. p. 247-307.

How to cite this article: Griesser MJ, Wakely PE, Mayerson J. Intraarticular fibroma of tendon sheath. Indian J Orthop 2011;45:276-9.

Source of Support: Nil, Conflict of Interest: None.

Announcement

Android App

A free application to browse and search the journal’s content is now available for Android based mobiles and devices. The application provides “Table of Contents” of the latest issues, which are stored on the device for future offline browsing. Internet connection is required to access the back issues and search facility. The application is compatible with all the versions of Android. The application can be downloaded from https://market.android.com/details?id=comm.app.medknow. For suggestions and comments do write back to us.