Abstract. Localized forms of giant cell tumors are defined as giant cell tumors of the tendon sheath (GCTTS). GCTTS arises from the synovium of a joint, bursa or tendon sheath, and 85% of the tumors occur in the fingers. GCTTS in the knee is extremely rare. We report an unusual case of a 15-year-old boy who presented with an occult growing swelling and a 2-month history of infra-patellar pain in the left knee. Magnetic resonance imaging (MRI) demonstrated a well-circumscribed soft tissue mass in the infra-patellar fat pad posterior to the patella tendon. Excision biopsy was performed by surgical removal. Histopathological examination revealed that it was GCTTS. During the follow-up period, his recovery was propitious and there was no recurrence. Owing to its few and non-specific symptoms, and local recurrence varying from 9 to 44%, its proper, early diagnosis and appropriate treatment is necessary.

Introduction

Giant cell tumor of the tendon sheath (GCTTS) is a benign lesion of uncertain etiology that involves inflammation, trauma, toxin, allergy, clonal chromosomal abnormalities and aneuploidy (1-2). Jaffe et al originally described the condition in 1941 (3). Approximately 85% of GCTTS occurs in the fingers, while 12% of tumors are located in the knee, elbow, hip and ankle (4). GCTTS may occur at any age but most typically occurs between the ages of 30 and 50, with a 2:1 female predominance (5). GCTTS in the knee is extremely rare, particularly in a 15-year-old boy. Most often, GCTTS is small (average tumor size, 2.0 cm) (5), although the mass in this case, measuring 3.8x2.0x1.7 cm was much larger than average. There are few reports in the literature of cases occurring in males, in younger age groups, tumor located in the knee and of greater tumor size. Antecedent trauma has been reported, but only 15% of cases have a definite history of trauma (6). Our patient reported a trauma while playing basketball at school 6 months previously. Treatments consist of careful local excision, particularly microscopic excision, and postoperative radiotherapy. GCTTS may recur following excision.

One of the most important clinical features is that the symptoms of GCTTS are non-specific. We report a valuable case that was not initially regarded as GCTTS. Following excision biopsy, the diagnosis of GCTTS was made. We believe that this study may assist clinicians in making a proper diagnosis and administering correct treatment.

Case report

The research findings are presented with the consent of the boy's parents and ethics committee approval. A 15-year-old boy presented with an occult growing swelling and a 2-month history of pain in the left knee at the Provincial Hospital Affiliated to Shandong University, China. Strenuous exercise aggravated the pain, which was felt at the infra-patellar region of the left knee. The skin temperature of the left knee was higher than the right one. The patient reported a trauma when playing basketball at school 6 months previously. Physical examination revealed local tenderness of the fibular and tibial collateral ligament. The range of motion was limited. Hyperextension and hyperflexion tests of the left knee were positive. There were no constitutional symptoms such as night sweats, fever or weight loss. The patient had previously been healthy, and no enlarged inguinal lymph nodes were detected. Analysis of the patient's blood was normal.

Plain radiography and magnetic resonance imaging (MRI) were obtained. Plain anteroposterior and lateral radiographs revealed no bone erosion of the left knee joint (Fig. 1). MRI demonstrated an encapsulated soft tissue lump in the infra-patellar fat pad, posterior to the patella tendon (Fig. 2). Initially, we regarded the neoplasm as an intra-articular lipoma, synovial sarcoma or fibroma of the tendon. Excision biopsy was performed by surgical removal, and an irregular shaped mass
measuring 3.8x2.0x1.7 cm was removed. Histopathological examination of the specimen revealed prominent mononuclear cells, accompanied by a variable number of foamy histiocytes, hemosiderin-laden macrophages and multinucleated giant cells (Fig. 3). No mitotic activity or malignant feature was reported. A diagnosis of GCTTS was finally made.

The recovery following surgery was propitious. There was no local tenderness of the left knee and the skin temperature of the knee was normal. The patient had full range of motion of the knee joint and returned to school life with no difficulty. During the 5 month follow-up period there was no evidence of local tumor recurrence.

**Discussion**

The diffuse and localized extra-articular forms of pigmented villonodular synovitis (PVNS) are presently referred to as diffuse-type giant cell tumor and giant cell tumor of the tendon sheath (GCTTS), respectively (7), and were originally alluded to by Jaffe et al in 1941 (3). Most localized giant cell tumors are small (average tumor size, 2.0 cm) (5), although lesions of greater size may be found in large joints such as the knee and elbow. Tumors are well-circumscribed and typically lobulated, and white to gray in color, with yellowish and brown areas. Most diffuse-type giant cell tumors are infiltrative and grow as diffuse, expansile sheets. These entities are now classified as ‘fibrohistiocytic tumors’ in the World Health Organization system of classification of bone and soft tissue tumors (4). GCTTS occurs in the digits of the hands and feet more frequently than in the large joints of the knee, elbow and ankle (8); 85% of GCTTS is observed in fingers, while 12% is observed in large joints (4). GCTTS in the knee is rare.

The etiology of GCTTS remains uncertain. A variable number of factors have been described as possible causes of GCTTS. These include inflammation, trauma, toxin, allergy, clonal chromosomal abnormalities and aneuploidy (1-2). GCTTS was initially regarded as an inflammatory disease; however, the finding of aneuploidy in certain cases and the demonstration of clonal chromosomal abnormalities strongly support a neoplastic origin (4).

The diagnosis of GCTTS may be difficult. Patients typically present with a painless mass in the digits or large joints. The lesions are usually well-circumscribed and localized, and infrequently erode or infiltrate the nearby bone (6). In large joints, the symptoms are non-specific and few, thus the diagnosis is difficult to make. The soft tissue mass grows and expands into areas of least resistance, such as the tendon sheath (8). Plain radiographs are usually not helpful in the diagnosis of the disease. Bone erosions or soft tissue swelling is occasionally found in plain radiographs.

Magnetic resonance imaging (MRI) is an effective and highly sensitive tool for diagnosis. Typically, MRI features T1- (T1WI) and T2-weighted (T2WI) images, which may present with a homogeneous low signal intensity in GCTTS. The reason for this is that tumors contain dense collagen and hemosiderin-laden macrophages (9).

Treatments consist of careful and complete local excision and postoperative radiotherapy. Surgical removal is the standard treatment of GCTTS. Adequate initial local excision may effectively limit the risk of local recurrence (10). Recently, microscopic excision has been prescribed to prevent recurrence. Ikeda et al (11) treated 18 patients with microscopic excision, and of the 18 patients which only one case recurred. The reason for this recurrence was that the microscope
excision was not used in this diffuse-type patient. As an adjuvant therapy, postoperative radiotherapy had been indicated to prevent recurrence following surgery. Kotwal et al (12) published a prospective study in which the recurrence rate was only 4%. This rate was much lower than previously reported incidences of between 25 and 45%.

Although GCTTS is a benign type of tumor, it has a high incidence of recurrence. Recurrence following resection has been a consistent problem, and occurs in 9 to 44% of cases (13). Lowyck and De Smet (13) retrospectively reviewed 49 patients and reported the rate of recurrence to be 16%.

In conclusion, the concurrence of GCTTS occurring in a younger male, occurring in the knee and being of greater than average size is unique to our patient, making this an exceptional case to those reported in previous literature. It is extremely difficult to make a proper diagnosis during the earlier stages of GCTTS. As the condition always presents with a painful lump, it should be considered as a differential diagnosis if the mass is found in the knee, particularly in younger males and when the tumor is of greater size. MRI may be useful for diagnosis when the symptoms of GCTTS are non-specific and few. To make a definite diagnosis, histopathological examination is required. If a possible diagnosis of GCTTS is suspected, excision biopsy should be carried out as early as possible in the same manner that the present case was treated. Finally, the purpose for which we report this case is to emphasize the possibility of GCTTS, even when the painless or painful mass accompanied with non-specific symptoms are found in the knee of a younger patient, and to avoid misdiagnosis wherever possible.

References