Tenosynovial giant cell tumour of the posterior cruciate ligament and its arthroscopic treatment

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ABSTRACT
Tenosynovial giant cell tumours originate from synovial tissues of the joints, tendon sheaths, mucosal bursas or fibrous tissues adjacent to tendons. Tenosynovial giant cell tumours are rarely intra-articular. We report a giant cell tumour of the tendon sheath arising from the posterior cruciate ligament diagnosed by magnetic resonance imaging and resected arthroscopically in a 54-year-old woman.

Keywords: arthroscopic resection, giant cell tumour of the tendon sheath, pigmented villonodular tenosynovitis, posterior cruciate ligament, tendon sheath lesion, tenosynovial giant cell tumour

INTRODUCTION
Tenosynovial giant cell tumours, also known as pigmented villonodular tenosynovitis, arise from the synovial tissue of the joint, tendon sheath, mucosal bursa or fibrous tissue adjacent to the tendon. These tumours predominantly involve the palmar side of fingers and toes, and seldom larger joints like the knees and ankles. Tenosynovial giant cell tumours of the tendon sheath are rarely intra-articular. In the literature, there has only been one reported case of tenosynovial giant cell tumour arising from the posterior cruciate ligament (PCL). In this report, we present a rare location of a tenosynovial giant cell tumour arising from the PCL and its treatment with arthroscopy.

CASE REPORT
A 54-year-old woman was referred to our clinic for right knee pain of two years. She did not feel any pain at rest but in walking and climbing up and down the stairs. She did not have knee trauma. She had active and full range of motion of the knee. On physical examination, there was no lateral and medial joint space tenderness. The knee was found to be stable in Lachman, anterior drawer, posterior drawer and medial-lateral stress tests.

No bone pathology was detected in anteroposterior and lateral knee radiographies. All laboratory tests were within normal ranges. On magnetic resonance (MR) imaging, a regular contoured mass localised in the PCL was detected (Figs. 1 & 2).

In the light of these findings, we performed an arthroscopy from standard portals. At the arthroscopy, a soft tissue mass that was yellow, pink and partially shiny white in colour was localised in the femoral attachment of the PCL. There was no meniscal and articular cartilage lesion. The mass was totally resected with a shaver.
and thermal ablation device. The histopathological examination revealed that the lesion composed of sheets of round or polygonal cells that blended with hypocellular collagenised zones. Multinucleated giant cells were scattered throughout the lesion. Xanthoma cells and haemosiderin granules were present (Fig. 3). The lesion was diagnosed as tenosynovial giant cell tumour. The patient had no more complaints after the operation, and there was no recurrence at 36 months' follow-up.

**DISCUSSION**

Tenosynovial giant cell tumours are generally differentiated into subgroups according to the region of occurrence and growing properties. These types have different clinical properties and biological behaviour. Tenosynovial giant cell tumours are classified as localised and diffuse tumours. Localised tumours are benign soft tissue tumours that predominantly involve the palmar side of the fingers and toes. These tumours are typically painless and grow slowly. The incidence in males is two times that of females, and the mean age of patients is 30–50 years. These tumours typically do not involve the larger joints and are very rarely placed intra-articularly. They are generally small (< 4 cm in diameter) and well-contoured masses, although tumours of up to 8 cm in diameter have been reported. A 10%–20% recurrence rate has been reported after local excision. As giant cell tumours do not have characteristic clinical features, they are usually diagnosed by investigations based on suspicion. 10%–14% of patients show cortical bone erosions.

The nature of this lesion is still controversial. Jaffe et al considered it a reactive process, hence the name, nodular tenosynovitis. Most authors currently regard it as being neoplastic, a hypothesis supported by the presence in this lesion of clonal chromosomal aberrations. MR imaging has been reported to be the best diagnostic technique for this entity. In this case, we report a histopathologically-proven giant cell tumour of the tendon sheath arising from the PCL, that was diagnosed by MR imaging and totally resected arthroscopically. This is the second case in the literature. We conclude that localised giant cell tumour of the tendon sheath can be placed as a unique lesion in the PCL, and a total arthroscopic resection is possible.

**REFERENCES**