Pigmented Villonodular Synovitis with Gross Extraarticular Extension and Intraarticular Infiltration

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Abstract
Pigmented villonodular synovitis (PVNS) is an unusual condition of unknown etiology. It is divided into localized and diffuse forms. Once diagnosed, treatment usually consists of surgical synovectomy. However, postoperative recurrence of PVNS occurs relatively frequently. This is a case report of a 78-year-old woman who suffered PVNS recurrence in her knee twice. Gross extraarticular extension and intraarticular infiltration is very rare. It is assumed that the lesion that originated in the extraarticular bursa or peritendineum infiltrated the intraarticular site. The patient was recurrence-free at 12 months after the operation, but six months later a few tumors were noted in her knee. The subsequent recurrences were due to incomplete excision of the previous tumors resulting from technical difficulties.

Keywords: Pigmented villonodular synovitis; Knee joint; Recurrence; Extra articular site

Introduction
Pigmented villonodular synovitis is a disease of unknown etiology that mainly develops in the synovial membrane of the knee joint [1,2]. It is said that the estimated incidence of PVNS is around 1.8 per million population [3]. It rarely involves multiple joints and often recurs after surgery. Gross extraarticular extension and intraarticular infiltration in the knee joint is very rare. We report a case of diffuse pigmented villonodular synovitis (DPVNS) that occurred in the popliteal region and that recurred after several surgical procedures.

Case Presentation

Fourteen years ago, a 78-year-old female, noted that she had a tumor on the medial side of the popliteal region of her knee. As there were no notable symptoms (such as pain), she left it untreated. However, over the years the tumor gradually grew, and due to restricted knee flexion, she became unable to sit with her knees bent. Four years later, she underwent an open synovectomy for PVNS, which was performed by another physician. After the operation, she was symptom-free, and her postoperative course was generally satisfactory. However, three years after surgery, another tumor developed on the lateral side of the left popliteal region, which caused her pain in her left knee joint during ambulation. Under a diagnosis of recurrent PVNS, a second synovectomy was performed. Four years after the second surgery, a tumor developed in the left popliteal region, which gradually caused numbness in the lower left leg and a heavy sensation in the same leg during walking. The patient was referred to our department for another surgical procedure.

A physical examination revealed swelling and slight local heat in the left knee joint. The range of motion of the left knee was 0 to 125º. She has no muscular weakness and sensory disturbance in her leg. A giant tumor, which was soft and elastic and displayed no adhesion to the skin, was found to extend from the medial to the lateral side of the left popliteal region.

A radiograph showed an imaging feature that was indicative of a soft tissue tumor on the medial side of the popliteal region of her knee. As there were no notable symptoms (such as pain), she left it untreated. However, over the years the tumor gradually grew, and due to restricted knee flexion, she became unable to sit with her knees bent. Four years later, she underwent an open synovectomy for PVNS, which was performed by another physician. After the operation, she was symptom-free, and her postoperative course was generally satisfactory. However, three years after surgery, another tumor developed on the lateral side of the left popliteal region, which caused her pain in her left knee joint during ambulation. Under a diagnosis of recurrent PVNS, a second synovectomy was performed. Four years after the second surgery, a tumor developed in the left popliteal region, which gradually caused numbness in the lower left leg and a heavy sensation in the same leg during walking. The patient was referred to our department for another surgical procedure.

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Arthroscopic inspection did not detect synovial proliferation in the supra-patellar pouch or
the tibiofemoral joint on the medial or lateral side; however, in the intercondylar fossa, villous synovial proliferation, as well as the growth of a yellowish-brown nodular synovium measuring 5 mm in diameter was noted around the anterior and posterior cruciate ligaments (PCL) (Figure 2).

Upon macroscopic examination, a brownish tumor measuring 15 x 10 x 7 cm on the medial side and 7 x 7 x 5 cm on the lateral side was found in the popliteal region (Figure 3). On the medial side, the tumor had displaced the popliteal artery and vein, pushing them in the lateral direction. It had also infiltrated the medial head of the gastrocnemius, semimembranosus, semitendinosus, and gracilis muscles and the area around the associated tendons. The tumor, the medial head of the gastrocnemius muscle, and the semimembranosus muscle were excised en bloc. The tumor had also proliferated in the region where the PCL is attached to the tibia.

A histopathological examination revealed the diffuse proliferation of synovial cells, a conglomeration of foam cells with iron deposition, and a scattering of multinucleated giant cells. There were no mitotic figures (Figure 4). These histopathological findings were consistent with those of pigmented villonodular synovitis.

As a postoperative treatment, the patient’s leg was immobilized with a plaster cast for 3 weeks. After the cast was removed, the patient wore an orthosis with a support and underwent weight-bearing training while taking care not to cause herself pain. One year after surgery, the pain at her left knee joint was still associated with slight swelling and fever. The range of motion of the joint was 5º during extension (hyperextension) and 130º during flexion. Her muscle strength was slightly reduced (flexion G, according to the manual muscle test). However, the patient was no longer dependent on orthotic devices, and there were no restrictions on her activities of daily living. The patient remained recurrence-free 12 months after the operation, but six months later a few tumors were noted on her knee.

**Discussion**

PVNS, a condition that mainly develops within the knee joints of young and mature adults, is a tumor-like proliferative disease of the synovium that involves chronic swelling and articular hematomas as its major symptoms. There are various theories concerning its etiology (e.g., inflammation, tumor formation, abnormal lipid metabolism,
and trauma) but none have been conclusively proven. However, it is believed that a synovium- or tendon-sheath-originating giant cell tumor and an associated non-specific inflammatory reaction are most likely the cause of this disease.

Gross extraarticular extension is very rare in this condition. Flandry noted that pigmented villonodular synovitis penetrates compartmental boundaries by mechanical means when the capacity of a given compartment no longer accommodates the expanding synovial mass [4]. One may consider the following two routes for the progression of the disease in this case: the lesion that originated at the intraarticular synovium infiltrated the extraarticular region or it originated in the extraarticular bursa or peritendineum and then infiltrated the intraarticular site. Enzinger et al. stated that PVNS that develops outside of a joint often represents tumor proliferation from the intra- to extraarticular regions [5]. However, the findings in the present case did not include a history of intraarticular hematoma; a tumor that was contiguous with the articular capsule at the initial synovectomy; the localization of the intraarticular lesion in the intercondylar region, despite the extensive spreading of an extraarticular lesion; or infiltration of the extraarticular lesion into intra- and intermuscular sites. Based on these findings, it was concluded that in the present case the lesion that originated in the extraarticular bursa or peritendineum had infiltrated the intraarticular site, while the subsequent recurrences were due to incomplete tumor excision.

The following have been employed for the surgical treatment of PVNS: endoscopic synovectomy for the localized type, synovial resection or therapy combining synovectomy and radiotherapy for the diffuse type, and arthrodesis or joint replacement with a prosthesis for the progressive type with osseous infiltration or joint deformation [2,4,5]. Despite improvements due to advances in surgical technology and in contrast to the localized type, synovectomy for diffuse PVNS (as was performed in the present case) is still reported to be associated with relatively high recurrence rates, even when the excision is complete [6,7]. Due to problems, e.g., infiltration of the lesion into the muscles, incomplete excision due to conditions such as adherions, damage to the articular components, and compromised articular functions, the efficacy of synovectomy is limited when applied to a diffuse-type PVNS that recurs at an extraarticular site. Diffuse PVNS is an inflammatory disease when classified according to its histopathology. However, it has been suggested that PVNS might be a benign synovial neoplasm rather than an inflammatory disorder [8]. The approach combining surgical synovectomy and radiotherapy could be an interesting option for the diffuse type at a high risk of recurrence [7,9-11]. However, little known about the long-term effect of radiation used for knee joint PVNS, and long-term complication should be considered after radiotherapy to prevent recurrence.

**References**