Recurrent spontaneous haemarthrosis of the knee associated with a synovial and juxta-articular haemangiohamartoma

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Abstract
Several spontaneous episodes of haemarthroses of the knee in a young man led to repeated arthroscopic examinations and one arthrotomy of the affected joint until a juxta-articular haemangiohamartoma was indicated by angiogram. Excision of the malformation stopped the bleeding tendency.

Traumatic haemarthrosis of the knee joint is common and currently the intra-articular causes of bleeding can be accurately diagnosed by arthroscopy. A spontaneous haemarthrosis of the knee joint is rare and is usually due to haemophilia. Vascular tumours and malformations are other, more uncommon causes. There are two forms of vascular tumour of the legs—namely, true haemangiomas and arteriovenous malformations or haemangiohamartomas, which may both involve the synovium and cause spontaneous intra-articular bleeding.

In this article a case of juxta-articular haemangiohamartoma with a synovial extension associated with a haemorrhagic synovitis and recurrent spontaneous haemarthroses is reported.

Case report
A 24 year old man sustained a haemarthrosis of the right knee while playing volleyball without any accident (November 1984). He had had varicose veins and hypertrophy of his right lower leg since childhood. At the age of 17 he underwent a successful operation for varicose veins. Apart from haemarthrosis the knee joint was normal clinically, radiologically, and in an arthrographic examination. The leg was lengthened by 2-5 cm and the circumference of the calf increased by 2 cm.

Between January 1986 and March 1986 the patient sustained three spontaneous haemarthroses of the knee. In January 1985 he underwent arthroscopy of the knee. The lateral part of the suprapatellar recess was thickened and uneven. Haemangiomatous tissue was suspected in this area, which was resected by an open arthrotomy. In the histological examination the synovium was recently thickened and covered by loose granulation tissue, under which were found both recent and old haemorrhagic changes, particularly around a large artery-type vessel. In the basal layer the synovium was rich capillary haemangiomatous tissue (fig 1A). At this stage the histological diagnosis was haemorrhagic synovitis.

The patient’s blood was analysed for any coagulation disorder. All tests were normal. Liver function tests were also normal. In October 1985 arthrography and arthroscopy were performed again, but the bleeding point could not be verified.

In February 1986 an angiographic examination around the knee was performed. An arterial phase was normal (fig 2A), but in the venous phase a collection of dye in small loculi was noted in the proximal and medial part of the knee (fig 2B). In June 1986 the vascular tumour was excised. It was located subcutaneously in an area between the vastus medialis and hamstring muscles, which consisted of a 3 cm collection of venous plexuses. The histopathology of the tumour showed non-neoplastic partially angiovenous and partially capillary vascular malformation (figs 1B–E). Recovery from the operation was smooth and uneventful. The patient has been able to continue with normal military exercises and sports activities up to the present.

Discussion
In 1990 Klippel and Trenaunay described a syndrome characterised by osseous and soft tissue hypertrophy, varicose veins dating from infancy, and haemangiomata of the skin. Since then there has been a confusing nomenclature for the vascular malformations of the legs: Parkes-Weber syndrome, muscle–skin angiomata, diffuse angiomatosis, congenital arteriovenous fistula, and genuine diffuse phlebectasia. In the last reports no clear difference was made between true haemangiomata and haemangiohamartomas, both were reported under the name: haemangiomata.

Goidanich and Campanacci studied 94 patients and classified the vascular malformations of the lower legs into six groups, the most common being localised deep vascular hamartomas in about half of the cases. The patient they presented had a typical subcutaneous vascular hamartoma as shown by its localisation, angiographic study, and histological findings.

The first case of synovial haemangiomata was described by Bouchul in 1856. Since then over 100 cases have been reported. Most cases have been haemangiohamartomas and only a few true haemangiomas.

True haemangiomas are small, sessile or stalked lesions arising near the infrapatellar fat pad and must be distinguished from villonodular synovitis. Their maximum size is 0·8 cm. Sometimes phleboliths are seen. Microscopic
Non-infectious, benign causes of spontaneous haemarthrosis of the knee

| Haemophilia and associated conditions |
| Synovial haemangioma |
| Synovial and juxta-articular haemangiomartoma |
| Kasabach-Merritt syndrome |
| Pigmented villonodular synovitis |
| 'Chronic rupture of anterior cruciate ligament' |
| Chondrocalcinosis |

examination shows that the haemangioma is usually cavernous, and occurs most often in childhood or in young adults.

In addition to pain, swelling, and palpable mass, both the haemangiomartoma and synovial haemangioma may cause spontaneous haemarthroses. Haemangioma was suspected in our patient, but despite two arthroscopies, two arthrogroms, and one arthrotomy no synovial vascular tumour could be confirmed because the synovial extension of the tumour was basal and capillary and therefore not visible. Excision of the extra-articular part of the tumour cured the bleeding, however. The tumour may cause venous congestion of the synovium and haemorrhagic synovitis.

Arthroscopy has been recommended as a diagnostic tool. Without it small, stalked synovial haemangiomas especially, may not be diagnosed. Arthroscopy should be performed without tourniquet to detect the blood filled tumour and without a television which may change the characteristic dark colour of the haemangioma. With small haemangiomas the angiogram may be negative. A tourniquet was used in our case, which possibly made a proper diagnosis more difficult.

Spontaneous haemarthrosis itself is rare. It may be caused by infections or malignant tumours. The table lists benign, non-infectious causes.

The most common cause of spontaneous intra-articular bleeding is haemophilia, and the common site the knee joint. Resnick and Oliphant described a case of Kasabach-Merritt syndrome associated with haemophilia-like arthropathy of the knee joint. This syndrome consists of extensive capillary haemangiomas, leading to consumption coagulopathy and bleeding tendency.

Pigmented villonodular synovitis is another rare cause of spontaneous haemarthrosis, sometimes accompanied by vascular malformation.

Recurrent, post-traumatic, spontaneous haemarthroses of the knee may be due to a partial tear of the anterior cruciate ligament. When an initial trauma causing a partial tear of this ligament is followed by poor scar formation the ligament remains insufficient. Further tearing is accompanied by bleeding in the joint. Recurrent haemarthrosis is pathognomonic of a 'chronic cruciate ligament rupture'.

Menkès and Rondier reported a series of 28 patients with haemarthrosis in 28 joints, including 20 knees. The haemarthroses were associated with chondrocalcinosis and joint destruction.
When vascular malformation is suspected in the spontaneous haemarthrosis an angiographic examination is indicated before arthroscopy.

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