Multimodal ultrasonographic evaluation in a case with unossified primary synovial osteochondromatosis

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Abstract
Primary synovial osteochondromatosis is a rare disorder and its diagnosis remains a challenge to the physician. We present the case of a 36 year old patient with right knee monoarthritis in which ultrasound findings, corroborated with clinical and histopathological evidence confirmed the diagnosis of unossified primary synovial osteochondromatosis. The arthroscopy with synovectomy found multiple intra-articular loose bodies occupying the entire joint. The ultrasonographic findings were extensively evaluated.

Keywords: primary synovial osteochondromatosis; knee; ultrasonography

Primary synovial osteochondromatosis (PSO) (Reichel-Jones-Henderson syndrome) is a rare, benign, monoarticular disease, affecting the synovium. The etiology is uncertain; however, elevated levels of bone morphogenetic protein, interleukin-6 and vascular endothelial growth factor-A are found in the loose body lesions. The syndrome is characterised by multiple intra-articular cartilaginous loose bodies, uniform in size, that may grow, calcify or ossify. Cartilaginous bodies are a result of proliferation and synovial metaplasia; in severe cases they may occupy the entire joint space or infiltrate into adjacent structures. Patients usually are in the 4th or 5th decades of life and men to female ratio is 2:1 to 4:1. The knee is affected preferentially (up to 70%); however, the disorder can arise in any other joint [1-6].

The most frequent clinical presentation includes pain, swelling and limitation of motion, often with slowly progression (years) [3]. The ultrasonographic (US) features were not extensively evaluated. Recent studies describe hyperechoic foci, posterior acoustic shadowing, intra-articular fragments which may change in position during dynamic US examination. Power Doppler examination reveals avascular process [7]. We report the case of a young male with PSO, in which we used multiple US techniques in order to evaluate the specific features of this syndrome.

Case report

A 36 year old male patient, without previous medical history, non-smoker, was referred to our department for pain, swelling and limitation of motion at the right knee. The symptomatology started 3 years before presentation with diffuse, almost continuous, constrictive pain, intensified by effort, accompanied by progressive limitation of motion.

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In this period the patient had few medical examinations and the most frequent etiologies of knee monoarthritis (septic arthritis, spondylarthritis, Lyme arthritis, crystal arthritis) were excluded. The magnetic resonance imaging (MRI) and US examinations found only unspecific findings (effusion and proliferated synovia). Local
corticosteroid, analgesics and non-steroid anti-inflammatory drugs had no significant effect and the patient reported aggravated progressive limitation of motion. At admission a very painful, swollen knee with severe limitation of articular mobility was found but no other pathological signs. Biological exams, inclusive immunologic tests, showed no lab alterations. Knee conventional radiography was in normal range.

US examination was performed using GE Logiq S8 machine, with 6-15 MHz linear transducer. In the suprapatellar region of the right knee numerous echogenic and homogeneous masses surrounded by transonic effusion were identified (fig 1a). The masses had no vascularisation and at strain elastography a stiff appearance was found (fig 1b,c). Three-dimensional US revealed that the entire recess, including the parapatellar recesses, was filled by these masses (fig 1d,e) and contrast-enhanced US confirmed the absence of vascularisation (fig 1f). The aspirated fluid had a rust-colored aspect (high level of iron was biochemical detected in this fluid) (fig 1g). A US-guided biopsy was performed (fig 1h). A combination of MRI T1-weighted fast spin echo proton density (PD) imaging with and without frequency-selective fat suppression (fs) and MRI T2-weighted gradient echo sequences in the sagittal, axial, and coronal planes was realized. MRI confirmed the presence of the large masses (diameter up to 4.5 cm), with intermediate signal intensity on T1-weighted images (fig 1i) and heterogeneously high signal intensity on T2-weighted images (fig 1j) with no enhancement.

Arthroscopy was performed, with the removal of numerous loose bodies (fig 1k,l) and synovectomy. Histopathological the loose bodies were composed of hyaline cartilage lobules with slightly increased cellularity (fig 1m). The final diagnosis was of unossified PSO.

**Discussions**

PSO is an uncommon benign disorder in which hyaline cartilage nodules form in the synovial tissue of the joint, tendon sheath or bursa. These nodules are usually detaching from the synovium and can occupy the entire joint [1-3].
Our patient was a young male, with a long history of disease and progressive aggravation of joint mobility. Previous studies showed that synovial chondromatosis can be highly aggressive and destructive, so a late diagnosis could severely affect the prognosis of the patient [2-5]. Microscopic appearance of PSO is composed of lobules of hyaline cartilage, surrounded by synovial lining (a two-cell layer of cuboidal epithelium). Sometimes, because of the metaplasia and pleomorphic cells, it is very difficult to make the differential diagnosis between this benign pathology and chondromyxoma. Diagnosis between this benign pathology and chondromyxoma may equal or exceed that seen in low grade chondrosarcoma [4].

Imaging findings depend on the stage of the disease and the presence of calcification or ossification of the cartilaginous bodies. For example, in 20-30% of the cases, wherein calcification is absent, there are no/non-specific radiographic findings (e.g. early osteoarthritis, erosions of adjacent structures, soft-tissue mass around the joint involved). Typical radiographic pattern includes multiple intraarticular chondral bodies with “ring-and-arc” chondroid mineralization and extrinsic erosion of bone. Our case had no calcification/ossification so there were no specific radiographic findings [5].

There are a few cases in literature in which the US findings in PSO are detailed. The presence of numerous small echogenic foci, representing the fronds of the synovium that have undergone metaplasia into cartilage were described [6,7]. We used all the US techniques (grey-scale, Doppler, elastography, three-dimensional US, elastography, contrast-enhanced US) in order to characterize this pathology. The homogeneity, stiffness, and lack of vascularization were the main US findings in our case of un-ossified PSO.

MRI can be used for better evaluation of the extension of PSO within the joint and surrounding tissue. The pattern was characterized as lobulated, homogeneous, intermediate signal intensity similar to muscle appearance on T1-weighted images, with high signal intensity on T2-weighted images. MRI can also evaluate marrow invasion, which is specific for malignant lesions, unusual for synovial chondromatosis [8].

Pigmented villonodular synovitis, synovial hemangioma and lipoma arborescens are conditions that can mimic synovial osteochondromatosis [9]. The necessity for a proper differential diagnosis and the impossibility of excluding a malignant process, were the reasons why we decided to perform a pre-surgical guided biopsy.

Management for PSO is mainly surgical and arthroscopic procedures are preferred. In our case, synovectomy was also realised, previous studies showing that this procedure gives better results compared to loose bodies removal alone [10]. Synovial chondromatosis can recur and when there is a recurrence it includes a fast growth and destruction of joints and a malignant degeneration must be considered [11,12]. It is well known that progression of synovial chondromatosis to chondrosarcoma is very rare. Distinction between these two conditions may be difficult using histology alone (the degree of cellularity and nuclear atypia may equal or exceed that seen in low grade chondrosarcoma), and should be based on clinical, imagistic and microscopic evidence [13].

In conclusion, US should be the first choice imaging technique in the evaluation of a clinically swollen or painful knee. Using the whole arsenal of US methods, the diagnosis and the plane for management of the local pathology can be generally accurately established. In unclear cases the need for other imaging techniques should be considered. Despite its lower frequency, in monarticular pathologies, the PSO should be taken in consideration for differential diagnosis.

References
