Primary tumors of the patella

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Abstract

The patella is an uncommon location for cancerous occurrence and development. The majority of tumors of the patella are benign, with a significant incidence of giant cell tumors and chondroblastoma. With the development of modern diagnostic technologies, there appear however many other histological types which raise challenges of diagnosis and treatment. In this article, we review the reported histological types of primary patellar tumors. Specifically, epidemiology, symptomatology, imageology, histopathology, and treatment options for these histological lesions will be discussed, respectively. As there is an increasing focus on the diagnosis and the treatment of these lesions, the availability of the integrated information about primary patellar tumors becomes more significant.

Keywords: Patella, Primary tumor, Symptom, Diagnosis, Surgery

Review

Introduction

A low incidence of tumors is found in the patella with the vast majority being giant cell tumor and chondroblastoma [1]. Diagnosis and treatment for these histological types of tumors are typically straight forward. However, some other notable, yet uncommon histological types which do not fall within these conventional patellar tumor types can cause both diagnostic and treatment dilemmas. The purpose of this manuscript is to discuss these current reported types of patellar tumors, their clinical manifestations, and appropriate treatment options. This list of patellar benign tumors and malignancies is not meant to be exhaustive. Many other, even rare factors that affect the condition of the patella are expected to be reported in the future.

Benign patellar tumor

Giant cell tumor

Giant cell tumor (GCT) is the most common type of the diagnosed patellar tumors (33% of all the patellar tumors) [1].

Clinically, the patients with the patellar GCTs usually complained about knee pains and/or swelling [2-5]. But there may be no inducement to the initiation of relevant diseases and the enhancement of such symptoms. Physical examination may show redness, local heat, swelling, effusion, tenderness, lump, crepitus, and the decrease of motional range [4-8].

The laboratory findings on some severe patients showed an increase of serum alkaline phosphatase (AP) [6,7] and erythrocyte sediment rate (ESR) [5,7].

Radiographs (Figure 1) revealed an osteolytic lesion of the patella with destruction of the bone [2-5,7,8], and sometimes, soap bubble appearance [7], sclerotic and radiolucent lesion [6], fracture of the femur and the tibia [3,6], and pathologic fracture [3] may also be found. Magnetic resonance imaging (MRI) showed abnormal extension and lesion of the patella, and there may be some evidences of adjacent tissues and sclerotic [2-4,8,9]. Additionally, chest radiography and bone scintigraphy are necessary for GCT patients to determinate the possible metastasis on the lung and other bones.

A needle biopsy or incisional biopsy of the patella is recommended for the diagnosis. The histopathological features of the patellar GCT, which mainly include numerous giant cells, short spindle-shaped cells, bone tissue calcification, and a few mitotic figures [2], should be distinguished from aneurysmal bone cyst or chondroblastoma. The histological examination of the aggressive GCT showed a compact stroma with nuclear atypia, frequent mitotic figures, hyperchromatism, and the unevenly distributed giant cells [5,7].
Surgery is the primary mode of treatment. GCT is a locally bony lesion, a tumor considered as having low recurrence rates using standard meticulous surgical technique according to Meyerding’s outline [10]. For small, benign GCT (Enneking stage 2), intralesional curettage or partial patellectomy with filling of the bone void and reconstruction of the extensor mechanism is the first choice in treating GCT of the patella. But if the patellar GCT is extensive or malignant (Enneking stage 2 to 3), patellectomy combined with adjuvant treatment and resection of metastasis is recommended.

Chondroblastoma
Chondroblastoma (CB) can also be found in the patella but at a very low occurrence rate (2%) [11]. It is the secondary common benign patellar tumor (16% of all the patellar tumors) [1].

CB appears as lesions on the patella that caused pain either with or without swelling [12-16]. The pain is aggravated by physical exercises and alleviated when at rest. Upon examination, CB patients had slight tenderness on the bad patella and some swelling with no erythema or warmness. The movement gave a slight feeling of pain when in extreme flexion. The patients who had the patellar CB with pathological fracture were found to move in pain, which heavily affected the range of knee motion [12,17].

Most of routine laboratory tests were within normal ranges. The increased serum calcium and AP were only found in the patients with patella fracture [12].

Plain radiographs (Figure 2) demonstrated a radiolucent lesion of the diseased patella with a well-defined sclerotic margin, lobulated rims, and thinned cortices. Patellar expansion and multicystic lesion of a ballooned patella was indicated in the patients with CB and aneurysmal bone cyst [16]. The fracture line appeared on the radiograph of the patient with pathological fracture [13,18]. Computed tomography (CT) scan revealed an osteolytic lesion with septation, sclerotic margins, and some calcifications in the patellar matrix [14-16]. MRI for the patella showed a lobulated lesion, expanded patella, and fluid-fluid levels. Meanwhile, the margins of the lesion were well defined [14-16]. Chest radiography was no anomalous change.

Histological examination revealed sheets of proliferating chondroblast with chondroid matrix and some multinucleated giant cells in places admixed with focal areas of coarse calcification [14-17]. Within the fibrous septum

Figure 1. Patellar giant cell tumor. (A, B) An enlargement of the cystic lesion in the patella and a bone translucency with peripheral rim change can be seen at the lateral down part of the patella on radiographs. (C, D) CT indicates that there is a well-defined lytic lesion with thin cortex occupying two-thirds of the patella.

Figure 2. Chondroblastomas of the patella. (A) On plain radiographs, the lesion appears as geographic changes with lobulated margins, thinning cortex, sclerotic rim, and reactive bone. (B) Calcifications inside the lesion are seen on CT images.
and certain multinucleated giant cells, there may be several walled cystic zones full of blood and immature bone formation, which are compatible with aneurysmal bone cysts [15,16].

CB treatment included curettage, curettage with burring, and filling of the lesion cavity with bone graft and/or bone cement [19]. For the patellar CB (Enneking stage 2), curettage by the anterior longitudinal bone window was performed, followed by the filling of the cavity with bone graft [14-16,18]. Partial patellectomy was used for the CB patients with a pathological fracture [17]. In addition, curettage followed by chemical adjuvant treatment (Enneking stage 2) and excision (Enneking stage 1) were reported in some cases [1,14-16,20]. For large-sized patellar CBs with aneurysmal bone cyst, curetting with bone grafts may be suitable; for large-sized or recurrent ones, curettage and bone graft or patellectomy may be a better option [16]. Postoperatively, immobilization of the affected limb and then static quadriceps exercises are necessary for the patients to recover normal joint activities [16].

Osteoid osteoma
Although the patella is a rare site for osteoid osteoma (OO) with only an incidence of 5% in all patellar tumors [1], a few cases of such kind have been reported [21-24].

Long-term pain was the main complaint of the patellar OO patients, which was presented throughout the daytime and exacerbated at nighttime and after physical exercises; which was well localized at the patella and could be relieved by analgesic drugs [21-24]. Swelling could occur in the diseased knee as well [24]. There was no previous trauma in most cases [21-23]. Physical examination revealed mild effusion, a tender point at the patella, and slight atrophy of the affected thigh; no redness or local temperature rise. Range of motion was confined due to pain [21-24]. Sometimes, clinical examination could not be entirely completed due to severe pain [22].

Laboratory studies of reported cases were normal [21-24]. X-ray, CT, MRI, and bone scintigraphy were meaningful to the determination of the patellar OO, for they showed a well-defined lesion with nidus and mineralization in the affected location of the patella [21-24] (Figure 3).

Several researchers suggested that the natural history of osteoid osteoma be a process of spontaneous healing. But studies indicated that it would take a long time to fully resolve symptoms if the osteoid osteoma was not excised. Non-steroidal anti-inflammatory drugs can reduce the time of recovery period. OO was traditionally treated with excision of the nidus, so did patellar OO. Curettage nidus or resection was performed to resolve symptoms (Enneking stage 1) [21,23-25].

Aneurysmal bone cyst
Less than 1% aneurysmal bone cyst (ABC) of all the known cases occurs in the patella [26]. Patellar ABC has an incidence of 5% in all patellar tumors [1].

Patellar ABC patients frequently presented complaints of pain in the anterior aspect of the affected knee and/or swelling of the knee joint [26-30]. The clinical assessment could not find any recent trauma or signs of local inflammation. Palpation of the patella was painful and/or swelling compared to the healthy side [26-29]. Additionally, physical examination revealed full active and passive ranges of motion of the affected knee [26-29].

Hemogram, c-reaction protein (CRP), and ESR were completely within normal ranges [27].

Radiographs of the knee showed a ‘geographic osteolysis’ all over the affected patella, that is, cortical thinning, smooth borders, and intact articular surface [26-28]. Mild expansion and a multiloculated appearance could also be observed [28]. CT scan showed a fluid-filled multiseptate cavities without intralesional calcifications in the patella suggesting an aneurysmal cyst [27,28]. MRI showed a hyperintense lobulated mass with fluid-filled cyst in the lesion of the patella [26,29]. Doughnut-shaped patella was found on bone scintigram [30].

![Figure 3](https://via.placeholder.com/150)

**Figure 3** An intracortical osteoid osteoma of the patella. (A) On plain radiographs, a clear lesion with small fuzzy periosteal reaction is observed. (B) CT demonstrates the sclerosis surrounding the lesion.
Histopathologic examination of the biopsy material revealed empty bloody cavities or the presence of red blood cells [27-29]. The cavities were separated by those fibrous connective tissues composed of spindle cells, giant multi-nucleated cells, some macrophages, masses of hemosiderin, and trabecular bone [27-29].

The best choice of ABC treatment is surgical saucерization and curettage with removal of the cyst lining followed by bone grafting. Bone grafting or bone cement following curettage with burring by a mid-line surgical approach was employed to remove the lesion (Enneking stage 2) [27-29]. Patellectomy was also performed in some patients (Enneking stage 2) [1,20]. Reddy reported an agressive ABC case (Enneking stage 3) that received a treatment consisting of curettage, chemical and thermal cautery of the bed, and autogenous bone grafts [26].

**Chondroma**

Chondroma (CH), including both enchondroma (EC) and juxtacortical chondroma (JC), is a rare lesion with approximately 5% of incidence in patellar tumors [1]. Clinically, CH patients would state their ache, swelling, and patellar mass [31,32]. One case had a history of struck injury on the knee due to an automobile accident [32]. After physical examination, such symptoms as mild motion pain, mass, swelling, tenderness, heat of the pre-patellar bursa, and extension were detected [31,32]. Patients usually had full active and passive range of motion of the affected knee [32].

Routine laboratory studies and ESR were within normal ranges [31].

Radiographs of the knee revealed an oval lytic defect within the patella in EC patients [31]. The mixed lesions were showed on the edge of patella in JC patients. Well-defined margins, normal cortex, and small calcifications were presented, without any periosteal shell or reaction, sclerotic rim, septation, fluid levels, pathologic fracture, or joint involvement [33]. MRI of patellar JC patient showed a hyperintense lesion in T2 sequences on the edge of the patella [32].

Histologic examination revealed diseased tissue of hyaline cartilage containing chondrocytes with small, condensed nuclei, which conformed to characteristic of chondroma [31]. Lamnott [32] reported that patellectomy was performed in the patellar EC patient (Enneking stage 2), who then recovered a stable knee without any further complaints. Excisional surgery was used to resect the JC lesion (Enneking stage 2), and the patient was finally capable of full range of motion [31]. For the Enneking stage 1 CH, excision and flap was used [1].

**Osteochondroma**

Osteochondroma (OC) in the patella is rare with an approximate 2% incidence of patellar tumors [1]. Clinically, patellar OC might cause the symptoms of knee pain and mass [34,35]. Upon physical examination, a hardened mass was found adhering to the patellar bone plane, without any inflammation [34]. Limitation of flexion was presented in the case of patellar intra-articular OC [35].

Routine laboratory studies were within normal ranges [35].

Upon radiographs, an irregular calcification neoformation that stretched from the edge of the affected patella was detected [33-35]. Some lesions projected over the pole of the patella, while others extended toward the joints. Sometimes, the 'bone bridge' might be present [35]. Both tomography and MRI were helpful to a meticulous observation of the tumor and the patellofemoral joint surface [34,35].

Anatomic pathologically, the lesion was a pale polyloid tissue with the cartilage cap. Microscopically, the outer layer was composed of connective tissues and the deep layer of cartilage cells. Between these two layers, there were normal spongy bone tissues [35]. The results accorded with the diagnosed characteristics of osteochondroma.

Although the surgery to excise the tumor is not essential to all OC cases, it is still a rational method for patients who suffered from some OC symptoms and malignant transformation. The OC surgery involved curettage, burring, bone grafting, and bone cementing. The simple resection was often found to inhibit the tumor growth and to resolve the symptoms in Enneking stage 2 cases [34,35]. During the operation, the periosteum and perichondrium covering the lesion were generally removed to reduce the chances of a recurrence [36].

**Osteoblastoma**

Osteoblastoma (OB) in the patella is very rare (about 2% of all patellar tumors) [1], and only a few cases of this kind were reported [37-39].

The common symptoms of the patellar OB patients are pain and swelling [38,39]. Local tenderness might be observed during physical examination [38]. The motion of the affected knee was usually of full range.

Radiograph of the patella revealed an osteolytic lesion surrounded by a sclerotic rim without evidence of extra-articular involvement [33,37,38]. OB might cause a patellar pathological fracture which could be detected by radiograph [39]. Both tomography and CT scans confirmed a well-defined lesion of the patella with calcifications [37,38]. A solitary hot spot of the affected patella could be revealed by bone scintigram [39].

The pathologic examination demonstrated that the lesion comprised irregular trabeculae of immature woven bone, lined by plump osteoblasts within a loosely textured fibrovascular stroma containing numerous giant cells [37,38]. In the case with a pathological fracture,
one hemosiderin deposit was presented, due to previous hemorrhage [37].

Generally, both conservative surgery and complete resection (marginal or wide en bloc) are the options for the OB treatment. Intralesional curettage can be effective but is associated with a high rate of recurrence. The treatment method for local recurrences should be wide en bloc resection [37]. Allogeneic bone grafting following intralesional curettage was conducted to treat the patellar OB in different cases (Enneking stage 1 to 2) [37,38]. Patellectomy could be the remedy for a failure after wide curettage [39]. After the surgery, the patients were asymptomatic, capable of full functioning of the knee, and complete healing of the lesion without evidence of recurrence [37-39].

**Osseous hemangioma**

Patellar primary osseous hemangioma (OH) is very rare with only an incidence of 2% in patellar tumors [1].

Clinically, complaints of the patellar OH patients could include aching, swelling, and limitation of movements of the knee [40].

Routine laboratory findings, such as blood count, Mantoux test, ESR, blood calcium, blood phosphorus, and AP, proved to be normal [40].

Upon radiographs of the knee joint, there is a multilocular cystic lesion all over a part of patella with possible surroundings of sharp margin, thinned cortex, and sclerotic rim [33,40]. A secondary fracture could be found in the radiographs [40]. CT could determine the radiographic results [40] (Figure 4).

Pathological findings showed that the loculated lesion containing fluid and clotted blood was traversed by thin bony septa. Existence of a fracture could be judged by visual inspection. Microscopically, the cyst consisted of large and small spaces filled with endothelium and blood. Reactive bone formation in the bony trabeculae running across the lesion could be indicated, as well [40].

Patellectomy was performed in the two patellar OH cases (Enneking stage 1 to 2), and the treated patients were finally capable of full movements of the knee joints [40].

**Simple bone cyst**

Simple bone cyst (SBC) comprises about 3% of all biopsied primary bone tumors and accounts for 4% of patellar tumors [1].

The patellar SBC patients usually complained about knee pain and swelling [41-43]. The pain could be related to traumatic history and aggravated by motion and climbing [41,42]. Upon physical examination, tenderness, swelling, flexion deformity, and the result of a positive patellar grinding test could be detected [41-43].

Radiographs showed a radiolucent lesion with well-defined margins in the affected location of the patella [41-44]. There might be a cortical destruction arising from a fracture [42]. CT could determine the radiographic results, indicating the elaborate changing of the cortex and patellofemoral joint [41,44]. Bone scintigram was normal [41].

Laboratory investigations did not reveal any abnormal findings [41].

Macroscopically, the lesion included a cyst lined with a thin membrane and a smooth inner surface, full of clear or yellowish fluid [41,42]. Microscopically, the histological findings about the simple cyst were consistent with the diagnosis of SBC [44].

The patellar SBC lesions were eliminated by autogenous grafting following curettage (Enneking stage 1) [41] or patellectomy (Enneking stage 1 to 2) [42,43]. Restoration of the extensor mechanism was performed after patellectomy [43]. Postoperative cases were asymptomatic, capable of full motion of the affected knee with no recurrence [41-43].

**Other benign patellar tumors**

Other benign patellar tumors that have been reported include lipoma, osteitis fibrosa cystica, ganglion, osteoma,
and non-ossifying fibroma. Because of the limited case reports, more details cannot be obtained. Known information thereof is shown in Table 1.

**Malignant patellar tumor**

**Osteosarcoma**

Osteosarcoma (OS) is the most common type of primary malignant osseous tumor. It mainly affects the long bones of the extremities but also affects other bones like patella. The patellar OS has an incidence of 6% in patellar tumors [1].

The patellar OS patients complained about the increasing anterior knee pain and swelling in the affected knee [45-49]. Most of them did not have history of knee injury [45-48], while some had a history of trauma or radiation [6,49]. Symptoms of a few cases might disappear after the conservative management [46]. Physical examination could reveal tenderness, swelling, redness, limited flexion, and muscle atrophy around the affected thigh [45-48].

Plain radiographs (Figure 5) showed lytic lesion, osteoblastic lesion, shape changing, and cortical thinning/breath in the patella [45-49]. OS mimicking GCT presented the multilocular lytic lesion [46]. Most of the cases had no cortical wall destruction in the patellofemoral joint [45-47,50]. The telangiectatic OS cases demonstrated a lytic lesion of the surrounding bone and osteoid matrix in the soft tissues extended from the patella [48]. CT scans could demonstrate soft tissue extension and an intraosseous lytic lesion with cortical thinning/breath [46,47]. MRI was performed to determine the diagnosis as well as to detect changing of the adjacent soft tissues and bones [45-47,51]. The mass showed low signal intensity on T1-weighted images, and then T2-weighted images presented high signal intensity. Contrast-enhanced fat-suppressed T1-weighted images demonstrated strong enhancement of the lesion [48,51]. Tc-99m bone scintigraphy only showed an uptake in the affected patella [45,46]. The CT scan of the chest was necessary for identifying pulmonary metastatic nodules [48].

Only in one case, ESR and white blood cell (WBC) count were increased [49].

Photomicrograph of histology revealed the diffuse proliferation of malignant cells that led to abundant osteoid formation. The cells were spindle shaped, atypical with large, hyperchromatic nuclei and prominent nucleoli [46-50]. Irregular osteoid formation, pleomorphic malignant fibrous histiocytoma-like areas, and telangiectatic areas were also found in some special cases [46].

Surgery is still the main therapeutic approach. For patellar OS, surgical methods, including patellectomy (Enneking stage IIB, III) [45,50], wide resection (Enneking stage IIA) [46], amputation (Enneking stage IIB, III) [47,49], arthrodesis with an allograft and total joint replacement (Enneking stage III) [48], were performed in patellar OS cases. Before and after surgery, all of the patients experienced chemotherapy. The surgeon tried the above-knee amputation, due to the poor results of preoperative chemotherapy or previous surgery [47,49]. The excision was performed with safe surgical margins, including for patella, the quadriceps tendon, a patella tendon, intrapatellar fat pad, and a medial and lateral retinaculum [45,46]. Cho [45] reported that an allograft patella was used for restoration of extensor mechanism function which was a difficulty after wide resection. Artificial knee joint tumor prosthesis replacement was used in one case because of the lesion spreading to the proximal tibia and adjacent soft tissue [48]. Most of the cases had no recurrence or metastasis on the patella within short postoperative period [45,46,48]. Two patients presented metastatic disease [47,49].

**Chondrosarcoma**

Originating from cartilage tissues, chondrosarcoma (CS) is the second most common primary bone tumor. It may occur at any endochondral ossification of the whole body bones, especially at the pelvis, femurs, humeruses, and scapulas [52]. CS occurrence in the patella is still scarce, with few cases being reported [33,53-55].

Clinically, increased mass, pain, swelling, and lameness of the affected knee might be the main complaint of the patellar CS patients [53,54]. The history of rheumatoid arthritis or a history of struck injury on the knee maybe caused patellar CS [53,54]. Upon examination, the patient might have swelling and tenderness over the affected patella [55]. Mass on the patella could be touched and felt [54]. The passive range of knee motion was limited in two cases [53,54].

Radiographs of the affected knee joint showed a lytic lesion shadow on the patella [53-55]. The tumor had a well-defined sclerotic rim and indicated some cortical continuity [54]. Meanwhile, the fracture shadow on the patella could be found by X-ray [55]. Sometimes, CS could mimic GCT with a ‘soap-bubble’ presentation and a ‘trumpet-shaped’ extension of the cortex [54]. MRI of the tumor showed a comparatively clear boundary around the lesion. T1-weighted images demonstrated low intensity, while heterogeneous low-high combined intensities were found on T2-weighted imaging with fat suppression [53]. Bone scan showed an increased focal activity on the lesion of the affected patella [53]. Chest X-ray or CT should be performed to determine lung metastasis.

Hematological tests of one case [53], including WBC count and AP, were within the normal range. Rheumatoid factors and CRP levels were slightly increased in one case, due to the history of rheumatoid arthritis. In another case, only the increased ESR was found [55].
| Type                        | Symptoms                                      | History                                    | Clinical examination | Imaging                                                                 | Laboratory test | Pathology                                                                 | Treatment                                                                 |
|-----------------------------|-----------------------------------------------|--------------------------------------------|----------------------|---------------------------------------------------------------------------|-----------------|---------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Lipoma [44]                 | -                                             | -                                         | -                    | R: a well-defined multilobulated lytic lesion replacing part of the patella | -               | The specimen demonstrated a lobulated lesion                             | -                                                                         |
| Ganglion [1,20,63]          | Immediate pain and swelling of the affected knee after twisting it during a fall | Injury of history                          | A knee joint effusion, point tenderness, limited range of motion, severe pain on flexion-extension of the knee | R: a well-defined lytic lesion, a pathological fracture, joint effusion, lobulated margins, septa, sclerotic rim, thick trabeculae. BS: a focal solitary area of intense activity in the affected patella | -               | A cyst-like lesion containing fibromyxoid and 'fatty'-appearing material, a mild amount of chronic inflammatory cells and reactive bone | Immobilization to treat the pathological fracture (stage 3), curettage and allograft bone graft (stage 3), curettage (stage 1) |
| Osteitis fibrosa cystica [62]| -                                             | A history of thyroid-related problems       | -                    | R: osteitis fibrosa cystica with multiple irregular cystic areas          | -               | -                                                                         | Excision of the lesion (stage 1)                                        |
| Leiomyosarcoma [64]         | Patellar pain and swelling                    | Be treated with curettage and bone graft of the patella due to uncertain diagnosis | Swelling, local heat, elastic hard mass, decreased range of motion, thigh muscle atrophy | R: a mixed lytic and sclerotic lesion with ill-defined margin in the patella. CT: sclerotic rim and osteolytic lesion with cortical disruption. MRI: extraneous high signal intensity infiltration on T2-weighted imaging. BS: isolated increased activity in the affected patella | WBC count, AP and CRP were within normal ranges | -                                                                         | Extra-articular wide resection with total patellectomy and reconstruction by Howmedica modified resection system (Stage IIIIB) |
| Angiosarcoma [1,20]         | -                                             | -                                         | -                    | R: multicentric lesion, permeative bone with cortex destruction and ill-defined margins, joint involvement. CT: destroyed cortex and determination of the grade of the tumor. | -               | -                                                                         | Radiotherapy (stage III, IIIIB)                                         |
| Hemangioendothelioma [1,20,33]| -                                             | -                                         | -                    | R: a lobular contour with ill-defined margins, septation 'soap-bubble appearance', thinned and piercing cortex, mimicking giant cell tumor | -               | -                                                                         | Patellectomy (stage IB), radiotherapy (stage IA), patellectomy and radiotherapy (stage IA, IB) |
| Ewing’s sarcoma [65]        | Increasing patellar pain and swelling, limp, impairing function, atrophy of quadriceps, weight loss, night sweats | A motor vehicle accident some months prior | A warm and swollen knee, painful limitation of flexion, patellofemoral irritability at compression, | R: mass-like change, sclerosis, permeative change. MRI: marrow replacement by tumor in the patella, soft tissue mass, lesion of proximal tibial, widespread involvement of bone marrow in the vertebral bodies and sacrum epiphysis. CT: pulmonary metastases. | Straw-colored joint fluid with 850 white cells with 94% mononuclear predominance, increasing of ESR. | Extensive necrosis of the lesion, neoplasm composed of nests of small round cells, scant amphophilic cytoplasm, cytoplasmic glycogen (+), membranous CD99 (+), nuclear FLI-1 (+) | Chemotherapy (stage IIIIB)                                               |

R = radiographs; CT = computed tomography; MRI = magnetic resonance imaging; BS = Bone scan.
Macroscopically, part of the patella had been replaced by the lesion, and certain necrosis of its surrounding bones and soft tissues could be found [53,54]. The lesion consisted of fragments of gelatinous tissue and bony spicules [54]. Yellow jelly-like material was drained from the joint in some case [55]. Upon microscopic examination, trabecular bone was admixed with proliferating chondroid tissue. Whereas, the arrangement of the cells and their relation to the intercellular substance resembled the structure of embryonal cartilage [54]. At higher magnification, a small number of cells with two nuclei and some single-nucleus cells with larger or smaller than average-sized nuclei were found [53].

Adjuvant chemotherapeutic drugs or radiotherapy is useless to CS due to its poor blood supply and lymph circulation. The wide surgical resection is still regarded as a mainstay method for treating CS [52]. The surgeons performed excision plus partial patellectomy or amputation to eliminate the tumor (Enneking stage IB), while amputation was also used for the worse case (Enneking stage IIIB) [53-55]. The two patients who accepted postoperative follow-up were found to have no recurrence in several years [53,55].

**Primary osseous lymphoma**

Generally, lymphomas arise in lymph nodes, while their formation in extranodal sites is less common. The patellar POL accounts for approximately 4% of all the patellar tumors.

One patient with the patellar POL presented pain, due to a pathologic fracture [56]. A physical examination of the affected knee could reveal swelling, painful flexion-extension motion, active arthritis, and tender subcutaneous nodules [56,57].

The patella exhibited moth-eaten appearance and multiple lytic lesions on knee through X-ray inspection [44,56,57]. A shadow of the fracture was shown as well (Figure 6). Bone scans and imaging of the chest, abdomen, and pelvis failed to indicate any other involvement [56,57].
Hematological and biochemical tests, bone marrow biopsy, and synovial fluid were unremarkable [56,57].

Excised lesion revealed the diffuse infiltration of tumor and/or surrounding tissues [44,57]. In the case of T-cell lymphoma, differently sized lymphoid cells with less cytoplasm, vesicular nuclei, single or double conspicuous nucleoli, and mitotic figures were found through the microscope. The cells could strongly stain CD3 and CD45 but not CD19 or CD30 [57]. Large B-cell lymphocytes were positive to CD45 and CD20, having intermediate proliferative activity; they were chased in the case of B-cell lymphoma [56].

The common treatments for primary bone lymphoma included the following: a combination of chemotherapy and radiotherapy, chemotherapy alone, and radiotherapy alone [58]. The POL patients (Enneking stage IIB) were treated with cyclophosphamide, doxorubicin, vincristine, and prednisone (CHOP) chemotherapy. But for the patients who suffered from severe POL (Enneking stage III) had received radiotherapy following CHOP chemotherapy [56,57]. Surgery was performed to repair the pathologic fracture [56]. Finally, both of them recovered from the disease. Only one recurrent case (Enneking stage III) was found. That patient treated with radiotherapy died 1 year later [1].

Malignant fibrous histiocytoma

Malignant fibrous histiocytoma (MFH), which arises in both the soft tissues and the bones, is the most frequent sarcoma of malignant soft tissue that is characterized by aggressive biological behavior. Patellar MFH, which originates in some unusual location, accounts for 1% of all the patellar tumors [1].

The patient’s complaints include pain, swelling, decreased range of motion, anorexia, and weight loss [59,60]. Physical examination could show swelling, tenderness, cutaneous erythema, and restricted knee motion [60]. One of the cases had a knife wound before his admission [60].

The radiographs showed multiple radiolucent lesion and permeating destruction in the patella [59,60]. Septated lesion and sclerosing areas were also found [60]. Plain films of knee joint could detect a fracture line [60]. CT scans demonstrated a mass and involvement of surrounding tissue. Bone scans were used to determine metastasis and other imaging results.

The only abnormal laboratory finding was the increased level of ESR [59,60]. The knee fluid test demonstrated 1+ WBC in one case [60].

Generally, specimens showed grayish, whorled tissue with thin, calcified septa and thinned cortex [60]. There were nodules which consisted of polymorphic cells with large eosinophilic cytoplasmic that were epithelioid-like in appearance [60]. MFH cells possessed prominent nuclear pleomorphism, hyperchromasia, and nuclei irregularity [59,60]. In some other areas, fusiform cells with a marked storiform pattern were found. Nontumoral bone trabeculae within the neoplasm were present, some lacking osteocytes. A positive intracellular stain with acid phosphatase was detected by histochemical method. The electron microscopy proved fusiform and polygonal structure of cellular elements [60].

The treatments of osseous MFH included wide resection, additional radiotherapy, adjuvant chemotherapy, systemic treatment approach, or the combination [61]. The first patellar MFH case (Enneking stage III) was reported to have undergone curative resection plus prosthesis but eventually died of the disease [50]. Amputation plus prosthesis fitting followed by chemotherapy had been employed to treat another patellar MFH patient (Enneking stage IIB). And no recurrence was found within 18 months after the treatment [59].

Other malignant patellar tumor

Other sporadic malignant cases that were ever reported, such as leiomyosarcoma (Figure 7), angiosarcoma, hemangiendothelioma (Figure 8), and Ewing’s sarcoma, have been summarized in the Table 1.

![Figure 7](image_url) Patellar leiomyosarcoma. (A) Plain radiograph shows a mixed lytic and sclerotic lesion of patella. The margin of the lesion is ill defined and associated with cortical breach. (B) CT scan reveals multiple lytic lesions of the patella with a sclerotic rim and cortical disruption.
Discussion

The patella is defined as a sesamoid originating in the quadriceps tendon. It develops from a cartilaginous precursor in the third month of gestation and ossifies about 3 years of age [1]. Its ossification is similar to that of a long bone’s epiphysis or apophysis. This makes the patella a possible location of bony lesions [62]. Due to limited published literatures, we still know a little about patellar neoplasms. As the patellar neoplasms are such a rare etiology as anterior knee pain, their determination is often postponed. However, a majority of primary patellar neoplasms usually cause a dilemma in diagnosis. Therefore, physicians urgently need more evidences to distinguish primary patellar tumors from other kind of lesions through integrated analysis of epidemiology, pathogenesis, symptomatology, imageology, and histopathology and in reference to the reported treatments.

The most common symptom of primary patellar tumors is knee pain that usually lasts a long period of time from pain onset to determination upon diagnosis [50]. For benign tumors, the feeling of pain is gaining slowly; for malignant ones, the feeling of pain enhances more rapidly. Sometimes, the immediate pain will present after injury history, resulting from a pathological fracture. Swelling - the secondary common complaint - occurs in several cases of both benign and malignant tumors. Upon physical examination, the impaired function of flexion and extension is easier to be determined than the patient’s complaint is. Affected by pain and swelling, the motion range of the diseased knee is forced to decrease. With the limitation of motion and limp, patients may find muscle atrophy of the quadriceps. Mass is another main complaint which is often indicated through the process of palpation. Noticeably, primary malignancies and aggressive tumors of the patella can raise local redness, heat, effusion, and other inflammatory representations. When signs of weight loss, night sweats, and other cachexia occur, physicians should mark a red flag for the possible malignant tumors. Moreover, there may be a relationship between the lesion site and the tumor type.

In patients with anterior knee pain, plain radiograph is the most common way to find patellar anomalous changes and can accelerate the diagnosis of a patellar tumor [50]. In most cases, radiographs of the patellar tumor accord with those of large bones. However, depending on the peculiarity of dimension, shape, and structure, radiographic reports of patellar tumors are still distinctive from one to another. More considerations should be made to imaging changes, such as bone destruction, extent, margins, cortex, sclerotic rim, septation, calcifications, ossification, periosteal reaction, pathologic fracture, and surrounding involvement. For benign tumors, ‘geographic aspect’ osteolytic lesion with well-defined margins and sclerotic rims are typical characteristics. Ossification, calcifications, and pathologic fracture are normal, as well (Table 2). Malignant and aggressive benign tumors are frequently accompanied by permeative destruction, total patellar extension, ill-defined margins, ossifications, and pathologic fracture (Table 2). Remarkably, involvement of surroundings is of utmost importance to the knee X-ray as well as to a visual measure to differentiate the malignant from the benign. Nevertheless, radiographic diagnosis is not absolutely correct because metastasis cancer, plasmacytoma, Paget’s disease, osteomyelitis, hyperparathyroidism, gout, and tuberculosis often mimic lytic neoplastic lesions of the patella [1]. Primary patellar tumors are often not studied through imaging results, which are otherwise very helpful to determining the staging of the lesion and the appropriate surgical course. Radiographic results combined with those of the CT and MRI are often needed to determine former diagnosis, surrounding involvement, and precise presentation of the tumor site. With bone scan, part of the primary patellar tumors is indicated as having only increased focal activity on the lesion of the affected patella. Among the existing case reports, the difficulty to obtain different diagnoses based
on radiographs happens at intervals [16,17,47,54]. Co-
existence of CB and ABC, CS-mimicking GCT, and
OS-mimicking GCT are the problems that should be
resolved by analyzing the results of histological exami-
nation. In addition, chest radiograph or CT is recommended
for all the patients with primary patellar tumors, especially
with malignant ones.

The past reviews on the topic of patellar tumors are
short of supports by laboratory tests, due to the insuffi-
cient reports and unusual changes. But as relative cases
are increasing, a certain amount of data has been made
available. Although the routine laboratory tests are usu-
ally within the normal ranges, a pathological fracture
may lead to the increased levels of serum calcium and
AP. A rise in levels of WBC and ESR can be detected in
some malignant cases after laboratory tests. This may
pertain to the secondary inflammatory reaction. Synovial
fluid examination has been used, which is of certain sig-
nificance to some cases. Therefore, laboratory tests must
not be overlooked.

Radiographic features of primary patellar tumors are
atypical of a specific histotype. Histological examination
is preferred, but in view of the risk of contamination, a
needle biopsy is a better option. An incisional biopsy is
recommended if the diagnosis and the radiographic fea-
tures are paradoxical. Two key points should be noticed:
(1) In needle biopsy, the joint must not be entered; (2)
in incisional biopsy, the patellar and quadriceps tendon,
the joint cavity, and the synovial membrane must not be
contaminated [1]. Histological examination is still a ‘gold
standard’ to determine the specific type of tumors, espe-
cially of the malignant ones. To our delight, pathological
features will not change as the site of the tumor changes.
Different types of patellar tumors still exhibit their in-
trinsic pathological characteristics. For instance, typical
spindle-shaped cells with large, hyperchromatic nuclei
and prominent nucleoli were found in the biopsied tissue
of the patellar OS [47-49].

Surgery is the most effective treatment for most of the
symptomatic patients who have a patellar tumor. Sur-
geons prefer to outline a course of treatment according
to Enneking stages [1]. Some advices are provided as fol-
lows: (1) Stage 1 tumors: only curettage with or without
grafts or excision of the lesion is recommended. (2)
Stage 2 tumors: curettage with adjuvants and grafting
are a better option. (3) Stage 3 tumors: total patellect-
omy is widely used. (4) Stage IA tumors: total patellect-
omy or wide excision can keep the affected knee in good
function. (5) Stage IB tumors: a patellectomy and wide
excision of the tendons and skin concerned is made
available. (6) Stage IIA tumors: resection with a wide
margin is recommended. Total patellectomy with re-
moval of the distal part of the quadriceps tendon, of the
proximal part of the patellar tendon and of the over
patellar fascia, is necessary. (7) Stage IIB tumors: in the
case with minimal invasion of adjacent soft tissue, a wide
resection and adequate reconstruction is preferred. Am-
putation is performed in the case with severe involve-
ment of surrounding soft tissue. (7) Stage 1 or stage 2
tumors: a pathologic fracture can be healed by conser-
native treatment. (8) Stage 3 or stage 1A tumors with a
pathologic fracture and involvement of synovium: a syno-
vectomy or a knee resection is recommended. (9) Stage IB
or stage IIA tumors with a pathologic fracture: a total
patellectomy with wide margins is preferred. (10) Stage
IIB tumors: a pathologic fracture does not change treat-
ment indications. For stage III tumors, the appropriate
cancer surgery intervention combined with radiotherapy or/and
chemotherapy are rational because of regional invasion or
metastases. Moreover, according to different characteri-
Zation of patients, suitable surgical options may range from
simple curettage with bone grafting, excision, patellectomy,
extensor mechanism reconstruction, and knee resection to above-knee amputation. If curettage is performed, prevention of tumor spread in the joint should be considered. If the external shell of reactive bone cannot be kept according to preoperative radiographic results, a patellectomy is better than a curettage. In order to preserve functional function, surgeons should keep intact the extensor apparatus or reconstruct extensor structure. Neoadjuvant chemotherapy has been used as a potential treatment for some primary malignant tumors. Radiotherapy should be chosen to treat highly radiosensitive tumors and benign tumors after incomplete excision. For certain postoperative tumors (stage 3), radiotherapy was unsuitable because of radiation-induced malignant degeneration [6]. Now, radiotherapy was only used for multiple malignant lesions, especially hematologic and vascular malignancies [1,20].

Conclusions
Although anterior knee pain seems simple, the primary tumors originating in the patella as a possible lesion cannot be neglected. More types of unreported primary patellar tumors will be confirmed. Due to the insufficient availability of the patellar characteristics, the integrated diagnosis of primary patellar tumors should be seriously considered. The extended application of MRI and CT is necessary. And noninvasive biopsy is worth studying in order to avoid contamination. While experts are inclined to outline a course of treatment for primary patellar tumors according to Enneking stages, new adjuvant therapy and improved methods of functional reconstruction may do more good to the patients.

Abbreviations
ABC: aneurysmal bone cyst; AP: serum alkaline phosphatase; CB: chondroblastoma; CH: chondroma; CHOP: cyclophosphamide; doxorubicin, vincristine, and prednisone; CRP: C-reactive protein; CS: chondrosarcoma; CT: computed tomography; EC: enchondroma; ESR: erythrocyte sediment rate; GCT: giant cell tumor; JC: juxtacortical; M: malignant fibrous histiocytoma; MRI: magnetic resonance imaging; OB: osteoblastoma; OC: osteochondroma; OH: osseous hemangioma; OD: osteoid osteoma; OS: osteosarcoma; POL: primary osseous lymphoma; SBC: simple bone cyst; WBC: white blood cell.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
SMZ searched for the papers, extracted the research information from those papers, and drafted the manuscript. LM and WYX revised the manuscript. MK and ZZ checked the structure of the manuscript. All the authors read and approved the final manuscript.

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References
1. Mercuri M, Casadei R. Patellar tumors. Clin Orthop Relat. 2001;163:35–46.
2. Yoshida Y, Kojima T, Taniguchi M, Osaka S, Tokuhashi Y. Giant-cell tumor of the patella. Acta Med Okayama. 2012;66:73–6.
3. Olufogun O, Donthnierei R. Iatrogenic seeding of a giant cell tumor of the patella to the proximal tibia. Clin Orthop Relat Res. 2007;465:260–4.
4. Malhotra R, Sharma I, Kumar V, Natarej A. Giant cell tumor of the patella and its management using a patella, patellar tendon, and tibial tubercle allograft. Knee Surg Sports Traumatol Arthrosc. 2010;18:167–9.
5. Chakraverty G, Chakravety U. Late presentation of G.C.T. of the patella. Internet J Orthopedic Surgery. 2009;12:1.
6. Okada K, Sato K, Abe E, Kabaoka Y, Miyakozki N, Ishikawa N, et al. Case report 858: postradiation osteosarcoma of the patella. Skeletal Radiol. 1994;23:471–4.
7. Achmad Kamal F, Erol U, Hugatalang, Sankani Gormay, Yogi Prabowo, Yanuanso Yanuanso. Primary malignant giant cell tumor of the patella: report of a rare case. Med J Indonesia. 2013;32:238–42.
8. Qureshi SS, Puri A, Agarwal M, Desai S, Janhikhar N. Recurrent giant cell tumor of bone with simultaneous regional lymph node and pulmonary metastases. Skeletal Radiol. 2005;34:225–8.
9. Macdonald D, Foranier V, Cameron J. Multicentric giant cell tumor involving the patella. Can J Surg. 2001;44:222–3.
10. Meyeirding HW. Treatment of benign giant cell tumors by resection or excision and bone grafting. J Bone Joint Surg Am. 1945;27:196–207.
11. Ben Maitigue M, Osman W, Dini F, Boughamroua H, Nawar N, Mtsouni M, et al. Chondroblastoma of bone. Tunis Med. 2012;90:781–8.
12. Rischke B, Engels C, Pietts E, Werner M, Delling G. Chondroblastoma of the patella with pathological fracture. Unfallchirurg. 2000;103:898–902.
13. Mosier JR RP, Brockmole DM, Vinh TN, Knardsford MJ, Aoki J. Chondroblastoma of the patella. Skeletal Radiol. 1988;17:413–9.
14. Rhodanmor J, Wehdi Ramanez S, Mohammad GH. Chondroblastoma of the patella treated with curettage and bone graft: a case report. Med J Islamic Republic of Iran. 2011;25:106–10.
15. Ghekere J, Geuens E, Lateur L, Samson I, Sciot R, Baert AL. Chondroblastoma of the patella with a secondary aneurysmal bone cyst. Eur Radiol. 1998;8:992–5.
16. Tan H, Van M, Yue B, Zeng Y, Wang Y. Chondroblastoma of the patella with aneurysmal bone cyst. Orthopedics. 2014;37:e87–91.
17. Gudi N, Venkatesh Reddy VR, Chhanadan KJ. Chondroblastoma patella presenting as a pathological fracture. Indian J Orthop. 2008;42:1–100.
18. OMarA JWI, Keeling J, Montgomery EA, Aaron AD. Primary lesions of the patella. Orthopedics. 2000;23:328.
19. Ranappa AJ, Lee FY, Tang P, Carlson JR, Gebhardt MC, Mankin HJ. Chondroblastoma of bone. J Bone Joint Surg. 2000;82-A:1140–5.
20. Mercuri M, Casadei R, Ferraro A, de Cristofaro R, Balladelli A, Picci P. Tumours of the patella. Int Orthop. 1991;15:115–20.
21. Kooz Z, Than P. Rare localization of osteoid osteoma in the patella. Pediatr Radiol. 2005;35:929–30.
22. Vailanatos PG, Tilenzoglou AC, Seleitaris SV, Mahera HJ. Osteoid osteoma of the patella: a case report. Knee Surg Sports Traumatol Arthrosc. 2006;14:161–4.
23. Mehdisasab SA, Sarrafan N, Alavieinnejad P. Osteoid osteoma of the patella presenting as chronic anterior knee pain: case report and review of the literature. Pakistan J Med Sci. 2010;26:490–3.
24. Ma K, Zhao HT, Niu XH, Zhang Q. Osteoid osteoma of the patella: report of two cases. Chin Med J (Engl). 2011;124:4096–8.
25. Bulas RV, Hayes CW, Conway WF, Loughran T. Case report 783: osteoid osteoma of the patella. Skeletal Radiol. 1992;21:326–9.
26. Reddy NS, Sathi VR. Primary aneurysmal bone cyst of patella. Indian J Orthop. 2009;43:216–7.
27. Traoré A, Doukouari B, Sie Essoh JB, Mobiart C, Soumaro K. Primary aneurysmal bone cyst of the patella: a case report. Orthop Traumatol Surg Res. 2001;97:221–4.
28. Arroud M, Attaraf K, Chater L, Affifi MA. Article patellae aneurysmal bone cyst in children: a case report. Pan Afr Med J. 2012;1:77.
29. Balke M, Dedy N, Mueller-Huebenthal J, Liem D, Haders J, Hoehler J. Uncommon cause for anterior knee pain - aggressive aneurysmal bone cyst of the patella. Sports Med Arthrosc Rehabil Ther Technol. 2010;2:9.
30. Kumar R, Dasan B, Malhotra A. Aneurysmal bone cyst of the patella: three-phase bone scintigraphic findings and differential diagnoses. Clin Nucl Med. 2000;25:1833–4.

31. Kaymaz B, Ergolu M, Kaymaz N, Ucar M. An uncommon cause of anterior knee pain: patellar chondroma. J Pediatr Hematol Oncol. 2014;36:561–2.

32. Lammot TR. Enchondroma of the patella. A case report. J Bone Joint Surg Am. 1968;50:1230–2.

33. Casadei R, Kreshak J, Rinaldi R, Rimondi E, Bianchi G, Alberghini M, et al. Imaging tumors of the patella. Eur J Radiol. 2013;82:2140–8.

34. de Barra FM, de Melo FC, Rocha AV, dos Santos MR. Patellar osteochondroma: case report. Revista Brasileira de Ortopedia (Engl). 2014;49:206–9.

35. Kim JH, Nam SO, Byun YS, Cho HS, Yoo HS. Intra-articular osteochondroma of the patella. J Korean Orthopaedic Assoc. 2012;47:648–72.

36. Khurana JS, McCarthy EF. Benign cartilage tumors. Diagn Histopathol. 2014;20:190–9.

37. De Coster E, Van Tiggelen P, Shahabpour M, Osteaux M, Opdecam P. Osteoblastoma of the patella. Case report and review of the literature. Clin Orthop Relat Res. 1989;243:216–9.

38. Shen CI, Shih HN, Hsu RW, Hsueh S. Osteoblastoma of the patella: case report. Chang Gung Med J. 2001;24:269–73.

39. Sicard A, Touzard RC, Squalli S. Osteoblastoma of the patella (author’s transil). J Chir (Paris). 1979;116:47–50.

40. Bansal VP, Singh R, Grewal DS, Bannerjee AK. Haemangioma of the patella. Acta Orthop Scand. 1979;50:1230–2.

41. Saglik Y, Uçar DH, Yıldız HY, Dogan M. Unicameral bone cyst of the patella. A case report. Int Orthop. 1995;19:280–1.

42. Wientroub S, Salama R, Baratz M, Papo I, Weissman SL. Unicameral bone cyst of the patella. Orthopedics. 2000;23:1285.

43. Chaudhary D, Bhatia N, Ahmed A, Chopra RK, Malik AC, Singh AK, et al. Osteoblastoma of the patella: a case report. Ups J Med Sci. 2012;117:342.

44. Chida S, Nagasawa H, Okada K, Shimada Y. Osteosarcoma of the patella: a review of 42 cases. Skeletal Radiol. 1989;18:365–71.

45. Cho Y, Kim JD, Chung SH. Osteosarcoma of the patella: biologic reconstruction with allograft. Orthopedics. 2009;32:10.

46. Aoki M, Nishio J, Iwasaki H, Masaki M, Kawakami Y, Nishino T, et al. Osteosarcoma of the patella mimicking giant cell tumor: imaging features with histopathological correlation. Anticancer Res. 2014;34:2541–5.

47. Chida S, Nagasawa H, Okada K, Shimada Y. Osteosarcoma of the patella: a case report. J Orthop Sci. 2012;17:342.

48. Shehadah AM, Haiba MA, Henshaw RM, Lack E. Telangiectatic osteosarcoma of the patella. Orthopedics. 2008;31:808.

49. Goodwin MA. Primary osteosarcoma of the patella. J Bone Joint Surg. 1961;43B:338–41.

50. Ferguson PC, Griffin AM, Bell RS. Primary patellar tumors. Clin Orthop Relat Res. 1997;336:199–204.

51. Singh J, James SL, Kroon HM, Woertler K, Anderson SE, Jundt G, et al. Tumour and tumour-like lesions of the patella - a multicentre experience. Eur Radiol. 2009;19:701–12.

52. Yang ZM, Tao HM, Ye ZM, Lin N, Yang DS. Multivariate analysis of the prognosis of 37 chondrosarcoma patients. Asian Pac J Cancer Prev. 2012;13:1171–6.

53. Kumagai K, Abe K, Tomita M, Osaki M, Taura T, Tsukeyoshi M, et al. Chondrosarcoma of the patella. J Orthop Sci. 2008;13:556–60.

54. Halpert B, Davis JB. Chondrosarcoma: report of two cases. Am J Cancer. 1935;23:784–90.

55. Boyle A, Walton N. Malignant anterior knee pain. J R Soc Med. 2000;93:639640.

56. Chandra A, Eilender D. Uncommon sites of presentation of hematologic malignancies. Case 1: localized primary non-Hodgkin’s lymphoma of the patella. J Clin Oncol. 1999;17:1640–2.

57. Agarwal V, Pal L, Aggarwal A, Misra R. Primary patellar T cell lymphoma: an unusual cause of mononorthritis. J Rheumatol. 2001;28:876–7.

58. Beal K, Allen L, Yahalom J. Primary bone lymphoma: treatment results and prognostic factors with long-term follow-up of 82 patients. Cancer. 2006;106:3652–6.

59. Roger DJ, Uhl RL, Carl A. Malignant fibrous histiocytoma of the patella. Orthopedics. 1994;17:189–92.

60. López-Barea F, Rodríguez-Peralto JL, Burgos-Lizáide E, González-López J, Sánchez-Herrera S. Case report 639: malignant fibrous histiocytoma (MFH) of the patella. Skeletal Radiol. 1991;20:125–8.

61. Belack SS, Schroeder G, Fuchs N, Bacci G, Bauer HC, Mapelli S, et al. Malignant fibrous histiocytoma of bone: a retrospective EMSOS study of 125 cases. Acta Orthop Scand. 1999;70:533–60.

62. Bhagat S, Sharma H, Bansal M, Reid R. Presentation and outcome of primary tumours of the patella. J Knee Surg. 2002;15:212–6.

63. Tam W, Resnick D, Hayhig P, Vaughan L. Intracapsular ganglion of the patella. Skeletal Radiol. 1996;25:588–91.

64. Inoue S, Tanaka K, Sakamoto A, Matsuda S, Tsuneyoshi M, Iwamoto Y. Primary leimyosarcoma of the patella. Skeletal Radiol. 2001;30:530–3.

65. Gorelik N, Dickson BC, Wunder JS, Bleakney R. Ewing’s sarcoma of the patella. Skeletal Radiol. 2013;42:729–33.