Giant Cell-Rich Variant of Extraskeletal Osteosarcoma Above the Patella – A Rare Case Study and Review of the Literature

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Abbreviations: MRI: Magnetic Resonance Imaging; ESOS: Extraskeletal Osteosarcoma; GCRO: Giant Cell-Rich Osteosarcoma; GCR-ESOS: Giant Cell-Rich Variant of Extraskeletal Osteosarcoma

ABSTRACT

Background: The giant cell-rich variant of extraskeletal osteosarcoma is a rare subtype of osteosarcoma, occurring in the extremities of adults prevalently in their sixth decade of life. We report an exceptional giant cell-rich variant of extraskeletal osteosarcoma that was detected above the patella in an adolescent patient. The exact histological criteria for giant cell-rich osteosarcoma is not clearly defined, which makes it very challenging to identify the key characteristics for preoperative discrimination between malignant giant cell-rich osteosarcoma and benign soft tissue giant cell tumors. This exceptional case can be a useful example for differential diagnosis of tumors with similar giant cell morphology for further appropriate choices of treatment.

Case Presentation: An 11-year-old Thai girl presented at our hospital with a 10-cm painless swelling in her right knee for three months. On examination, an irregular mass measuring 10x5 cm² in size was detected above her right patella bone along with two 4-cm right inguinal lymph nodes. Radiological investigations and MRI showed the heterogeneous soft tissue mass beneath the subcutaneous fat without any patella bone invasion. Histopathology displayed numerous scattered multi-nucleated giant cells among the medium-size round cells to spindle cells in the background, together with osteoid formation. Abnormal mitoses, multi-nucleation and bizarre nuclei were found. One inguinal lymph node out of the two was positive for metastatic sarcoma. Our final diagnosis was giant cell-rich variant of extraskeletal osteosarcoma. The patient underwent neoadjuvant chemotherapy (EUMOS-1) and a wide resection with a rotational flap. Histopathology reported less than 1% tumor necrosis. The patient remains alive without local recurrence or distant metastasis during the past 13 months.

Conclusion: Giant cell-rich subtype of extraskeletal osteosarcoma has an extremely rare incidence, as well as considerable variations in histological entities. As a result, it has a very poor prognosis and possesses a high risk of misdiagnosis. Our report illustrates an unprecedented rare case of the giant cell-rich variant of extraskeletal osteosarcoma detected above the patella in an adolescent patient. Using a combination of radiography, magnetic resonance imaging, and histological examination, while carefully noticing all points of the clear cutting-criteria for differential diagnosis proposed in this study, a diagnosis of the giant cell-rich subtype of extraskeletal osteosarcoma can be reached. The preoperative differentiation of this exceedingly rare entity is crucial and needs to be examined by musculoskeletal pathologists for early accurate diagnosis and further effective treatment.
Bathurst et al. [1], is a rare variant of osteosarcoma, which accounts for 1-3% of conventional osteosarcomas [2-4]. To date, there have been 22 cases of GCRO reported, and the most common sites found are the femur and tibia [5,6]. Only 2 cases of the giant cell-rich variant of extraskeletal osteosarcoma (GCR-ESOS) were found (Table 1). Considering historical appearance, GCRO contains a large number of benign giant cells similar to those found in benign giant cell tumors. Due to its rarity and histologically close resemblance to giant cell tumors, the interpretation of lesions can be easily mistaken for giant cell tumors [7-9]. The GCRO needs to be differentiated from benign soft tissue giant cell tumors, due to the different treatment strategies that should be provided to patients between the two neoplasms. Preoperative discrimination between GCRO and other tumors with similar giant cell morphology is essential for giving proper diagnoses and further applying proper choice of treatments. In this study, we first report a giant cell-rich variant of extraskeletal osteosarcoma detected above the patella in an adolescent patient that may mimic benign giant cell tumors. The clear-cutting criteria for preoperative discrimination are also discussed.

**Table 1**: Previously reported 22 cases of giant cell-rich osteosarcoma (GCRO) and only 2 cases of giant cell-rich variant of extraskeletal osteosarcoma (GCR-ESOS).

<table>
<thead>
<tr>
<th>Age/Gender</th>
<th>Extraskeletal area</th>
<th>Location</th>
<th>Symptoms</th>
<th>Final diagnosis</th>
<th>Treatment</th>
<th>Recurrence/Metastasis</th>
<th>Outcome/Follow-up</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>19/M</td>
<td>No</td>
<td>Meta-diaphysis of left femur</td>
<td>Motion pain without swelling</td>
<td>GCRO</td>
<td>Chemotherapy, surgical resection, limb salvage surgery with an autogenous autoclaved bone graft, vascularized fibular graft</td>
<td>None/None</td>
<td>NED, 72 months</td>
<td>Sato et al., 1996 (15)</td>
</tr>
<tr>
<td>19/M</td>
<td>No</td>
<td>Diaphysis of right femur</td>
<td>A fracture (6 months before), hypertrophy of the right leg muscles, no peripheral vascular or neurologic deficit</td>
<td>GCRO</td>
<td>Resection, reconstruction of femur with a metallic prosthesis and autogenous fibular graft</td>
<td>Yes (after 13 months)/multiple bone metastasis</td>
<td>NED, 12 months; DWD, 21 years</td>
<td>Bertoni et al., 2003 (2)</td>
</tr>
<tr>
<td>17/M</td>
<td>No</td>
<td>Distal end of the radius</td>
<td>Continuous pain of left wrist joint</td>
<td>GCRO</td>
<td>Operation, Wide resection followed by a vascularized fibular graft, Upper arm amputation</td>
<td>Yes/Yes</td>
<td>DWD, 8 months</td>
<td>Shinozaki et al., 2004 (16)</td>
</tr>
<tr>
<td>29/F</td>
<td>No</td>
<td>Proximal metaphysis of right tibia</td>
<td>Right knee pain</td>
<td>GCRO</td>
<td>Resection, limb salvage operation</td>
<td>None/None</td>
<td>NED, 11 months</td>
<td>Hong et al., 2005 (6)</td>
</tr>
<tr>
<td>29/M</td>
<td>No</td>
<td>Left femur</td>
<td>Knee pain for 2 months without swelling</td>
<td>GCRO</td>
<td>Surgical resection, adjuvant chemotherapy</td>
<td>None/None</td>
<td>NED, 36 months</td>
<td>Gambarotti et al., 2011 (4)</td>
</tr>
<tr>
<td>16/F</td>
<td>No</td>
<td>Left proximal tibia</td>
<td>Left knee pain for 1 year with swelling</td>
<td>GCRO</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>Imran et al., 2012 (8)</td>
</tr>
</tbody>
</table>
An 11-year-old Thai girl presented with a painless swelling (Figure 1) in the right knee for three months. She was eventually diagnosed with a giant cell-rich variant of extraskeletal osteosarcoma. On examination, a 10-cm irregular mass was detected over the right patella along with two 4-cm right inguinal lymph nodes. Radiological investigations showed no bone lesion (Figure 2) and bone scintigraphy also displayed no bone metastases. The magnetic resonance imaging (MRI) of the right knee displayed heterogeneous soft tissue mass beneath the subcutaneous fat without any patella bone invasion (Figure 3). According to the pelvic MRI scan, the MR images showed two enlarged lymph nodes which displayed similar intensity to the soft tissue above the patella, postulating that the two lymph nodes might be positive for malignancy (Figure 4), although histological evidence obtained afterwards suggested that only one lymph node out of the two was indeed positive for metastatic sarcoma. Preoperative open biopsy of the soft tissue above the patella was performed, and the soft tissue was examined by musculoskeletal pathologists.
Histopathology showed numerous scattered multi-nucleated giant cells among the medium-size round cells to spindle cells in the background, along with the osteoid matrix (Figure 5). As a result, the final diagnosis was a giant cell-rich variant of extraskeletal osteosarcoma (GCR-ESOS). Neoadjuvant chemotherapy, including cisplatin, doxorubicin and methotrexate was given prior to a wide resection with a rotational flap, shown in Figures 6 & 7. Histopathology reported less than 1% tumor necrosis. Adjuvant chemotherapy, including cisplatin, doxorubicin, methotrexate with the addition of ifosfamide, and etoposide was given. The patient remains alive without local recurrence or distant metastasis during the past 13 months Figures 8 & 9.
Discussion and Conclusion

Giant cell-rich variant of extraskeletal osteosarcoma (GCR-ESOS) is an exceptional subtype of osteosarcoma, considering that it possesses the combined histopathological features of extraskeletal osteosarcoma and giant cell-rich osteosarcoma. The possibility of the occurrence of either extraskeletal osteosarcoma or giant cell-rich osteosarcoma alone is rare, below 5% of overall osteosarcoma occurrence [2,4,10]. Our report on the GCR-ESOS, which is the combination of both types of osteosarcoma, is therefore considered extremely rare. Due to this extreme rarity, the GCR-ESOS has a poor prognosis and is easily misdiagnosed, which may lead to inefficient treatment. We thus recommend that the interpretation of lesions should be examined by musculoskeletal pathologists. Considering our case, radiological investigation showed a soft tissue mass with mineralization and without any patella bone attachment. The MRI demonstrated nonspecific T-1 hypointense and T-2 hyperintense lesions.

Differential diagnosis can include any other tumors, such as extraskeletal chondrosarcoma, undifferentiated pleomorphic sarcoma, post-traumatic myositis ossificans, sarcomatoid carcinoma and synovial sarcoma. Histologically, the GCR-ESOS contains numerous osteoclast-like giant cells that may obscure the malignant cells in the background. In such cases, the interpretation of lesions may be mistaken for a benign giant cell tumor. For a reasonable decision concerning treatment, GCR-ESOS needs to be clearly differentiated from benign soft tissue giant cell tumors. We recommend that the GCR-ESOS should be best distinguished by histological features: osteoid formation, atypical mitotic activity and multi-nucleation. Treatment strategies can include a wide resection, or alternatively amputation, which will not cause any difference in overall survival. Adjuvant chemotherapy (Euromos-1) resulted in a 5-year survival rate of 66-77%. Radiation is recommended when the resection is marginal or can be used as a palliative treatment.

In this study we discovered, for the first time, an unusual case of giant cell-rich subtype of osteosarcoma detected above the patella in an adolescent patient. This subtype can be characterized by the striking features of an abundance of osteoclastic giant cells and the paucity of the tumor osteoid. Although it is easily confused by benign soft tissue giant cell tumors, due to its possible overlapped histological features, we suggest clear cutting criteria for preoperative discrimination through the observation of histological structure. From the histological point of view, scattering of osteoclast-like giant cells, extensive spindling, atypical mitotic activity, multi-nucleation, and most importantly osteoid formation are the key characteristics for efficient differential diagnosis. On the side of immunohistochemical analysis, mesenchymal tissue markers such as Vimentin have been noted as a potential diagnostic tool for the GCR-ESOS [11]. While MDM2 and CDK4 can be used to differentiate low-grade osteosarcoma from benign mimics [12, 13-21].
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References