INTRODUCTION

Extraskeletal chondrosarcomas are rare tumors counting 2% of all soft-tissue sarcomas. Myxoid and mesenchymal chondrosarcomas are the histologic subtypes. Extraskeletal mesenchymal chondrosarcomas (EMC) are rare than myxoid subtype and commonly involve head and neck. If it involves the extremities, it affects particularly thigh. Wrist is a rare location for EMC. EMC has two peaks. First peak is in 3rd decade, commonly involving head and neck. Second peak is in the 5th decade and it occurs most frequently in the thigh.

Published reports about EMC are mostly based on conventional radiography, tomography and conventional magnetic resonance imaging (MRI) findings. The largest studies evaluating imaging features of EMC were reported by Hashimoto et al in 10 patients and Shapero et al in 7 patients. To the best of our knowledge we did not recognize any report of MRI with dynamic contrast-enhanced MRI so far.

CASE

A 53-year old man presenting with painful swelling of the right wrist was admitted to our hospital 3 years ago. He had been operated for Kienbock’s disease six months before the administration. Physical examination revealed a soft-tissue swelling on the volar surface of the right wrist. Then we performed X-Ray, conventional, dynamic enhanced
and diffusion weighted MRI. The mass did not contain calcification on X-Ray (Fig. 1). T1 and T2 weighted MRI revealed a well defined, lobulated mass within the carpal tunnel, arising from the distal forearm, extending to the carpometacarpal joint, measuring 2.5x3.4x5.7 cm (Figs. 2 and 3). On T1 and T2 weighted images the tumor did not cause any change in the adjacent osseous bony cortex and the medulla. On T1 weighted images the lesion was isointense to muscle (Fig. 2). On T2-weighted images the lesion had heterogenous intermediate signal intensity (Fig. 3). The heterogenous signal intensity on T2 weighted images and deep location of the tumor suggested malignancy. There weren’t any necrosis in the tumor. The T1-weighted fat-saturated volumetric interpolated gradient echo sequence was obtained for dynamic contrast enhanced MRI. Contrast enhanced images were subtracted from the precontrast scan on the console. Subtracted images depicted peripheral enhancement at early arterial phase (Fig. 4). In the late phase the central portion of the mass did not enhanced (Fig. 5). Time-signal intensity (TSI) curve showed a steep rise to an early peak followed by slightly washout and it suggested malignancy (Figs. 6A and 6B). Single shot spin echo planar imaging sequence at b values of 0, 500, and 1000 s/mm (2) were obtained for diffusion-weighted MRI and the apparent diffusion coefficient (ADC) map of the lesion. The peripheral portions of the mass were hyperintense at all b values and the mass showed low ADC values on ADC maps corresponding with restricted diffusion of the peripheral portions of the mass (Figs. 7A and 7B). Mean ADC value was 0.98 ± 0.06 x10^-3 mm^2/sec. On diffusion-weighted MRI, the areas showing restricted diffusion corresponded with the enhancing areas on dynamic-contrast enhanced images.

The differential diagnosis of the tumor includes; synovial sarcoma, malignant fibrous histiocytoma which has more recently classified as pleomorphic undifferentiated sarcoma. Extraskeletal chondrosarcoma and osteosarcoma were less likely in the differential diagnosis because no matrix calcification was found in the mass. Patient underwent gross total resection of the tumor. The specimen demonstrated 2.3x3x6 cm lobulated solid mass including hemorrhage. Microscopically, lesion showed atypical pleomorphic mesenchymal cells with increased mitotic activity, chondroid matrix and necrosis (Figs. 8 and 9). The mass did not contain calcification on X-Ray and histology. Non-calcified chondroid matrix was found on histology. The final diagnosis was a grade 3 EMC. No bony involvement was identified at histological assessment. He received radiotherapy after surgery. After three years of surgery local recurrence was occurred.
Figure 3. On a coronal T2-weighted MRI, the mass had heterogeneous intermediate signal intensity. The low T2 signal intensity area within the central portion of the mass may due to chondroid matrix with high collagen components.

Figure 4. Subtracted images at early arterial phase showed peripheral enhancement pattern of the mass.

Figure 5. Subtracted images at delayed phase showed no central enhancement pattern.

Figures 6A and 6B. Time to signal intensity curve showed a steep rise to an early peak followed by slightly washout.
Figures 7A and 7B. Diffusion weighted images at 1000 b value and ADC maps shows restricted diffusion of the peripheral portions of the mass

Figure 8. (H&E, X200) grade 3 chondrosarcoma characterized by atypical pleomorphic mesenchymal cells with prominent nucleolus

Figure 9. (H&E, X400) High power view reveals increased mitotic activity, pleomorphism and necrosis in the dedifferentiated areas of chondrosarcoma

DISCUSSION

EMC are high grade malignant tumors that most frequently affect head and neck and also thigh. Wrist is a very rare site for extraskeletal mesenchymal chondrosarcomas. X-Ray findings of these lesions are nonspecific soft tissue masses which may include chondroid matrix mineralization. Imaging findings of EMC were rarely reported and there weren’t any specific findings. EMC shows typically intermediate signal intensity on T2 weighted MR images. Some of the cases showed lobulation in the previous reports similar with this case. Peripheral enhancement pattern on contrast-enhanced CT images were also described. The gadolinium enhancement pattern of EMC was very rarely reported. Inhomogeneous enhancement pattern and contrast enhancement except low signal intensity areas on T1 weighted MRI were previously described. So far to the best of our knowledge
the dynamic enhancement pattern of EMC have not been reported previously. According to our case early arterial rapid enhancement and wash-out pattern of the mass may suggest a malignant process. The differential diagnosis of the tumor includes; synovial sarcoma and malignant fibrous histiocytoma which has more recently classified as pleomorphic undifferentiated sarcoma. Synovial sarcomas showed typical intermixed areas of low, intermediate and high signal intensity defining as ‘triple sign’ on T2 weighted images. Our case shows predominantly intermediate signal intensity on T2 weighted images. Pleomorphic undifferentiated sarcoma is more likely in the differential diagnosis with low or intermediate T1 and high T2 signal intensities. Razek et al proposed using a threshold mean ADC value of 1.34×10−3 mm²/sec to distinguish benign soft tissue neoplasms of extremities from malignant neoplasms. We found a mean ADC value of 0.98 ± 0.06 x10−3 mm²/sec which is highly suspicious for malignancy.

In conclusion, EMC is a rare soft tissue tumor that can be unusually origin in wrist and might show peripheral enhancement at early arterial phase and restricted diffusion on MRI. The knowledge of the imaging spectrum of the EMC might help us suggesting the diagnosis.

REFERENCES