Case Report

Osteosarcoma or Ewing’s sarcoma? Radiologist’s Dilemma
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Abstract: Misdiagnosis of osteosarcoma is usual with the condition commonly known as Ewing’s sarcoma. It is the second most frequent primary malignant bone cancer, after osteosarcoma. We report a case of 13 year old boy who radiologically presents all features of Ewing’s sarcoma but histopathologically comes out to be osteosarcoma.

Keywords: Osteosarcoma, Ewing’s sarcoma, Distal femur, Bone neoplasias.

INTRODUCTION
There was a broad spectrum of misdiagnoses between ewing’s sarcoma and osteosarcoma. Symptoms that might aid diagnosis include early local volume increase and the presence of fever. Early symptom of osteosarcoma and Ewing’s sarcoma is local pain. This case report is helpful in identifying signs, symptoms and imaging features of both osteosarcoma and Ewing’s sarcoma that could be used in differentiating the two types of tumors, providing information and impetus for shorten the delay to diagnosis which ultimately results in improved prognosis.

CASE REPORT
A 13-year old boy with limping, staggering gait and pain and soft tissue swelling in the lower thigh region for 20 days i.e. distal femur, referred by an orthopaedician, presented in our department. X-ray (Fig. 1) and CT scan (Fig. 2) revealed diffuse circumferential cortical thickening and associated presence of saucerization of the cortical margin and interrupted type of periosteal reaction along the posterolateral aspect of distal femur as evidenced by sunburst/onion-peel type of reaction with associated Codman’s triangle. MR imaging of the right thigh was performed and T1- T2- and IR images were obtained in the axial and coronal planes (Fig. 3a & 3b). MR study revealed altered signal intensity of the bone marrow of right femoral diaphysis exhibiting hypointense signal on T1W1 & hyperintense signal on T2W1/ STIR images with aggressive sunburst periosteal reaction, codman’s triangle & large soft tissue component.

DISCUSSION
Ewing’s sarcoma and Osteosarcoma are the most common primary malignant bone tumors in children [1]. For tumor diagnosis, focus should be on early signs and symptoms in order to distinguish it from other conditions, such as osteomyelitis and tenosynovitis, and to distinguish different bone tumours from one another [2]. Prognosis is improved by early diagnosis and treatment. The prognosis of Ewing’s sarcoma and osteosarcoma is mainly depend on the local extent of the tumor and presence of metastases [3]. During the last 20 years, new knowledge and studies on bone sarcomas have been presented, and great advances in the treatment of these diseases have been achieved [4]. Limb salvage treatment allows tumor resection while preserving the limb, moreover, modern prostheses preserve limb function also. [5]. Primary osteosarcoma is classified according to site as central (conventional
high-grade and low-grade), intracortical or surface (parosteal, periosteal or high-grade). Osteosarcoma may sometimes be multicentric (osteosarcomatosis) or arise within the soft tissues (extraskeletal). Intramedullary component sometimes shows dense sclerosis. Extraskeletal component occurs after destruction of cortex resulting in typical osseous ‘cloud-like’ matrix mineralization. The extraskeletal mass appears dense and prominent due to periosteal reaction, which typically shows ‘sunburst’ appearance. Reactive Codman’s triangles may be seen at the margin of the lesion [6]. Although MRI contributes little help in the diagnosis of the variety of the tumour, it is useful for local staging of the tumour. Both intramedullary and extraskeletal extension are clearly demonstrated, as is extension into the adjacent articular joint. Skip metastases, if any can be identified and therefore it is very essential to image the entire bone [7]. Ewing’s sarcoma accounts for approximately 5% of biopsied primary tumours and, along with osteosarcoma, represents 90% of primary malignant bone tumours in children [8]. 75% of patients are between 5 and 15 years of age [9]. Localized pain and swelling are the most common chief complaints. Diaphysis (35%) or the metadiaphyseal region (59%) of a long bone are the most common sites. Bones most commonly affected are the femur and humerus (together 31% of cases), the pelvic bones (21% of cases, most commonly the ilium) and ribs (6.5-8% of cases) [10]. A wide zone of transition with permeative lytic bone destruction is seen. The tumour rapidly extends through the cortex producing a large extraskeletal, sub-periosteal mass. Soft tissue component is clearly appreciated on imaging. There is erosion of the outer cortex, producing so-called ‘saucerization’ of the cortex. Occasionally Codman’s triangle is also seen. A vertical ‘hair-on-end’ type of periosteal reaction is also classical of Ewing’s sarcoma. Other reported features include cortical thickening and rarely bone expansion or pathological fracture. Ewing’s sarcoma may occasionally show a mixed or mainly sclerotic lesion, especially in the flat bones and spine. These lesions may resemble osteosarcoma [11]. CT and MRI are helpful in demonstrating a large extraskeletal component. Intramedullary component is accurately demonstrated and ‘skip’ metastases may be identified [12].

**Fig. 1**: X-ray AP (A), Lateral view (B) and 3D (C) image of right thigh shows interrupted, sunburst periosteal reaction seen involving the distal femoral diaphysis with saucerization along the lateral femoral aspect, obliteration of medullary cavity and soft tissue swelling.

**Fig. 2**: CT A) Axial & B) Coronal image shows aggressive, interrupted, sunburst/onionpeel periosteal reaction & codman’s triangle involving the femoral diaphysis with associated diffuse circumferential cortical thickening & obliteration of medullary cavity with soft tissue extension.
CONCLUSION
Lower femoral diaphyseal permeative destructive, lytic bone lesion with extraosseous extension into the surrounding soft tissue in young child could be Ewings or osteosarcoma. As the age of patient, presenting signs and symptoms and radiological (x ray, CT and MRI) features are common in both the tumors and could be proved only on histopathological diagnosis. Therefore it could be diagnostic dilemma for the radiologist and osteosarcoma should always be put in the differential diagnosis of Ewing’s sarcoma.

REFERENCES
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