Atypical Presentation of Primary Ewing’s Sarcoma

Abstract
Ewing’s sarcoma is the second most common bone tumor in children and adolescents. Its annual incidence is 0.6/million total population worldwide, and in some populations in India, its incidence can account for 15%–19.4% of the primary bone tumors. Ewing’s sarcoma is a highly malignant round cell tumor, affecting the metaphyseal and the diaphyseal regions of the long bones. Spine involvement, as a primary focus in Ewing’s sarcoma is uncommon, but when present, usually affects the sacrum. Origin from non-sacral region is extremely rare. We report a case of acute onset of paraplegia progressing to quadriplegia in a child with metastatic involvement of the spine, primary focus being the nonsacral spine which is inherently challenging to an early diagnosis. Ewing’s sarcoma though has improved survival with chemotherapy and radiotherapy, primary spine focus has poor prognosis.

Keywords: Acute onset paraplegia, cord compression, Ewing’s sarcoma, malignant bone tumors of childhood, spine involvement

Introduction
Ewing’s sarcoma is the second most common bone tumor in children and adolescents with its incidence being 15%[1,2] of the primary bone tumors.
Ewing sarcoma, a malignant tumor, though has improved survival postadvent chemotherapy, certain primary sites of involvement especially the spine carry poor prognosis. Nonsacral origin of primary Ewing’s sarcoma has an inherent low survival, which is further complicated by its rarity and late presentation presenting challenges to diagnosis and treatment.

Case Report
A 16-year-old boy presented to us with history of acute onset of weakness of bilateral lower limbs progressing over 24 h. There was no history of prior trauma or associated fever. On examination, there was weakness of both lower limbs with extensor plantar response. Magnetic resonance imaging spine revealed multiple space-occupying lesions of the spine with loss of cervical lordosis and significant spinal cord compression at the thoracic D4-D7 level [Figures 1 and 2].
Emergency decompression of the spinal cord was done with laminectomy at D4-D7 level and excision of the thoracic lesion. Histopathology of the lesion showed small round cell tumor suggestive of primitive neuroectodermal tumor of the bone/Ewing’s sarcoma. Immunohistochemistry confirmed the diagnosis of Ewing’s sarcoma. His lower limb weakness after the decompressive laminectomy did not show any improvement, and over the next 2 days, he further progressed to develop upper limb weakness as well. Local radiotherapy was started followed by adjuvant chemotherapy with four-drug regimen. However, despite this multidisciplinary approach, he succumbed to the illness after the first course of chemotherapy, within 2 months of the onset of symptoms.

Discussion
Ewing’s sarcoma accounts for 15%–19% of primary bone tumors mainly occurring between 10 and 20 years of age and affecting males more than females.[3]
Ewing’s sarcoma is a tumor of the growing bones, common location being the metaphyseal plate and the diaphyseal regions of the long bones. Long bone involvement classically presents with local pain and swelling. It rarely involves the spine with a 4% incidence;[4] where then, the most common location is that of the sacrum. Nonsacral location is rare, with silent multifocal involvement of the spine being rarer. In these cases, features of

Address for correspondence:
Dr. Nancy V Jeniffer,
Assistant Professor,
Department of Paediatrics,
M S Ramaiah Medical College and Hospital, Bengaluru, Karnataka, India.
E-mail: dollyvijiyan@yahoo.com

Access this article online
Website: www.ijmpo.org
DOI: 10.4103/ijmpo.ijmpo_71_17
Quick Response Code:
spinal cord compression usually occur late in the course of the disease and predispose to unfavorable outcomes due to delayed diagnosis.

This child presented in the later stages of the disease with metastatic lesions of the spine with compression and cord changes at various levels. He was asymptomatic before the onset of his weakness of lower limbs sparing nonspecific complaints of fatigue and neck pain after long hours of reading.

Multidisciplinary approach in the treatment involving surgical decompression, chemotherapy, and radiotherapy are necessary to improve chances of survival.[9]

Laminectomy is an effective approach for tumors invading spinal cord from paravertebral soft tissues through the intervertebral foramen as was done in this case. Anterior decompression is the preferred intervention for tumors involving spinal cord from the body of the vertebra.

Radiotherapy is administered in cord-tolerant low doses and should always follow decompression in such cases to prevent posttreatment edema of the spinal cord and worsening of deficits.

Although chemotherapy has advanced in providing up to 75% survival in localized lesions, metastatic lesions still do have poor prognosis.[9] Chemotherapy which is usually given in three–four-drug regimen is to help restrict further spread, prevent micrometastasis, and also decrease the bulk of the tumor. The four-drug chemotherapy classically followed consists of vincristine sulfate, dactinomycin, and cyclophosphamide and doxorubicin hydrochloride.[7]

The other differential diagnosis of a small round cell bone tumor includes Ewing’s sarcoma, rhabdomyosarcoma, neuroblastoma, malignant lymphoma, and primitive neuroectodermal tumor of bone.[7]

Ewing’s sarcoma/malignancy should be considered in the differential diagnosis of any child who presents with acute onset of paraplegia due to its poor outcome. Spine involvement, as a primary focus in Ewings sarcoma is uncommon, but when present, usually affects the sacrum. Origin from non sacral region is extremely rare. Multidisciplinary approach is crucial to good prognosis and survival.

Financial support and sponsorship
Nil.

Conflicts of interest
There are no conflicts of interest.

References