Case Report

A Fatal Case of Primary Osteogenic Sarcoma of Cervical Spine: A Rare Entity

Authors
Dr Rachana Swapnil Binayke, Dr Shantilal. M Sisodia, Dr Banasri Devi
Department of Pathology, Grant Government Medical college And Sir JJ Group of Hospitals
Byculla, Mumbai-400008 Maharashtra, India
Corresponding Author
Dr Rachana Swapnil Binayke
Address- 1504, Shree Datta Tower, Dr.Walimbe Road, Behind KEM Hospital, Parel.Mumbai-400012
Email: rbinayke@gmail.com, Telephone:+91-9987595300

ABSTRACT
Primary osteosarcoma of cervical spine is a rare condition with poor prognosis. The clinical, radiological as well as histopathological features of osteosarcoma vary widely, often leading to diagnostic difficulties. The report highlights the rarity of osteosarcoma of the cervical vertebra wherein the radiological and clinical findings suggested an infective pathology and histological findings remained the gold standard for final diagnosis of neoplastic etiology. A 21-year-old female presented with pain in the nape of the neck and tingling numbness in left upper & lower limb. On examination patient had quadriplegia. Radiological investigations were suggestive of infective etiology most likely Koch’s causing involvement of C3-C5 vertebrae with prevertebral, paravertebral, parapharyngeal and intraspinal infective soft tissues abscess. She underwent C4 laminectomy with excision of the lesion and C1-C7 fixation with titanium Hartshil. The patient eventually died in the post-operative period. The intraoperative squash diagnosis was reported as a spindle cell tumor with malignant potential. The subsequent neuropathological diagnosis of C4 vertebral tumor on paraffin sections revealed characteristic features of osteogenic sarcoma (osteoblastic variant) showing osteoblastic giant cells and neoplastic growth of atypical spindle shaped cells arranged in sheets and trabeculae laying down mineralized osteoid and permeating the adjoining host bone. Early detection and accurate diagnosis is important for improving not only patient prognosis but also the quality of life. We should always consider this rare entity, particularly in young patients who present with intractable neck pain.

Keywords: Primary osteogenic sarcoma, cervical vertebra, young female, death.

Introduction
Osteogenic sarcoma is one of the most common malignant tumors of bone arising frequently in extremities and only rarely involve spine. Primary vertebral osteogenic sarcoma accounts for only 0.85-3% of all osteogenic sarcoma and 3.6-14.5% of all primary spinal tumors. Most osteogenic sarcoma arising denovo are located in metaphysial area of long bones particularly lower end of femur, upper end of tibia and upper end of humerus. The mean age of occurrence is 38 years for axial osteogenic sarcoma than its appendicular counterpart which occurs predominantly between 10-25 years of age. Osteogenic sarcoma is a high grade tumor with poor prognosis. Overall the prognosis of cervical osteogenic sarcoma is worse...
when compared to osteogenicsarcoma at conventional sites. It is a locally aggressive tumor treated with chemotherapy, radiotherapy and surgical resection. Due to rarity of spinal osteogenicsarcomas we report a case of 21 years old female who presented with primary osteogenic sarcoma of cervical spine.

Case Summary
A 21 years old female presented with pain in the nape of neck and tingling numbness in left upper and lower limb since 7 months. On examination patient had quadriplegia. Radiological investigations were suggestive of infective etiology most likely Koch’s causing involvement of C3-C5 vertebrae with pre vertebral, paravertebral, parapharyngeal and intraspinal soft tissue abscess. [Fig 1&2]. She underwent C4 laminectomy with excision of the lesion. The patient eventually died during postoperative period.

Pathological Findings
The intraoperative squash diagnosis was reported as spindle cell tumor with malignant potential. The subsequent neuropathological examination of C4 vertebral tumor on paraffin sections revealed characteristic features of osteogenic sarcoma showing neoplastic growth of atypical spindle shaped cells arranged in sheets and trabeculae lying down mineralized osteoid and permeating the adjoining host bone. [Fig 3(a, b, c, d)]

Fig 1: Xray cervical spine showing Osteolytic lesion of C4 vertebra causing cord compression

Fig 2: MRI spine showing collapse of C4 vertebra and intraspinal extension

Fig 3(a)

Fig 3 (b)
DISCUSSION
Primary spinal osteosarcomas are rare, mostly located in the sacral region followed by lumbar and thoracic segment of spine (3). Cervical segment being the least common site needs high degree of suspicion for diagnosis. Clinically osteosarcoma almost always present with insidious onset neck pain which progressively become intractable and is associated with neurological deficit in more than two third of patients. Age distribution of primary vertebral osteosarcoma is bimodal. The mean age of occurrence is 4th decade for axial osteosarcoma which is two decade later than its appendicular counterpart. A second smaller peak in incidence of the tumor occurs in patient older than 50 years (4). Patients presenting with non mechanical back pain along with rapid neurological deterioration should be thoroughly evaluated for primary spinal malignancy. Clinical examinations and radiological investigations are complimentary but histopathological examination remains gold standard in the establishment of diagnosis. Though radiology plays an important role in establishment of diagnosis, however imaging studies alone may be misleading at times. Plain radiograph, computed tomography, magnetic resonance imaging, angiography and dynamic bone scintigraphy are used for diagnosis and evaluation of tumor extent. Plain radiograph and CT shows a mixture of osteoblastic and osteolytic areas. However purely osteolytic (moth eaten appearance) and pure sclerotic osteosarcoma (ivory, dense cloudlike appearance) are also been reported. The differential diagnosis for a lytic expansile lesion on a cervical spine radiograph in this age group include aneurysmal bone cyst, giant cell tumor, eosinophilic granuloma, osteoblastoma, fibrous dysplasia and osteosarcoma (5). So conclusive diagnosis is based on histopathology. Tumor commonly involve spinal canal as a soft tissue mass which makes cross sectional imaging like CT and MRI essential. In many parts of developing world tuberculosis is the most common cause of vertebral body infection with majority of cases aged around 20 years, and it is difficult to differentiate atypical tuberculous lesion of vertebra from neoplastic process radiographically and clinically (as in our case). Osteogenicsarcoma represent a heterogeneous group of tumor with different histologic features. Conventional osteogenicsarcomas are intramedullary and high grade besides this another group of low grade osteogenicsarcomas consisting of low grade intramedullary and various forms of surface osteogenicsarcoma. (6) According to World health organization bone osteosarcoma is currently classified as follows; conventional, telengectesic, small cell, low grade central, secondary, parosteal, periosteal, high grade surface (7). Majority of tumor shows osteoid matrix mineralization and few shows lytic lesion. Conventional osteosarcoma is the classic form of
osteosarcoma and traditionally according to the predominant type of extracellular matrix produced it is further subdivided in to osteoblastic, chondroblastic and fibroblastic subtypes\(^{(3)}\). The tumor cells are often highly anaplastic with pleomorphic and hyperchromatic nuclei and are spindle shaped. Poor prognosis is due to location and proximity of vital structures.

**Conclusion**

Early detection and accurate diagnosis is important for improving not only patient prognosis but also quality of life. This rare entity should be considered particularly in young patients who present with intractable neck pain.

**Disclosure** – The authors report no conflict of interest.

**Funding** - Nil

**References**