Enchondroma of the cervical spine in young woman: A rare case report

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ABSTRACT

Enchondroma is a type of benign cartilaginous bone tumor. Enchondroma of the spine is very rare. There are only a few cases of enchondromas located in the lamina of the cervical spine have been reported. Therefore, we report a case of enchondroma in the cervical spine. A 24-year-old female patient presented with a history of neck pain, restriction of neck movement, pain and numbness along the right scapula, and weakness accompanied by wasting of the right hand. Presumptive diagnoses included bony tumors such as aneurysmal bone cyst or a giant cell tumor. Radiologic examinations revealed a round tumor in the right lamina of C4 with extracortical extension and foramen of C4-5. C4 right hemilaminectomy and facetectomy were performed with near complete removal of the tumor. On histological examination, the tumor was confirmed to be an enchondroma. At the 6-month follow-up, a computed tomography scan showed no recurrence with good alignment.

Key words: Bone tumor, cervical spine, enchondroma, lamina

Introduction

Chondromas are benign cartilaginous tumors. These are classified as enchondromas that arise within the medullary cavity and periosteal chondromas that arise on the surface of the bone. Enchondroma is a rare benign bone tumor where it commonly involves the long tubular bones. Involvement of the spine is also rare that enchondroma of the spine has rarely been reported. To our knowledge, only 11 cases of chondromas in cervical spine have been reported in the literature since 1960 [Table 1]. We report a case of enchondroma that was located in the right lamina of the fourth cervical vertebra in a young woman.

Case Report

A 24-year-old female patient presented with a history of neck pain, restriction of neck movement, pain, and numbness along the medial border of the right scapula, and weakness accompanied by wasting of the right hand.

In imaging studies, a computed tomography scan revealed a lobulated margined soft-tissue lesion measuring approximately 15 mm in the right lamina of C4 with extracortical extension to the right epidural space and neural foramen of C4-5. The lesion showed widening of the right neural foramen of C4-5 and indentation to the thecal sac along calcifications. On magnetic resonance imaging, the lesion showed slightly heterogeneous signal intensity on T2-weighted images, and iso-signal intensity on T1-weighted images [Figure 1a and b]. The radiologist diagnosed a suspicious aneurysmal bone cyst or a giant cell tumor. Surgery was performed. The fascia was incised along the skin incision and the paravertebral muscles were stripped away from the spinous processes and lamina of C4 on the right side. A hard irregularly shaped tumor mass was seen. C4 right hemilaminectomy and partial facetectomy were performed. The tumor mass was almost completely removed [Figure 2a and b].

Microscopic examination revealed multiple fragmented masses that were composed of multiple cartilage islands separated by fibro-osseous septa. The tumor cells were arranged in small clusters in nodules. The chondrocytes, with bland-looking nuclei, were situated within sharp-edged lacunar spaces (inlet) [Figure 3a and b]. The histological diagnosis was enchondroma.
Chondroma is a type of benign bone tumor that is a well-differentiated hyaline cartilage tumor. It occurs more frequently in patients between third and fifth decades of life. Enchondroma of the vertebral body with neural compression has rarely been reported. Therefore, enchondromas are frequently asymptomatic and are discovered incidentally. When they are symptomatic, spinal enchondromas present with signs of nerve root or spinal cord compression, local pain, palpable masses, and/or pathological fractures. For bones, enchondromas typically arise in metaphysis or diaphysis and may appear roducent, partially mineralized, or heavily mineralized. The pattern of mineralization is characteristic and includes a combination of rings or arcs and stippled radiodensities. The tumor is typically lobular in contour with intermediate signal intensity on T1-weighted images and high signal intensity on T2-weighted images, consistent with hyaline cartilage. Spinal chondromas may be derived from a hyperplasia of immature spinal cartilage with migration outside the vertebral axis or from metaplasia of the connective tissue in contact with the spine or the annulus fibrosus. Enchondroma of the vertebral body with neural compression has rarely been reported. Therefore, enchondromas are frequently asymptomatic and are discovered incidentally. When they are symptomatic, spinal enchondromas present with signs of nerve root or spinal cord compression, local pain, palpable masses, and/or pathological fractures. For bones, enchondromas typically arise in metaphysis or diaphysis and may appear radiolucent, partially mineralized, or heavily mineralized. The pattern of mineralization is characteristic and includes a combination of rings or arcs and stippled radiodensities. The tumor is typically lobular in contour with intermediate signal intensity on T1-weighted images and high signal intensity on T2-weighted images, consistent with hyaline cartilage.
deformity may be present, and the neural foramen may be wide similar to our case if the tumor is intraforaminal.\[7\]

The treatment of choice is excision of the tumor, preservation of vital neural and vascular structures, and maintenance of spinal stability.\[12\] Surgical resection is used to establish a histologic diagnosis, prevent sarcomatous degeneration, and preserve neurologic function. Following complete resection of an enchondroma, the recurrence rate is <10%.\[1,11\] Regarding adjuvant treatments for spinal enchondroma, chemotherapy is ineffective, and radiation therapy is used only for patients with tumors that are not amenable to resection or in cases where surgical margins have been histologically positive for tumor. These lesions are either only slightly responsive or resistant to radiotherapy, and the required high doses of radiation can be hazardous to adjacent nervous tissue.\[1,11,17\] Sarcomatous degeneration is possible, but spontaneous remissions have been reported, particularly in elderly patients.\[16\] Cartilage tumors present challenges in the differentiation of benign and malignant tumors through clinical symptoms or imaging studies. Surgical resection is the treatment of choice and is required for histologic diagnosis. In enchondroma, if only partial excision has been performed, close long-term follow-up is mandatory due to the facts that these tumors have been known to recur. If necessary, adjuvant radiation therapy should be considered to prevent malignant change or recurrence.

**Conclusion**

Enchondroma involving the lamina of the cervical spine rarely occurs. In our case, the tumor was located in the lamina, and this mass invaded the area around the soft tissue and compressed the spinal cord. We initially thought that the lesion was most likely a benign bone tumor and suspected aneurysmal bone cyst or giant cell tumor. However, the pathologic diagnosis was enchondroma. The mass was most likely derived from metaplasia of the connective tissue in contact with the lamina of the spine. Furthermore, enchondromas should be best treated with an aggressive operation to prevent recurrence.

**References**


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