Ossifying Fibroma of the Lumbar Spine: Case Report

Lumbar Spinanın Ossifiye Fibromu: Olgu Sunumu

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ABSTRACT

Ossifying fibroma of the vertebra is a very rare condition. We report a 16-year-old boy with ossifying fibroma of the third lumbar vertebra. Neurological and laboratory examinations demonstrated normal findings. Low-back pain, in the absence of radicular pain, was the presenting symptom. Plain radiography and computerized tomography scanning revealed an osteolytic lesion. The tumor in the third lumbar vertebra was curetted and filled with bone allograft. The histopathological diagnosis was consistent with ossifying fibroma. There was no postoperative complication. The case is the second to be reported. The clinical and radiological aspects of this rare tumor are discussed with the pertinent literature.

KEYWORDS: Ossifying fibroma, Lumbar spine, CT of spine, Fibro-osseous lesion

ÖZ


ANAHTAR SÖZÜKLER: Ossifiye fibrom, Lumbar spina, Spinal CT, Fibro-osseöz lezyon

INTRODUCTION

Ossifying fibromas are fibro-osseous lesion. The true incidence and common locations vary widely among reports. They most commonly occur in the mandible and maxilla (4,5) and also in the long bones of the lower extremities (2,5). The most common sites in the cranium are the frontal and temporal bones (4). Ossifying fibroma of the vertebra is a very rare condition (5). Here we report the second case of an ossifying fibroma of the lumbar vertebra in a 16-year-old boy. The clinical significance and management of this rare lesion are discussed with a review of the pertinent literature.

CASE REPORT

A 16-year-old boy was admitted to our department with low back pain. The patient had low back pain for 18 months and the pain had increased in the last three months. Neurological and laboratory examinations demonstrated normal findings. The plain lumbar radiography revealed an osteolytic lesion involving the left part of the L-3 vertebral body. Lumbar CT scans revealed a soft-tissue density mass of 3.5x2x2 cm. in size at the L3 vertebral body involving the median and posterior part of the corpus and destructing the posterior cortex, and showing minimal extension into the spinal canal with perilesional sclerosis (Figure 1A). Bone window CT scan showed vertebral body destruction (Figure 1B, C). We recommended en bloc resection of the L3-vertebral body but the parents of this young boy rejected any aggressive surgical intervention. The tumor was then resected subtotally and the curettage cavity filled with allograft bone chips.

Histopathological examination revealed a lesion characterized by fibroblastic stroma involving trabecular bone tissue surrounded by osteoblastic cells. Fibroblastic cells formed a storiform pattern in some areas. Hemosiderin-laden macrophages and a few osteoclastic multinuclear giant cells were also present in the lesion. The histopathological diagnosis was consistent with ossifying fibroma (Figure 2).

The postoperative period was uneventful, and the patient was discharged on the seventh postoperative day. The follow-up lumbar CT scan taken 18 months later showed the residual tumor and microcalcification located at the posterior portion of the L3 vertebrae corpus. No dural sac compression was observed (Figure 3).
Primary spinal neoplasms account for less than 10% of all tumors seen in the spinal column (1). The mean age at diagnosis is approximately 21 years for benign lesions of the spine, and approximately 49 years for malignant lesions (6).

Ossifying fibroma is a very rare lesion, and is more common in females. It appears in patients before the age of 40, with a peak incidence in the third and fourth decades (4). Ossifying fibroma most commonly occurs in the mandible and maxilla (4,5). The frontal and temporal bones are the most common sites in the calvarium (4). It also grows in the long bones of the lower extremities (2,5). These lesions present as a slowly-growing mass that is initially asymptomatic. Symptoms and signs depend on the location of the tumor.

Radiographically, ossifying fibromas are initially radiolucent lesions. They gradually become radiopaque with calcifications. Expansion of involved bone with the preservation of a thin rim of cortex is seen in ossifying fibromas. Dilated vascular channels are seen in skull films. CT scans reveal a well-delineated dense lesion. Scalp and dural feeding vessels are occasionally revealed by angiography. Radionuclide 99m Tc scans also may be positive (4).

Computerized tomography plays a major role in the differential diagnosis of ossifying fibroma from aneurysmal bone cyst, giant cell tumor, osteoid osteoma and osteoblastoma (3).

Osteoid osteoma is a small benign, primary bone tumor composed of atypical bone and an osteolytic nidus surrounded by a thick sclerotic reaction. CT findings of a smooth, round lucent nidus less than 15 mm in diameter, surrounded by a well-defined sclerotic area are pathognomonic. Osteoblastoma generally attacks the neural arch and affects the spinal column in 25-50% of cases. The vertebral corpus is rarely involved. The nidus of the osteoblastoma is often poorly circumscribed and is larger than that of the osteoid osteoma. Surrounding sclerosis is not prominent but may be seen. Paraspinal extension and extension into spinal canal may occur. Aneurysmal bone cyst is an uncommon, benign,
markedly expansile bone lesion that affects a single vertebra. CT shows characteristically marked “aneurysmal” cortical “eggshell” expansion and thinning, enclosing a noncalcified cyst. Giant cell tumors are rare, destructive lytic bone tumors that may involve the vertebral body, pedicle, or neural arch, and pathological fractures may cause pain. CT demonstrates a lytic noncalcified expansile mass that may have a benign appearance, similar to an aneurysmal bone cyst, but the lesion may also expand into surrounding soft tissues in a more aggressive fashion.

Ossifying fibroma is a slow-growing sharply marginated benign tumor. It is composed of a fibrous stroma with varying amounts of woven bone, but has areas of mature lamellar bone at the periphery. It is a part of the spectrum of fibro-osseous lesions that includes fibrous dysplasia. The only feature that separates ossifying fibroma from fibrous dysplasia is the presence of a “capsule” of lamellar bone, surrounding an ossifying fibroma (4).

Histopathological examination shows a cellular fibrous spindle cell growth in whorled or matted pattern. Bony fragments can be observed as in meningioma psammoma bodies. Woven bone spicules circumscribed by lamellar bone development are present. Woven bone appears first matures into lamellar bone. Polarization of widely-spaced parallel birefringent lines and osteoblastic and osteoclastic rim of trabeculae in the lamellar bone are some distinguishing characteristics of lamellar bone. Fibrous dysplasia may be recognized by random birefringence under polarized light. A giant cell reaction, hemorrhage and inflammation may also be observed in fibrous dysplasia (4).

The differential diagnosis of fibro-osseous lesions must include not only ossifying fibroma and fibrous dysplasia but also reactive bone formation. Spicules of bone appear along the lines of trauma with woven bone undergoing complete maturation to lamellar bone in reactive bone formation, and additional endochondral bone formation is also present.

Juvenile ossifying fibroma is a rapidly growing destructive lesion of the maxilla occurring before the age of 15. Extension into the paranasal sinuses and orbit is common. This lesion is histologically identical to the adult tumor, but its aggressive local invasion can be fatal.

Our patient is the second case in the literature with ossifying fibroma of the spine. The first one was reported by Ohyama et al. in 1992 (5). The follow up of our patient is eighteen months, and this is a short follow-up period for these patients.

Total surgical excision is the preferred treatment of ossifying fibroma. Subtotal removal should be considered in extremely large lesions and the lesions of the skull base with excellent outcomes of more than ten years. Radiation therapy for the treatment of these tumors has also been reported but malignant transformation limits this application (4).

REFERENCES