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Fibrous Dysplasia of the Spine—A Case Involving the Polyostotic Form Isolated to the Thoracolumbar Spine

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ABSTRACT

Fibrous dysplasia of the spine in a polyostotic form is very rare, with fewer than 36 cases discussed in the literature and there is no such case in Indonesia that has been reported. The aim of this report is to present a case from Indonesia of polyostotic fibrous dysplasia isolated in the spine. We report a case of a 38-year-old Sundanese man with a 1-year history of progressive back pain and weakness of both lower extremities. There was no history leading to infection and no previous trauma. A physical examination revealed kyphoscoliotic deformity, a café au lait spot, tenderness at the thoracolumbar region, and neurological deficits. Laboratory studies were within normal ranges. Plain radiographs showed lytic lesion and kyphoscoliosis. Magnetic resonance imaging showed an endosteal scalloping, infiltrative process, expansion, and destruction in the vertebral bodies from T2 to L5. The findings of an aggressive destructive process was highly suspicious of a malignant process, relying on differential diagnosis and metastases, plasma cell myeloma, bone tumor and chronic infectious spondylitis. Histology revealed an irregularly oriented osteoid without osteoblastic rimming but surrounded by fibroblastic proliferation with a C-shaped sign. Investigations revealed a diagnosis of polyostotic fibrous dysplasia of the thoracolumbar spine in isolation. The patient underwent T5-S1 stabilization and bone grafting. At 1 year postoperative, the patient was asymptomatic; there was no recurrence and minimal neurological deficit with grade II on the modified McCormick scale. A case of the polyostotic form of fibrous dysplasia of the spine in isolation has never been reported in Indonesia. The extreme rarity of this type of presentation can pose a diagnostic dilemma, and in cases isolated to the spine, surgical treatment with posterior stabilization, decompression, and bone grafting gives a good functional outcome.

Tumors

Keywords: fibrous, dysplasia, polyostotic, spine

INTRODUCTION

Fibrous dysplasia is a benign fibro-osseous lesion of the bone, characterized by the replacement of the medullary component of one (monostotic) or, less commonly, several bones (polyostotic) with fibroblastic tissue.^{1,2} Such tumors are predominantly found in males. Most patients with fibroblastic dysplasia are asymptomatic, but some may present with a pain or swelling on the back. Fibroblastic dysplasia accounts for 7% of benign bone tumors and 2.5% of all bone lesions. It may occur in any bone but most commonly in the proximal femur, tibia, ribs, and skull.³ Polyostotic disease commonly affects the sphenoid, frontal, maxillary, and ethmoid bones; occipital and temporal bone involvement is rare.^{4,5} The spine is involved in 1.4% to 5.5% of fibrous dysplasia lesions.^{6–8} Fibrous dysplasia of the spine in a polyostotic form is very rare, with fewer than 36 cases discussed in the literature, and no such case has

previously been reported in Indonesia. The aim of this report is to present a case from Indonesia of polyostotic fibrous dysplasia isolated in the spine.

CASE REPORT

We report a case of a 38-year-old Sundanese man with a 1-year history of progressive back pain and weakness of both lower extremities. The pain was a stabbing pain, radiating to the trunk and both lower extremities. Initially, the pain was intermittent, then slowly increasing in frequency and severity so that it was interfering with daily life activities. His visual analog scale score was 7 out of 10. The patient was unable to stand and tended to lie down. There was no history of infection or previous trauma. There was no history of family with the same conditions. A physical examination revealed kyphoscoliotic deformity, a large and irregularly bordered café au lait spot, and tenderness at the thoracolumbar



Figure 1. Clinical pictures of a 38-year-old man with kyphoscoliotic deformity and café au lait spot.

region. (Figure 1). His spinal range of motion was limited due to pain. Neurological examination found hypesthesia at levels below the T6 dermatome. Extremity strength examination indicated good strength in all key muscles, except for weakness (4 out of 5) in both lower extremities, especially in dorsiflexion, great toe extension, and plantarflexion. Reflex examination indicated bilateral 3+ hyperreflexia throughout the lower extremities. The Babinski sign was positive in both feet. The straight-leg-raise test was positive at the right leg. He had normal bowel and bladder function.

Laboratory studies were within normal ranges. Plain radiographs showed a lytic lesion and

kyphoscoliosis at the thoracolumbar spine. Magnetic resonance imaging of the thoracolumbar area was carried out and the results showed endosteal scalloping, an infiltrative process, expansion, destruction in the multiple vertebral bodies from T2 to L5 (Figure 2a), and multiple disk bulging with mild cord compression at levels T6-7, T11-12, L1-2, L3-4, and L-L5 (Figure 2b).

The finding of an aggressive destructive process was highly suspicious of a malignant process, relying on differential diagnosis and metastases, plasma cell myeloma, bone tumor, and chronic infectious spondylitis. Histology from core biopsy examination at the lumbar region revealed an

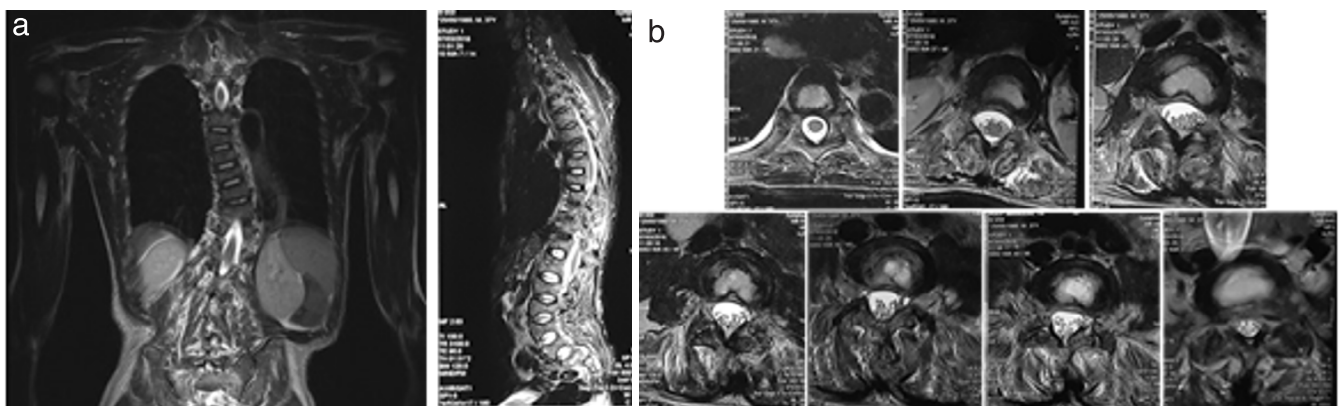


Figure 2. (Left) Magnetic resonance imaging of thoracolumbar spine showed endosteal scalloping, infiltrative process, expansion, and destruction in the vertebral bodies from T2 to L5. (Right) Multiple disk bulging with mild cord compression at levels T6-7, T11-12, L1-2, L3-4, and L4-5.

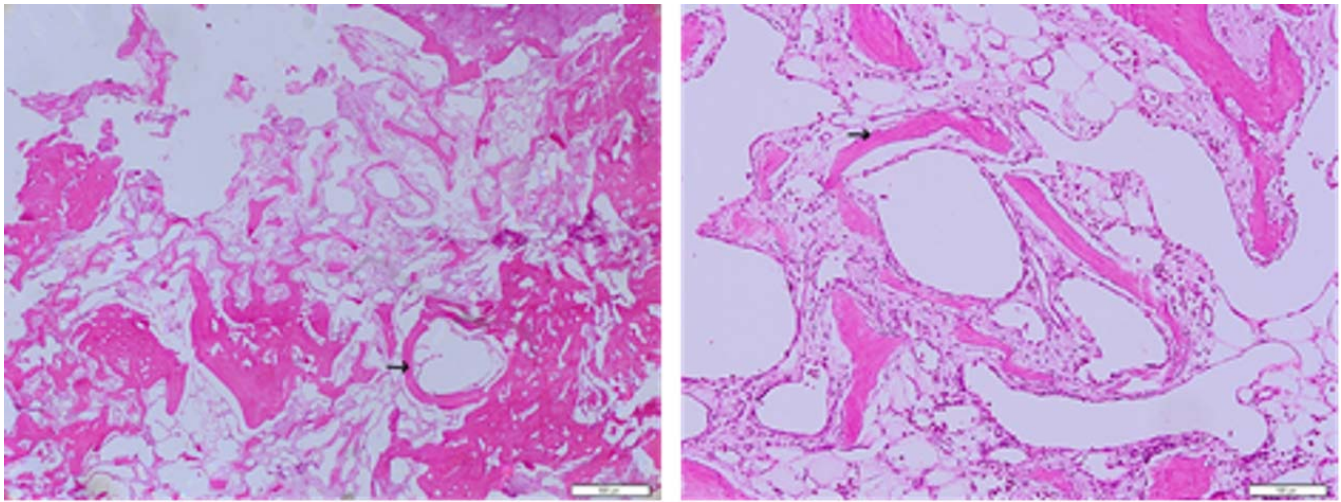


Figure 3. Irregularly oriented osteoid with fibroblastic proliferation and C-shaped sign (black arrow).

irregularly oriented osteoid without osteoblastic rimming but surrounded by fibroblastic proliferation with C-shaped sign (Figure 3). A bone scan investigation revealed an isolated hot spot at the spine. Investigations revealed a diagnosis of polyostotic fibrous dysplasia isolated to the thoracolumbar spine (Figure 4). The patient underwent decompression with partial laminectomy, pedicle screw fixation, and fusion with bone grafting. Partial laminectomy was performed at levels T6-7, T11-12, L1-2, L2-3, L4-5, and L5-S1 under general anesthesia (Figure 5). Pedicle screws were inserted with biplanar fluoroscopy guidance (left T5, T7, T9,



Figure 4. Bone scan revealed isolated hot spot at spine.



Figure 5. The patient treated with T5-S1 posterior stabilization and bone grafting.

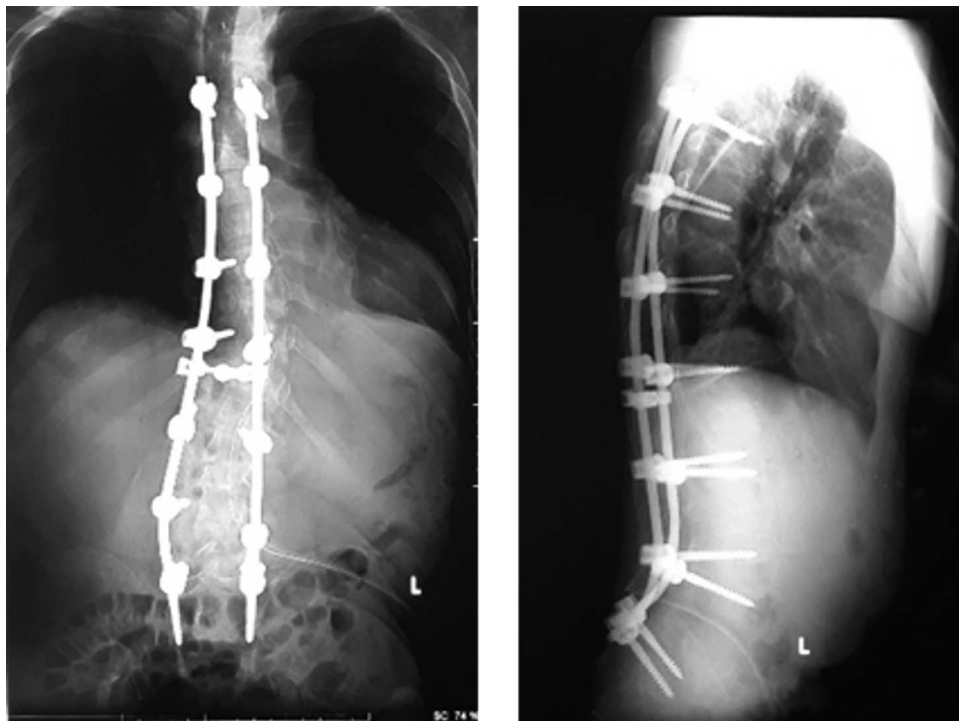


Figure 6. Anteroposterior and lateral radiographs showing the final result after decompression and pedicle screw fixation from T5 to S1.

T11, L1, L3, L4, and S1 pedicles; right T5, T7, T9, T11, L1, L3, L5, and S1 pedicles). The operative time was 210 minutes, and the estimated blood loss was 2000 mL. Two weeks after the operation, the visual analog scale was 3. The pathologic diagnosis of fibrous dysplasia was confirmed by histologic examination. Risedronate sodium was administered orally, 35 mg once a week, in combination with vitamin D and calcium. The patient had recovered from neurologic deficit and back pain at the 3-month follow-up. At 1 year postoperative, the patient was asymptomatic, with no recurrence and

minimal neurological deficit with grade II on the modified McCormick scale (Figures 6 and 7).

DISCUSSION

Fibrous dysplasia was first described by Von Recklinghausen in 1891.⁹ It is a slow, progressive bone disorder characterized by replacement of normal cancellous bone in the medullary canal with immature fibro-osseous tissue.^{10–12} According to the involvement of the bones, it is classified into 2 forms: monostotic and polyostotic. The polyostotic form is rarer than the monostotic variant.¹³ The prevalence of spinal involvement is very low, also the presence of multiple lesions of fibrous dysplasia in the thoracolumbar spine in isolation is rare.³ In the previous study conducted by Leet et al,¹⁴ none of the cases from a large cohort of patients had isolated spinal localization. The clinical features of fibrous dysplasia are usually painless and asymptomatic; when symptoms occur they usually manifest as pain, swelling, and deformity. In our case, the patient had a progressive back pain with weakness in both lower extremities and kyphoscoliotic deformity in thoracolumbar region.^{15–18}

On plain radiographs, it appears as a lytic lesion in the metaphysis or diaphysis with a “ground glass” appearance. In our case, the plain radiographs showed lytic lesions along with kyphoscoliotic



Figure 7. Clinical pictures after 1 year postoperative.

deformity. Magnetic resonance imaging was done to evaluate the soft tissue component and the extent of a lesion. In our case, the results showed endosteal scalloping, infiltrative process, expansion, and destruction in the vertebral bodies from T2 to L5. However, fibrous dysplasia cannot be diagnosed only through radiographic findings. Histopathological examination is needed to rule out the differential diagnoses, which include metastases, plasma cell myeloma, bone tumors, and chronic infectious spondylitis. The lesion may also be complicated by fractures and reparative changes. Hemangioma, giant cell tumor, and aneurysmal bone cyst are also some of the cases of fibrous dysplasia with lytic lesions. Histology revealed an irregularly oriented osteoid without osteoblastic rimming but surrounded by fibroblastic proliferation with a C-shaped sign which corresponded to the previous case reported by Joyce et al.¹⁹

The treatment of fibrous dysplasia was determined by the presence of the symptoms. In the present case, the spinal cord was compressed and stretched by the kyphoscoliotic deformity. Surgery is indicated for a patient with persistent pain, neurologic impairment, vertebral collapse, instability, and/or cord compression.²⁰ We performed a surgical treatment regarding this condition in order to achieve following goals: decompression of the spinal cord, maintenance of spinal stability, eradication of symptomatic lesions, and prevention of further pathological fractures. In our case, the patient underwent a surgical treatment by performing T5-S1 stabilization, decompression, and fusion with bone grafting. One year after the surgery, the patient was asymptomatic, showed no sign of recurrence and minimal neurological deficit with a grade II on the modified McCormick scale.

CONCLUSION

A case of the polyostotic form of fibrous dysplasia isolated to the spine has never been previously reported in Indonesia. The extreme rarity of this type of presentation can pose a diagnostic dilemma, and in cases that are isolated to the spine, surgical treatment with posterior stabilization, decompression, and bone grafting gives a good functional outcome.

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