Case Report
Percutaneous kyphoplasty for vertebral fracture second to polyostotic fibrous dysplasia: a case report and review of the literature

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Abstract: A 26-year-old female patient suffered from intermittent back pain for 4 years and a fall from bicycle aggravated the symptom. Physical and imaging examinations revealed the diagnosis of polyostotic fibrous dysplasia in McCune-Albright syndrome. No neurological symptom was found. A vertebral fracture was seen at the eighth thoracic body. Percutaneous kyphoplasty was performed at the fractured level. The patient got a remarkable pain relief after surgery. As a suggestion, percutaneous kyphoplasty may be a usable choice to treat the patients with vertebral fracture second to polyostotic fibrous dysplasia.

Keywords: Percutaneous kyphoplasty, polyostotic fibrous dysplasia, vertebral fracture, McCune-Albright syndrome

Introduction
Fibrous dysplasia (FD) is an uncommon disorder of bone characterized by progressive replacement of normal bone with fibro-osseous tissue, which may results in pain, deformity and pathological fracture [1, 2]. The disease could affect single bone (monostotic) or several (polyostotic) bones. Polyostotic fibrous dysplasia (PFD) can occur combining with café-au-lait skin spots and precocious puberty, which is known as McCune-Albright syndrome (MAS) [3, 4]. In most cases, FD preferentially affects long bones and craniofacial bones are the most frequently affected [5]. As reported, FD rarely affects the spine in about 2.5% of cases [6]. Herein, we present an even rare case of PFD in MAS involving with thoracic spine, ribs and ilium.

Case report
The patient was a 26-year-old female patient with a history of intermittent back pain. She reported that the pain had lasted for 4 years until a tumble accidently occurred on her two weeks ago, which aggravated the symptom. On questioning, we learned the history of twice vaginal bleeding at the patient’s age of 5. Two light brown-pigmented patches were observed on the back and medial side of her left thigh, which were snatchy and asymptomatic (Figure 1A, 1B). Physical examination revealed conspicuous tenderness and percussion pain in the thoracic region, within T8-9. Neurological examination demonstrated normal status. Visual analog scale (VAS) was 6 before treatment and painkiller had little help.

Computed tomography (CT) of thoracic spine revealed multiple lytic expansile lesions involving several ribs, T2, T3 and T8 (Figure 2). Sagittal and coronal CT showed obvious collapse in T8 (Figure 2B, 2C). Magnetic resonance imaging (MRI) of short tau inversion recovery (STIR) sequence showed abnormal hyperintense of the T8 vertebral body (Figure 3). Fresh vertebral fracture of T8 was confirmed, in line with the location of back pain. Positron emission tomography-computed tomography (PET-CT) suggested hypermetabolism in several ribs, T2, T3, T8 and left ilium. Additional MRI of sacrum demonstrated the lesions of left ilium (Figure 4). A needle biopsy of T8 showed
cellular fibrous matrix, which is consistent with fibrous dysplasia (Figure 5).

All the findings above appeared to fit the classical triad of symptoms of MAS. In case of endocrinopathies, the patient received an all-round endocrine evaluation, which revealed no abnormality.

With diagnosis established, the treatment was discussed. After explaining benefits and risks of different surgical methods, the patient and her family chose percutaneous kyphoplasty, consistent with our option. The bilateral PKP was performed under general anesthesia. After T8 was located with help of fluoroscopy, a trocar and cannula was inserted into vertebral body through each pedicle. Then the trocar was removed and a balloon was inserted and inflated through each cannula. After that, cement was introduced into the fractured vertebral body. No cement leaked during the whole procedure.

After the surgery, VAS was reduced to 0. Patient got a complete pain relief. Radiography after 1 week showed the recovery of height of T8 (Figure 6). In 1 month, 3 months and 6 months, the patient’s follow-up visits showed a good status of her.

Discussion

Fibrous dysplasia is a non-inheritable disorder due to the activating mutations of codon 201 in the GNAS1 gene that encodes for the alpha-subunit of protein Gs [7]. From the perspective of cytology, it is caused by abnormal proliferation and differentiation of bone marrow stromal cells, which leads to the replacement of normal bone with matrix, trabeculae and collagen [8, 9].

Biological variability results in the multiformity of clinical manifestation. In general, FD is asymptomatic, which may be a chance finding by radiography [10]. However, pain, deformity or even fracture could happen in many cases. The typical MAS is classically defined by the triad of PFD, precocious puberty and café-au-lait skin spots [11]. In patients with MAS, there is a high-

Figure 1. Light brown pigmented patches on the back of truncus (A) and inner of left thigh (B).
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er rate of fractures, which is 0.29 versus 0.08 fractures per year, compared with patients with PFD alone [12].

FD has a typical matrix “ground glass” appearance in radiography [13]. Vertebral involvement often presents with lytic expansile lesions. With the help of needle biopsy, the diagnosis can be established. If a vertebral fracture is suspected in FD case, MRI is necessary to check the abnormal hyperintense in STIR sequence [14].

Conservative treatment for FD aims at optimizing function and minimizing morbidity. Conventional medicines consist of calcitonin, etidronate, mithramycin, and bisphosphonates [15]. But all these options are palliative. Surgical treatment is challenging. Methods like curettage, grafting and screws are frequently ineffective [16, 17]. However, aggressive resection with rigid fixation and fusion for FD of spine are supported and proved to be effective by some scholars [18, 19]. Deen et al suggest that balloon kyphoplasty may be an option in patients with painful vertebral compression fracture caused by fibrous dysplasia [20]. Above all, the treatment for FD is complex and individualized.

In our case, the patient was diagnosed with MAS and PFD was involved with her thoracic spine, ribs and ilium. Her main symptom was back pain due to the fractured eighth thoracic body. Previous medications were ineffective. According to radiography, no other fracture was found except T8. Taking the principle of minimal invasion and rapid pain relief into consideration, we chose percutaneous kyphoplasty as a surgical method [21]. With bone cement injected into the vertebra, height of T8 got a remarkable recovery. After the surgery, the patient gained significant pain relief and the VAS score was improved to 0 from 6. No complication sets in during 6 months’ follow-up.

Conclusion

This report presents a rare case of polyostotic fibrous dysplasia in McCune-Albright syndrome involving with thoracic spine, ribs and ilium. The

Figure 2. CT of thoracic spine revealed multiple lytic expansile lesions involving several ribs, T2, T3 and T8. (A) Cross section. Sagittal (B) and coronal (C) CT pictures showed obvious collapse in T8.

Figure 3. MRI of STIR sequence showed abnormal hyperintense of T8.
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A patient suffered from back pain and a vertebral fracture of T8 was found after examinations. By performing PKP to relieve the pain, we suggest it as a usable choice to treat this kind of patients.

Disclosure of conflict of interest

None.

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