Aim: The purpose of the study was to analyze and summarize the computed tomography (CT) and magnetic resonance imaging (MRI) findings of spinal monostotic fibrous dysplasia (MFD) as well as evaluate the clinical value of CT and MRI in MFD diagnosis. Materials and Methods: CT ($n = 4$) and MRI ($n = 5$) images of six patients with pathologically confirmed spinal MFD were examined. The assessed image features included location, shape, rib involvement, vertebral collapse, margin, attenuation, and sclerotic rim on CT, as well as signal intensity, dark signal rim, and enhancement pattern on MRI. Results: In total, four of six patients underwent CT scanning. The most common findings on CT scanning were expansile lesions ($n = 4$), sclerotic rims ($n = 4$), and ground-glass opacity (GGO) ($n = 4$). In total, five of six patients underwent MRI. The lesions were low-signal intensity ($n = 2$), low-to-isointense signal intensity ($n = 1$), and low-signal intensity with several isointense portions ($n = 2$) on T1-weighted imaging (T1WI). The lesions were low-signal intensity ($n = 1$), isointense to high intensity ($n = 1$), and isointense signal intensity with several high portions ($n = 3$) on T2WI. A dark signal rim was found in most cases on T1WI and T2WI ($n = 4$). The lesions ($n = 2$) showed obvious enhancement. Conclusions: The CT and MRI manifestations of spinal MFD have the following characteristics: expansile lesion, GGO, sclerotic rim, and no obvious soft-tissue mass. The combined use of CT and MRI examinations is necessary for patients with suspected spinal MFD. Keywords: Computed tomography, magnetic resonance imaging, monostotic fibrous dysplasia, spine

INTRODUCTION

Fibrous dysplasia (FD) is a benign bone lesion originally described by Lichtenstein and Jaffe in 1942.\(^1,2\) According to the literature, FD represents approximately 7% of all benign tumor-like bone lesions.\(^3,4\) Depending on the extent of involvement, FD can divided into two types: monostotic FD (MFD) and polyostotic FD (PFD).\(^5\) FD is usually found in the ribs, the proximal femur, tibia, humerus, and craniofacial bones.\(^6\) Only 2.5% of FD occurs in the spine.\(^7\) Spinal involvement occurs mostly in PFD, and it is unusual for it to occur in MFD.\(^8,9\) Treatment of spinal MFD depends on the presence and severity of symptoms. Asymptomatic patients with stable lesions only need clinical observation.\(^7,9\) Therefore, a correct preoperative diagnosis of spinal MFD is important to avoid unnecessary surgery.

The purpose of this study was to assess the characteristic computed tomography (CT) and magnetic resonance imaging (MRI) findings of spinal MFD and to evaluate the clinical value of CT and MRI in MFD diagnosis.

MATERIALS AND METHODS

Patients

This study was approved by the institutional ethics committee. Since it was a retrospective study, the sample size was limited by the number of patients with spinal MFD who had undergone imaging. The study included six patients who had undergone CT and/or MRI examinations between 2015 and 2017 at our institution. All patients had been diagnosed with spinal MFD by a combination of clinical symptoms, imaging findings, and pathological confirmation. The study included four male and two female patients, with a mean age of 32 years (range: 16-50 years).

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informed consent was not required. In April 2010 to May 2017, a total of six patients with pathologically confirmed spinal MFD were retrospectively reviewed. The patients included three males and three females (age 22–77 years, median age 48.7 years). The clinical symptoms included backache (n = 3) and numbness of the limbs (n = 1). The remaining lesions were found by routine examination (n = 2). The duration of symptoms ranged from 1 month to 20 years (median, 7 months).

**Imaging techniques**

CT examinations (n = 4) were performed using the standard CT protocol. The imaging parameters were as follows: field of view (FOV) of 200–400 mm, matrix of 512 × 512, thickness of 1.5 mm, and a reconstruction interval of 1 mm.

MRI examinations (n = 5) were performed using a 1.5T MRI scanner (Signa Advantage Horizon; GE Medical Systems, Milwaukee, WI). Both axial and sagittal T1-weighted imaging (T1WI) (500–700 ms repetition time [TR], 15–30 ms echo time [TE], 200–360 mm FOV, 256–512 × 208–512 matrix) and turbo spin echo T2WI (3000–5000 ms TR, 60–90 ms TE, 200–360 mm field of view, 256–512 × 208–512 matrix) were performed on MRI.

Two cases underwent enhanced MRI. Contrast-enhanced T1-weighted spin-echo images with fat saturation were obtained after the intravenous injection of 0.1 mmol/kg of gadolinium dimeglumine.

**Imaging analyses**

All the images were reviewed by two senior radiologists. The reviewers did not know the clinical and histopathological findings of the patients. The criteria for the evaluation of the disease included the lesion location (cervical vertebra, thoracic vertebra, or lumbar vertebra) (vertebral body, vertebral pedicle, or posterior element), shape (circular or irregular), rib involvement (yes or no), and vertebral collapse (yes or no). Margin (well-defined or ill-defined), attenuation (high, low, intermediate, or mixed density), and the sclerotic rim (yes or no) of the lesion were examined on CT. Signal intensity (high-, low-, or intermediate-signal intensity), dark signal rim (yes or no), and the enhancement pattern (mild or obvious) of the lesion were examined on MRI. The signal intensity of the lesion was compared with the spinal cord on MRI. Any differences of opinion were resolved by discussion.

**RESULTS**

**General features of the lesions**

The CT and MRI imaging features of six patients with spinal MFD were summarized in Table 1. In all six cases, the lesions occurred in the cervical vertebra (n = 2), thoracic vertebra (n = 3), and lumbar vertebra (n = 1). In addition to one lesion involving vertebral body, vertebral pedicle, and posterior element, at the same time, the rest of lesions were confined in the vertebral body. The shape of the lesions was regular (n = 4) or irregular (n = 2). One case occurred in the thoracic vertebra with rib involvement. Two cases had vertebral collapse [Figures 1a-c and 2a-d]. The morphology of the lesion was assessed in each case.

**Computed tomography findings**

The most common findings on CT scanning were expansile lesions (n = 4), sclerotic rims (n = 4), and ground-glass opacity (GGO) (n = 4). All lesions were well defined (n = 4) [Figure 3d-e]. All lesions showed low density, and a sclerotic rim (n = 4) [Figure 2a]. The bone cortex was thin (n = 4). The adjacent bone was not involved in any cases (n = 4). No obvious soft-tissue masses were found in any cases (n = 4).

**Magnetic resonance imaging findings**

On MRI, the lesions were low-signal intensity (n = 2) [Figure 2b], low-to-isointense signal intensity (n = 1) [Figure 1a], and low-signal intensity with several isointense portions (n = 2) [Figure 3a] on T1WI. The lesions were low-signal intensity (n = 1) [Figure 1b], isointense-to-high intensity (n = 1) [Figure 2c], and isointense signal intensity with several high portions (n = 3) [Figure 3b] on T2WI. A dark signal rim was found in most cases on T1WI and T2WI (n = 4) [Figure 3a and b]. The lesions (n = 2) showed obvious enhancement on enhanced MRI [Figures 3c and 2d].

**Pathological changes**

Histologically, the lesions had the delicate trabeculae of immature bone, with no osteoblasts. The delicate trabeculae floated in immature mesenchymal cells. Cellular fibrous tissue containing a proliferation of spindle cells could be found, and there were no features of malignancy [Figures 3f and 4].

*Figure 1: A 22-year-old male with monostotic fibrous dysplasia in L3 (original). (a-c) Sagittal T1-weighted imaging, T2-weighted imaging, and T2-weighted imaging–fast spin magnetic resonance imaging showed severe collapse of the vertebral body. Note that the adjacent intervertebral space was not involved.*
Discussion
Overview
FD occurrence in the spine is rare. Although MFD represents 70% of these lesions, involvement of axial bone is mostly seen in PFD. MFD involving the axial bone is extremely rare.[5,6,10,11] To the best of our knowledge, the characteristic radiographic appearances of MFD in long bones have been reported many times in the literature. However, only a few reports have described the imaging features of spinal MFD.[5,7,10,12,13]

The present study reports the characteristic CT and MRI findings of six cases of MFD, with the aim of improving diagnostic accuracy.

Table 1: The results of computed tomography/magnetic resonance imaging findings of spinal monostotic fibrous dysplasia in 6 patients

| Patient/age (years)/sex | CT/MRI image | Site | Shape | Rib damage/vertebral collapse | CT Margin | Attenuation | GGO | Sclerotic rim | TIWI* | T2WI* | Dark signal rim | Enhancement degreea |
|-------------------------|--------------|------|-------|-------------------------------|-----------|-------------|-----|--------------|-------|-------|----------------|---------------------|
| 1/36/female             | Yes/yes      | C6   | Circular | No/no                         | Well defined | Low | Yes | Yes | Low SI with several isointense portions | Isointense with several high SI portions | Yes | Obvious |
| 2/59/male               | Yes/yes      | C7   | Circular | No/no                         | Well defined | Low | Yes | Yes | Low SI | Isointense to high SI | Yes | — |
| 3/77/female             | Yes/yes      | T5   | Irregular | No/yes                        | Well defined | Low | Yes | Yes | Low SI | Isointense to high SI | Yes | Obvious |
| 4/59/male               | Yes/no       | T11  | Circular | Yes/no                        | Well defined | Low | Yes | Yes | — | — | — | — |
| 5/39/female             | No/yes       | T12  | Circular | No/no                         | — | — | — | — | — | — | — | — |
| 6/22/male               | No/yes       | L3   | Irregular | No/yes                        | — | — | — | — | — | — | — | — |

*aCompared with signal intensity and attenuation of the spinal cord. bCompared with signal intensity of the lesion on precontrast T1WI. CT: Computed tomography, MRI: Magnetic resonance imaging, GGO: Ground-glass opacity, SI: Signal intensity, —: nil

Figure 2: A 77-year-old female with monostotic fibrous dysplasia in T5 (original). (a) Axial computed tomography showed an irregular and expansile lesion. (b) Sagittal T1-weighted imaging magnetic resonance imaging showed low-signal intensity and partial vertebral collapse. (c) Sagittal T2-weighted imaging-FS magnetic resonance imaging showed isointense to high signal intensity. (d) Sagittal T1 weighted imaging-FS + C magnetic resonance imaging showed obvious and homogeneous enhancement.

Figure 3: A 36-year-old female with monostotic fibrous dysplasia in C6 (original). (a) Sagittal T1-weighted imaging magnetic resonance imaging showed low-signal intensity with several isointense portions. (b) Sagittal T2-weighted imaging magnetic resonance imaging showed isointense with several high-signal intensity portions. (c) Sagittal T1-weighted imaging + C magnetic resonance imaging showed obvious and heterogeneous enhancement. (d) Axial computed tomography showed an expansile lesion and sclerotic rim. (e) Reformatted sagittal computed tomography showed the sclerotic rim more clearly. (f) Histopathology examination (H and E) showed delicate trabeculae of immature bone with no osteoblasts.
Etiologic and pathological changes

At present, the exact etiology of MFD is not clear. It is well-recognized that the etiology of MFD has been linked to an activating mutation of the Gs alpha gene on chromosome 20, which leads to an increase in cyclic adenosine monophosphate availability.[12] MFD develops during skeletal development.[8] However, the clinical presentation depends on the location of the lesion.[8,12] Pain is usually proportional to the degree of vertebral body involvement.[7] Regarding pathology, MFD is characterized by the replacement of bone marrow with poorly organized spicules of immature bone.[8,10] The combination of a lack of stress alignment and insufficient mineralization results in substantial loss of mechanical strength, leading to the development of pain, deformity, and vertebral collapse.[8] In this study, two cases had vertebral collapse. With the medical history of the patients, the backache both appeared 2 years ago without treatment.

Analysis of computed tomography imaging findings

The CT imaging features of spinal MFD depend on the underlying histopathology of the lesion.[3,5,17] Radiolucent areas are composed of fibrous elements, and radiopaque areas are composed of woven bone.[5] The normal bone is replaced by more radiolucent tissue and has a “ground-glass opacity (GGO)” pattern.[8] Another imaging feature of the lesion is the sclerotic rim. The lesion is bounded by a sclerotic rim of reactive bone. The sclerotic rim is defined more sharply on its inner border than its outer border.[8] In this study, the sclerotic rim and GGO were found in all cases. Spinal MFD has benign biological behavior, and the adjacent intervertebral disk can maintain its normal form. In this study, the adjacent bone was not involved in all cases.

Based on previous descriptions in the literature and the findings of the present study, spinal MFD has the following CT imaging findings: (1) expansile lesion, (2) well-defined margin and sclerotic rim, (3) GGO, (4) thinning of cortical bone, and (5) normal adjacent intervertebral disk.

Analysis of magnetic resonance imaging findings

MRI findings of spinal MFD in this study were nonspecific. Most lesions show low-signal intensity or low-signal intensity with several isointense portions on T1WI and isointense signal intensity with several high portions on T2WI. The cause of the heterogeneous signals may be related to the trabecular bone, cellularity, collagen fibers, cystic changes, and hemorrhage.[3,18] MFD may be surrounded by a layer of thick, sclerotic reactive bone, called a rind.[3] The rind can be seen as a dark signal rim on T1WI and T2WI.[3] Dark signal rims were found in four cases in this study. Two cases underwent enhanced MRI. The lesions showed obvious enhancement. Enhanced MRI can provide useful information regarding the quality of the blood supply and help to predict whether the lesion is benign.

Choice of imaging method

CT and MRI imaging have respective advantages in the diagnosis of spinal MFD. CT imaging can

Figure 4: Histopathology examination (H and E) of a 59-year-old male with monostotic fibrous dysplasia in T11 (original). Histopathology examination showed delicate trabeculae of immature bone with no osteoblasts. The mesenchymal stroma surrounding the dysplastic trabeculae was relatively hypocellular and was composed of spindle-shaped primitive mesenchymal cells.
help evaluate the destruction of bone and provide accurate anatomical information to guide surgery.[3,5,7] MRI imaging can show the spinal cord and internal structures of the lesion more clearly. In summary, the combined use of CT and MRI examinations is necessary for patients with suspected spinal MFD. It should be mentioned that it will be quite difficult to diagnose spinal MFD with imaging studies when the vertebral body is collapsed.[3,11] In these cases, pathological examination should be taken into account.

### Differential diagnosis

The imaging findings of MFD are similar to other benign bony tumor-like lesions.[3,7,19,20] Differential diagnoses should include benign bony lesions such as (1) giant cell tumor (GCT); (2) vertebral hemangioma (VH); and (3) aneurysmal bone cyst (ABC). Table 2 described the differential diagnosis of spinal MFD.

GCT of the spine often shows expansile lesions and a “soap bubble” appearance.[21,22] In contrast to MFD, only a small number of lesions have a sclerotic rim. Cortical destruction was commonly seen in GCT.[21] On MRI, GCT often shows a heterogeneous or homogeneous signal intensity on T1WI. The solid areas of the tumor show heterogeneous low-to-intermediate signal intensity on T2WI.[23]

The “corduroy cloth” appearance is an important sign for VH, which represents the transverse cuts through the thickened vertical trabeculae.[24] On CT, the lesion is often nonexpanded and circular-like shape. On MRI, the lesion often shows high-signal intensity on T1WI and T2WI. Enhanced MRI imaging shows obvious enhancement.[25,26] Some scholars have suggested that the VH which causes nerve damage always shows a specific signal intensity (low signal intensity on T1WI and high signal intensity on T2WI).[25] This specific signal intensity is mainly due to the prominent hypervascular stroma.[26]

ABC of the spine often shows expansile lesion. The inner edge of the bone shell has different sizes of trace.[27] Lesions can breakthrough the cortical bone to form soft-tissue masses. Due to the different components of the lesion, the lesion has various signal performances on MRI.[28] The presence of a “fluid–fluid” level on MRI is characteristic for ABC.[29] This feature is mainly due to the presence of blood in different stages of evolution.[29]

### Treatment

The treatment of spinal MFD still remains controversial. Some scholars have suggested that the treatment should focus on symptom relief and prevention of lesion progression rather than a simple resection.[5] It is well-recognized that bisphosphonates as a first-line drug have successfully relieved the pain.[5,10,12] However, once medical management fails, surgical management should be taken into consideration. The purpose of surgery is to reconstruct the stability of the vertebral body, and the most frequently used methods are curettage, internal fixation, and bone grafting.[11]

### CONCLUSIONS

The CT and MRI manifestations of spinal MFD have the following characteristics: expansile lesions, GGO, sclerotic rims, and no obvious soft-tissue masses. These features can provide a highly suggestive diagnosis of spinal MFD. In addition, the combined use of CT and MRI examinations is necessary for patients with suspected spinal MFD.

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### Conflicts of interest

There are no conflicts of interest.

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