Giant Polyostotic Fibrous Dysplasia: F-18-fluorodeoxyglucose Positron Emission Tomography/Computerized Tomography and Radiologic Findings

Dev Poliostotik Fibröz Displazi: F-18-florodeoksiglukoz Pozitron Emisyon Tomografi/Bilgisayarlı Tomografi ve Radyolojik Bulgular

Melis Baykara Ulusan1, Tevfik Fikret Çermik2
1University of Health Sciences Turkey, Istanbul Training and Research Hospital, Department of Radiology, Istanbul, Turkey
2University of Health Sciences Turkey, Istanbul Training and Research Hospital, Department of Nuclear Medicine, Istanbul, Turkey

ABSTRACT

A 40-year-old man with polyostotic fibrous dysplasia underwent F-18-fluorodeoxyglucose (FDG) positron emission tomography/computed tomography imaging to rule out a possible malignancy. It showed lytic, expansive and moderate to high hypermetabolic bone lesions with “ground glass” pattern and surrounded by a distinct rim of reactive bone in the right temporal bone, 8th and 9th ribs on the left hemithorax, T8 vertebra and sacrum. As some lesions had high F-18-FDG uptake, it was recommended to repeat the histopathological examination with suspicion of sarcomatous pathology. A second biopsy of the mass on the 8th rib was confirmed the diagnosis of fibrous dysplasia.

Keywords: Polyostotic fibrous dysplasia, F-18 FDG PET/CT, CT, MRI

ÖZ


Anahtar Kelimeler: Poliostotik fibröz displazi, F-18 FDG PET/CT, CT, MRI

Introduction

Fibrous dysplasia (FD) is a benign congenital non-neoplastic condition of children and young adults characterized by the replacement of normal cancellous bone by abnormal fibrous tissue. FD can affect one bone (monostotic, 80%) or multiple bones (polyostotic, 20%). When polyostotic, it tends to be unilateral. It can literally affect all bones but is mainly seen in long bones, craniofacial bones, and ribs. The polyostotic form can be seen in McCune-Albright and Mazabraud syndrome with endocrinopathies (1,2).

Lesions of FD continue to grow until bone maturation occurs; they regress in adulthood or remain silent. FD is usually an incidental finding in adults and asymptomatic but it may become symptomatic with pathological fractures, secondary aneurysmatic bone cysts and very rarely with malignant transformation (0.5%) (3,4).

The diagnoses of FD and its complications are based on physical, radiological, and histopathological examination. The use of the whole body F-18-fluorodeoxyglucose (FDG) positron emission tomography/computed tomography (PET/CT) imaging is not rare because FD shows various FDG uptake and can mimic multiple bone metastases, especially in polyostotic form (5,6).

This article presents a case of polyostotic FD in a patient with the imaging and histopathological features, diagnosed during the investigation of a giant rib mass.

Case Report

A 40-year-old man presented with chest pain and a big, firm swelling on his left flank area. Magnetic resonance imaging (MRI) examination images showed a heterogenous, expansive bone mass involving the left eighth and ninth ribs and T8 vertebra. The lesion was expanding from
the bottom of left scapula level posteriorly to the midclavicular line anteroinferiorly. The bone mass showed heterogeneous intermediate signal on T2W images with mainly heterogeneous avid enhancement with concomitant mild enhancement areas on postcontrast T1W images (Figure 1). The lesion was limited to the bone without a soft tissue component and mostly well-demarcated. However, giant cell bone tumor could not be excluded due to vertebral destruction and signal similarities. Aneurysmal bone cyst findings and some bone-forming regions were reported in the first histopathological examination. No primary tumor was found but FD and giant cell tumor of bone were highlighted as differential diagnosis.

The patient was referred for F-18-FDG PET/CT imaging to rule out a possible malignancy. It showed lytic, expansile and moderate to intense hypermetabolic bone lesions with “ground glass” pattern and surrounded by a distinct rim of reactive bone in the right temporal bone, left hemithorax, left T8 vertebra and sacrum (Figure 1-2). The F-18-FDG uptake in the right temporal bone was variable, with the maximum standardized uptake value ($SUV_{max}$) of 11.9. The right hemithorax lesions appeared to arise from the left eighth and ninth ribs ($SUV_{max}$: 15.1), with the involvement of the T8 vertebra ($SUV_{max}$: 9.6). FDG-avid lesions were also noted in bilateral sacral wings with a maximum $SUV_{max}$ of 9.8 in the right sacral wing. As the heterogeneous and intense F-18-FDG uptake in the lesions raised concern of sarcomatous degeneration, secondary histopathological examination was recommended.

On microscopic examination, the tissue revealed irregular, bony trabeculae in a collagenous stroma with no evidence of osteoblastic rimming and lamellar bone were evident. The mesenchymal stroma surrounding the dysplastic trabeculae was relatively hypocellular. Both features were characteristics of FD. A diagnosis of polyostotic FD was made with a second biopsy of the rib mass. Informed consent was obtained from the patient to use his data in this study.

Discussion

FD accounts for approximately 7% of benign bone tumors and is caused by an activating mutation of $GNAS$ gene, encoding $\alpha$-subunit of stimulatory G protein seen in both mono and polyostotic forms, resulting in abnormal osteoblastic differentiation and increased bone turnover (2,6).

The gross histological picture of FD constitutes a firm solid tan-gray mass which gradually replaces the medullary cavity and the surrounding cortical bone. It consists of uniformly cellular fibrous tissue, cytologically bland spindle cells with sparse mitotic activity. Irregular curvilinear woven bones are also present without any significant osteoblastic rimming (7).

A sudden increase in the size of a previous FD maybe due to a superimposed aneurismatic bone cyst or malignant transformation. Malignant transformation is rarely seen with an incidence of 0.5% in patients with FD. The incidence increases in patients with McCune-Albright syndrome by nearly 4%. It may develop after radiation therapy. Although the most common malign transformation is osteosarcoma, fibrosarcoma and chondrosarcoma can also be seen. Radiographic changes suggesting malignancy are lytic regions, intralesional calcification, periosteal reaction, and a cortical disruption (4,8-13). Radiologic differential diagnoses of FD include paget disease (mostly skull), simple bone cyst, giant cell tumor (mostly pelvis), non-ossifying fibroma, neurofibromatosis, and osteoblastoma. In tibia, when originates in cortex, FD is indistinguishable from adamantinoma. This distinction can be made pathologically (5,11,12,14).

X-ray imaging may show not specific but characteristic features. The density of lesions varies from lytic to sclerotic. Mostly central lesions show...
enlargement of medulla, deformity with endosteal thinning, increased trabeculation to a characteristic “ground-glass” appearance which is formed by a mixture of immature bone and fibrous tissues (2,4,8).

The conventional three-phase bone scan shows markedly increased uptake of tracer in both perfusion and the delayed phase. This high uptake is related to the bone turnover characteristic for the disorder. CT is still the best technique for delineating the extent of involvement with the typical ground glass pattern of the bone. The lesion is surrounded by a reactive sclerotic bone (5,14).

F-18-FDG PET/CT imaging is also used to make a differential diagnosis. Benign bone lesions like giant cell tumor of bone can show high F-18-FDG uptake, and the $S_{UV \text{max}}$ values vary according to the different stages of disease and remodeling (15).

In cases of FD, a highly variable F-18-FDG uptake was identified, between none and avid, especially in cases mimicking metastatic disease (3,16-18). The highest $S_{UV \text{max}}$ value of 15.1 for a benign FD in our study was higher than most of the benign and some of the malignant lesions reported in the literature (18,19). The report of Su et al. (19) and many other publications have discussed the necessity and usefulness of PET/CT. The lesion diversity among the body parts and the distribution of hyper- and hypometabolic areas inside the lesions made it impossible to diagnose a lesion by PET/CT. They argued that the correct diagnosis could only be made by biopsy (17).

In patients who were followed up with FD, focal increase in $S_{UV \text{max}}$ was evaluated in favor of early malignant transformation, while some authors reported that patients with benign proven lesions showed F-18-FDG activity changes in follow-up examinations (15).

CT and MRI imaging features in our case suggested FD in the first differential diagnosis when multiple lesions were considered. The 40-year-old patient presented with a growing lesion, and high F-18-FDG uptake of the lesions compared to the literature increased the suspicion of malignant transformation. When both pathology results are evaluated together, this patient with undiagnosed polyostotic FD is likely to become symptomatic due to aneurysmal bone cyst secondary to the left hemithorax lesion. In addition, repeat biopsy results from different locations were not found in favor of malignant transformation. This case shows the potential pitfalls in the interpretation of F-18-FDG PET/CT and when multiple increased uptake is seen in the skeleton, particularly in the context of suspected malignancy.

In conclusion, on F-18-FDG PET/CT imaging, the quantity of radiopharmaceutical uptake of FD lesions does not distinguish between a benign lesion and a lesion that has undergone malignant transformation. However, PET/CT imaging can lead to biopsy by identifying the maximum F-18-FDG-avid bone lesions and it may show multifocality of an undiagnosed polyostotic FD. And also, PET/CT may be useful for long-term follow-up of low F-18-FDG-avid FD lesions.

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References


