Diagnosis of solitary eosinophilic granuloma by CT, MRI, and 18F-FDG PET/CT: two clinical cases

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ABSTRACT
This paper presents two clinical cases of eosinophilic granuloma of bone diagnosed by CT, MRI, and 18F-FDG PET/CT. In both cases the patients were admitted to the clinic with suspected primary malignant bone tumor and the diagnosis of a solitary eosinophilic granuloma was made based on the results of comprehensive radiological diagnostic examination and histological verification. Solitary eosinophilic granuloma of bone is an infrequent condition, occurring in less than 1% of cases of skeletal tumor masses. The most common eosinophilic granuloma is found in the parietal and frontal bones of the skull and is an osteolytic volumetric mass that gradually increases in size. Although most bone tumors can be detected by radiography, computed tomography is preferred, primarily because of its superior ability to detect cortical bone destruction. The diagnostic accuracy of computed tomography and magnetic resonance imaging may be different. The combined use of radiological and radionuclide methods allows us to narrow the spectrum of differential diagnosis. Unfortunately, relatively low specificity of existing radiological diagnostic studies in most cases does not allow to establish a precise diagnosis, and biopsy with subsequent pathological examination remains the method of choice. These clinical observations demonstrate the need to include eosinophilic granuloma in the differential diagnosis when a solitary osteolytic focus is detected.

Keywords: eosinophilic granuloma; osteolytic focus; computed tomography; magnetic resonance imaging; positron emission tomography; case report.

To cite this article
Gelezhe PB, Bulanov DV. Diagnosis of solitary eosinophilic granuloma by CT, MRI, and 18F-FDG PET/CT: two clinical cases. Digital Diagnostics. 2021;2(1):75–82. DOI: https://doi.org/10.17816/DD59690
Два случая верифицированной солитарной эозинофильной гранулёмы: визуализация методами КТ, МРТ и 18F-ФДГ ПЭТ/КТ

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АННОТАЦИЯ

В работе представлены два клинических наблюдения эозинофильной гранулёмы кости, диагностированной методами компьютерной, магнитно-резонансной и позитронно-эмиссионной томографии с 18F-фтордезоксиглюкозой, совмещённой с компьютерной томографией. В обоих случаях пациенты поступили в клинику с подозрением на первичную злокачественную опухоль кости, по результатам комплексного лучевого диагностического исследования и гистологической верификации установлен диагноз солитарной эозинофильной гранулёмы. Солитарная эозинофильная гранулёма кости — достаточно редкое (менее 1% случаев всех опухолевых объёмных образований скелета) заболевание. Наиболее часто эозинофильная гранулёма обнаруживается в теменной и лобных костях черепа и представляет собой остеолитическое объёмное образование, постепенно увеличивающееся в размерах. Несмотря на то, что большую часть опухолей костной ткани можно выявить при помощи рентгенографии, предпочтительно применение компьютерной томографии, в первую очередь из-за её превосходной способности визуализировать деструкции кортикального слоя кости. Диагностическая точность компьютерной и магнитно-резонансной томографии может быть различна. Комплексное применение методов лучевой и радионуклидной диагностики позволяет сузить спектр дифференциального диагноза. К сожалению, относительно низкая специфичность существующих лучевых диагностических исследований в большинстве случаев не позволяет установить точный диагноз, и методом выбора остаётся биопсия с последующим патоморфологическим исследованием. Данные клинические наблюдения показывают необходимость включения эозинофильной гранулёмы в дифференциальный диагноз при обнаружении солитарного остеолитического очага.

Ключевые слова: эозинофильная гранулёма; остеолитический очаг; компьютерная томография; магнитно-резонансная томография; позитронно-эмиссионная томография; клинический случай.

Как цитировать
2例证实的孤立性嗜酸性肉芽肿
CT、MRI和18F-FDG PET/CT成像

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简评:
本文介绍计算机诊断嗜酸性骨肉芽肿、磁共振和18F氟脱氧葡萄糖正电子发射断层扫描以及计算机断层扫描的两个临床观察。根据综合的放射学诊断研究和组织学证实，确诊为孤立性嗜酸性肉芽肿，两例患者都因怀疑原发性恶性骨肿瘤而入院。孤立性嗜酸性肉芽肿是一种相当罕见的疾病（不到1%的骨骼肿瘤体积形成病例）。最常见的是，嗜酸性肉芽肿见于头骨的顶骨和额骨，是一种溶骨体积的形成，逐渐增大。虽然大多数骨肿瘤可以通过X线摄影发现，计算机断层扫描是首选，主要是因为它能很好地显示骨皮质层的破坏情况。计算机断层扫描和磁共振成像的确诊准确性可能不同。辐射和放射性核素诊断方法的复杂应用使能够缩小鉴别诊断的范围。在大多数病例中，现有的放射学诊断研究的特异性较低，不能做出准确的诊断，选择的方法仍然是活检后进行病理形态学检查。这些临床观察表明，当发现孤立的溶骨性病灶时，鉴别诊断需要包括嗜酸性肉芽肿。

关键词：嗜酸性肉芽肿；溶骨性病灶；计算机断层扫描；磁共振成像；正电子发射断层扫描；临床病例。

引用本文:
DOI: https://doi.org/10.17816/DD59690
BACKGROUND

The newly diagnosed osteolytic focus in young patients leads inevitably to an extensive differential diagnosis, which involves a variety of pathological processes. Under conditions of oncological alertness of radiologists and general practitioners, the osteolytic focus is often unambiguously interpreted as a manifestation of a malignant tumor. It should be remembered that benign and inflammatory processes can also cause the emergence of an osteolytic focus.

The paper presents two clinical cases of solitary eosinophilic granuloma of the bone, which is a rare pathological process that must be included in the differential range when a solitary osteolytic focus is detected.

DESCRIPTION OF THE CASES

Clinical case 1

A 30-year-old female patient considers herself sick since August 2016, when pain began in the lumbar region on the left, progressing over the course of a year. She visited the clinic in August 2017 after experiencing severe pain.

The studies of the pelvis were performed using magnetic resonance imaging (MRI) and computed tomography (CT). MRI revealed a cystic formation measuring 2.2 × 1.4 × 2.0 cm on the gluteal surface of the upper sections of the left iliac crest, along with edema of the musculus gluteus medius with a vertical length of up to 7 cm. A trabecular edema of the iliac crest on the left with a length of 5.0 cm was determined. The CT scan showed an osteolytic focus of the upper sections of the wing of the left iliac bone of up to 1.8 × 1.2 × 1.2 cm with clear uneven contours, destructed cortical layer of the bone, and signs of generalization beyond its limits (Fig. 1).

Mono-mode positron emission tomography (PET) with 18F-fluorodeoxyglucose (18F-FDG) was performed, which revealed a single focus of the radiopharmaceutical agent hyperfixation in the area of the wing of the left iliac bone (SUVmax1 13.1) (Fig. 2); therefore, the widespread metastatic process was ruled out.

CT-guided 18G needle biopsy was performed from the wing formation of the left iliac bone (Fig. 3). The histological study (No. 2017-10802-01) concluded on the morphoimunohistochemical presentation, which is most consistent with Langerhans cell histiocytosis (eosinophilic granuloma, histiocytosis X) (Fig. 4).

Clinical case 2

A 12-year-old boy, during his football practice, he hit the ball with his head. According to his parents and the child himself, after which, they noticed swelling in the forehead area, which gradually increased in the subsequent days. A CT scan was performed, as recommended by the doctor of the primary health care facility, which revealed an osteolytic rounded defect of the frontal bone with a diameter of about 3.5 cm, punch-type destruction of the external and internal cortical laminas, and soft tissue parasosseous formation. MRI was recommended as further examination.

According to the brain MRI data, a subcutaneously located space-occupying lesion was detected in the frontal region parasagittally, with a mild right-sided priority, with a non-uniformly increased MR signal in the T2-WI and T2-dark fluid modes, with signs of diffusion restriction, of

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1 SUV (standardized uptake value) — стандартизированный уровень накопления радиофармпрепарата.
an ovoid shape with indistinct uneven boundaries, sized 47 × 17 × 35 mm. It was widely adjacent to the squama of the frontal bone, with destruction of the external and internal cortical plates with a minimal intracranial soft tissue component, limited by the brain dura mater (Fig. 5).

Lymphocytosis of up to 49.5%, neutropenia of 39.3%, and thrombocytosis 491 were noticeable in the general blood test, as well as an increase in the erythrocyte sedimentation rate of up to 29 mm/h and an increase in C-reactive protein up to 10.65 mg/L.

According to the results of the studies, a neoplastic lesion of the frontal bone was suggested. Differential diagnostics was made between lymphoma, plasmocytoma, and sarcoma. In order to search for a primary tumor focus, 18F-FDG PET/CT was performed. In the frontal region, along the midline, an ovoid lesion of 30 × 15 mm in size was revealed, with a significant accumulation of the radiopharmaceutical agent (SUVmax up to 11.2), destruction of the external and internal cortical lamina of the frontal bone (Fig. 6). For the rest of the 18F-FDG foci, no positive neoplastic process was detected; therefore, a widespread metastatic process was ruled out.

An incisional biopsy of the subcutaneous lesion of the frontal region was performed for histological verification. Percutaneous puncture biopsy of the frontal region lesion was performed in the supine position with a thick needle. When aspirated with a syringe, no tissue was obtained. A 1-cm transverse linear incision was made along the hairline of the frontal region soft tissues. A biopsy sample of the pathological tissue, which was represented by gray soft tissue masses, was taken using a Volkmann curet and Royce forceps. According to the histological conclusion No. 2015-11688-01, the changes are more consistent with Langerhans cell histiocytosis (eosinophilic cell granuloma).

**DISCUSSION**

Solitary eosinophilic granuloma of bone is a relatively uncommon occurrence, accounting for less than 1% of tumor-like bone lesions. A histological sign of histiocytosis X,
including eosinophilic granuloma, is the presence of proliferating histiocytes (Langerhans histiocytes) [1]. The histiocyte contains puffed oval nuclei and eosinophilic cytoplasm. Birbeck granules are cytoplasmic organelles found in Langerhans cells, but their function is still unclear. The granuloma also comprises a large number of eosinophils and giant multinucleated cells.

In their work, K. M. Herzog et al. [2] reported that the skull is the most common site of eosinophilic granuloma (43%), whereas the femur is the second most common site. C. Arseni et al. [3] reported the lesion of the skull was solitary in 80% of the patients with eosinophilic granuloma, as in our clinical case 1.

Eosinophilic granuloma of the skull is manifested as an osteolytic space-occupying lesion, gradually increasing in size, often with localization in the parietal and frontal bones. On the basis of 25 patients with a total of 41 eosinophilic granulomas, L. Ardekian et al. [4] found that pain, often accompanied by local edema, was the most common symptom (92% of cases).

Although X-ray can accurately identify and distinguish most bone foci, CT is the preferred method, mainly due to its excellent ability to detect bone cortical layer destruction.

The radiographic characteristics vary significantly depending on the lesion location. The lesion in the skull is usually 1 to 4 cm in diameter, demonstrating punch-type clear boundaries, with frequent destruction of the external and internal cortical lamina. At the same time, there can be sequestrum inside the focus. Flat bone lesions are characterized by a periosteal reaction, thinning of the cortical layer, and local swelling of the bone. A hole within a hole may form when multiple small foci merge. A marked destruction of bone tissue, imitating a malignant process, occurs in rare cases.

The most significant in differential diagnostics with eosinophilic granuloma, in the range of the destructive benign and malignant lesions of the cranial vault, are osteomas (benign tumors), plasmacytoma, epidermoids, dermoid cysts, vascular tumors, osteosarcoma (malignant sarcoma), metastatic disease, meningiomas, as well as infectious and pathological conditions [5].

Bone eosinophilic granuloma may look like osteomyelitis, Ewing’s sarcoma, or lymphoma on X-ray. Other skeletal lesions, such as neuroblastoma metastases, intraosseous hemangiomas, and fibrous dysplasia should also be considered in differential diagnostics. In adults, eosinophilic granuloma can mimic osteolytic metastases, multiple myeloma, and hyperparathyroidism.

The most common finding based on MRI data is a mild diffuse decrease in the signal according to the T1-WI data, combined with an increase in the signal according to the T2-WI. Edema is also visible in the soft tissues surrounding the lesion, as shown by an increased signal on the spin-echo inversion-recovery sequence. The focus of eosinophilic granuloma of the skull bones limits diffusion compared with the white matter of the brain [6]. The described changes are not specific and can occur in a number of conditions, including osteomyelitis, traumatic changes, and avascular necrosis [7].

The sensitivity indicated in the literature for 18F-FDG PET scanning is greater than 90%, whereas the specificity remains low and varies considerably, from 65% to 80% [8, 9]. False negative results are most commonly caused by low-grade tumors, which often show low levels of 18F-FDG fixation. False positive results can be caused by some benign diseases, including fibrous dysplasia and aneurismal bone cyst, as well as acute inflammation [10].
Treatment for eosinophilic granuloma is determined on the degree of the disease progression. Surgery, radiotherapy, and chemotherapy are all possible treatment methods, which can be used in isolation or combination. Surgery is usually indicated for isolated lesions when an appropriate cure can result in complete elimination of the lesion. Despite the fact that eosinophilic granuloma is considered to be a benign condition, there have been reports of spontaneous regression and relapses after surgical excision in the literature [11, 12]. Because local relapses are often registered in a series with longer follow-up periods, it is recommended that subsequent follow-up studies be performed for at least 10 years [11–13].

CONCLUSION

Differential diagnostics of a solitary osteolytic focus can be difficult. The use of an integrated approach in radiation diagnostics, which includes CT, MRI, and 18F–FDG PET/CT, enables to narrow the range of possible pathological conditions. At the same time, the specificity of existing radiological diagnostic studies does not always enable to establish a precise diagnosis, so histological verification remains the preferred method. When identifying a solitary osteolytic focus in young patients, an eosinophilic granuloma should be considered.

ADDITIONAL INFORMATION

Funding source. This study was not supported by any external sources of funding.

Competing interests. The authors declare that they have no competing interests.

Author contribution. The authors confirm contribution to the paper as follows: material collection and article writing. P.B. Gelezhe; histological examination data. D.V. Bulanov. All authors contributed significantly to the work’s conception, acquisition, analysis, interpretation of data, drafting and revising, final approval of the version to be published, and agree to be accountable for all aspects of the work.

Patient’s permission. The patient and the legal representative of the juvenile signed a voluntary informed consent to the publication of medical information in an anonymized form.

REFERENCES


СПИСОК ЛИТЕРАТУРЫ


DOI https://doi.org/10.17816/DD59690

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