INTRAOSSEOUS HAEMANGIOMAS ON HYBRID IMAGING: A PICTORIAL REVIEW

Mairah Razi¹, Humayun Bashir¹, Saima Riaz¹, Zia S. Faruqui²
¹Department of Nuclear Medicine, Shaukat Khanum Memorial Cancer Hospital and Research Centre, Lahore, Pakistan,
²Department of Radiology, Shaukat Khanum Memorial Cancer Hospital and Research Centre, Lahore, Pakistan

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Abstract

Osseous haemangiomas are benign skeletal tumours, usually identified as incidental findings on different imaging modalities. Bone scan is the most frequent radionuclide procedure performed as metastatic workup in patients with various malignancies. Not every hotspot on a staging bone scan is malignant. Haemangiomas with variable degree of radiotracer uptake on technetium-99m (Tc-99m) bone scintigraphy may be falsely labelled as metastases in background of known malignancy. The addition of single-photon emission computed tomography-computed tomography (SPECT-CT) enhances the specificity of bone scan which allows accurate detection and anatomical localisation of scintigraphic findings. We present a case series as pictorial review of osseous haemangiomas identified on Tc-99m methylene diphosphonate SPECT-CT at our department.

Key words: Haemangioma, methylene diphosphonate, bone scan, single-photon emission computed tomography-computed tomography

Introduction

Haemangioma is a slowly growing non-malignant bone lesion. It is characterised by vascular spaces lined with endothelial cells usually located in the medullary cavity. It constitutes approximately 1% of all primary bone tumours. Spine and skull are the most common sites for haemangioma, representing 75% of the lesions. The remaining sites may involve tibia, femur and humerus. The sternum is a highly uncommon location.[1] Mostly, these are multiple in up to one-third of the cases and the peak incidence is in the fourth or fifth decade. There is hamartomatous proliferation of vascular tissue within the endothelium which may also contain fat, smooth muscle, fibrous tissue and thrombus. Histological subtypes include intraosseous cavernous or capillary haemangioma, however, may also occur as arteriovenous and venous subtypes. Vertebral haemangiomas are generally capillary type and may cause neurological symptoms if they extend into the epidural canal. Cavernous haemangiomas are common in the skull.[2] Haemangiomas are largely asymptomatic, usually found incidentally on radiography or autopsy.

Haemangiomas mostly appear as round or oval radiolucent lesions on radiographs. A characteristic finding on X-ray will be fine spiculae emanating from its centrum giving a corduroy, spoked wheel or honeycomb appearance.[1] On axial computed tomography (CT) scan, vertebral body lesions have a typical spotted ‘polka dot’ pattern due to thickened trabeculae appearing as small punctate areas of sclerosis in cross-section. Calvarial haemangiomas are usually lytic and trabecular thickening resemble as radiating, web-like or spoked wheel pattern. In long bones, the metaphyseal or epiphyseal lesions appear lytic that gives a spiculated array like an ‘Irish lace.’ Trabecular pattern of vascular malformations of the bone usually displays a high signal intensity on magnetic resonance imaging (MRI). However, these features may vary; largely depend on the proportion of fat and vascularity of the lesions. T1-weighted MRI scans diverge from low to high intensity depending on the content of adipose tissue present. T2-weighted MRI scans display lesions with high intensity due to the vascularity. Bone scintigraphy is usually normal but may show increased or decreased uptake in haemangiomas.[3]

Image Acquisition

Planar bone scintigraphy acquired after 20 mCi of technetium-99m (Tc-99m) methylene diphosphonate (MDP).
Standard single-photon emission computed tomography-CT (SPECT-CT) acquisition parameters were used.

SPECT images acquired using low-energy parallel-hole collimators with large field-of-view gamma detectors (range, 40 cm), 180° arc, 6° view angle, zoom of 1.0 and 30 s/stop. Images were acquired with a 128 × 128 matrix and then reconstructed using a three-dimensional ordered-subset expectation maximisation iterative technique (eight subsets and four iterations). CT images acquired using single-detector step-and-shoot technique, 10-mm slice interval, current of 2.5 mA, voltage of 140 kV, 256 × 256 matrix and a Gaussian filter. Total imaging time for SPECT-CT was approximately 25–30 min. [Figures 1-6] represent cases of haemangioma identified with SPECT-CT.

Discussion

Skeletal scintigraphy has the advantage of entire skeletal visualisation in oncological patients who are at high risk for osseous metastasis. $^{99m}$Tc-MDP is the most commonly available tracer for skeletal imaging. Despite the high sensitivity, not every hotspot on bone scan is malignant.

Variable causes of increased tracer uptake mimicking metastatic disease impair the specificity of the bone scintigraphy. Solitary finding on bone scan often requires further radiologic correlation with CT to improve the limited specificity of bone scan. This limitation of planar scintigraphy has been overcome by the introduction of hybrid SPECT/spiral CT since 2001.

Osseous haemangiomas are usually asymptomatic and the diagnosis is incidental most of the time. Rarely, they appear symptomatic constituting approximately 1% of all cases.

Appearance of haemangioma on bone scan has been well documented in literature. Variable degree of radiotracer uptake is seen in haemangiomas; increased, decreased or equal to adjacent bone. Vertebral haemangiomas which are smaller than 3 cm in size generally show normal uptake of a radiotracer. However, larger lesions may demonstrate either increased or decreased uptake.

They may be erroneously labelled as metastases on bone scan in background of primary malignancy which have predilection for skeletal (lytic) metastases. In addition to post-external radiotherapy, haemangioma is also one of the causes of ‘cold’ vertebrae on skeletal scintigraphy.

Han et al. evaluated case series of 15 vertebral haemangiomas in 10 patients on bone scan. Planar images showed normal findings throughout the skeleton with the exception of only one. SPECT images also displayed normal findings in 11 vertebral haemangiomas which were smaller than 3 cm in diameter. However, three of four vertebral haemangiomas were 3 cm or larger and demonstrated variably increased or decreased uptake on SPECT images.

The role of SPECT-CT has been reported in a 20-year-old woman with upper back pain by Bhoil et al. Planar bone scintigraphy showed focal uptake in the seventh thoracic vertebra. SPECT-CT showed uptake in the vertebral body and transverse process at D7 with CT findings typical of haemangioma. Furthermore, MRI of the thoracic spine confirmed the findings of haemangioma in the same vertebra.

A case of increased tracer uptake in haemangioma has also been reported in a patient of renal cell carcinoma who underwent bone scintigraphy to assess for skeletal
metastases. SPECT-CT localised the MDP uptake in D8 vertebra with vertical trabecular thickening typical of haemangioma ruling out metastasis. Furthermore, on review of thoracolumbar spine on CT, similar lesions were identified in D10 and L3 without any significant uptake.[10]
Intraosseous haemangiomas show variable Tc-99m MDP avidity. When combined SPECT-CT is used, the benefit of precise anatomic localisation of the Tc-99m MDP uptake and the corresponding CT appearance help characterise the nature of these benign lesions.

Conflict of Interest

The authors declare that they have no conflict of interest.

References


Figure 5: A 43-year-old patient with follicular carcinoma of thyroid. (a) Planar whole body and (b) right anterior oblique views show mild tracer uptake in the tip of the sternum. (c) axial and (d) sagittal views of the conventional computed tomography revealed trabecular thickening at the tip of the sternum – a rare site for haemangioma.

Figure 6: A 55-year-old female patient with left breast carcinoma. (a) Planar images of bone scan showing increased tracer uptake in D11 vertebra. Sagittal and axial fused single-photon emission computed tomography-computed tomography (CT) and CT images (b-e) show typical polka dot appearance at D11 hemivertebra with increased tracer uptake. (f and g) T2-weighted sagittal magnetic resonance imaging (MRI) of the thoracic spine shows multifocal haemangioma in this patient with hyperintense signal in D2 and D11 vertebrae and (h) axial T2-weighted MRI.