Imaging approach to unifocal sclerotic bone lesions – A pictorial essay

Subbarao Kakarla¹,²,*

¹KIMS Foundation and Research Centre, Minister Road, Secunderabad - 500003, Telangana, India
²Kakarla Subba Rao Radiological and Imaging Educational Sciences Trust (KREST), Shaikpet Nala, Hyderabad - 500008, Telangana, India

Abstract
Sclerotic bone lesions are occasionally encountered in routine practice a few of which may not be clinically significant. Whenever a single focus is noted, it is mandatory to examine the entire skeleton to rule out concomitant lesions by way of limited skeletal survey or radionuclide bone scan. The etiological factors of unifocal sclerotic bone lesions are many. The aim of this essay is to identify the lesions and to find out the etiology and to arrive at a final diagnosis. The sclerotic lesions include developmental/congenital, post traumatic, inflammatory, infective, metabolic, hemopoietic, neoplastic, and miscellaneous groups. The radiological characteristics are described in detail so that a final diagnosis can be made. Often, conventional radiology is adequate but CT/MRI may be helpful. The final diagnosis rests on histopathology.

Keywords: Sclerotic bone lesions; Unifocal; Conventional CT; MRI; PET CT

Introduction
Unifocal sclerotic bone lesions are either symptomatic or asymptomatic. Evaluation of the lesion is important in the diagnosis and management of these lesions. The radiological characteristics are described in detail so that a final diagnosis can be made [1-3]. A sclerotic bone lesion radiologically may appear as homogenous and dense lesion. The entire lesion may not be sclerotic but if three fourths of the lesion is sclerotic, it can be called as a sclerotic lesion. The etiology of this lesion is varied and various lesions are listed below (Table 1).

The first modality that detects the sclerotic lesion is by conventional radiology. Computerized tomography (CT), magnetic resonance imaging (MRI), radionuclide scan and positron emission tomography (PET) CT are advanced technics that help to pinpoint the final diagnosis. Ultimately, histopathology report is necessary.
Table 1: Etiology - The differential diagnosis of a solitary sclerotic bone lesion is heavily influenced by the age of the patient, and includes.

- Bone island
- Enostosis
- Fibrous dysplasia
- Post traumatic/ stress, callus
- Inflammatory/ infection
- Sclerosing osteomyelitis
- Osteoma
- Osteoid osteoma
- Osteoblastoma
- Osteosarcoma
- Enchondroma
- Healed fibrous lesions
- Ewing's sarcoma
- Lymphoma
- Plasmacytoma
- Paget's disease
- Healed brown tumors

Unifocal sclerotic lesions with discussion on imaging - Congenital/ developmental bone island

A bone island, also known as an enostosis, is a focus of compact bone located in cancellous bone (Figure 1a-i). This is a benign entity that is usually found incidentally on imaging studies; however, the bone island may mimic a more sinister process, such as an osteoblastic metastasis (for example, from prostate cancer). Bone islands demonstrate characteristic radiographic findings. In the correct clinical context, findings on radiographs are considered diagnostic. However, if the lesion is large or demonstrates increased scintigraphic activity, or if the patient is symptomatic or has a history of malignancy, clinical follow-up and/or biopsy may be warranted.

Bone islands typically appear as sclerotic, round-to-ovoid intramedullary foci. The long axis of the bone island is aligned parallel to the long axis of the bone [4-6].

Enostosis

An enostosis is a small area of compact bone within the cancellous bone. They are commonly seen as incidental findings on radiographs or CT scans. They are typically very small and do not cause any symptoms (Figure 2a,b). Their radiodensity is generally similar to cortical bone. Most of the authors treat enostosis as same as bone island. However, enostosis is linear, oblong, and close to the endosteum [3, 4].
Fibrous dysplasia

Fibrous dysplasia (FD) is a developmental disorder with replacement of the osseous part with fibrous tissue. Three major forms are described but unifocal sclerotic FD is quite rare. Often biopsy is required to make a final diagnosis. Skull and facial bones are the most common sites. Polyostotic involvement should be ruled out before it is called to be unifocal [7, 8]. Radiologically, solitary sclerotic area is noted with a ground glass type of matrix (Figure 3a-c). The margins are well outlined without any soft tissue swelling or periosteal reaction.

Post traumatic/ Stress

Healed fractures may show dense sclerotic lesion simulating osteoid osteoma. Hypercallosis is noted in some clinical conditions such as osteogenesis imperfecta, and neurological disorders. History of trauma is essential to make a diagnosis. Bone grafts may also show unifocal sclerotic lesion during healing stage and history is essential to make a diagnosis (Figure 4a-e). Stress lesions including fractures show sclerosis subsequent to healing. Stress and ischemic lesions produce sclerosis (Figure 4f) [9, 10]. Stress lesions need not show the fracture line all the time even with CT. Radionuclide scan identifies the lesion.

Osteonecrosis - Avascular necrosis

It often occurs in the articular ends of the long bones and there are many causes. Head of the femur is the most common site. Avascular necrosis may be the result of the following:

- Injury / Fracture
- Damage to blood vessels
- Long-term use of drugs, such as corticosteroids
- Excessive, long-term use of alcohol
- Specific chronic medical conditions
- Radiation
Radiologically several grades are observed depending upon the stage of the disease. In the final stage sclerosis is predominant (Figure 4f).

Figure 4a: (i) Healing fracture 3rd metatarsal with callus; (ii, iii) Healed stress fracture.

Figure 4b, c: Healed stress fracture in typical location.

Figure 4d, e: Freibergs disease dense 2nd metatarsal head.

Figure 4f: Stress lesion tibia.
Inflammatory/ Infection

Inflammatory lesions such as Langerhans cell histiocytosis and sarcoidosis may produce sclerotic lesions. Langerhans cell histiocytosis (LCH) represents a spectrum of rare disorders characterised by idiopathic infiltration and accumulation of abnormal histiocytes. It may be unifocal or multifocal [11]. When it is unifocal it is called eosinophilic granuloma. In the healed phase of a single lesion, sclerosis is noted which may be mistaken for other lesions. Osteomyelitis of Garre's type presents as sclerotic lesion and so also does healed osteomyelitis.

LCH is a nonspecific granulomatous lesion and generally gives lytic and mixed type of bone lesions. Eosinophilic granuloma can be unifocal but sclerotic appearance is extremely rare except when it is healed.

Bone infections – Osteomyelitis

Except in Garre's type of osteomyelitis, unifocal sclerotic lesion is not seen in osteomyelitis, whether acute or chronic.

Sclerosing osteomyelitis of Garré

In 1893 Garre described a peculiar form of chronic low grade, diffuse, non-purulent osteomyelitis characterized by a striking absence of viable pathogens on attempted tissue culture. Radiologically the bone involved is uniformly dense with or without minimal periosteal reaction [12]. It can be diaphyseal or metaphyseal (Figure 5a-d).
Idiopathic – Condensing osteitis, Osteitis condensans ilii

Condensing osteitis of the clavicle is a benign, often painful disorder, marked by bony sclerosis at the sternal end of the clavicle. Sclerosis of the inferomedial part of the medial end of the clavicle is noted (Figure 6a) [13].

Neoplastic – Benign, malignant and metastatic

Benign bone lesions appearing as unifocal sclerotic lesion

Osteoma

Osteoma is a benign bone tumor commonly encountered in skull and sinuses. Occasionally osteoma may be noted in the long bones also. These are painless benign tumors and the imaging characteristics vary from ivory dense to slightly trabeculated appearance [15].

Osteomas are frequently present in paranasal sinuses and occur in the frontal as well as the outer and inner tables of the skull (Figure 7a-c). Further imaging is not generally needed.

Osteoid osteoma

Osteoid osteomas are benign bone-forming tumors that typically occur in children (Particularly adolescents). They have a characteristic lucent nidus less than 1.5 or 2 cm and surrounding osteosclerotic reaction, which classically causes night pain that is relieved by the use of salicylate analgesia, e.g. aspirin [16]. Although there is a lucent nidus in the lesion,
it may be obscured by abundant reactive new bone (Figure 8a-e). CT/ MRI are essential to demonstrate the nidus (Figure 8f).

Figure 7a, b: Osteoma of the right frontal sinus.

Figure 7c: CT osteoma of the skull.

Figure 8a: Osteoid osteoma 3rd metacarpal.

Figure 8b-d: (b, c) Osteoid osteoma of the distal phalanx of the thumb, (d) thumb showing clubbing.

Figure 8e: Osteoid osteoma femur, nidus is hidden by new bone.

Figure 8f: CT Osteoid osteoma skull with nidus.
**Osteoblastoma**

Osteoblastoma has a nidus more than 2 cm in diameter and matrix may be heavily calcified (Figure 9a). The posterior elements of vertebra are commonly involved both in osteoid osteoma and osteoblastoma [17] (Figure 9b).

**Osteosarcoma**

Osteosarcoma (OS) is a common primary malignant tumor of bone that produces osteoid matrix. Conventional OS is the most common subtype of OS and is readily identified at radiography as an intramedullary mass with immature cloudlike bone formation in the metaphyses of long bones. The most common radiographic appearance is expansile lytic bone lesion with coarsely thick or thin incomplete trabeculation (61% of cases). A dense sclerotic pattern is less common (<30% of cases). Variable rates of periosteal reaction (22%–50%) at radiography are also reported. Radiologically unifocal sclerotic lesion without periosteal new bone or soft tissue swelling is quite rare [18].

Malignant lesion of bone generally manifests as a predominantly sclerotic lesion except in telangiectatic osteosarcoma (Figure 10a, b).
Other malignant lesions of bone [2, 19]

**Ewing’s sarcoma**

Ewing’s sarcoma is a small cell sarcoma with heterogeneous appearance. A permeative lytic appearance with onion peel type of periosteal reaction is a common manifestation. However, purely sclerotic with ivory appearance also may be encountered (Figure 11a-c). In the latter instance differentiation from osteogenic sarcoma and lymphoma is difficult as all of them appear sclerotic.

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**Lymphoma**

Hodgkin’s lymphoma or other lymphomas may present as an isolated ivory focus more commonly in the spine (Figure 12) [20].

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**Plasmacytoma**

Plasmacytoma is a unifocal lesion of multiple myeloma. Plasmacytoma presenting as a unifocal sclerotic lesion is quite rare. In Poem Syndrome multiple lesions are noted associated with other clinical findings [21].

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**Metastasis**

Unifocal sclerotic metastasis is quite rare but the presence of other lesions must be ruled out before mentioning that it is unifocal. Radiologically no specific characteristic is noted (Figure 13a-c), although they may be lytic, expansile, sclerotic and mixed. A primary lesion in prostate, breast, GI and GU tracks must be identified to say it is unifocal metastasis [2]. A PET CT scan is quite useful in making a proper diagnosis and identify the extent of lesions.

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**Cartilaginous lesions**

**Enchondroma**

The tumors arising from cartilage generally contain calcifications which are distinct with circular, arc like, nugget shapes. Occasionally a homogenous dense calcification is noted simulating a unifocal sclerotic lesion. Enchondroma when it is unifocal involves femur and other long tubular bones (Figure 14a, b). The hands are also common sites but heavy calcification is rare.
Healed fibrous lesions

Fibrous lesions of bone are generally not dense but healed non ossifying fibroma may show as a unifocal sclerotic lesion (Figure 15a-d). When they are limited to cortex and endostemeum they may simulate enostosis (Figure 15e).

Miscellaneous

Paget’s disease

Paget disease is rare but when present it may be lytic, lytic and sclerotic, and purely sclerotic. Radiologically a single ivory vertebra is one of the manifestations of Paget (Figure 16a). Occasionally other single bones may be involved (Figure 16b) [22, 23].

Healed brown tumors

Brown tumors of primary hyperparathyroidism may show sclerosis subsequent to parathyroidectomy (Figure 17ab). However, single brown tumor without other skeletal changes is quite rare [24].
Conclusion

Unifocal sclerotic bone lesions constitute a wide spectrum of disorders. Conventional radiology is the initial imaging method and is best in making an early diagnosis. Radiological appearances of some of these lesions are similar but by analysis and with the help of clinical and laboratory studies, most of the lesions can be differentiated. Advanced imaging is quite unnecessary in most of the cases. However, the unifocality should be established by eliminating multiple lesions with the help of other imaging methods such as Spect CT and PET CT.

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Conflicts of interest

Author declares no conflicts of interest.

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