Benign tumors of the bones of the hands - Pictorial essay

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Abstract

Bone tumors of the hands are uncommon. However, cartilaginous tumors such as enchondromas are occasionally encountered. Other tumors such as osteoma, osteoid osteoma and osteoblastoma may also occur but are uncommon. Other fibrous, vascular and neural tumors are rare. The radiological characteristics of these tumors including giant cell tumor and aneurysmal bone cysts are described. This pictorial essay is mostly restricted to conventional radiographs.

Keywords: bones of the hand; benign tumors; tumoral lesions

Introduction

Hand is the distal functional tool of the upper limb, an important organ for day-to-day functions. Hand anatomy is complex and intricate. This enables hands to do gross as well as precise functions. A total of 27 bones constitute the basic skeleton of the wrist and hand [8 carpals, 5 metacarpals, 14 phalanges and 3 for each digit except thumb which has 2]. The classification of benign tumors occurring in the bones of the hands is mentioned in Table 1.

Table 1: Classification of benign tumors.

<table>
<thead>
<tr>
<th>Benign tumors</th>
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<tbody>
<tr>
<td>Osseous</td>
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<tr>
<td>Cartilaginous</td>
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<tr>
<td>Fibrous</td>
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<tr>
<td>Vascular</td>
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<tr>
<td>Miscellaneous – Giant cell tumors,</td>
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<tr>
<td>epidermoid cyst, aneurysmal bone cyst</td>
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</tbody>
</table>

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Received 22 January 2018; Revised 27 February 2018; Accepted 16 March 2018; Published 26 March 2018


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Diagnosis with radiological characteristics

Conventional radiography plays a major role in the diagnosis of tumors of the bones of the hand. Benign tumors of the bones of the hands are more frequent than malignant lesions [1]. As with any bone tumor in any part of the body, it is critical to know the precise location of the tumor as well as the age of the patient to narrow the differential diagnosis. Also, the margin of the lesion and the zone of transition between the lesion and normal bone are key factors in determining the aggressiveness of the lesion. A lesion with well-defined, sharp margins on radiographs or CT images is considered nonaggressive, whereas a broad zone of transition and poorly defined borders suggest an infiltrative osseous process, such as an aggressive tumor or a destructive metabolic or infectious process. Mineralization and opacity of a lesion can be appreciated at radiography, although they are better characterized with CT, and can help determine the origin of the lesion. Identification of a matrix with a "rings-and-arcs" appearance indicates a chondral origin (e.g., enchondroma or chondrosarcoma). On the other hand, an opaque, cloud-like matrix is suggestive of osseous mineralization (osteoblastoma). The presence and type of periosteal reaction can also help in the differential diagnosis. In general, periosteal reaction is uncommon in most of the benign tumors. A smooth, solid-appearing periosteal reaction denotes a slow, nonaggressive process and a disrupted or "sunburst" appearance suggests an aggressive lesion [2-4]. More aggressive indicate malignant tumors [5]. Presence of soft tissue swelling without mineralization does not help in arriving at the diagnosis. However, mineralization either cartilaginous or osseous would determine the nature of the lesion.

Imaging characteristics - Osseous tumors

- Osteoma
- Osteoid osteoma
- Osteoblastoma

Osteoma: This lesion is more common in skull bones and paranasal sinuses. However, osteoma can be encountered in the bones of the hand also. The imaging appearance is the same as is encountered in the skull. It is an uniformly dense structure attached to the cortex (Figure 1). A bone island should be distinguished as the latter is located in the medulla (Figure 2).

Osteoid osteoma: It may be intra cortical, periosteal, intramedullary or subarticular. Intracortical osteoid osteoma is a round lucent lesion with a nidus (Figure 3). Intense new bone formation is noted in both cortical and periosteal lesions. Associated soft tissue swelling is present. Intramedullary lesion does not usually produce reactive new bone. Subarticular lesion produces synovitis simulating arthritis without any periosteal reaction.
Osteoblastoma: A benign osteoid producing tumor similar to osteoid osteoma but has a larger nidus. Moderate degree of new bone formation is also present (Figure 4a,b). Morphologically, it may simulate a large osteoid osteoma, aneurysmal bone cyst and when aggressive, an osteosarcoma.

**Cartilaginous tumors**

- Enchondroma
- Enchondroma protuberans
- Periosteal chondroma
- Exostosis – Single, multiple
- Ollier’s enchondromatosis
- Chondromyxoid fibroma
- Chondroblastoma
- Trevor’s

**Enchondroma**: It is a benign expanding cartilaginous neoplasm that is usually solitary in intramedullary bone. The typical rings and arcs of calcification may not be present all the time. However, punctuate or micro nodular calcification may be seen sometimes [6]. Endosteal scalloping is a characteristic feature.
(Figure 5a-d). The primary significant factors of enchondromas are related to their complications, most notably pathologic fracture (Figure 5e-h). A small incidence of malignant transformation is noted, which may be associated with a pathologic fracture (Figure 5i-k). It is not infrequent to realize that a radiologically benign enchondroma showing a low grade chondrosarcomatous change histologically.

**Enchondroma protuberans**: It is a rare tumor that arises from an intramedullary enchondroma with an exophytic growth pattern (Figure 5l,m).

**Periosteal chondroma**: It is also called ‘Juxtacortical chondroma’ and is a chondromatous lesion arising from the periosteum. It may grow into the medullary cavity. A characteristic cortical hook, similar to the hook noted in gout indicates the cartilaginous nature.

![Figure 5a-d: Enchondroma spectrum; (a-c) phalanges, (d) metacarpal.](image)

![Figure 5e-h: Enchondroma with a fracture; (h) MRI of the same.](image)

![Figure 5i-k: Enchondroma with malignant transformation.](image)
(Figure 6a-e). MRI is rarely necessary. Nodular calcification may be noted in the matrix (Figure 6c,f).

Exostosis (Osteochondroma): Isolated osteochondroma is rare in the bones of the hand (Figure 7a,b). A subungual exostosis may be occasionally encountered [6] (Figure 7c). However, in multiple exostoses bones of the hand may also be involved (Figure 7d).
Ollier's disease: In Ollier disease, two types have been described. One is multiple enchondromas which often appear to be larger than they do in other conditions. Because enchondromas occur in young patients and can be large, growth of the affected limbs may be adversely affected, and pathologic fractures may occur (Figure 8a-d). Second type is Ollier's dyschondroplasia where the appearance of linear lucencies, in which the chondrocytes appear to line up in a vertical orientation along the length of the bone. These lesions may be unilateral or bilateral and may shorten the bones (Figure 8e).

Maffucci syndrome: In Maffucci syndrome, multiple enchondromata with associated soft-tissue hemangiomas are seen. Soft-tissue hemangiomas typically have numerous rounded calcifications with central luencies, which are consistent with phleboliths on plain radiograph (Figure 9a,b).
**Metachondromatosis:** Enchondromas and associated osteochondromas are noted, which differ from conventional osteochondromas in that they point toward rather than away from the joint (Figure 10). Metachondromatosis is the only one that is hereditary, which is by autosomal dominant transmission.

**Chondromyxoid fibroma:** It is very rare as compared with other cartilaginous lesions. Although it is listed under cartilaginous tumor, calcification is rarely seen on radiographs. It is a lytic, eccentric and expanding lesion with lobulation and surrounding sclerosis (Figure 11).

**Chondroblastoma:** Chondroblastoma of the hand bones has been reported. However, we have not encountered in our series.

**Trevor’s:** It is also called dysplasia epiphysealis hemimelica (DEH). It is a rare cartilaginous lesion arising from the epiphysis or ossification centre. It may involve the adjacent bone and generally has an unilateral distribution (Figure 12).

**Tumors of fibrous origin**

- Non ossifying Fibroma
- Fibroxanthoma
- Desmoplastic fibroma
- Cortical desmoid

These fibrous tumors are encountered in the long bones, but are rare in short bones.

Vascular tumors of the hand bones are rare. However, glomus tumor is relatively common in the distal phalanges (Table 2).

**Table 2: Vascular tumors.**

<table>
<thead>
<tr>
<th>Tumor Type</th>
<th>Capillary</th>
<th>Cavernous</th>
<th>Venous</th>
<th>Arteriovenous malformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemangioma</td>
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<tr>
<td>Cystic angiomatosis</td>
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<tr>
<td>Glomus tumor</td>
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<tr>
<td>Gorham disease</td>
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<tr>
<td>Hemangiopericytoma</td>
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**Hemangioma:** Radiological appearance of hemangioma is that of an expansile, lytic, fine lacy network or honey combing (Figure 13a-c). Radiating coarse linear strands diverging from center also may occur, simulating a sunray appearance. Angiogram demonstrates the type of hemangioma [7] (Figure 13d). Occasionally, hemangioma may produce reactive new bone and is sclerotic on radiographs (Figure 13e). On occasion it produces a multiloculated lytic lesion with a soft tissue swelling (Figure 13f).
Cystic angiomatosis involves multiple sites of skeleton. Bones of the hands may also show hemangiomas. However, the radiological pattern is the same as solitary hemangioma.

**Glomus tumor:** It is a rare benign vascular tumor occurring in the terminal phalanx mostly of the fingers. Radiologically a lytic lesion is noted in the terminal phalanx with large soft tissue swelling (Figure 14a,b). In the differential diagnosis epidermoid inclusion cyst [8] may be considered. Epidermoid cysts also most commonly occur in the distal phalanges of the hands. A lucent lesion with surrounding sclerosis located in the distal phalanx is a classical finding (Figure 14c).

**Gorham disease:** This is also known as vanishing bone lesion. Radiologically massive osteolysis of adjacent bones is noted without any sclerosis [9]. Generally, a single bone or multiple adjacent bones are affected (Figure 15).

**Giant cell tumors**

Giant cell tumour of the small bones of the hands and feet have also been described. Giant cell tumor is a most common benign tumor occurring at the ends of tubular bones [10]. It extends subarticularly with septae formation. The aggressive nature simulates malignant bone tumor. It occurs in the 3rd and 4th decades of life. They are mostly benign lytic lesions with a thin transitional zone without any matrix mineralization. No periosteal reaction is noted unless there is a fracture (Figure 16a-c). Sometimes it is difficult to differentiate this from aneurysmal bone cyst (ABC). Secondary ABC may be present in some of the giant cell tumors.
Aneurysmal bone cyst

It can be primary or secondary. Radiologically, it is an expanding osteolytic lesion with thinning of the cortex with a peripheral shell. The zone of transition is narrow but can be wide and aggressive (Figure 17a,b). The lesion may cross the growth plate in children [11]. Fluid fluid levels may be shown on CT/MRI [12]. These are often encountered in younger age groups.

Conclusion

Benign tumors of the bones of the hands are very rare, although cartilaginous tumors are common. Other tumors of osseous nature and vascular origin are described with radiological characteristics. Giant cell tumors and aneurysmal bone cysts are also included. This pictorial essay deals mostly with conventional radiographs of the hands.

Acknowledgements

NIMS (Hyderabad), KIMS (Secunderabad) and KREST Museum (Hyderabad).

Conflicts of interest

Author declares no interests of conflicts.

References


