“Soap Bubble” Lesion of the Middle Phalanx: Enchondroma or Epitheloid Hemangioma

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Abstract

Introduction: Epitheloid hemangioma, a benign vascular tumor that arises in skin and soft tissues can also involve the skeletal system. Occasionally this has been reported from small tubular bones of the hand.

Case Report: Authors report a case of epitheloid hemangioma of the middle phalanx in a young girl without any cutaneous manifestations. The lesion presented as a swollen middle finger, and plain radiographs showed a geographic area of destruction with cortical thinning and intralesional calcifications. The case was managed by curettage and bone grafting. Histology confirmed this as a case of epitheloid hemangioma.

Conclusion: Epitheloid hemangioma should be considered in the differential diagnosis of hand masses with expansile lytic lesions with cortical thinning.

Keywords: Hand; Vascular tumor; Enchondroma; Epitheloid hemangioma.
pain of one month duration. There was progressive increase in size. The middle phalanx was diffusely swollen, with mild tenderness. She did not have any lesions over the skin. Plain films of the right middle finger showed an expansile lesion with thinning of the cortex and intralosomal calcifications. [Fig 1]

Her complete blood count was normal. Considering her age and the radiological features a diagnosis of enchondroma was made. Computerized tomographic scan (CT scan) showed a central oval round area of geographic destruction in the medullary canal at the metaphyseal region of the middle phalanx of the right middle finger. There were dystrophic calcifications with bulbous expansion of the bone with cortical thinning. There was no soft tissue involvement. As shown in the plain films and the CT scan. [Fig 2]

With a provisional diagnosis of enchondroma she was operated under general anesthesia. The mass was approached through a curvilinear dorsolateral approach under tourniquet. Dark-brown jelly-like contents of the tumor were curetted out. There was break in the volar cortex. The defect in the bone was packed with cancellous grafts harvested from the proximal tibia.

Histopathology showed grey and dark brown tissue, with sections showing fragments of tissue composed of many perforating capillary sized blood vessels that are lined by plump endothelial cells with abundant cytoplasm and vesicular nuclei. Some cells showed vacuoles. At places there was hobnail appearance of the endothelial cells. These were admixed with numerous eosinophils, lymphocytes and few plasma cells. Occasional scattered giant cells also were seen. Immunohistochemistry for vascular markers showed CD31 positive in cords of cells and cells lining blood vessels, many CD68 positive cells were also seen. Features were consistent with epitheloid hemangioma. [Fig 3 & 4]

At one-year follow up, there was no recurrence of the lesion. The patient regained full movements of the finger, but the finger remained swollen. [Fig 5 & 6].

**Discussion**

The differential diagnosis of expansile multi-loculated phalangeal lesions includes enchondroma, giant cell tumor of bone (GCT), aneurysmal bone cyst (ABC), and chondromyxoid fibroma [4]. Enchondromas are the most common primary tumor of the hand, presenting as expansile lesion usually of the proximal phalanx [10]. Aneurysmal cysts are occasionally reported in the phalanx, and when it occurs in the phalanx it is practically impossible to differentiate it from GCT or enchondroma [10].

In GCT of the phalanges, there is bone destruction and diaphyseal and soft tissue extension [11].

In our case because of her age and the clinical and radiological findings, a diagnosis of enchondroma was made, as enchondromas are the second most common benign cartilaginous tumor in her age group, second only to osteochondroma [12].

In small tubular bone tumors like GCT and enchondroma there is significant amount of bone loss due to matrix resorption by osteoclasts [3]. A tumor with similar pattern in the plain films is epitheloid hemangioma [3].

Presentation of EH is pain in the involved bone; however it may be seen as an incidental finding [3]. There is equal sex predilection [13].

EH has been reported from the femur, humerus, tibia, scapula, and vertebrae, which are the usual sites [4-9]. It is a tumor predominantly of the long bones; however lesions in the small tubular bones have been reported [4]. In an analysis of 50 cases of EH, 8% of cases were with lesions of the small bones of the hand [3]. In most of the cases it presents as a solitary lesion [7].
Lesions in the hand are expansile, with endosteal scalloping and thinning of the cortex in the meta-diaphyseal region, and rarely with matrix mineralization[12]. There may be soft tissue extension as well[3]. Endosteal scalloping was seen in 100% of cases in a report by Bierry et al; however none of their cases showed matrix mineralization[12]. There are reports of EH presenting as well defined lytic lesions of bone with sclerotic margins and intralesional calcifications[13], giving the appearance of soap bubble[2]. The expansile nature of the tumor and the cortical thinning may suggest aggressive malignancy[5]. EH is known to be associated with skin lesions, and the absence of skin lesions made us think of enchondroma as the only diagnosis[13]. Histologically there are large polyhedral cells in sheets or cords or lining numerous well defined vascular spaces. Cells are oval, with vesicular nuclei and with different sized nucleoli. There is a lobular growth pattern, with cellular lobules separated by loose connective tissue containing well formed vessels which are of capillary size[5,13].

Cytological atypia is not a feature. Vessels are surrounded by multiple epithelial endosteal cells. The endothelial cells lining the vessels have abundant eosinophilic cytoplasm. The sheets of epithelial cells lining the vascular structures have “tomb stone” like arrangement [3].

Management of the lesion is by curettage and bone grafting or by marginal excision [5]. Spontaneous remission of the tumor has also been reported suggesting a benign nature of the tumor, however a case of recurrence was reported by Nielson et al.[3, 5].

Conclusion
Authors report a case of epitheloid hemangioma in a 16-year-old girl, which was clinically and radiologically diagnosed as enchondroma. Subsequent histopathological studies proved it to be a case of epitheloid hemangioma.

Clinical Message
This is a rare presentation of phalangeal tumor, and the diagnosis can be confirmed by immunohistochemistry for vascular markers.

References


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