A case report of osteoblastoma on the distal phalanx of the ring finger successfully treated with curettage and polymethylmethacrylate filling

Seyit Ali Gumustasa, Talat Cagırmazb,*, Olcay Gulerb, Onder Ofluoglu, Sibel Kayahan
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Orthopedics and Traumatology Department, Yavuz Selim Bone and Joint Disease State Hospital, Trabzon, Turkey
Orthopedics and Traumatology Department, Medipol University, Medical Faculty, Istanbul, Turkey
Orthopedics and Traumatology Department, Bahçeşehir University, Medical Faculty, Istanbul, Turkey
Pathology Department, Dr. Lutfı Kırdar Kartal Training Hospital, Istanbul, Turkey

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BACKGROUND: Osteoblastoma is an aggressive benign tumor whose presentation varies with location and size. This rare bone tumor is thus difficult to diagnose particularly when it occurs outside its most common location – the vertebral column and long bones.

CASE: We report a case of osteoblastoma of the fourth distal phalanx of the left hand in an 18-year-old male, presented with pain and swelling and treated with curettage and polymethylmethacrylate filling followed by immobilization by a cast, which was opened 10 days later to start physical therapy. Patient was pain-free, recovered full function of his finger, and remained without pain at one month post-surgery. The finger was monitored closely for two years; sequential films showed a radiopaque interface and no evidence of local recurrence.

CONCLUSION: This is the first report of osteoblastoma on the distal phalanx. The possibility of osteoblastoma should be considered in cases of pain and swelling of phalanx, and if diagnosed, curettage and polymethylmethacrylate filling may be the treatment of choice.

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1. Introduction

Osteoblastoma is a rare primary neoplasm of the bone. Despite its benign nature, the tumor may sometimes exhibit aggressive behavior and is typically treated with curettage; then the resulting space is packed with bone or a bone substitute.

Osteoblastomas account for only 0.5–2% of all primary bone tumors and 3% of benign bone tumors [3]. While this tumor can occur in any part of the skeleton, the most common sites are the vertebral column and long bones. The tumor rarely presents in the hand; the phalanx is involved in less than 5% of cases [3,14]. Metaphyseal lesions are slightly more common than diaphyseal, with very few lesions in epiphyseal locations [6].

Osteoblastoma can occur at any age, and predominantly affects younger persons, with about 80% of these tumors occurring before 30 years of age. Osteoblastoma affects males more often than females, at a ratio of 2–3:1 [3].

Symptoms and signs are variable, but the characteristic symptom is a localized, dull pain that does not get worse at night and cannot be relieved by salicylates [14]. We present a case of osteoblastoma located at the rare site of the distal phalanx of the hand.

2. Case report

An 18-year-old male student was referred to our clinic for pain and swelling of the fourth distal phalanx of the left hand. He complained of progressive distal phalanx pain and swelling in the area for two months that was not activity-related and only partially relieved by non-steroidal anti-inflammatory medications. In contrast to characteristic pain pattern of osteoblastoma [14], which is not worse at night, the pain in the present case was more intense at night. There was no history of trauma.

Upon referral to a medical center, an X-ray revealed a lytic lesion containing some calcification in his distal phalanx (Fig. 1).

The rheumatologist referred him to an orthopedic surgeon. Physical examination identified a tender, firm swelling over the distal phalanx. Movements were slightly restricted by pain. Blood work data, including complete blood count, chemistry, coagulation analysis, erythrocyte sedimentation rate, and C-reactive protein were all normal.
Based on clinical and radiological findings, the lesion was considered a benign primary tumor. An excisional biopsy was performed through a lateral approach and the resulting gap was filled with polymethylmethacrylate (PMMA). Macroscopic biopsy findings included a cavity containing spongy bone and reddish soft mesh. Histopathological appearance of the osteoblastoma consisted of a well-vascularized connective tissue stroma actively producing osteoid and primitive woven bone (Fig. 2).

Then the digit was immobilized by a cast, which was opened 10 days later to start physical therapy. At that time, the patient was pain-free; he recovered full function of his finger and remained without pain at one month post-surgery. The finger was monitored closely for two years; sequential films showed a radiopaque interface and no evidence of local recurrence (Fig. 3).

3. Discussion

Osteoblastoma is an uncommon primary bone tumor with clinical and histological manifestations similar to those of osteoid osteoma. The most helpful single imaging technique for diagnosing osteoblastoma is radiography, on which the tumor appears as an expansile radiolucent lesion with an intact surrounding shell of bone and may contain varying degrees of calcification [13,14]. Osteoblastoma may have features similar to those of malignancy, such as cortical destruction and extra-osseous soft tissue expansion [13,19].

Other relevant imaging methods for this tumor type include computed tomography (CT), bone scan, and magnetic resonance imaging (MRI). CT can detect small mineralization in the tumor, which can help with diagnosis. CT also provides information about the size and extent of the lesion in the cortical bone, and aids preoperative evaluation and planning for surgery. Bone scans are also useful for localizing lesions, which appear as increased activity areas, but the scintigraphic appearance of the tumors is nonspecific. MRI findings of osteoblastoma are not specific, with a low signal in T1-weighted images and a high signal in T2-weighted images. However, MRI can reveal the extent of intra-osseous and soft tissue extension more accurately than other methods [4,13].

Although typical for an osteoblastoma, cortical attenuation may also be seen in other benign lesions such as fibrous dysplasia, aneurysmal bone cysts, and giant cell tumors [19].

The pathological features of osteoblastoma are similar to osteoid osteoma; both are bone-producing lesions arising from osteoblasts. Usually, the lesion is reddish-brown, even pinkish; friable with a gritty consistency; and hemorrhagic in appearance [9].

Osteoblastoma can be aggressive if associated with large epithelioid osteoblasts, and carries risk of local recurrence or malignant transformation into osteosarcoma [5]. These “aggres-
sive osteoblastomas" are more mitotically active than conventional osteoblastoma [14].

Benign osteoblastoma has a reported recurrence of approximately 10–20% [9,12,19]; therefore, close follow-up monitoring of patients is required. The size, location, and ability to resect the entire lesion are important factors influencing the risk of recurrence, morbidity, and mortality, although metastasis has rarely been documented [14]. The patient in this case study was followed-up closely for two years and sequential films revealed a radiopaque interface and no evidence of local recurrence.

Simple curettage with or without bone grafting results in less local morbidity but a higher rate of recurrence [2,15,20]; total removal is required to eliminate recurrence. Complete excision may be possible by intra-lesional curettage aided by high-speed burr and adjuvant cryotherapy [2], or in expendable bones such as ribs, fibula, or distal ulna, by en bloc resection [8]. Cementation in conjunction with excision may be helpful for extending the surgical margin. Postoperative radiation is usually not recommended; however, it may be considered for tumors that have recurred multiple times, particularly around the spine [11,16].

The majority of patients are cured by initial therapy. Recurrences usually occur within two years of initial surgery because of growth of remaining tumor cells; recurrences after two years are very rare. In the present case, we successfully applied curettage (excisional biopsy) and filled the resulting gap with polymethylmethacrylate filling without any complication or recurrence for two years postoperatively. Depending on the size of tumor or preference of surgeon, the curettage cavity is filled with bone graft or polymethylmethacrylate [20]. The treatment approaches and outcome in previous reports [2,16,20] and for the present case were summarized in Table 1.

Case reports of osteoblastoma in the small bones of the hands describe swelling as the main feature of metacarpal and phalangeal lesions and curettage and bone grafting as sufficient treatment [18]. Mosher and Peckham [17] described a case of osteoblastoma of the fifth metacarpal in a 10-year-old boy that was treated with en bloc excision and fibular grafting. Although usually a solitary bone tumor, Adler [1] described a case with multiple osteoblastomas occurring in multiple bones of the right hand in a nine-year-old child. All lesions identified radiologically were treated by curettage.

The phalanx is involved in less than 5% of cases, making this localization exceptional [3,14]. At least one case of benign osteoblastoma, localized on the proximal phalanx of the fifth digit of the left hand, was treated by en bloc surgical resection of the lesion and curettage [10]. In a summary of six cases of osteoblastoma of the hand by Farzan and Mortazavi [7], half were on the metacarpals and the other half were on phalanges; recurrence was observed in one case.

In conclusion, to the best of our knowledge, this is the first report of osteoblastoma on the distal phalanx. Although rare, osteoblastoma can be localized on phalanx. Unless the lesion is not very large, excisional biopsy followed by cementation with polymethylmethacrylate may effectively treat osteablastoma without causing recurrence for up to two years.

**Conflict of interest**

There is no conflict of interest.

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None.

**Consent**

In our study informed full consent was taken from the patient, and we did not need any detailed information about the patient.

**Author contribution**

Gumustas, Seyit Ali: surgeon and one of the author.
Cagırmaz, Talat: study designer, data collector.
Guler, Olcay: study designer, one of the author.
Öfluoglu, Onder: surgeon, supervisor.
Kayahan, Sibel: pathologist.

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