Parosteal lipoma of humerus with a medical history of 24 years: a case report

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Abstract: Parosteal lipoma is an extremely rare benign tumor, responsible for less than 0.1% of primary bone neoplasms and 0.3% of all lipomas, which mainly constitutes of mature adipose tissue with a bony component. Bony lesions are often discovered in patients with this tumor (59.2%), creating the significance of differential diagnosis of malignant tumors. The tumor is known to be related with underlying bony changes including focal cortical hyperostosis, pressure erosion of the underlying bone, and bowing malformation. Here we state a case of a parosteal lipoma appearing in the humerus with a bony excrescence in a 70-year-old male patient, which had the longest medical history (24 years) in documents of the last 50 years. Twenty-four years after the mass was found, he presented with complaints of night pain. MRI presented a large 11 cm × 6 cm × 5 cm well defined, lobulated mainly fat intensity lesion with a small part of chondroid component measuring 2.7 mm × 1.9 mm × 0.9 mm in proximal left humerus. The mass was excised, and the pathology demonstrated ossifying parosteal lipoma without features of malignancy. The patient had no recurrence and no complaints for 2 years after the operation. In conclusion, orthopedic surgeons should bear the diagnosis of parosteal lipoma in mind, thus providing appropriate treatment. Surgery is usually ideal treatment, which requires that any periosteal involvement is removed completely.

Keywords: Parosteal lipoma; humerus; case report; diagnosis; treatment

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Introduction

Parosteal lipoma is an extremely rare benign tumor, being responsible for less than 0.1% of primary bone tumors and only 0.3% of all lipomas (1). This tumor consists mainly of mature adipose tissue and shows contiguity to periosteum of the underlying bone. It is believed to be one of the rarest primary bone tumors, which comprised mostly of mature adipose tissue with a bony component. The most common sites of this tumor are femur followed by proximal radius. This has become great importance of differential diagnosis of malignant tumors, due to bony lesions are found in almost half of patients with this tumor. We hereby describe a case of parosteal lipoma of the humerus in a 70-year-old male patient, which had the longest reported medical history (24 years) in the last 50 years. We present the following article in accordance with the CARE reporting checklist (available at http://dx.doi.org/10.21037/aoj-19-152) (2).

Case report

A 70-year-old male patient appeared with a 24-year history of mass on his left upper arm. In 1991, the mass firstly appeared with the size of a bean and asymptomatic. The mass enlarged slowly over the years. In April 2015, the mass displayed the size of a fist and caused night pain. No interventions had been done before. The patient had no history of trauma or other prior similar tumors. There was no drug history or family history.

On clinical examination On April 27, 2015, an approximately 7 cm × 5 cm sized mass was revealed on
left upper arm, with an oval shape, elastic, immobile and nontender. Pain was complained while the left shoulder joint moved. There was no sensory deficit in any part of the left arm.

X-ray image showed a radiolucent soft-tissue mass contiguous with left proximal humerus and irregular osseous protuberance of subjacent bone (Figure 1A). On the magnetic resonance imaging (MRI), a 11 cm ×6 cm ×5 cm fat intensity lesion with underlying bony excrescence measuring 2.7 cm ×1.9 cm ×0.9 cm. (D) Postoperative X-ray. Left humerus was normal and showed no bony excrescences.

On May 5, 2015, total tumor resection was performed under brachial plexus anesthesia. Tumor was discovered to be situated on the humerus with tight continuity, with minute areas of spiculation palpable on the bone surface. On histopathology (Figure 2), a close relationship has been demonstrated between the lesion composing of mature lipocytes and the periosteum, consistent with parosteal lipoma.

The patient was satisfied and experienced neither movement disorder nor dysesthesia after the surgery. Postoperative X-ray (Figure 1D) of left humerus was normal and presented no bony excrescences. He had no recurrence and no complaints at latest follow-up (June 2020, 5 years after the operation).

All procedures performed in studies involving human participant were in accordance with the ethical standards of the institutional and national research committees and with the Helsinki Declaration (as revised in 2013). Written informed consent was obtained from the patient for publication of this manuscript and any accompanying images.

**Discussion**

Parosteal lipoma is a rare type of lipoma, “Periosteal lipoma” had been its name once, while it was replaced by “parosteal lipoma” to indicate that the tumor adjacent to

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**Figure 1** Timeline of the patient’s X-ray images. (A) Pre-operative X-ray showed a radiolucent soft-tissue mass contiguous with left proximal humerus and irregular osseous protuberance of subjacent bone. (B,C) Magnetic resonance imaging (MRI). A 11 cm ×6 cm ×5 cm fat intensity lesion with underlying bony excrescence measuring 2.7 cm ×1.9 cm ×0.9 cm. (D) Postoperative X-ray. Left humerus was normal and showed no bony excrescences.
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Figure 2 Histopathology (HE, ×100). The lesion was composed of mature lipocytes.

The bone but not necessarily originate from periosteal (3). These lesions are typically obtained on the diaphyses of long bones in middle-aged people (4). Followed by the location of proximal radius, the most common sites are femur. The lesion has also been reported originating in scapula, ribs, clavicle, metacarpals, pelvis, metatarsals, mandible and skull (3), perhaps almost all the bones in the body are affected.

According to the degree of chondroid modulation and endochondral ossification, these tumors are classified into 4 subtypes: (I) no ossification; (II) pedunculated exostosis; (III) sessile exostosis; and (IV) patchy chondro-osseous modulation. As revealed in our case, the focal hyperostotic change mimicking malignancy is the most common finding in the underling bone.

The radiographs clearly illustrated that parosteal lipoma is a well-defined area of lucency located adjacent to a long bone. About 60% of parosteal lipomas may have potential bony alterations, mostly hyperostotic reactive changes (fine linear densities, calcification, cortical thickening or undulation, or frank excrecences of bone), but these lipomas also have cortical bowing (in patients with growing bones), smooth cortical erosions or underlying osteochondroma (5). Bone destruction was absent.

On computed tomography, parosteal lipomas usually present as well-defined fat density mass with lobulated appearance adherent to underlying bone. The presented osseous excrecences can be distinguished from an osteochondroma, since it lacks of contiguity of the marrow space with the adjacent bone. The CT images are helpful in evaluating the relationship of the mass with the adjacent bone, which is important for surgical planning.

MRI is considered most valuable for evaluation of parosteal lipoma. On MRI, despite of pulse sequence, the tumor is recognized as a juxtacortical mass with same signal intensity to that of subcutaneous fat. Low-signal-intensity strands on T1-weighted images in the lesion, corresponding to fibrovascular strands, can be differentiated from those of well-differentiated liposarcoma, as these are thin and lack postcontrast enhancement (6). Adjacent muscle atrophy, caused by associated nerve entrapment, is identified on MRI as general pattern of increased striations of fat in the affected muscle. This finding is best valued on T2-weighted images because of the reduced signal intensity of normal muscle relative to fat. Cartilaginous components in parosteal lipoma showed intermediate signal intensity on T1-weighted images but high signal intensity on T2-weighted images (7). From the above, MRI is the best approach to exhibit the relationship between the tumor and the underlying bone, which is crucial for surgical planning.

Pathologically, a parosteal lipoma is usually a multilobulated mass circumscribed by a thin, fibrous capsule, and it is well encapsulated with a broad-based attachment to the underlying bone (8). Under the microscope, the adipocytes of a parosteal lipoma occur identical to fat cells found in the subcutaneous tissues. There are no reports that this tumor undergoes malignant transformation. The differential diagnosis with low-grade liposarcoma may be difficult, despite there are no previous reports of either primary parosteal liposarcoma or degenerated benign parosteal liposarcoma with these parosteal lipomas of patients (9).

The treatment of parosteal lipoma is intact surgical excision with further subperiosteal dissection, osteotomy, or segmental resection of the bone in cases with hyperostosis (10,11). The prognosis is good and local recurrence is unusual. Due to its negligible malignant potential, parosteal lipoma can be followed conservatively. In our case, it would be better if an X-ray or MRI was performed at the time of 2-year follow-up.

Conclusions
Parosteal lipomas are benign with an excellent prognosis. Surgery, which is ideal treatment, requires particular attention to ensure that any periosteal involvement is removed completely. For purpose of providing suitable treatment, parosteal lipoma should be incorporated in the differential diagnosis of soft tissue tumors.
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Footnote

Reporting Checklist: The authors have completed the CARE reporting checklist. Available at http://dx.doi.org/10.21037/aoj-19-152

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/aoj-19-152). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All procedures performed in studies involving human participant were in accordance with the ethical standards of the institutional and national research committees and with the Helsinki Declaration (as revised in 2013). Written informed consent was obtained from the patient for publication of this study and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

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References


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