Phosphaturic mesenchymal tumor

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ABSTRACT

Phosphaturic mesenchymal tumors (PMTs) are very unusual tumors, commonly associated with tumor induced osteomalacia (TIO). The tumor cells produce a peptide hormone-like substance known as fibroblast growth factor 23 (FGF23), which regulates phosphate levels. When FGF23 is over expressed by the tumor cells it leads to phosphaturia, which in turn causes mobilization of calcium and phosphate from bones, and decreased osteoblastic activity, which finally leads to severe osteomalacia. Usual presentation of PMT will be gradual muscular weakness and diffuse bone pain from pathologic fractures. The majority of PMTs are benign, malignant PMTs are extremely rare, but exist. However, very rarely PMT present without phosphaturia and TIO i.e., the nonphosphaturic variant. The challenge in diagnosis of PMT is due to the vague nature of the symptoms and lack of clinical suspicion. The diagnosis is often missed due to the rarity of PMTs. The usual location for PMT involves the extremities followed by head and neck sites. We report a case of a 37 year old male who presented to the outpatient department with a swelling in the left parotid region. He had no associated symptoms. CT scan showed a well-defined heterogeneous mass measuring 2.3 × 2.0 cm with calcification and soft tissue density seen superficially in the left pre auricular region. Per operatively the tumor was found to be intramuscular, embedded in the left masseter muscle. The final histopathology report was confirmative of phosphaturic mesenchymal tumor.

Keywords: Phosphaturic mesenchymal tumors, Tumor induced osteomalacia, Nonphosphaturic

INTRODUCTION

Phosphaturic mesenchymal tumors (PMTs) are very unusual tumors which are commonly associated with tumor induced osteomalacia (TIO), the paraneoplastic syndrome associated with PMTs present as phosphaturia. The tumor cells produce a peptide hormone-like substance known as fibroblast growth factor 23 (FGF23), which regulates phosphate levels.1 When FGF 23 is over expressed by the tumor cells, it leads to phosphaturia, mobilization of calcium and phosphate from bones, and decreased osteoblastic activity, which finally leads to extensive osteomalacia. Patients usually present with gradual muscular weakness and diffuse bone pain from pathologic fractures.2 The majority of PMTs are benign, malignant PMTs are very rare, but exist.1 However, rare cases of PMT present without phosphaturia and TIO i.e., the nonphosphaturic variant.4 The diagnosis is often demanding due to the non-specific nature of the symptoms and lack of clinical suspicion. The pathologic diagnosis is often missed due to the rarity of PMTs and histologic overlap with other mesenchymal neoplasms such as enchondroma and hemangiopericytoma.5-6 The most common location for PMTs includes soft tissues, bones of the extremities followed by head and neck sites, with the paranasal sinuses and nasal cavity accounting for more than half of the cases in the head and neck region.7,8

CASE REPORT

In this article, we report the case of a 37 year old male who presented to the outpatient department with a swelling in the left parotid region. He had no associated symptoms. Clinically he was suspected to have a lesion
arising from the left parotid gland (Figure 1). CT scan showed a well-defined heterogeneous mass measuring 2.3 × 2.0 cm with calcification and soft tissue density seen superficially in the left pre auricular region. CT was suspicious of a soft tissue tumor or lymph node (Figures 2 and 3).

Figure 1: (A) Lesion, (B) zygoma, (C) angle of mandible.

Figure 2: (A) Lesion, (B) masseter muscle, (C) branch of facial nerve, (D) parotid gland.

Figure 3: (A) Lesion, (B) facial nerve, (C) parotid gland.

Figure 1 (A-H): CT head axial view showing lesion.

Figure 3 (A and B): CT head coronal view, lesion in left zygomatic region

Figure 6 (A-C): Histopathological pictures of PMT.
A left sistrunk incision was placed. Per operatively the tumor was found to be intramuscular and deep to the superficial lobe of parotid gland, embedded in the left masseter muscle. To minimize the injury to facial nerve we proceeded with a left superficial parotidectomy. A 2 × 2 cm white firm lesion was excised out by splitting the left masseter muscle (Figures 4 and 5). The lesion was sent for histopathology examination. Postoperatively patient had neuropaesthesia of the orbital branch. Histopathology showed hyalinised tissue with scattered blood vessels, foreign body giant cell, lymphoid nodule and focal areas of osseous metaplasia which was suggestive of PMT (Figures 6 A, B and C). After a diagnosis of PMT was made serum phosphorus, alkaline phosphatase and Vit D3 was done on follow up. Serum phosphorus and alkaline phosphatase was normal. Serum VitD3 was low, hence VitD3 supplementation was given.

DISCUSSION

PMTs are rare neoplasm associated with tumor induced osteomalacia and is often under recognized by clinician and pathologist.4 This problem is aggravated by the fact that several variants of PMTs exist to date no consistent or defining histopathological and immune-phenotype has been established.5 This can be reemphasized in the series by Folpe et al, even after expert review five cases remain unclassified as PMTs.9

The paraneoplastic syndrome associated with PMTs manifest as phosphaturia. The tumor cells produce a peptide hormone-like substance known as fibroblast growth factor 23 (FGF23), a physiologic regulator of phosphate levels. Over expression of FGF23 by the tumor cells leads to phosphaturia, mobilization of calcium and phosphate from bones, and the lessening of osteoblastic activity, eventually resulting in extensive osteomalacia. Patients normally present with gradual muscular weakness and diffuse bone pain from pathologic fractures.10 The majority of PMTs are benign, malignant PMTs are exceedingly rare, but exist.11 PMTs without phosphaturia i.e., non phosphaturic variant are even rarer. The non phosphaturic variant is even harder to diagnose for its lack of symptoms because symptoms of osteomalacia more or less always precedes the clinical presentation of PMTs.12

Phosphaturic mesenchymal tumors are of the mixed epithelial, connective tissue type and mixed connective tissue type [MCT], of which MCT is the most common one.5,6 Histopathologically, fibrous capsule, calcification and giant cell reaction is observed in soft-tissue PMT-MCT, while PMT-MCT of bone and multiple PMT-MCT showed an infiltrative growth pattern. Non-phosphaturic PMT also shows histopathological features of mixed connective tissue type.13,15

The most common location for PMT involves soft tissues, bones of the extremities followed by head and neck sites. Roughly 95% of PMTs is seen in the extremities and appendicular skeleton, only 5% of the tumor is found in the head and neck region. Localization in the head and neck is very rare, with the paranasal sinuses and nasal cavity representing more than half of the cases.7,8,16

Regrettably, preoperative phosphate and serum FGF23 levels are not available for patients with “non-phosphaturic” variant of PMT so far. There have been incidence wherein clinically noteworthy osteomalacia was diagnosed only after the biopsy analysis of PMT was done, emphasizing for the pathologist the significance of identifying the histopathological features of this tumor, even when a history of TIO is not given.12 Many other benign mesenchymal tumor types have been associated with TIO, such as enchondroma and hemangiopericytoma.10 However, majority of the reports reviewed has shown that PMTs are misdiagnosed, due to the uncommonness of these tumors and lack of acquaintance and experience of the pathologists with their histopathological features.

Even though PMTs are rare tumors and even if there is no clinical symptoms of Osteomalacia it should be considered as differential diagnosis for tumors in head and neck region.

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REFERENCES