Case Report

Dorsal vertebral body tumor and non-compressive quadriparesis – A rare case report of a phosphaturic mesenchymal tumor

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INTRODUCTION

Quadriparesis is a symptom of varying etiology and is investigated, diagnosed, and treated by the neurologist, the neurosurgeon, the spine surgeon, and the physician, alike. Proximal muscle weakness narrows down the spectrum which ranges from infectious agents including influenza, Epstein-Barr virus, and Human immunodeficiency virus in patients with associated risk factors to the use of alcohol or steroids. Neurologic conditions that can cause proximal weakness include cerebrovascular disease, brain as well as spinal tumors, myopathies, demyelinating disorders (i.e., multiple sclerosis, Guillain-Barré syndrome), and neuromuscular disorders (i.e., myasthenia...
Localizing neurologic deficits can help the physician to focus the diagnostic work-up; however, one always encounters the odd case where despite all efforts, no answer is found amidst the common causes.

A phosphaturic mesenchymal tumor (PMT) is a neoplasm most commonly of mesenchymal origin, arising from and involving bone or soft tissue. The tumor produces a phosphaturic substance, fibroblast growth factor-23 (FGF23) that causes renal phosphate loss and subsequent systemic phosphate depletion, commonly leading to oncogenic osteomalacia, which is its most common presenting feature. Rarely hypophosphatemia may present as proximal muscle weakness and must be considered as a differential for the same. The diagnosis is often delayed due to the lack of clinical suspicion and nonspecific nature of the symptoms and, hence, it is important to be aware of this entity as one of the easily reversible causes of hypophosphatemia and resultant quadriplegia.

Surgery for spinal tumors is usually performed to decompress the neural elements and/or for spinal stabilization. We present a rare case where a hemivertebrectomy was performed to excise a small vertebral body PMT that was the cause of a paraneoplastic syndrome resulting in non-compressive quadriplegia.

CASE PRESENTATION

A 34-year-old gentleman presented to the clinic with complaints of weakness of all limbs, proximal greater than distal, for the past 3 years along with nonspecific mild back pain. His weakness had been gradually progressive, and for the past 1 year, he could no longer walk independently and needed support for daily activities. For the past 10 months, he was bedridden and severely functionally restricted. Examination revealed findings corroborated with the history. While the tone was normal, muscle power in the shoulders and hips bilaterally was ⅗ (MRC Grading), bilateral elbows and knees ⅘, and bilateral ankles, wrists, and grip 5/5 suggesting a predominant proximal muscle weakness. Sensations over the body and deep tendon reflexes were normal. An magnetic resonance imaging (MRI) whole spine was performed that revealed a normal result with no structural pathology that could explain the weakness.

Laboratory tests revealed hypophosphatemia (serum phosphorus levels of 1.84 mg/dl), which explained his proximal muscle weakness, with >5% fractional phosphorus excretion in the urine. An avid focus on DOTATOC positron emission tomography (PET) scan [Figure 1] and a corresponding hyperdense focus on a plain computed tomography (CT) [Figure 2] suggested the possibility of a PMT of the left side of D10 vertebral body. Even on revisiting the MRI no lesion could be identified at D10. Correction of hypophosphatemia was commenced with phosphorus supplements and calcitriol; however, there was no improvement in the serum phosphorus levels despite correction due to persistent phosphatura and no clinical improvement before surgery was observed. He underwent a right D10 costotransversectomy and hemivertebrectomy where the sclerotic lesion was drilled and excised. The entire area corresponding to the hot spot in the PET scan was drilled, along with a 3–4 mm margin [Figure 3]. Within the third postoperative day, he had subjective improvement in his symptoms and by 14 days made a near-full recovery. His phosphate levels gradually normalized [Figure 4], muscle power returned to normal in all muscle groups and he could walk with minimal support. Histopathology revealed bony fragments infiltrated by a cellular tumor, composed of spindle cells in fascicles having an oval nucleus with moderate clear to pink cytoplasm and minimal atypia.
with no mitosis [Figure 5]. Immunohistochemically, the tumor cells stained positive for Vimentin, thus confirming its mesenchymal origin [Figure 6]. A PET scan performed 1 year later confirmed no locoregional or distant recurrence [Figure 7]. On a 5-year follow-up, he is fully functional with no residual neurological deficit and no evidence of clinical recurrence.

**DISCUSSION**

PMTs are rare tumors of mesenchymal origin. Most tumors are found in soft tissues, with bone being the second most common site.\(^1\)

**Presenting features**

They are termed “phosphaturic” as a factor produced by the tumor cells, FGF23 also termed “phosphatonin,” promotes renal phosphate excretion, leading to chronic and rarely acute hypophosphatemia, in turn, leading to generalized weakness, malaise, and in severe cases hypotension, congestive heart failure, rhabdomyolysis, respiratory failure, seizures, altered mental status, and even death.\(^1\) As the initial symptoms of acute hypophosphatemia are nonspecific, it may not initially

**Figure 3:** Post-operative computed tomography scan showing complete excision of the hyperdense focus with the surrounding bone, amounting to a right D10 hemivertebrectomy.

**Figure 4:** Normalization of serum phosphorus levels after tumor resection.

**Figure 5:** Histopathological examination of the tumor showed spindle cells with an oval nucleus, minimal atypia, and no mitosis.

**Figure 6:** Immunohistochemical staining showing a positive stain for Vimentin, a marker for mesenchymal cells.

**Figure 7:** 18-FDG positron emission tomography computed tomography on follow-up shows no evidence of any locoregional or distant hot spot suggesting no recurrence or metastasis.
be suspected or recognized when a patient first presents to the clinic. Chronic hypophosphatemia on the other hand results in defective bone mineralization, ultimately leading to osteomalacia in adults and rickets in children.\[^3\] Tumor-induced osteomalacia, as this is known, occurs usually with PMTs but may also be seen in some other mesenchymal tumors (e.g., hemangiopericytomas). The symptoms of osteomalacia and rickets include weakness, fractures, bone pains, and bowing of long bones,\[^2\] which constitute the most common presenting symptoms of PMTs.

Due to the rarity of this condition, most cases, as in our case, usually suffer for years before being diagnosed. Most are treated nonspecifically and symptomatically for long periods at primary and secondary care centers, before finally being diagnosed.

Pathogenesis and the role of FGF23

A high FGF23 level in a patient with normal renal function, and hypophosphatemia, with or without osteomalacia, is suggestive of a PMT.

FGF23 was first discovered in 2000, in autosomal dominant hypophosphatemic rickets by White \etal[^5]. The best-established receptor of FGF23 is FGF receptor-1 (FGFR1). Agaimy \etal. 2017[^1] analyzed a series of 22 PMTs using an extended immunohistochemical marker panel, examined them by fluorescence \textit{in situ} hybridization, and found that FGFR1/FN1 gene fusions were detected in roughly half of the cases. Mutations in FGF23 that render the protein resistant to proteolytic cleavage lead to increased activity of FGF23. Its main function seems to be the regulation of phosphate concentration in plasma. It is secreted by osteocytes in response to elevated calcitriol and inhibits renal Pi reabsorption by reducing the apical expression and activity of NaPi-IIa in the proximal tubule epithelium.\[^3\] FGF23 also reduces intestinal absorption of dietary Pi through a Vitamin D receptor-dependent decrease in NaPi-IIb activity. Hence, significant hypophosphatemia ensues.

Histopathology

Histologically, PMTs are characterized by a proliferation of bland spindle cells associated with a variable amount of a “smudgy” calcified matrix. Mitosis and atypia are extremely rare and most tumors are benign. However, a very small percentage does exhibit nuclear atypia and a high Ki-67 index, and this subset might also be clinically malignant. There is also a small percentage of tumors that may be non-phosphaturic, and these pose a diagnostic challenge. Agaimy \etal. in their series showed consistent expression of CD56 (100%), ERG (90%), and SATB2 (90%) on immunohistochemistry of PMTs.\[^1\]

Management

In phosphaturic or “classical” PMTs, before surgery or when surgery is not indicated, oral phosphate and calcitriol can alleviate symptoms and metabolic imbalance.\[^6\]

Complete tumor resection confers a good prognosis in most patients. However, surveillance for recurrence and metastasis is necessary, although most tumors do not recur or metastasize.

CONCLUSION

PMTs must be included in the differential diagnosis of patients presenting with quadripareisis with low serum phosphate levels. Elevated serum FGF-23 levels are diagnostic with a fair level of accuracy. Locating the tumor might prove challenging due to its small size at presentation and the possibility of involving almost any mesenchyme-derived structure in the body. DOTATOC PET CT can help locate these tumors when clinical and laboratory findings strongly suggest the same. Once diagnosed, complete surgical excision is the recommended treatment. Phosphate supplementation can hasten symptomatic recovery in operated cases and might alleviate the symptoms to a certain extent in inoperable or high-risk cases. The return of phosphate levels to normal usually provides excellent results and patients returning with recurrence are not common.

Declaration of patient consent

Patient’s consent not required as patient’s identity is not disclosed or compromised.

Financial support and sponsorship

Nil.

Conflicts of interest

There are no conflicts of interest.

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How to cite this article: Bhatjiwale MM, Chandrachari KP, Kannan S. Dorsal vertebral body tumor and non-compressive quadriplegia – A rare case report of a phosphaturic mesenchymal tumor. Surg Neurol Int 2022;13:452.