A case of oncogenic osteomalacia detected by In-pentetreotide total body scan

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**ABSTRACT**

A case of tumor-induced osteomalacia in a 35-year-old woman suffering from severe bone pain and muscle weakness is described. This uncommon disease is characterized by a reduced serum phosphorus level with elevated urinary phosphate excretion, normocalcemia, high serum bone alkaline phosphatase and a deficiency of 1,25 dihydroxyvitamin D3. The tumors responsible for oncogenic osteomalacia are usually small, benign and commonly located in bone or soft tissues of the head and the limbs, so the diagnosis can often be difficult. In this case a 111In-pentetreotide scintigraphy was able to detect a hemangiopericytoma located in the right maxillary sinus. Removal of the tumor resulted in the reversal of clinical and biochemical abnormalities.

**Introduction**

Tumor-induced osteomalacia (TIO) is a rare and insidious paraneoplastic syndrome characterized by acquired hypophosphatemia, hyperphosphaturia and low serum levels of 1,25 dihydroxyvitamin D3. Usually, patients complain of muscle weakness, bone and muscle pain with gradual onset. These symptoms commonly involve weight-bearing areas such as the legs, ankles, hips and back. Following surgical removal of the tumor the clinical and biochemical abnormalities disappear, indicating that they are due to a humoral factor secreted by the tumor (1, 2).

The responsible tumor, often a benign mesenchymal tumor, may be located in almost any part of the body, with a large proportion occurring in the head and in the upper and lower extremities. To date there is no standard method for detecting these tumors. The finding of somatostatin receptors expressed by human mesenchymal tumors (3-5) opens a new pathway for the diagnosis of this disease, as demonstrated in some other cases (6).

Here we report a case of TIO in which the lesion was finally detected by 111In-pentetreotide scintigraphy and successfully removed.

**Case report**

A 35-year-old Caucasian woman was admitted to our hospital because of severe bone pain, polyarthralgia and muscle weakness. The patient had been healthy until two years before admission, when she began to complain of pain in the ankles and knees on weight bearing but gradually subsiding with rest. These symptoms seriously worsened in the last nine months. Her gait was anserine, the weakness and bone pain became so severe and disabling that the patient was not able to walk unassisted. No response to nonsteroidal antiinflammatory and steroidal medications (prescribed by the general practitioner) was reported. She had lost 9 Kg in weight in the last year and 5 cm in height. There was no family diagnosis of bone or renal disease. The patient did not have a history of risk factors for bone disorders. On physical examination there was no evidence of synovitis, but the motion of the hips was limited. There were no neurological deficits. Laboratory tests showed extreme hypophosphatemia 0.7 mg/dl (normal range, 2.5-4.6 mg/dl), mild hypocalemia 7.8 mg/dl (normal range, 8.4-10.2 mg/dl), an increase in total alkaline phosphatase of 194 U/L (normal range, 32-92 U/L) with bone isoenzyme as high as 72% (normal range, < 35%). Urinary phosphate excretion was 458 mg/24h (normal range, 400-1000 mg/24h), the renal clearance of phosphate was 26.3 ml/min (normal range, 6-15 ml/min). Urinary calcium excretion was 59 mg/24h (normal range, 100-300 mg/24h). The level of 1,25(OH)2D3 was low (7 pg/ml; normal range, 14-50 pg/ml) with normal plasma concentrations of 25(OH)D3. Other laboratory investigations, including parathyroid hormone, protein electrophoresis, creatinine, liver and thyroid function tests, muscle enzymes, erythrocyte sedimentation rate, blood cell count were in the normal range.

On plain radiographs many thoracic and lumbar vertebrae showed bowing of the endplates without abnormalities in the remainder of the skeleton, except for a diffuse loss of radiodensity. Bone densitometry (DXA; Hologic 4500) revealed decreased bone mass in the spine and normal values in the femoral neck. The value of the lumbar measure-
ment (L2-L4) was 0.779 g/cm² corresponding to -2.72 T score.

A bone scintigraphy was performed: the acquisition was done 3 hours after the injection of 555 MBq (15 mCi) of ⁹⁹mTc metilene-diphosfonate, whole body technique, with a double head gamma camera PRISM 2000 XP (Picker) matrix acquisition 1024 x 256 pixel (2000 KC count). Diffuse increased uptake corresponding to the meta-epiphysal region of both femurs, tibiae, radii and humeri was discovered. An enhanced radiotracer uptake was also shown by six ribs and four vertebrae, probably due to recent fractures (Fig. 1).

The clinical, laboratory and radiological features were consistent with the diagnosis of hypophosphatemic osteomalacia. A bone biopsy of the iliac crest showed a thickened layer of unmineralized osteoid consistent with osteomalacia. Further investigations excluded primary renal and bowel diseases.

In the search of an underlying tumor a total body scan was performed with octreotide 4 and 24 hours after injection of 111 MBq (3 mCi) of ¹¹¹In-octreoscan (Byk Gulden-Italia). Two large field of view digital SPET cameras (General Electric-Millennium VG) with medium energy, general purpose collimator and two energy peacks of 171 and 245 keV (20% window) were used. Total body images in anterior and posterior views (scan speed 12 cm x min−1, matrix acquisition 128x512) were supplemented with appropriate planar views. Abnormal ¹¹¹In-octreoscan tumor uptake was defined as any focal or diffuse area of increased activity in a location incompatible with normal anatomy. In this case an increased uptake in the right maxillary sinus was disclosed (Fig. 2). Magnetic resonance imaging (MRI) confirmed the presence of a solid round mass, 2 cm in diameter, in the right maxillary sinus. There was no destruction of the bone wall (Fig. 3).

While waiting for surgery, the patient received daily supplementation with phosphate 2 g, calcium carbonate 2 g, 1.25(OH)₂D₃ 1.5 mcg. The serum phosphate level increased up to a maximum value of 2.3 mg/dl. The urinary phosphate excretion increased as well.
and reached a maximum value of 2145 mg/24h. Removal of the tumor resolved the biochemical abnormalities in a few days. Two weeks after surgery and withdrawal of oral therapy, her serum phosphate level increased up to 4 mg/dl and there was a rapid decrease in urinary phosphate excretion to 609 mg/24h. The value of 1.25(OH)₂D₃ returned to the normal range. As shown in Figure 4, histologically the tumor was composed of epithelioid cells, range from round or oval to slightly spindled and are arranged in a various patterns including nests and cords, with elaborated and ramifying blood vessels (pattern hemangiopericytoma-like). Cytological atypis was minimal and mitoses were rare. The cells are separated by osteoid matrix and focally by hyalinized collagen.

Bone densitometry performed 8 months after surgery showed a value for the lumbar scan of 0.950 g/cm² corresponding to a percentual increase of 21.9%.

The patient showed progressive improvement in her symptoms with a complete recovery in three months.

Discussion
After the description of an acquired osteomalacia resistant to vitamin D supplementation by McCance in 1947 (7), Prader et al. in 1959 first described a case of oncogenic osteomalacia in which a tumour was recognised as the cause of osteomalacia (8), and to date about 120 cases of TIO have been reported in the literature. The majority of patients are adults over 30 years (range 1-74), with a male:female ratio of 1.2:1. The duration of symptoms before diagnosis ranges from 2.5 months to 19 years, with an average of more than 2.5 years (1). The musculoskeletal symptoms of TIO (gradual onset of muscular weakness and bone pain on weight-bearing areas such as legs, ankles hips and back; fatigue and gait disturbances) are non-specific and unrevealing. The radiological features of osteopenia and pseudofractures and the bone histology are highly suggestive for osteomalacia. The biochemical profile is characterized by an impressive

Fig. 3. Coronal SE T2-weighted MR image of the skull reveals opacification of the right maxillary sinus. A round soft tissue mass, 2 cm in diameter, can be seen along the inferior aspect of the sinus. The mass shows homogeneous, intermediate signal intensity. No destruction of the bone wall is detected.

Fig. 4. The histology of the tumor: pattern hemangiopericytoma-like (x10, x25) (see text).
reduction of serum phosphorus level and an elevated urinary phosphate excretion related to a low tubular reabsorption of phosphate (TRP). Serum calcium in the normal range or slightly decreased, high serum bone alkaline phosphatase and a deficiency of 1,25(OH)D$_2$ are typical biochemical features as well. Serum levels of 25(OH)D$_3$ and PTH are usually in the normal range.

The tumors responsible for oncogenic osteomalacia have been described as ‘strange tumors in strange places’ (9); these neoplasms are often small in size (1 to 7 cm), and are commonly located in the bone or soft tissues of the lower extremities and craniofacial regions.

About 40% of these tumors are vascular, with hemangiopericytoma features being the most common. Although most tumors are benign, several malignant cases have been reported (10, 11).

The way in which these tumors lead to hypophosphatemia is not yet understood. Useful observations have come from the study of X-linked hypophosphatemic rickets (HYP), which is phenotypically similar to TIO. Although there are some differences in clinical presentation, both diseases are characterized by an excess in circulating phosphatonin which inhibits sodium-dependent phosphate reabsorption in the kidney and decreases 1α-hydroxylase activity, resulting in hypophosphatemia and reduced 1,25 (OH)$_2$D$_3$ levels. Excess of phosphatonin in HYP is due to impaired degradation caused by mutation in the gene PHEX, which encodes for the endopeptidase responsible for phosphatonin catabolism. In TIO a large amount of phosphatonin, overwhelming endogenous PHEX activity, is likely to be generated by the tumor itself (12-14). Phosphatonin is a low molecular weight (8-25 KD) heat sensitive substance (15); its inactivation by trypsin supports a likely peptide structure (16). Recent studies suggest that FGF-23 (a member of the fibroblast growth factor family that is overexpressed in tumors associated with TIO) could be identified with phosphatonin (17).

The diagnostic challenge is related to the identification of these insidious neoplasms. In several patients TC scan did not identify the tumor (18) and MRI scan, especially the whole-body one, is an expensive method and not always a certain diagnostic tool.

The discovery of somatostatin receptors on mesenchymal tumors have disclosed a new diagnostic path. Somatostatin receptors have been found in many normal tissues such as the gastrointestinal tract, central nervous system, anterior pituitary gland, endocrine and exocrine pancreas, activated immune cells, but have also been observed in high concentration in human neuroendocrine tumors stemming from the neural crest. Five different subtypes of somatostatin receptors have been identified: the subtypes 2, 3 and 5 show a high responsiveness to a somatostatin analogue such as octreotide (4). In this regard, the 111In-pentetreotide body scan can be considered a useful tool not only for detecting these tumors, but also for a therapeutic approach. As demonstrated in this case, scintiscan using radiolabelled somatostatin analogues can play a pivotal role in detecting a mesenchymal tumor related to an osteomalacia that is inexplicable in other way (4, 5).

References
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