Diagnosis and Clinico-Radiological Presentation in an Aggressive Maxillary Brown Tumour

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ABSTRACT

Brown's tumour is an uncommon focal giant cell lesion which arises as a result of the effect of increased parathyroid hormone on bone tissues in hyperparathyroidism. The mandible is the predominantly affected site in the maxillofacial area and a maxillary involvement is rare. The severity of the lesion, caused by a Brown's tumour, may lead to evident osteolysis and gross deformity in the maxillofacial region, which suggests the need for making an early diagnosis and giving prompt treatment. We are reporting a male patient who presented with a massive painful swelling in the right maxilla as the first manifestation of primary hyperparathyroidism, caused by a parathyroid adenoma.

CASE REPORT

A 34-year-old male patient reported to the Department of Oral Medicine and Radiology, Government Dental College and Research Institute, Bangalore India, with the chief complaint of a swelling on the right side of the face of 4 months duration and pain of 2 months duration. The patient had difficulty in chewing food since 4 months, after which he had noticed a swelling on the right side of the face, which was insidious in onset, initially small in size and had gradually progressed to the present size. His medical history revealed that he suffered from epilepsy since 25 years and that he was under medication. On physical examination, the patient was found to be short stunted and his vital signs were in normal range. Extraoral examination revealed a diffuse swelling on the right side of the face, which led to gross asymmetry and the skin over the swelling appeared to be unaltered. On palpation, the swelling was found to be tender and firm in consistency. The submandibular lymph nodes on the right side of the face were examined, ten and on palpation and mobile. On intraoral examination, a diffuse swelling was seen on the right side of the hard palate, which roughly measured about 4 x 3 cm, with anterior extension from the distal aspect of the canine, posteriorly to the tuberosity area, which caused buccal displacement of the maxillary molar teeth and Grade III mobility, which caused hindrance in speech and mastication. The mucosa present over the swelling appeared to be inflamed, with indentation of the lower molar teeth [Table/Fig-1]. On palpation, the swelling was found to be tender and soft in consistency. The patient indicated that these lesions had been present since about 1 year, but that they had rapidly enlarged in size since 4 months. Based on the clinical findings, a provisional diagnosis of a benign soft tissue neoplasm was made, because of the site of occurrence, history and character. On routine radiographic examination, IOPA of maxillary molars and orthopantamograph revealed a generalized loss of lamina dura around teeth, with change in occlusal plane and a diffuse radiolucency above the apex of right maxillary molar teeth, which were suggestive of bony destruction [Table/Fig-2-4]. A radiographic differential diagnosis of primary hyperparathyroidism, osteomalacia and Paget's disease of bone was given. Computed tomography (CT) scan of head and neck which was done, revealed a well defined irregular heterogenous soft tissue attenuation, which measured 5.4 x 5.9 x 5.2 cm, which was an osteolytic, bone-expanding lesion seen in the anterior part of the right maxillary sinus, with invasion of the adjacent floor of the orbit, anterior ethmoids and nasal cavity. Post contrast studies showed an early intense enhancement with few non enhancing areas within the lesion [Table/Fig-5-7]. The laboratory findings were as follows: intact PTH 314 pg ml-1 (15–65 pg ml-1), calcium 12.0 mg dl-1 (8.5–10.5 mg dl-1), phosphate 6.7 mg dl-1 (2.7–4.5 mg dl-1) and alkaline phosphatase 283 U L-1.
Ultrasoundogram (USG) of the neck showed a well defined, homogenous, hypoechoic circular lesion which measured 1.5 x 2 cm, which was noted to lie inferior to the lower pole of left lobe of thyroid [Table/Fig-8]. The lesion was placed medial to the mid left common carotid artery and lateral to the cervical oesophagus, just deep in the subcutaneous plane, which showed minimal internal vascularity and was suggestive of a parathyroid adenoma. There was no evidence of significantly enlarged cervical lymph nodes. The thyroid gland appeared to be normal in size, shape and location. Ultrasound guided fine needle aspiration cytology (FNAC) revealed on microscopic examination; many multinucleated giant cells which were arranged in groups, which were adjacent to haemosiderin granules within a fibrovascular haemorrhagic stroma. These findings were compatible with a diagnosis of Brown’s tumour of the maxilla, which was associated with a parathyroid adenoma. Finally, to locate and define the extent of the spread of the tumour, a parathyroid Scaniscan was done after giving an IV injection of technetium-99m sestamibi, which showed a focus of abnormal accumulation of tracer in the location of the left inferior parathyroid gland, left submandibular region and right maxillary region [Table/Fig-9]. It was interpreted as a parathyroid adenoma which involved the left inferior parathyroid gland. Finally, after a thorough diagnostic work up, a diagnosis of Brown’s tumour of the maxilla, which was associated with primary hyperparathyroidism was given.

**DISCUSSION**

Hyperparathyroidism (HPT) was identified concurrently in Europe and United States in the mid 1920s. This disorder was first described in the setting of severe bone disease which resulted in significant morbidity. Incidence of primary hyperparathyroidism (PHPT) is 2-3/1000 in women and it is 1/1000 in men. The disease is most frequent in postmenopausal women during the sixth decade of life, although it is seen in all age groups. Among patients with PHPT, 80–85% develop the disease due to parathyroid adenoma, 15–20% develop it due to hyperplasia of the parathyroid gland, while in less than 0.5%, the cause is parathyroid carcinoma [1-3]. Brown’s tumours are rare, focal giant-cell lesions which arise secondary to both primary and secondary HPT and they have been reported to occur in 4.5% of patients with primary HPT and in 1.5 to 1.7% patients with secondary disease [4]. Some parathyroid adenomas and hyperplasias are familial (5% of cases), and others are a part of multiple type I, IIa, and IIb endocrine neoplasias [5]. The ribs, femur and pelvis are more commonly involved sites of Brown’s tumours [6]. This bony lesion of HPT is caused by increased circulating levels of PTH, which result in increased osteoclastic bone resorption, primarily in cortical bone [7]. This may explain as to why the mandible, a cortical bone, is the most commonly affected site, and why a maxillary involvement is less common in the maxillofacial area [8].

In some cases, blood calcium levels which are within normal limits may also be observed. Although many systems are affected in PHPT, the most pronounced alterations are observed in the bone. With new and more objective diagnostic PTH radioimmunoassay techniques, along with the successful treatment of the disease, bony complications of HPT have started to decline rapidly. The earliest and most reliable change which is seen in hyperparathyroidism is subtle erosion of bone on the subperiosteal surfaces of the phalanges [9]. The lesion usually presents as a slight swelling in the jaw bones. However, our case was similar to the case which was presented by Dinkar et al., [6] and this case differed from previously reported cases with respect to its gross maxillary destruction. Radiographically, Brown’s tumour appears as a well defined unilocular or multilocular radiolucency and is very similar to the other giant cell lesions like cherubism and aneurysmal bone cysts [10]. On clinical examination and on using only routine
panoramic radiography, the lesions may resemble bone metastases of a carcinoma, multiple myeloma, Langheran’s cell histiocytosis, osteosarcoma, Paget’s disease, osteomyelitis or osteonecrosis which is secondary to bisphosphonate therapy [8].

As it is difficult to distinguish Brown’s tumours from other giant cell lesions on the basis of histopathological and radiological examinations, the confirmatory diagnosis is made on the basis of hyperparathyroidism [11]. Brown’s tumours mainly result from secondary HPT in patients with renal insufficiency, but they have also been described as a rare manifestation of calcium malabsorption and some forms of osteomalacia [12]. Primary treatment of Brown’s tumour is surgical removal of associated parathyroid pathology. Brown’s tumours generally regress spontaneously over time and other treatment options include enucleation and curettage, radical resection and reconstruction, radiation therapy, and chemotherapy [13,14]. The most significant point about the case which has been described here, was the appearance of Brown’s tumours in the maxilla, which is rare and that the characteristic biochemical hallmark of primary hyper-parathyroidism was not seen. Thus, the patient whom we are describing to have normocalcaemic primary hyperparathyroidism was distinct from primary hyperparathyroidism patients who present with overt hypercalcaemia, in whom the serum calcium levels can occasionally be normal. From time to time, the serum calcium levels may be normal and most of the time, the serum calcium levels in these patients are elevated. Our case was similar to the case which was described by Bilizikian and Silverberg [15]. Maruani and Cols postulated that in normocalcaemic primary hyperparathyroidism, there was resistance to target tissues. After an oral calcium load, normocalcaemic subjects showed inadequate suppression of PTH when they were matched with a cohort of subjects who had hypercalcaemic primary hyperparathyroidism [16]. Surgical excisions of Brown’s tumours are indicated for large and disfiguring lesions and in cases where the affected bone is weakened, as it was seen in our case. However, other authors had initially treated such lesions with systemic corticosteroids and, when they had reduced in size, they had done surgical excisions [17]. The present report describes a rare case of a peripheral Brown’s tumour which was associated with primary HPT, which simulated a peripheral giant cell lesion of the jaws.

CONCLUSION
This case emphasizes the importance of making an early diagnosis of hyperparathyroidism by doing a thorough diagnostic work-up. This case should attract the attention of general practitioner dentists, oral and maxillofacial surgeons, endocrinologists and radiologists, whose consultations may be vital for patients with hyperparathyroidism, since this disease may result in nearly fatal results if it is neglected. Dentists should also be alert to the possible occurrence of Brown’s tumours in the jaws of patients who had previously been diagnosed to have hyperparathyroidism.

REFERENCES