Brown tumor of the palate as first manifestation of primary hyperparathyroidism: a case report

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(Reçu le 16 septembre 2011, accepté le 3 octobre 2011)

Abstract – Brown tumor is one of the lesions that develop in patients with hyperparathyroidism. Skeletal bones including maxillo-facial ones can be the site of this lesion. Owing to the improve methods of blood analysis most of cases of primary hyperparathyroidism are diagnosed early and asymptotically making advanced disease with bone lesions extremely rare. This article contains a case of a 43-year-old female patient who presented with palatal swelling as the first sign of primary hyperparathyroidism. The diagnosis was suggested by the histological findings and confirmed by the endocrinologic status.

Case report

A 43-year-old woman patient presented with a swelling in the anterior right region of the palate since 6 months. Extraoral examination revealed a swelling in the right middle third of the face (Fig. 1A). Intraoral examination a large palatal swelling measuring around 4 × 3 cm extending from the right premolar region to the left incisive region (Fig. 1B). Panoramic radiograph revealed an anterior radiolucency (Fig. 2A). Computed tomography scan showed a palatal destructive bone lesion extending to the floor of nasal cavity and expanding the cortical plate (Figs. 2B and 2C).

Incisionnal biopsy was made. Histopathological examination showed proliferation of multinucleated giant cells mixed
with mononuclear spindle shaped cells indicative of giant cell lesion (Fig. 3). Blood analysis revealed an elevated serum calcium 148 mg.l⁻¹ (normal: 86-105), decreased serum phosphorus: 17 mg.l⁻¹ (normal: 25-50), elevated parathyroid hormone level (PTH: 8608 pg.ml⁻¹; normal: 9-55) and elevated bone densitometry. These findings suggested the diagnosis of hypercalcemia and hyperparathyroidism.

Ultrasound and computed tomography scan of the neck showed a large mass in the left lower lobe of thyroid gland measuring about 3.5 × 1.7 cm (Figs. 4A and 4B). There were no renal stones on the abdominal ultrasonography. The diagnosis of primary hyperparathyroidism is confirmed. Other bones lesions related to hyperparathyroidism have been showed in the skull and phalanges (Figs. 5A and 5B).

Fig. 1. (A) Extrabuccal view: facial asymmetry. (B) Intrabuccal view: well circumscribed mass on the palate.

Fig. 1. (A) Vue extra-buccale : asymétrie faciale. (B) Vue intra-buccale : tumefaction palatine bien circonscrite.

Fig. 2. (A) Panoramic radiograph: radiolucency in the anterior of the maxillary (arrow). (B) Coronal computed tomography scan showing a lytic bone lesion extending to the floor of nasal cavity (arrow). (C) Axial computed tomography scan showing the expansion of cortical bones (arrow).

Fig. 2. (A) Radiographie panoramique : radiotransparence maxillaire antérieure (flèche). (B) CT-scan : coupe coronale montrant une ostéolyse s'étendant vers le plancher des fosses nasales (flèche). (C) CT-scan : coupe axiale montrant une soufflure de la corticale (flèche).
The treatment consisted of surgical removal of the parathyroid mass. The lesion was histopathologically diagnosed as a parathyroid adenoma (Fig. 6). Medical treatment by bisphosphonates was associated. No surgical treatment of the intraoral brown tumor has been achieved. Regression of the swelling was noticed months later.

Discussion

Primary hyperparathyroidism is more frequently seen in patients over 50 years old, with gender predilection toward females [5]. Classically, it is associated with two major sites of potential complications: the kidneys and the bones [6]. The renal manifestations (nephrolithiasis) are the most common symptom [3]. Bone involvement is the late manifestation [7]. In the past, bones lesions were recognized in 80%
to 90% of patients with primary or secondary hyperparathyroidism [4]. This rate has decreased to less than 5% of cases because of early diagnosis by routine biochemical screening and successful treatment of the disease [4,8]. Classic skeletal lesions are bone resorption, bone cysts, brown tumors and generalized osteopenia [6]. Brown tumors are non neoplastic lesions resulting from abnormal bone metabolism in hyperparathyroidism [9]. They have been described in both primary (4.5% of patients with primary hyperparathyroidism) and secondary hyperparathyroidism (1.5–1.7% of patients with secondary hyperparathyroidism) as resulting from an imbalance of osteoclastic and osteoblastic activity with bone resorption exceeding the bone formation [10,11]. The ribs, clavicles, and pelvis are the sites of predilection of this lesion [11]. Jaw involvement is extremely rare with mandible the most common site than maxillary [9,11].

Clinical symptoms caused by brown tumors depend on their seize and location [11]. They most commonly present as slowly growing, painful masses [11]. Asymptomatic lesions accidentally diagnosed by radiological examination are possible [11]. Radiographically, brown tumors of the jaws present as well-demarcated, monolocular or multilocular osteolytic lesions [4]. As for our patient, other radiographic symptoms related to the primary hyperparathyroidism are usually associated to the brown tumor such as subperiostal bone resorption of phalangeal tufts, loss of lamina dura around the teeth, generalized osteoporosis and “salt and pepper” radiologic appearance of demineralization of the skull [12,13].

Histologically, microscopic findings in brown tumor are non-specific showing classically population of mononuclear stromal cells mixed with multinucleated giant cells, among which recent hemorrhagic infiltrates and hemosiderin deposits are often found [14]. The hemorrhage and hemosiderin give the tumor a brownish color which gives rise to its name [4,14]. In our case, histological features alone cannot establish a certain diagnosis because of many giant cell lesions of the jaw bone (central giant cell reparative granuloma, cherubism, aneurysmal bone cyst). A certain diagnosis was confirmed by the endocrinologic status of the patient.

The treatment of hyperparathyroidism is the first step in the management of the brown tumor [14]. There is general consensus that the treatment of primary hyperparathyroidism is parathyroidectomy, but opinions are divided about the management of the bony lesions [5,11-15]. Most authors believe that brown tumor regression and healing are expected after the correction of hyperparathyroidism [5,14]. The time necessary for bone regeneration varies from several months in young patients to several years in older patients [5]. In the case reported here, no treatment of the palatal brown tumor has been done. However, several cases of brown tumor that grew after parathyroidectomy or normalization of hyperparathyroidism level have been reported [14]. In these cases, many authors have reported the surgical resection of remaining brown tumor [4,14].

Conclusion

Despite the improve methods of blood analysis that have led to early diagnosis of this endocrine disorder, there is still the possibility of patients presenting advanced bony lesions of primary hyperparathyroidism. We should therefore investigate all jaws giant cell lesions to exclude primary hyperparathyroidism.

Competing interests: none

References


