Brown tumour revealed: A literature review with a case study

Marcelo Pinto*, Gustavo Gaffrée, Roberto Santos, Fábio Ramoa, Jonathan Paixão, Hugo Melo and Célia Archer

Received: December 08, 2017; Published: December 13, 2017
*Corresponding author: Marcelo Pinto; Email: marcelodomingues1991@hotmail.com

Summary
The mandibular hyperparathyroid brown tumour is considered a rare metabolic disorder, also known as mandibular osteoclastome, therefore its therapy is not well defined yet. This lesion is triggered by the excess of the production of PTH parathyroid hormone, which are directly linked to mechanisms of control of levels of calcium, phosphorus and vitamin D, which confers the lesions the aspect of numerous osteoclastic cells, circumscribed and richly vascularized. Which causes their reddish-brown coloration, individuals affected by this disorder show signs and symptoms of weakness, nausea, fatigue, anorexia, excessive thirst, polyuria, constipation and consequently depression. The purpose of this study is to perform a literature review by describing the case of mandibular brown tumour, explaining the conduct performed according to the case by clinical, laboratory analysis and image studies that guided the case.

Introduction
This disorder, called brown tumour, has its determinant metabolic bone disarray caused by the excess production of parathyroid hormone, which is produced by the four parathyroid glands located posterior to the thyroid gland, which are responsible for the control of phosphorus (P), calcium (Ca) and vitamin D. The level of calcium present in the blood is considered a trigger factor of the disorder by the release and production of this hormone. Shetty “the hyperparathyroidism HPT is a disease in which there may be a complex, of biochemical anatomic and clinical abnormalities”. Therefore the Brown Tumor has its histopathological complexity described as multiple adenomas, numerous osteoclastic cells, of cystic format, and separated by a highly vascularized tissue which will confer the lesion a dark red or brownish coloration which characterizes its nickname. Some of the signs and symptoms observed in this disorder are fatigue, nausea, weakness, anorexia, excessive thirst, polyuria, constipation, pain, swelling and frequent urination. In severe cases, may present kidney stones, loss of bone mass and fractures, mental confusion and consequent depression.

The systemic scenario begins when the body presents hypocalcemia, and in the other hand will be inhibited in hypercalcemia, it will be stimulated the uptake of calcium to the extracellular environment, which leads to the increase of serum calcium concentration and decrease of the phosphate ion which is responsible to transport the phosphorous. Two sites of action in bone metabolic disorder are recognized, which are described respectively as actions on kidneys that reduce phosphate absorption and increase calcium absorption and the action on bones that stimulates bone mobilization with increased serum calcium concentration. Another important aspect and the issue of vitamin D which is a regulator of osteo mineral calcium physiology vitamin D is used in the treatment of the secondary form of the disorder as supplementation. The disorder may be divided into primary, where the treatment will be surgical removing the lesion, and secondary where the basic disorder that will affect the normal functioning of the parathyroid glands and vitamin D supplementation should be treated.

Literature Review
Brown tumor of Hyperparathyroidism it is a metabolic disorder that can affect the entire skeleton. This disorder is named due to macroscopic aspect to the intraossic lesions, presenting reddish-brownish hue, due to the intense bleeding that occurs in these lesions and the deposit of hemosiderin. Commonly, this disorder is a result of a framework of primary hyperparathyroidism, secondary rarely and tertiary. Brown tumor arises only in chronic cases of non-treated Hyperparathyroidism (PTH). The lesions are caused by the increased production and release of PTH, leading to increased re absorption of calcium in bones. With the constant calcium sequestration, it will start to grow a repair tissue, which increases gradually in size [1,2]. This disorder usually rise in individuals who are in third, fourth and sixth decade of life, especially in females. The most affected is in the head and neck area especially in mandibular area [1,2]. Clinical manifestations vary according to the location and extent of the Tumour. In the mandibular bone, there may be pain and hard tumefaction. When extensive it can lead to deformation of the region, impairing the function of the bone [1,2].

HTs symptoms are observed including fatigue, weakness, nausea, anorexia, excessive thirst, polyuria, constipation and depression [1,2]. The diagnosis is achieved through radiological exams, computerized tomography scans and Histopathological examination of the Tumour Histopathologically the Brown Tumor is characterized by a soft tissue mass composed of giant...
The Brown Tumour generally shows multiple occurrences, although it can also occur as a single one. When present in the mandibular bone, it usually appears at the hard blade [5,6]. In addition, one could observe bone trabeculae neoformed with osteoblasts tumefactions on its edges. Blood tests indicate the increased levels of calcium and alkaline phosphates, as well as the reduction in the level of phosphorus is useful for finding the correct diagnosis [1,2]. The Brown Tumor does not demand specific treatment in most cases because the correction of Hyperparathyroidism leads to the lesion disappearance. However, it can cause pathological fractures and bone marrow compression even when it involves the spine. When it reaches the face can cause breathing difficulties and facial deformities. In such cases, it is recommended that the patient undergoes surgical treatment [7-9].

According to radiographs and Tomography scans: it shows a radiolucent / hypodense diffuse, irregular image without bulging of the vestibular and / or lingual bone cortices was verified. Radiographic and CT scans did not show radiopaque halus, that is, descorticalized, involving the region of the element 36. Bone resurfacing in the furcation region in the same element. In computed tomography (CT), the medullary bone presents a granular aspect with bone and lingual cortical thinning, vertical bone loss located in the mesial and distal part of the 36 element, and increased ligament space with absence of hard lamina, apparent mobility, initial external root resorption in the Element mentioned above. Hypothesis: Bone tumor, endo-periodontal envelopment or giant cell lesion. According to the laboratory test of Parathyroid hormone (PTH), by the enzyme immunoassay method by chemiluminescence with a result of 502.8 pg/ml (normal 12 to 65 pg/ ml), creatinine level is high with 2.94 mg/dl (Normal for male patients is 0.70 to 1.30 mg/dl). Histopathological examination by Hematoxylin and Eosin staining method with diagnosis of Brown Tumour (osteoblast matrix with prominent osteoblasts and osteoclasts, amidst fibrous stromal fibroblastic proliferation: proliferated, ecstatic and congestive vessels, haemorrhage and hemostasis).

The parathyroid glands were increased in cervical echography. The findings of hyperparathyroidism were confirmed together with other exams to the diagnosis of Brown Tumour of Hyperparathyroidism. Patient was submitted to surgical intervention under general anaesthesia, naso-tracheal intubation, vestibule access in the region of the left jaw body, enucleation of the lesion, removal of the element 36, curettage and synthesis of the region. All material was collected and sent to the pathology center of the State University of Rio de Janeiro (UERJ) for histopathological analysis where the diagnosis of Brown Tumour of Hyperparathyroidism was confirmed. Post surgery evolution of the patient in question was compatible with the surgical procedure and without intercurrences.

Case Report

Patient male, 35 years W.L.M. R, Brazilian, born in Rio de Janeiro, attended the Oral clinic and traumatology maxillofacial surgery at Municipal Lourenço Jorge Hospital, Barra da Tijuca, capital of Rio de Janeiro. During the anamnesis reported as main complaint “volume increase in unilateral mandibular region being the left side, painful. Current history of the disease (HDA) with account of start approximately 2 (two) years, as a minor injury to the gums, which evolved gradually, resulting in lumps region mentioned above. Still reported that does not have habits like smoking and alcoholism, drug hyper sensibilities, morbid antecedents denies personal and family. Clinical and physical examination showed, increased mandibular unilateral left significant volume, pain and mobility the palpation of the 46 element associated with bulging Protuberance in the abovementioned causing bone, with absence of fungi signs and infectious in the oral cavity (Figure 1).
Figure 3: Diagnostic hypothesis: Bone tumor.

The imaging tests: X-rays and computed tomography verified a radiolucent image/hypoechoic diffuse, irregular without bulging of the vestibular cortical bone and/or lingual side. Image does not show halo Radiopaque, i.e. descriptolized, involving 36 element regions. Bone rarefaction in the furcation region on the same element. In computed tomography scan (CT) the medullary bone features granular aspect with tapering of the vestibular and lingual cortical bone loss mesial and distal vertical located on 36 and element increase ligament space with absence of the lamina dura, apparent mobility, external root reabsorption on the initial element mentioned above (Figures 2-4). As the laboratory examination of the parathyroid hormone (PTH), by the method of chemiluminescence enzyme immuno assay with result of 502.8 pg/ml (normal 12 to 65 pg/ml), creatinine high level observed with 2.94 mg/dl (normal for male patient is 0.70 to 1.30 mg/dl) (Figures 5 & 6).

Figure 4: Diagnostic hypothesis: Endo-or giant cell lesion periodontal.

Figure 5: Result of presence of parathyroid hormone, by the method of chemiluminescence enzyme immuno assay.

Figure 6: Brown Tumor diagnosis.

Histopathological examination, by coloring method by hematoxylin and eosin with Brown Tumor diagnosis (with prominent osteoid matrix osteoblasts and osteoclasts in fibrous stroma with fibroblastic proliferation; vases proliferated, entranced and congestion, pockets of deposition of hemosiderin and hemorrhage). Parathyroid glands were increased in cervical ultrasound. The findings of hyperparathyroidism confirmed together with the other tests the diagnosis of Hyperparathyroidism.
Brown Tumor (Figure 7). Patient underwent surgery under general anesthesia, naso-tracheal intubation, accomplished access in lobby Fund in the region of left mandibular body, enucleation of the lesion, dental extraction of 36 element, curettage and synthesis of the region. All material was collected and forwarded to the pathology Center of Rio de Janeiro State University (UERJ) for histopathological analysis where it was confirmed the diagnosis of Hyperparathyroidism Brown Tumor. Postoperative evolution of the patient in question was compatible with the surgical procedure and without complications (Figure 8).

Figure 7: Result of microscopic examination.

Figure 8: Patient underwent surgery under anesthesia.

Discussion

The Brown tumour is a lesion associated with the hyperparathyroidism and can be divided into primary or secondary. Affects more the mandible than the maxilla and your prevalence is by the feminine gender above 50 years. We can clinically identify a Brown tumour of hyperparathyroidism because of an increase volume in the region causing pain and mobility, presenting as an extroverted mass of slow growth and destructive character. Depending on where the tumour is located, they can cause: diplopia, deformity, bleeding, chewing trauma, among other adversities. Radiographically, we identified this Brown tumour of hyperparathyroidism, due to the presence of a uni or multicellular radiolucent image, descorticalized, besides that when the Computed Tomography was performed, described in the above case, absence of the hard blade was revealed.

Histologically, the Brown tumour of hyperparathyroidism manifests as a mass of soft tissue composed of a giant cells inside the fibro vascular stroma, presenting focus of bleeding and hemosiderin deposition as a friable red-brown mass.

Diagnostic errors can occur if we only evaluate their histological characteristics, since other lesions such as the giant cell lesion present similar aspects to this lesion, thus causing unnecessary radical therapeutics and severe sequel if the diagnosis is incorrectly concluded. Therefore, it is extremely important to analyze the history of the disease and laboratory tests. On laboratory examination, elevated creatinine level revealed parathyroid gland hyperplasia confirming the diagnosis, in addition with the other exams, of Brown tumour of hyperparathyroidism. The treatment accomplished in the relate of case was surgical with the objective was remove all the tumour and for this reason a naso-tracheal intubation was performed, with your purpose was allow a good surgical access, giving better tranquillity to the surgeons work, besides ensuring that the patient was treated in a single surgical time. Posteriorly, was done the enucleation of the tumour, which was the purpose of the surgery, the element 36 removal, curettage and suture of the region.

Final Consideration

The literature review on the mechanism of pathological action of the brown tumour associated with a clinical case study, it was possible to reaffirm that dentistry is on the correct path both to diagnose the disorder through the diagnostic exams, establishment of differential diagnostic and laboratory exams as long as long-term treatment, not only through aspects closely related to the dental surgeon, but also in the supplementation of vitamins and minerals, it is worth emphasizing that all cases should maintain long-term treatments booking appointments regularly. Although cases of Brown Tumour are rare, the treatment was performed with total efficiency.

References


