Parathyroid carcinoma is an extremely rare neoplasm which typically is associated with hardened nodules in the parathyroid region, hypercalcemia and renal failure. A 69 year-old woman presented with knee pain at an emergency unity. On clinical examination the patient also presented a 2.0cm fibroelastic and mobile node in the left neck, level II. Serum total calcium and parathyroid hormone were increased, and the node was also detected by CT and 99mTc-sestamib scintigraphy. The patient had been surgically previously treated of parathyroid carcinoma misdiagnosed as a thyroid adenoma (eleven years ago), and a hyperplasia of parathyroid chief cells (nine years ago), that after revision was diagnosed as cervical metastasis of parathyroid carcinoma. She was surgically treated by neck dissection, confirming the second regional metastasis of parathyroid carcinoma. The diagnosis of parathyroid carcinoma is a challenge, and although rare, it should be considered in the presence of hypercalcemia and palpable neck nodule.

**Key Words:** Parathyroid Carcinoma; Hypercalcemia; Hyperparathyroidism; Lymph Node Metastasis; Nodule.

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**Introduction**

Parathyroid carcinoma is an extremely rare neoplasm derived from parathyroid parenchymal cells, representing fewer than 4% of cases of parathyroid disease [1,2]. Primary hyperparathyroidism is mainly associated with adenomas (90%) and hyperplasias (7%) of the parathyroid, and less than 1% with carcinomas [3].

The main targets of PTH are the kidney and skeleton. Primary hyperparathyroidism is the most common cause of hypercalcemia, that when severe causes various symptoms, including fatigue, weakness, weight loss, anorexia, nausea, vomiting, renal insufficiency, nephro lithiasis, nephrocalcinosis, polyuria, and polydipsia, bone pain and pathologic fractures [4,5,6]. A palpable and hardened neck mass is a common finding in patients with parathyroid carcinoma, and it is reported in over 75% of cases [7,8].

Histologically it is very difficult to distinguish parathyroid carcinoma and adenoma, and parameters to be considered include fixation to the thyroid, metastasis, and vascular, perineural, capsular and adjacent tissues invasion. In parathyroid carcinomas hypercalcemia can reach very high levels [3].

This report describes a patient who was initially surgically treated of a possible nodular hyperplasia of the parathyroid. Later she presented high hypercalcemia associated with lymph node metastasis of carcinoma of the parathyroid. The clinical diagnostic reasoning was fundamental to clarify this case.

**Description of The Case**

A 69 year-old Caucasian woman was referred in 2010 to the Emergency Service of the Fornecedores de Cana Hospital, Piracicaba, Brazil, complaining of pain in her knee. She had been previously submitted to left partial thyroidectomy in 1999 with histological diagnosis of thyroid adenoma. In 2008 the patient was submitted to cervical node biopsy with the diagnosis of hyperplasia of parathyroid chief cells.

During clinical and laboratory examinations, it was found that the patient presented high levels of hypercalcemia and a 2.0 cm fibroelastic, mobile nodule in the left cervical region. In addition, the patient also showed evident hoarseness. Based on these informations, laboratorial investigations were carried out, showing high levels of PTH and hypercalcemia (Table 1). A 99mTc-sestamib scintigraphy showed intense uptake in the cervical node, and CT scan confirmed that the palpable node highlighted by the scintigraphy was a cervical level II lymph node (Figs. 1 and 2).

Considering the present clinical data, a review of the original...
surgical specimen diagnosed as parathyroid hyperplasia in 2008 prompted the diagnosis of metastatic parathyroid carcinoma in cervical lymph node. It was then considered that the initial surgery of a left partial thyroidectomy realized in 1999 with the diagnosis of thyroid adenoma, was probably due a parathyroid carcinoma misdiagnosed as a thyroid adenoma. On this way, the patient had in 1999 a parathyroid carcinoma, in 2008 the first cervical lymphnode metastasis and in 2010, when we saw the patient, the second cervical metastasis.

The patient did not show evidences of distant metastasis or bone lesions, and she was then treated by radical left neck dissection (Fig. 3). One out of 38 nodes removed was involved by the carcinoma. Levels of calcium and PTH rapidly came close to normal, and symptoms improved.

Eight months after surgery the patient showed slight increase in the PTH level, and a metastatic parathyroid tissue surrounded by fibrous connective tissue in the left side of the neck was surgically removed, followed by radiotherapy. The patient is under follow up for 10 months, without evidences of disease and with normal levels of calcium and PTH.

Histologically the review of the original surgical specimen from 2008 (first metastatic parathyroid carcinoma in cervical lymph node showed an encapsulated nodule, with areas of infiltration into the adjacent tissues. Most of the tumor was formed by solid sheets of polyhedral cells with well defined limits and eosinophilic cytoplasm, central nuclei and prominent nucleoli. The peripheral cells showed scant cytoplasm, with more hyperchromatic nuclei in relation to the central cells. An increased nuclear-cytoplasmic ratio and cytoplasmic inclusions also were found. Only a few cells showed positivity for Ki-67, indicating a low proliferative rate (Dako, clone MIB-1, dilution 1:100) (Figure. 4).

The second metastatic lymph node detected in 2010, was invaded by the malignant parathyroid cells, showing similar morphological and architectural features as above described. Residual lymphoid
tissue was found close to the lymph nodal capsule, which showed areas of invasion by the tumor cells. It was also observed areas of cystic degeneration and necrosis.

**Discussion**

Hypercalcemia occurs in approximately 10-30% of all cancer patients, and it is the most common life-threatening metabolic disorder associated with malignancies. Chemical mediators as RANKL, TGF-β, PTHrP, IL-1, IL-6, IL-7 and TNF-α can be activated by factors released by tumor cells and cause hypercalcemia [9].

Hypercalcemia in patients with cancer can be either PTH-independent or dependent. PTH-independent includes humoral hypercalcemia of malignancy (HHM), local osteolytic hypercalcemia, hypercalcemia due to increased production of 1,25 dihydroxyvitamin D3, hypercalcemia related to drugs, and immobilization-associated hypercalcemia. PTH-dependent is associated with parathyroid carcinoma or ectopic production of PTH by malignancies [9].

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The normal level of PTH (10-65pg/mL) is inversely proportional to the level of calcium in the blood, which is tightly regulated between 8.8 and 10.2 mg/dL. Our patient showed a serum level of PTH and calcium of 619pg/mL and 16.3 mg/dL respectively. High levels of hypercalcemia can be fatal, and determination of its causes is fundamental for the correct treatment. Rarely parathyroid carcinomas do not secrete PTH and consequently the patient does not show the signs and symptoms of hypercalcemia. On these situations the diagnosis of the tumor usually is late, and frequently the patients die due the large size of the tumor mass [10]. More frequently elevated levels of PTH can also be seen in non-neoplastic lesions of the parathyroid as hyperplasia, parathyroiditis and cysts, as well as in adenomas.

Hypercalcemia and its symptoms can be found in 86% of patients with parathyroid carcinoma, particularly bone and joint pain and renal stone [7,11,12]. Our patient looked for treatment because of pain on her knees, but she also showed a neck nodule, that is found in more than 75% of the patients with parathyroid carcinoma [1,7,8,13]. Our patient also showed evident hoarseness and slight loss of voice, and it is well known that recurrent palsy of the laryngeal nerve in a patient with primary hyperparathyroidism is indicative of parathyroid carcinoma [3,13].

It should be considered that patients with primary hyperparathyroidism can present bone brown tumors, histologically similar to giant cell lesions of the jaws. Also patients with hereditary hy-
perparathyroidism with jaw-tumor syndrome (HPT-JT) present a fibrous-osseous lesion indistinguishable of central ossifying fibroma, and up to 15% of these patients have parathyroid carcinoma [6,14,15]. Our patient was submitted to radiographic bone screening including the jaws, and no bone lesion was found.

The histological distinction between parathyroid adenoma and carcinoma is difficult. It is suggested that trabecular pattern, mitosis, Ki-67 index, nuclear hyperchromatism, thick fibrous bands, capsule and tissue invasiveness favors malignancies, but it should be also considered that some adenomas can show atypical morphology [6]. More reliable characteristics include metastasis, comedonecrosis and severe hypercalcemia. It is not uncommon that a parathyroid carcinoma is misdiagnosed as a benign lesion, as it happened initially on the present case.

Clinical and pathological staging is not yet well established for parathyroid carcinoma. Metastasis may occur in the later course of the disease, mainly in the lungs (40%), liver (10%) and also in regional lymph nodes (30%), as on the present case [7]. The clinical relevance of cervical node involvement is not well established [3]. Our patient showed two ipsilateral lymph node metastasis and one local recurrence, but a close follow up indicates that she is free of the disease and without clinical and laboratory alterations.

The main modality of treatment of parathyroid carcinoma is radical surgery, with removal of the lesion together with the ipsilateral thyroid, thyroid isthmus and lymph nodes. Surgery is also the treatment of choice for local recurrences and regional metastases [7]. Prior surgery, severe hypercalcemia should be controlled by drugs. Although the tumor is resistant to radiotherapy, it is indicated in cases of microscopic residual disease, and it was used on our patient, since she had two regional nodal metastases and one local recurrence involving fibrous tissue. The patient must have a long follow up, since metastases have been described later in the course of the disease, as happened on the case here described.

In short, the patient was treated 11 years before of an adenoma of the thyroid that probably was a non diagnosed parathyroid carcinoma. After 9 years a node on the neck was removed with the diagnosis of hyperplasia of the parathyroid, which on revision was in fact the first metastatic node of parathyroid carcinoma. After another 2 years she presented knee pain and high levels of hypercalcemia and a neck node that it was confirmed as the second metastatic parathyroid carcinoma. Finally after 8 months the patient presented a local recurrence, also surgically removed followed by adjuvant radiotherapy. Conclusively, clinical and histological diagnosis of parathyroid carcinomas can be a challenge, and severe hypercalcemia and its symptoms, associated with primary hyperparathyroidism and neck nodule are the main indicators of this disease.

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