Abstract

The columnar cell variant is a rare subtype of papillary thyroid carcinomas (PTC), characterized by its morphologic features and aggressive biologic course. Its diagnosis depends on the postoperative histological study of the surgical piece and its prognosis remains controversial. The treatment is mainly surgical while post-operative radio iodine therapy has proven its worth. We report the case of a 73 years old patient, with history of arterial hypertension, having presented during a medical checkup a cervical painless swelling that was mobile on swallowing, and the lymph node areas were free. The thyroid laboratory tests were normal. The ultrasound cervical examination found a hetero multi nodular goiter classified TIRADS III. The patient has undergone a total thyroidectomy. The anatomopathological study of the surgical piece revealed a papillary carcinoma of the right lobe of the thyroid in its cylindrical cells variant measuring approximately 4 mm, located in contact with the thyroid capsule which it invades focally with presence of a single neoplastic embolus. Postoperative recovery was marked by transient hypocalcemia that was controlled by calcium supplementation and the patient was discharged from hospital 5 days later. The patient received 131-Iodine therapy (3.7 GBq) followed by post therapy whole body scan 6 days and 6 months later that didn’t show any pathological focus. The columnar cell variant of papillary thyroid carcinomas represents an aggressive variant of PTC potentially portending and unfavorable prognosis. The encapsulated form or small tumors demonstrates a more favorable outcome with indolent clinical process, which shows relatively slow growth and low incidence of recurrence or metastasis. The surgery still remains the main treatment while the post-operative radio iodine therapy plays an important role allowing the sterilization of tumor residues and thus reduces the rate of recurrence.

Keywords: Columnar Papillary Thyroid Carcinoma.

INTRODUCTION

Papillary thyroid carcinomas are the most common form of thyroid cancers. Generally, of good prognosis, they present various clinical and evolutionary aspects according to their origin.

The columnar cell variant of papillary thyroid carcinoma (CCVPTC) is a rarer variant with a more reserved prognosis.

We report the case of a patient who presented a columnar cell variant of papillary thyroid carcinoma discovered in the anatomopathological study of the surgical piece.

CASE REPORT

73-year-old patient with history of arterial hypertension under treatment has presented during a medical check-up a cervical swelling justifying the realization of a cervical ultrasound then referred to our facility for medical care.

The cervical examination found a patient with a painless cervical swelling that was mobile on swallowing. The lymph node areas were free and the rest of the clinical examination was without abnormalities.

The thyroid laboratory tests were normal. A cervical ultrasound examination found a hetero multi nodular goiter; the largest nodule being on the left lobe measuring 3.5cm classified TIRADS III.

The patient has undergone a total thyroidectomy. The anatomopathological study of the surgical piece revealed a papillary carcinoma of the right lobe of the thyroid in its cylindrical cells variant...
measuring approximately 4 mm, located in contact with the thyroid capsule which it invades focally with the presence of a single neoplastic embolus.

The patient was discharged from hospital 5 days later. The decision of the multidisciplinary concertation meeting was Radioactive Iodine therapy.

The patient received 131-Iodine therapy (3.7 GBq). The post-therapy whole body scan was performed 6 days later, revealing the presence of a thyroid remnant without distant pathological focus with low level of thyroglobulin.

The patient was put on L-thyroxine-based hormonal braking treatment. An efficacy assessment was carried out 6 months later, the whole body scan didn’t show any pathological distant focus and the patient was declared cured.

**DISCUSSION**

Papillary thyroid carcinoma is the most common malignancy in the thyroid gland, which accounts for 90% of all thyroid cancers [1, 2]. It is considered an indolent malignant tumor with an overall mortality rate of 1% per year [3].

The World Health Organization classification of thyroid tumors recognizes a subcategory of thyroid papillary carcinoma termed the biologically aggressive variants that include the diffuse sclerosing variant, the tall cell variant, and the columnar cell variant [4].

The columnar cell variant of papillary thyroid carcinoma (CCVPTC) is a rare subtype, which accounts for 0.15−0.2% of all PTCs [5].

Studies showed that CCVPTC has a fast growth rate and a high incidence of recurrence, and this type of tumors is associated with local invasion and early lymph node metastasis [6-8]. However, the prognosis of CCVPTC remains controversial, because the encapsulated form has a more favorable outcome with indolent clinical process, which shows relatively slow growth and low incidence of recurrence or metastasis [9-11].

The presence or absence of extrathyroidal invasion represents the most important parameter in predicting the behavior of these tumors. Tumors confined to the thyroid gland are associated with an excellent prognosis and can be managed conservatively (less than total thyroidectomy with or without radioactive iodine therapy) whereas tumors that invade beyond the thyroid capsule with extension into perithyroidal soft tissues behave more aggressively, necessitating more aggressive management (total thyroidectomy with supplemental radioactive iodine therapy) and evaluation for disseminated visceral disease [12].

Ultrasonography may show hypoechoic nodules with or without calcifications and with at least more than one typical malignant sonographic feature [13].
The diagnosis of columnar-cell carcinoma largely depends on the postoperative pathologic findings:

The accurate diagnosis of CCVPTC is difficult upon FNA (fine needle aspiration) cytology, because the tumor cells in CCVPTC lack the nuclear features of the classic pattern of papillary carcinoma, which shows nuclear grooving and intranuclear pseudo-inclusions [14, 15]. Based on histology, CCVPTC is defined as papillae or gland-like structures lined by columnar cells that have prominent nuclear stratification [16].

Because the nuclear features of PTC are not well represented in the CCV, these tumors may be mistaken for metastatic intestinal, endometrial, or pulmonary adenocarcinomas, from which they can be distinguished because of immunohistochemical positivity for both thyroglobulin and thyroid transcription factor 1 (TTF-1). To the best of our knowledge, there is no clear consensus regarding the minimal percentage of columnar cells that confers a diagnosis of PTCCCV, with reported cases varying from 30% to 80% [17-19].

The management of CCV is controversial, and the current guidelines of the American Thyroid Association recommend total thyroidectomy or near total thyroidectomy and lobectomy, central lymphadenectomy, and postoperative RAI (radioactive iodine) as indicated [20], the rates of definitive hypoparathyroidism and recurrent laryngeal nerve injury following total thyroidectomy and total thyroidectomy associated with routine central lymph node dissection were higher than that after lobectomy and total thyroidectomy [21, 22].

External-Beam Radiation Therapy (EBRT) is typically reserved as a last resort after surgery and RAI have been attempted [25]. Some studies revealed that EBRT has been traditionally used to treat incompletely resected tumors, unresectable diseases, aggressive variants, and to reduce the risk of locoregional recurrence [23, 24].

CONCLUSION

The thyroid papillary carcinoma of columnar cell type is a distinct morphologic type of thyroid papillary carcinoma.

The biologic behavior of this tumor is predicated on clinical stage, with the presence or absence of extrathyroidal invasion being the single most important parameter.

The most useful features in the differentiation of CCVPTC from other PTC are the presence of columnar cells. Treatment for columnar-cell carcinoma of the thyroid is mainly surgical resection.

Due to the frequent recurrences and metastases, the prognosis for unencapsulated columnar-cell carcinoma is poor, whereas the encapsulated columnar-cell variety has been reported to have a favorable prognosis.

A post-operative treatment with radioactive iodine 131 allows to irradiate any persistent neoplastic focus decreasing the risk of subsequent recurrence, eradicate normal thyroid remnants and to perform a post therapy whole body scan to detect metastatic disease.

REFERENCES