Concomitant thyroid MALT lymphoma and papillary thyroid carcinoma

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SUMMARY
The objective of this study was to describe the rare thyroid MALT lymphoma concomitant with papillary thyroid carcinoma in a male patient who was submitted to total thyroidectomy. Treatment and follow-up issues are addressed. Male patient complains of fast thyroid enlargement without lymphadenopathy and normal clinical exams. Total thyroidectomy was indicated and performed without any complications. The pathology showed multicentric papillary thyroid carcinoma, concomitant thyroid MALT lymphoma and Hashimoto’s thyroiditis. The immunohistochemistry assay was positive for CD 20, CD 43, CD 79, AE1/AE3. The staging studies showed no evidence of both metastasis, Ann Harbor stage IE, without B symptoms. After RIT no further radiotherapy or chemotherapy was indicated. Nowadays the thyroglobulin is undetectable, without recurrences at two years of follow-up. It was concluded that primary thyroid MALT lymphoma is uncommon being the papillary thyroid carcinoma more frequent. Both occurring concomitantly is very rare and the treatment has to prioritize the tumor of worst prognosis at the discovery moment. Arq Bras Endocrinol Metab. 2010;54(4):425-8

SUMÁRIO

Primary thyroid MALT lymphoma is very uncommon, accounting for 0.6% to 5% of all thyroid malignancies, with a distinct elderly female predominance and incidence peak between 60 and 69 years. Some authors correlate it to auto-immune diseases like Hashimoto’s thyroiditis (1-3). Papillary carcinoma is the most frequent thyroid cancer and responsible by nearly 80% of all thyroid cancer in the USA, occurring predominately in women (3:1 ratio) in the 3-4 decade of life (4-6). The concomitance of both is very rare and the present case report has the objective of describe the rare thyroid MALT lymphoma concomitant with papillary thyroid carcinoma in a single old male patient who noted a sudden thyroid enlargement and was submitted to total thyroidectomy. Treatment and follow-up issues are addressed.
CASE REPORT

The present case report was previously approved by the Hospital Ethics Committee. A 61 year-old male patient was referred to this service due to fast painless thyroid enlargement during about the last three months without any other complaints. The clinical feature showed a multinodular thyroid enlargement, without neck lymphadenopathy, hoarseness, dysphagia or dyspnea. The thyroid ultrasound showed a 54 cm³ multinodular goiter with some nodules presenting gross calcifications, without other cervical anomaly, laboratory screening showed normal functioning of the thyroid and only serum thyroglobulin antibody was slightly increased. The patient was a non-smoker, denied alcohol consumption and has arterial hypertension. Although the preoperative USG guided needle aspirative biopsy is the mainstay in the current literature and should be done in every suspected nodule, we preferred the prompt surgical treatment in this particular case due to the fast multinodular growing in the male patient and the positive history of familial thyroid carcinoma (mother).

Total thyroidectomy was performed without any complications, preserving the inferior and superior laryngeal nerve and the four parathyroids, some technical difficulty occurred due to the extremely hard nodule at the superior left thyroid pole. Discharge was the next day following surgery without complaints until follow-up visits.

The pathology exam showed multicentric papillary thyroid carcinoma, classic variant, without vascular or lymphatic embolization in both lobes, accompanied of Hashimoto’s thyroiditis and a concomitant thyroid MALT lymphoma in a nearly 3.5 cm nodule surrounded by normal thyroid tissue and areas of papillary carcinoma at the left thyroid pole. The immunohistochemistry assay was positive for CD 20, CD 43, CD 79, AE1/AE3 (Figures 1 and 2).

Staging included neck CT, thorax, abdomen and pelvis, PET-CT, total body scintigraphy with MIBI, laboratory evaluation and bone marrow biopsy. All exams showed no evidence of both metastasis so the Ann Arbor (classification for lymphomas modified by Mussoff) staging was stage IE, without systemic B symptoms. Levothyroxine was initiated at 150 mcg per day and the patient was referred to the radioiodine therapy with 150 mCie demonstrating a negative whole body scan. No further radiotherapy or chemotherapy was indicated and nowadays, after two years of follow-up, the patient is stable, with undetectable serum thyroglobulin and without evidence of recurrences.

DISCUSSION

Primary thyroid lymphoma is very rare, accounting for 0.6% to 5% of all thyroid malignancies, and MALT lymphoma is the most common subtype. It occurs more frequently in the elderly female linked with chronic autoimmune thyroiditis (Hashimoto’s disease), being moderate to poor in degree of differentiation. Presentation in the male gender in this report is associated with poor prognosis when compared the WDTC (well differentiated thyroid carcinoma); however, no sufficient data exits regarding MALT lymphoma in literature (7-9).

Differential diagnosis between primary thyroid lymphoma and generalized lymphoma with thyroid involvement must be performed. Clinically the first has the presentation of painless rapid mass growth in the thyroid, sometimes with compressive symptoms which...
can be confused with intracystic hemorrhage or enlargement of a nodule (10). B symptoms like fever, weight loss more than 10% and nocturnal sweating suggests typical disseminated lymphoma either primary in thyroid gland or in another site. In this report, as other few cases, the disease was discovered only after surgery, once the patient related solely a globus sensation. Preoperative exams such as neck ultrasound, thyroid scintigraphy and even neck CT can be indistinguishable for compressive goiter.

In literature, the majority of MALT lymphomas occur in a previous autoimmune thyroid disease (Hashimoto’s thyroiditis), probably due an acquired pathological transformation of the intrathyroid lymphoid tissue. The level of concordance between Hashimoto’s disease and MALT lymphoma is 25%-75%, even though MALT lymphoma occurs in 0.5% of the cases (11,12). The malignant course is difficult to characterize, attributable to the indolent and long period of time in both clinical and morphological views, as the present case report. Although nearly 40% of patients present clinical or subclinical hypothyroidism in the primary thyroid lymphoma, this clinical case showed no evidence of laboratory alterations (11).

The present concomitance of MALT thyroid lymphoma and papillary thyroid carcinoma, very rare in the literature, make this case report a challenge in the literature, the patient has not any metast PET-CT and annual WBS as the standard follow-up from the literature, the patient has not any evidence of laboratory alterations (13). In this report the PET-CT has not evidenced disseminated disease or other primary sites for both thyroid neoplasms. Retrospective reports suggest an indolent behavior and excellent clinical prognosis for this subset of thyroid lymphomas. However, recent published observations in patients with other extranodal MALT-lymphomas have documented the dissemination of disease in about one-third of cases at the time of diagnosis (14). One must prove which tumor is more aggressive in behavior between the papillary thyroid carcinoma and MALT lymphoma to assure the ideal treatment for both.

The Ann Arbor classification for Lymphomas modified by Mussoff was used in this case (15): stage IE disease corresponds to disease confined to the thyroid, stage IIE corresponds to disease confined to the thyroid gland and the regional lymph nodes on the same side of the diaphragm, stage IIIIE corresponds to disease confined to the thyroid and lymph nodes on both sides of the diaphragm and/or spleen; and stage IV corresponds to disease in nodal and/or additional extranodal involvement. Localized disease was defined as IE and IIE disease, and disseminated disease as stage IIIIE and stage IV.

The present case was staged as IE (localized disease) where single treatment like surgery is evidence-based in the literature (7). However, multimodality (surgery plus radio- or chemotherapy) is still proposed in some institutions with curative intent (16). Radiotherapy, when indicated is performed in total dose of 40 Gy (10) and multiple chemotherapy (CHOP) is classically indicated for disseminated or locally aggressive lymphoma. Its role in localized disease is not clear (17-19).

The patient who presenting concomitant MALT thyroid lymphoma and papillary thyroid carcinoma must be judiciously evaluated, since the treatment has to prioritize the tumor which presents the worst prognosis at moment. In the present report the staging work-up showed no evidence, at PET-CT and WBS I-131, of disseminated disease for both tumors. Standard treatment with RIT at 150 mCie was then performed for the WDTC while no further treatment was proposed for MALT lymphoma.

Until this date, after trimestreal thyroglobulin, semestrial PET-CT and annual WBS as the standard follow-up from the literature, the patient has not any...
evidence of regional or distance recurrence loci during the two years of follow-up.

Despite the short follow-up of this rare presentation of concomitance of MALT lymphoma and papillary thyroid carcinoma, the patient has not evidenced recurrence. Based on the current literature showing few reports of MALT lymphoma associated to other tumors, this case report has the aim to add the experience of the service in this rare pathology to medical data.

A patient presenting the concomitant MALT thyroid lymphoma and papillary thyroid carcinoma must be judiciously evaluated, since the treatment has to prioritize the tumor with worst prognosis at the moment of discovery. Despite the rarity of thyroid lymphoma, is has to be part of the differential diagnosis when examining the elderly with sudden thyroid enlargement and who have clinical history of Hashimoto’s thyroiditis.

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REFERENCES