Primary thyroid MALToma– a rare diagnosis of an unassuming thyroid nodule

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ABSTRACT

Primary thyroid lymphoma, although a rare malignancy, can arise in common chronic inflammatory conditions such as Hashimoto’s thyroiditis. Incidental finding of a thyroid nodule with chronic thyroid inflammation warrants further investigation. Early detection of malignancy can play a vital role in improved outcomes. We report a case of a 60-year-old male who presented to the clinic for a routine visit. An enlarged, firm, non-tender thyroid gland was appreciated on exam with high thyroid stimulating hormone (TSH) level. Fine needle aspiration of the mass revealed nonspecific atypical lymphocytes. The pathology and immunohistochemical stains were consistent with histologic impression of extra nodal marginal B-cell lymphoma (mucosa-associated lymphoid tissue [MALT] lymphoma) and Hashimoto’s thyroiditis. Patient was treated with thyroxine after complete surgical excision of left thyroid lobe and remains in remission with close follow-up with his primary care provider. Primary thyroid MALT lymphoma follows an indolent process and remains asymptomatic in most patients. These are usually found to arise at sites of ongoing chronic inflammation with underlying autoimmune or infectious etiologies. Treatment modalities include surgical excision and/or radiation therapy for localized lesions, with both radiation and chemotherapy indicated for disseminated disease.

1. Background

Mucosa-associated lymphoid tissue (MALT) lymphoma is defined as extra nodal lymphoma composed of heterogeneous small B cells arising from marginal zone of MALT [1]. It most frequently occurs in the gastrointestinal tract mucosa (50%), head and neck (15%), lung (14%), skin (11%), thyroid (4%), and breast (4%) [1–3]. Primary thyroid lymphoma (PTL) is a rare form of malignancy, constituting about 2–8% of all thyroid malignancies and 1–2% of all extra nodal lymphomas [3,4]. Thyroid lymphomas are most commonly diffuse large B-cell lymphomas (60–80%), and only about 30% are extra nodal marginal zone lymphomas [3]. Primary thyroid MALT lymphoma is rare, accounting 6–28% of PTLs [5] and usually arises in the setting of chronic lymphocytic thyroiditis, such as Hashimoto’s thyroiditis [2]. The mean and median ages at diagnosis is between 65 and 75 years with 4:1 female predominance [5–7]. Co-occurrence of papillary thyroid carcinoma has also been reported [3,8]. Diagnosis requires a high degree of suspicion and is often difficult due to coexistence of reactive as well as neoplastic processes in the thyroid gland.

2. Case description

A 60-year-old male presented to our outpatient clinic for a routine visit. He had no complaints except for minimal hand dryness and denied fatigue, generalized weakness, heat or cold intolerance, constipation, diarrhea, hair loss, or any recent weight changes. His past medical history was pertinent for a basal cell carcinoma of scalp treated with extensive excision 3 years prior.

On physical examination, an enlarged, firm, non-tender thyroid gland was appreciated with about 3 cm left thyroid lobe mass felt on palpation. Xerosis of hands, arms, and legs were also noted with no pruritus or erythematous rash. No palpable lymphadenopathies or hepatosplenomegaly was noted. He denied any lump like sensation in his neck, neck pain, difficulty in swallowing, hoarseness, or dry cough. Signs of tracheal, esophageal, or neck vein compression were not found. He was an active smoker (35 pack-years) but did not have any history of radiation exposure, family history of thyroid-related disorders, or malignancy. His diet was normal with iodine-rich meals.

Complete blood count and basic metabolic profile were within normal limits. TSH was found to be elevated at 10.14 uIU/mL. All other lab values were within normal limit. Thyroid ultrasound was done which revealed large 5.8 × 3.1 × 2.5 cm hypoechoic mass occupying almost complete volume of the left thyroid lobe, and persistent elevation of TSH was found in repeat thyroid panel. Patient underwent...
Computed Tomography (PET–CT) is a common finding associated with Hashimoto’s thyroiditis, as observed in our patient. Nevertheless, association of H. pylori and thyroid MALT lymphoma has not been adequately addressed in the literature [3]. Thyroid gland does not contain any native lymphoid tissue; in fact, lymphoid tissues can only be seen in pathological conditions, mostly involving autoimmune processes like Hashimoto’s thyroiditis [2]. Histologically, lymphocytic infiltration of stroma with oxyphilic change of follicular epithelium is seen in autoimmune thyroiditis, as opposed to atypical lymphoid cells originating within marginal zone of lymphoid follicles and extending into interfollicular spaces and germinal centers (follicular colonization) which is seen in MALT lymphoma [2,3,6]. In our patient, FNA showed neoplastic proliferation of mostly singly dispersed small- to medium-sized cells with lymphoid cell proliferation, composed predominantly of small mature lymphocytes. As can be seen in Figure 1, histologic sections of thyroid parenchyma showing architectural effacement by an atypical lymphoid infiltrate with a focally nodular pattern. Also, demonstrated in Figure 2 is a focus of intraluminal lymphoma cells, representing a so-called ‘MALT ball.’ IHC stain and phenotype of neoplastic cells of MALT lymphoma are regarded to be virtually identical to that of non-neoplastic marginal zone B cells which are positive for CD20, CD 22, and CD79A and negative for CD5 and CD10.

Treatment of thyroid MALT lymphoma depends upon its stage and subtype. Localized thyroid MALT lymphoma can be treated with surgical resection with 100% survival at 5 years [5,8–10]. Review of 103 cases done by Tsang et al. with localized (stage IE/IIIE) extra nodal MALT lymphoma had overall 5 years survival of 98% in 85 patients with extra nodal MALT lymphoma treated with radiation therapy alone [11]. The optimal treatment regimen and follow-up for these patients remains controversial. Five-year disease-specific survival has been found to be variable (35–79%) in literature due to differences in population size, study design, lymphoma classification, and treatment modalities used [6]. The stage and histologic grade do play a major role in determining prognosis of the disease process. Diffuse large B-cell lymphoma has a poorer prognosis compared to localized primary MALToma. Stages I/II respond well to localized treatment as compared to stages III/IV which are more disseminated and need combination of chemotherapy and radiation therapy [6,10]. Patients who present with enlarging tumors or symptoms of compression also have poorer

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**3. Discussion**

Primary thyroid MALT lymphoma follows an indolent course and presents without any overt symptoms. Although most lymphomas present with typical ‘B’ symptoms such as fever, nocturnal sweating, and weight loss, this is usually not seen in this disease process, which make the diagnosis even more challenging [2,3]. Approximately 10% of patients may have these symptoms and 10% may have features of hypothyroidism, usually associated with Hashimoto’s thyroiditis [6].

MALT lymphomas are initially recognized at mucosal tissues, hence defined as tumors of mucosa-associated origin. Since first described by Isaacson and Wright [6], various extra nodal locations have been defined and studied. These were usually found to arise at the sites where ongoing chronic inflammation with underlying autoimmune or infectious etiologies was observed [2]. For example, MALT lymphoma of the stomach has been associated with chronic infection with Helicobacter pylori. A similar pathophysiology is seen with chronic inflammation of thyroid gland in the setting of ongoing Hashimoto’s thyroiditis, as observed in our patient.
prognosis [10]. Although this disease has excellent survival rate after treatment, the optimal follow-up remains controversial at present [5,10]. A usual practical approach is surgical excision for the localized disease, followed by radiotherapy and chemotherapy for disseminated disease [8]. However, some recent studies support radiation therapy as a primary modality of treatment in patients at Stages IE and IIE and adjuvant therapy for those who are suspected to have some residual disease even after thyroidectomy procedure [3,12]. Unfortunately, due to rare occurrence of PTL and lack of larger randomized controlled studies, definitive guidelines for treatment and follow-up on these patient groups are awaited.

Figure 1. Histologic sections of thyroid parenchyma on low power demonstrate architectural effacement by an atypical lymphoid infiltrate with a focally nodular pattern.

Figure 2. Histologic sections demonstrate intraepithelial lymphocytosis with involvement of the thyroid epithelium by the neoplastic lymphocytes. Shown is a focus of intraluminal lymphoma cells, representing a so-called ‘MALT ball’.
4. Conclusion

Primary thyroid MALT lymphoma follows an indolent process and remains asymptomatic in most patients with occasional nonspecific symptoms. It requires a high index of suspicion for diagnosis, and should be considered in patients with chronic inflammatory conditions of thyroid in presence of thyroid nodules. Treatment modalities include surgical excision and/or radiation therapy for localized lesions, with both radiation and chemotherapy indicated for disseminated disease.

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