Primary thyroid lymphomas constitute only 3% of non-Hodgkin Lymphomas (NHLs) and 5% of all thyroid neoplasms. Thyroid lymphoma is seen more frequently in women; the mean and median ages are 65 and 75 years, respectively. It is highly curable without surgery. We report a 34-year-old man with primary thyroid lymphoma arising from Hashimoto’s thyroiditis. The patient was admitted to the hospital for a growing mass on the left side of his neck within 2 months. MRI showed a 3 cm diameter, contrast-enhancing solid nodule at the left lobe adjacent to isthmus and bilateral cervical lymph nodes. The patient was referred to our hospital. A fine-needle aspiration biopsy (FNAB) showed a large number of atypical lymphocytes with necrosis. FNAB was suspicious for malignancy, and total thyroidectomy and left lateral neck dissection were applied. Pathology was reported as primary thyroid lymphoma. Surgery was followed by chemotherapy including rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisolone. No relapse has been detected during next 15-months follow-up. This is very rare occurrence of primary non-Hodgkin’s thyroid lymphoma arising from Hashimoto’s thyroiditis in a young man.
30 years from the beginning of thyroiditis. The majority of lymphomas are derived from B lymphocytes. Primary T-cell thyroid lymphomas are extremely rare. Clinical presentation is usually a rapidly growing thyroid mass. This mass grows rapidly which takes 1 to 3 months and it may cause obstructive symptoms such as dysphagia, dyspnea, and hoarseness.

CASE REPORT

Thirty-four-year-old male patient who had a rapidly growing mass on the left side of the neck was admitted to the State Hospital. He had no history of hoarseness, shortness of breath, hypothyroidism, and any drug usage. Neck MRI showed approximately 3 cm intense, heterogeneous, contrast-enhancing solid mass at the left lobe adjacent to the isthmus and bilateral cervical and submandibular 2.5 to 3 cm lymph nodes. Thyroid function tests showed subclinical hypothyroidism [TSH 9.43 mIU/ml (0.27-4.2) and FT4 1.14 ng/dl (0.93-1.7)]. Anti-thyroglobulin antibody level was high 218 IU/ml (0-115). A fine-needle aspiration biopsy (FNAB) was done, and a large number of necrosis and atypical lymphocytes were found. Total thyroidectomy and left lateral neck dissection were applied. The pathology was reported as primary thyroid lymphoma. The patient was referred to us for chemotherapy and further management. FT3 1.25 pg/ml (1.8-4.6), FT4 0.35 ng/dl (0.93-1.7), TSH, 64 mIU/ml (0.27-4.2), anti-thyroglobulin antibody 218 IU/ml (0-115) were found, postoperatively. At histopathological examination of surgical specimens the mass and lymph nodes, diffuse large cell lymphoma and Hashimoto’s thyroiditis were reported (Figure 1, 2). Positive immunohistochemical staining of B-cell marker (CD 10, 20) were detected. A L-thyroxine therapy was started (100 µg/day). At the postoperative period, positron emission tomography (PET) showed a weak FDG uptake at thyroid area probably due to postoperative inflammation, and paratracheal, right axillary lymph nodes. In this state, rituximab, cyclophosphamide, doxorubicin, vincristine and prednisone chemotherapy were given to the patient totally eight cycles of chemotherapy was applied. After chemotherapy, the patient has been followed without any recurrence for 15 months.

DISCUSSION

Primary thyroid lymphoma is usually seen in patients with advanced age and Hashimoto thyroiditis. Very rare cases were reported under the age of 40. Our case, who is 34-year old man is probably one of the extremely rare cases of thyroid lymphoma and Hashimoto thyroiditis. Most patients suffer from a rapidly growing neck mass; in 30% of patients, dysphagia, stridor, hoarseness and feeling of pressure around the neck were seen. This mass typically grew rapidly in 1 to 3 months.
Ten percent of patients diagnosed with thyroid lymphoma suffer from fever, night sweats and weight loss. Nine percent of patients are hypothyroid while 10% of them are euthyroid. A rapidly growing neck mass was the only complaint in our case and our patient was subclinically hypothyroid.

The vast majority of thyroid lymphomas are derived from B cells and the percentage from 60 to 80 percent of diffuse large B-cell lymphoma are considered to be. Thyroid lymphomas usually develop after 20-30 years from the beginning of Hashimoto’s thyroiditis. Although the relationship between thyroid lymphoma and Hashimoto’s thyroiditis is not understood clearly, Hashimoto thyroiditis is a well-known risk factor for thyroid lymphoma. Risk of thyroid lymphoma in patients with Hashimoto’s thyroiditis is 60 times more frequent than non-thyroiditis patients. In a study conducted in Japan, the risk of development thyroid lymphoma in patients with chronic lymphocytic thyroiditis is 70-80 times more than normal population. Lymphocytes in Hashimoto’s thyroid exposed to chronic antigenic stimulation were claimed to be caused by malignant transformation. Thyroid lymphomas may also occur after iodine supplementation and exposure to ionizing radiation. Although there are no specific and diagnostic laboratory abnormalities for thyroid lymphoma, approximately thirty percent of the patients has high serum IgA, IgM or IgG concentrations. In our case, there was subclinical hypothyroidism and high anti-thyroglobulin antibodies before the operation. Diffuse large B-cell lymphoma can be usually diagnosed after cytologic examination of material obtained by thyroid biopsy. An aspiration biopsy by a thick needle instead of fine needle was suggested to obtain enough material by some authors. Given the frequent association with Hashimoto’s thyroiditis, cytological diagnosis of small cell lymphomas are more difficult and may require immunohistochemical staining or flow cytometry. Unfavorable prognostic factors of thyroid lymphoma are increased tumor size (≥10 cm), advanced stage of disease (≥stage IE large), the presence of obstructive symptoms, the rapid growth of the tumor and the mediational involvement. Five-year survival rate is 80%, 50%, and lower than 36% for stage IE, stage IIE, stage IIIIE and IVE, respectively. Treatment plan depends on the histological type of thyroid lymphomas. Thyroid lymphomas are highly sensitive to both radiation and chemotherapy which consist of cyclophosphamide, doxorubicin, vincristine, and prednisone, but, the role of surgery in the treatment of this disease is limited, even in Stage IIE and IIIIE lymphomas. Surgical resection for thyroid lymphoma is not superior to any chemo- or radiotherapy modality and surgery has some known risks as well.

Finally, primary thyroid lymphoma is seen more frequently in women and elderly which is in contrast to our young male patients. In our case, although our patient has high probability of being evaluated by clinically and cytologically as lymphoma, non-thyroid malignancy has been suspected because of patient’s age and sex, and surgery have been performed and the lymphoma was diagnosed. If a patient presents with a rapidly growing mass in the thyroid, primary thyroid lymphoma should also be kept in mind, even among young and male patients, and means of diagnosis lymphoma from preoperative biopsy material should be investigated.

REFERENCES


