Case presentation – thyroid lymphoma

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Malignant tumors of the thyroid gland account for about 1% of the newly diagnosed malignant tumors each year, and their incidence in women is twice the incidence in men. According to the WHO classification (2004) thyroid tumors are divided into: carcinoma of the thyroid, adenoma and similar tumors, and other thyroid tumors which include: teratomas, angiosarcomas, paragangliomas and others, as well as primary lymphomas and plasmacytomas. Primary thyroid lymphomas are defined as lymphomas which originate in the thyroid gland. This study presents the case of a 68-year-old patient with a thyroid lymphoma, which caused compression of the airways. In the patient presented there was reduced activity of the thyroid gland. The dominant symptoms were: breathing difficulties, hoarse voice and the enlargement of the thyroid. An ultrasound examination was performed before surgery on the neck, which showed a multinodular thyroid, with compromised and compressed trachea to the right and rear. An emergency surgical procedure was performed to reduce the tumor. Pathohistological diagnosis confirmed diffuse large B cell lymphoma. The aim of the study was to present a patient with a thyroid lymphoma, who had previously not had any immunological changes to the gland, that is, she had not had any chronic lymphocyte thyroiditis, but due to the compressive syndrome it was necessary to perform an emergency surgical procedure to reduce the tumor.

Key words: Thyroid, Lymphoma, Malignant tumor.

Introduction

Malignant tumors of the thyroid gland account for about 1% of the newly diagnosed malignant tumors each year (1) and their incidence in women is twice the incidence in men (2). They most often occur in older patients where the average age of occurrence is 65 years. According to the WHO classification (2004), thyroid tumors are divided into: carcinoma of the thyroid, adenoma and similar tumors, and other thyroid tumors which include: teratomas, angiosarcomas, paragangliomas and others as well as primary lymphomas and plasmacytomas (3). In the group
of carcinomas the most common in the thyroid are papillary (80%), follicular (10%), medullary (5-10%) and the exceptionally rare anaplastic carcinoma (1-2%) (2), whilst the frequency of primary lymphoma is 5% (4). Primary thyroid lymphomas are defined as lymphomas which primarily originate in the thyroid, so this definition excludes lymphomas which affect the thyroid whether by metastasis or by direct spreading from the neighboring lymph nodes. Primary thyroid lymphoma in its late stages may spread to the lymph nodes and other organs, including the gastrointestinal tract, thereby representing a form of mucosa-associated lymphoid tissue (MALT) lymphoma (5).

Primary thyroid lymphomas are usually non-Hodgkin type, whilst primary Hodgkin's disease is very rare (6). Extranodal marginal zone B cell lymphoma (EMZBCL) and diffuse large B cell lymphoma (DLBCL) are lymphomas which most often occur in the thyroid with areas of morphology of a transient form between these two types of lymphoma. Other, especially follicular lymphomas are extremely rare (3). Diffuse large cell B lymphomas are lymphomas which occur most often in the lymph nodes, whilst in 30% of cases they can occur as extra-nodal. The most common extra-nodal site where lymphomas of this type occur is the thyroid (7).

The rapidly growing mass of the thyroid gland in an older woman should always arouse suspicion of a tumor. In the past it was necessary to perform an open biopsy to obtain enough tissue for pathohistological examination (6). However, today diagnosis is made on the basis of cytological analysis of aspirate changes in the thyroid gland, immunocytochemistry, flow cytometry, immunophenotypization of lymphocytes obtained by fine needle aspiration (FNA), and finally pathohistological analysis and molecular genetic analysis (8). If it is a DLBCL lymphoma, that is, a lymphoma with a high degree of malignity, there is usually no problem in establishing a diagnosis from smears obtained by FNA. In cases of unclear diagnosis, if a monomorphic population of small lymphatic cells is found in the smears, it is necessary to repeat the aspiration to obtain sufficient cells for immunophenotypization (6). Differentiating between EMZBCL and lymphocyte thyroiditis, as well as differentiating between other lower level lymphomas than Hashimoto thyroiditis can sometimes be difficult both from cytological smears and histological preparations. Although pathological verification is the gold standard for diagnosis of lymphoma, in rendering a diagnostic decision immunohistochemistry, flow cytometry or genetic molecular analysis are still necessary in addition (3).

Treating large cell lymphomas of the thyroid gland is no different from treatment of any other lymphoma occurring in the lymph nodes (9). Treatment is based on the sub-type of the lymphoma and the scope of the disease. Today, the trend in treatment of large cell lymphomas is for treatment to be selected on the basis of prognostic factors. Treatment of a disease which is limited to the thyroid gland is performed by local regional radiation or surgery. In most patients, treatment is mainly by a standard chemotherapy regime, which consists of Cyclophosphamide, Doxorubicin, Vincristine and Prednisolone (CHOP) in combination with radiation, and must always be under the supervision of an oncologist (11).

The goal of this paper is to present a patient with thyroid lymphoma, who had not had any previous immunological disease of the thyroid gland, that is, chronic lymphocyte thyroiditis, and in whom it was necessary to perform surgery to reduce the tumor due to compression syndrome.

Case presentation

The patient, aged 68, was sent to our institute for examination. A year earlier, the
diagnosis of hypothyroidism had been estab-
lished in another institution and therapy
with 1-thyroxin 100 micrograms once a day
prescribed. On clinical examination, a node
was palpitated in the left lobe of the thyroid
gland, 2 cm in size. The follow up finding
of the thyroid-stimulating hormone - TSH
was 9.9 mIU/l (0.27-3.75 mlU/l); Free thy-
roxine - FT4 17 pmol/l (10-20 pmol/l); Thy-
roglobulin - Tg 0.4 ng/ml (2.0-70 ng/l) and
thyrogblobulin antibodies - TgAt 9.0 IU/ml
(>100 IU/ml). Therapy was recommended
with 150 micrograms of l-thyroxin once a
day and control of hormonal status in four
months.

Three months later the patient was ad-
mitted to our Department of Ear, Nose
and Throat for breathing difficulties, loss of
voice and sudden increase in the size of the
thyroid gland. Physical examination estab-
lished that the thyroid gland was enlarged,
with multimodal changes, hard, painless
and immobile on swallowing, with bilateral
neck lymphadenopathy. Ultrasound showed
multimodal changes to the thyroid gland
with a compromised and compressed tra-
chea on the right and to the front. On both
sides of the neck there were pockets of en-
larged lymph nodes. The ultrasound finding
indicated a malignant tumor in the thyroid
gland. Since compressive syndrome and re-
spiratory insufficiency were present, it was
proposed following consultation that tumor
reduction and tracheotomy be performed.

The pathohistological diagnosis of the tu-
morous mass was Lymphoma malignum dif-
fusum centroblasticum B glandula thyreoide-
ae – diffuse large cell B lymphoma (figure 1).
Immunohistochemical analysis showed that
the neoplastic cells were positive for CD20
(figure 2) and bcl-6, whilst they were nega-
tive for CD 3, CD 10, CD 23, cyclin D1, TdT
and bcl-2.

Post-operative computerized tomography
of the neck, thorax and abdomen showed the
remains of the tumorous mass, which went
from the level of the hyoid bone to the left,
paralaryngeally and para- and retropharyn-
geally, right up to the level of the jugular fos-
sa. It infiltrated the left half of the supraglot-
tic and glottic regions of the larynx, reducing
the air space to a latero-lateral diameter of 2
mm. In the distal half of the neck to the left,
pathologically enlarged lymph nodes were
visible on the jugular chain about 15 and 24
mm in size, which compromised and, it was
suspected, also infiltrated the jugular vein
internally. In view of the stage of the illness,
further therapy was proposed and transfer
of the patient to the Oncology, Hematology
and Radiotherapy Clinic. During her hos-
pitalization she received a second cycle of

Figure 1 Diffuse large cell B lymphoma of thyroid
gland (HE, x 10)

Figure 2  Immunohistochemical analysis - CD20
positive, x 40.
chemotherapy according to the CHOP regime, with the addition of monoclonal CD 20 antibodies. On both occasions she had iatrogenic neutropenia which was treated with granulocyte growth factor and broad spectrum antibiotics. There were also raised levels of the enzymes lactate dehydrogenase, aspartate transaminase and creatinine present in the serum. Despite the therapy applied no regression of the tumorous mass was recorded, so the patient was still unable to swallow spontaneously, but was fed through a nasogastric tube. After two months from admission to hospital in the ENT clinic the illness progressed into the front of the upper mouth and gingiva of the upper jaw, there was loss of body mass and the patient died.

Discussion

Our patient was treated for one year for reduced activity of the thyroid gland, that is, hypothyroidism. The medical history and negative Tg-At finding excluded the existence of Hashimoto thyroiditis. In a relatively short period of time the thyroid gland increased rapidly in size due to the infiltration of neoplastic cells in the form of diffuse nodes, with compression of the airways and resulting respiratory insufficiency. This condition demanded urgent surgical intervention, that is, total thyroidectomy with bilateral dissection of the neck.

The incidence of primary thyroid lymphoma in patients with Hashimoto thyroiditis has increased significantly (7, 10). In fact lymphomas in the thyroid gland in almost all cases occur on the basis of chronic lymphocyte thyroiditis. Graff-Baker et al. believe that there is a pathophysiological connection between autoimmune disorder and thyroid lymphomas. The proposed theory supposes that chronic antigen stimulation is a secondary autoimmune disorder which leads to chronic proliferation of lymph tissue, with resulting mutation leading to the development of lymphoma (9).

Until recently there was concern whether the diagnosis of lymphoma could be established by analysis of a sample obtained by aspiration. Cha et al. (11) showed that this approach is successful in 63% patients with thyroid lymphoma. Aspirates of large cell lymphomas are typically hypercellular with visible individual lymphatic cells, which have a cytological appearance, similar or identical to lymphoma of other sites, and in that case it is simple to reach a diagnosis. In contrast to them, in aspirate smears of the lymphoma marginal zone a mixture is found of small atypical lymphocytes, centrocytes, monocytoid B cells, immunoblasts and plasma cells. As a result of the morphological appearance of the cytological smears, differentiating this type of lymphoma and reactive changes in the lymph node is almost impossible (3). The introduction of molecule technology has led to definite diagnosis and as a result open thyroid biopsy is no longer performed in modern diagnostics (6). Takashima et al. (12) and Daria et al. (13) described the use of chain reaction polymerase in strengthening the immunoglobulin heavy chain to establish the diagnosis of lymphoma. The presence of antigen CD-20 and heavy chain clonality confirm the diagnosis.

Large cell B lymphoma in two patients aged 64 and 50 years, without a previous history of Hashimoto thyroiditis, were described by Akcala et al. (14), saying that due to the sudden increase in size of the thyroid gland and compression of the respiratory path, total thyroidectomy was performed and CHOP chemotherapy applied.

Conclusion

This unusual case of thyroid lymphoma is interesting because it occurred in a patient who did not have Hashimoto thyroiditis.
The surgery was performed because of compression of the airways caused by the rapid increase in size of the thyroid gland due to infiltration by neoplastic cells. After two rounds of chemotherapy there was no regression of the disease and the patient died.

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