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2 **Synchronous thyroid and gastric mantle cell lymphoma.**

3 **Running title: thyroid and gastric mantle cell lymphoma.**

4 **Abstract:**

5 **Introduction:**

6 Mantle cell lymphoma (MCL) is a distinct entity within the World Health Organization
7 classification of lymphoid neoplasm and represents approximately 8% of lymphoma. Patients
8 with mantle-cell lymphoma typically present with extensive disease and involvement of
9 multiple lymph nodes as well as the spleen, bone marrow, blood, and gastrointestinal tract.
10 MCL of the thyroid occurs exceptionally. The MCL of the stomach is also an exceptional
11 occurrence.

12 **Observation:** we describe the case of a 58-year-old male who was diagnosed with thyroid and
13 gastric MCL. The patient was classified into high risk group according to the Mantle Cell
14 Lymphoma International Prognostic Index (MIPI). The R-CHOP (Rituximab,
15 Cyclophosphamide, Adriablastine, Vincristine and Prednisone) regimen was started and
16 complete remission was achieved after 8 courses. He currently receives a maintenance
17 treatment with rituximab every two months.

18 **Conclusion:** This case is a combination of two rather infrequent extranodal localizations of
19 the MCL.

20

21 **Keywords:** Mantle cell lymphoma- thyroid- stomach- chemotherapy.

22

23 **Introduction:**

24

25 Thyroid non-Hodgkin's lymphoma (TNHL) represents 2-8% of thyroid malignancies and 1-
26 2% of extranodal lymphomas [1]. Diffuse large B cell lymphoma is the most common
27 histological type, accounting for up to 70% of primary TNHL [2]. The mucosa-associated
28 lymphoid tissue lymphoma (MALT) accounts for 15-40% of primary TNHL [2]. Follicular
29 lymphoma of the thyroid is very rare. Mantle cell lymphoma of the thyroid (MCL) occurs
30 exceptionally. In the gastrointestinal tract, the MALT is the most common low-grade
31 lymphoma, arising mainly in the stomach (60%-70%) [3]. The MCL of the stomach is also an
32 exceptional occurrence. To our knowledge, this is the first report of a patient with
33 synchronous thyroid and gastric MCL.

34

35 **Case report:**

36 A 58-year-old male was admitted in the department of ENT for further evaluation of a mass
37 of the thyroid gland, associated with gradually increased pain and dyspnea. He had no family
38 or personnel history for thyroid pathology and gastric complaints. The local examination of
39 the thyroid revealed a painless palpable mass which was hard in consistency, fixed to the
40 musculature and invading the entire thyroid (Figure 1). The ECOG Performance Status was

41 equal to 2. The rest of the physical examination was normal (no palpable lymph nodes and no
42 hepatosplenomegaly). Serum laboratory values, including LDH, b2-microglobulin, fT4 and
43 TSH were within normal ranges. Anti-TSH receptor antibodies were absent. Viral serology
44 and particularly HIV, HBV, HCV and EBV tests were negative. Complete blood cell count
45 was normal. Ultrasound revealed a heterogeneous nodule involving almost the entire lobe of
46 the thyroid. The thyroid fine needle aspiration was not performed. After a biopsy of the
47 thyroid mass, histological examination demonstrated a diffuse lymphomatous infiltrate.
48 Lymphoepithelial lesions were characterized by neoplastic lymphocytes that infiltrated and
49 destroyed thyroid follicles, often showing regressive changes. Lymphoma cells appeared
50 monotonous and slightly larger than small lymphocytes. Their nuclei displayed variable
51 degrees of angulation with fairly condensed chromatin and their cytoplasm was very scanty
52 (Figure 2A). Immunohistochemically, the tumor cells were positive for CD20, cyclin D1 and
53 CD5 (Figure 3) and negative for CD23, CD10, and the epithelial membrane antigen. Few
54 CD3 positive lymphoid cells were detected. Ki 67 was identified in 80% of neoplastic cells.
55 In consequence of this finding, the tumor was diagnosed as MCL. The examination of the
56 ENT was normal. Computed tomography scans showed cervical lymph node associated with
57 two nodular thickening at the cardia and fundus regions of the gastric wall. The gastroscopy
58 showed a loss of substance of 15 mm in diameter at the gastric antrum whose biopsy revealed
59 the infiltration of the gastric mucosa by the same lymphoid cell proliferation (Figure 2B). The
60 cells were also positive for CD20, CD5 and cyclin D1 and negative for CD10. Ki 67 was
61 identified in 75% of neoplastic cells. Helicobacter pylori infection was not detected. In
62 consequence of this finding, the diagnosis of gastric MCL was confirmed. The colonoscopy
63 was not performed. The bone marrow biopsy revealed the absence of a medullary extension
64 of the lymphoma. Cytogenetic study of the bone marrow cells was normal. Cytogenetic
65 analysis was not performed on the fragments of the thyroid and gastric biopsy. The final
66 diagnosis was a double gastric and thyroid localization of MCL. After this staging,
67 lymphoma was classified as stage IV according to the classification of Ann Arbor. The
68 patient was classified into high risk group according to the Mantle Cell Lymphoma
69 International Prognostic Index (MIPI). The R-CHOP (Rituximab, Cyclophosphamide,
70 Adriablastine, Vincristine and Prednisone) regimen was started and complete remission was
71 achieved after 8 courses. Six intrathecal prophylaxes therapy with 12 mg methotrexate were
72 done. Control gastroscopy showed a cicatricial ulcer of the antrum whose biopsy was
73 negative. The autograft was refused by the patient. He currently receives a maintenance
74 treatment with rituximab every two months. Rituximab maintenance therapy will be applied
75 for 2 years. No relapse has occurred during a follow-up of 4 months.

76

77 Discussion:

78 MCL is an aggressive lymphoma of older adults, with a male preponderance and it represents
79 6% of all NHL [4] and just a minority of the extra nodal lymphomas [5]. Clonal plasma cell
80 differentiation may occur within germinal center in some cases of MCL [6]. Patients with
81 extra nodal MCL will be found, in the most of cases, to have lymphadenopathy or more
82 widespread disease on staging [5]. Lymphoproliferative disorders affecting the thyroid are
83 characterized by diverse clinical and pathologic spectrum and must be differentiated from

84 carcinoma and benign thyroiditis. **MCL of the thyroid** is an exceptional occurrence. The
85 clinical presentations include an enlarging neck mass, as in our case, but patients may also
86 present the symptoms of dysphagia, hoarseness and choking, or a cold thyroid nodule [7].
87 Since MCL of the thyroid is an uncommon malignancy, a misdiagnosis is possible. Other
88 malignant thyroid tumors, especially anaplastic carcinoma, and other lymphomas, such as
89 follicular lymphoma and marginal zone lymphoma must be differentiated from MCL because
90 of the subsequent management strategies. In such cases, diagnosis and subclassification can
91 be established using study of routine sections augmented by immunohistochemistry [8].
92 **Despite the absence of digestive clinical symptoms in our case, the gastroscopy showed a**
93 **gastric infiltration by the MCL.** In other cases, patients may have diarrhea and abdominal
94 pain [9]. By using additional immunological and molecular markers, lymphomas are
95 classified into subtypes according to the **World Health Organization** classification and that is
96 important for further decision making. For an adequate prognostic evaluation and appropriate
97 clinical decisions, histological diagnosis must be combined with IPI prognostic parameters.
98 The MCL international prognostic index has been proposed as a new prognostic index for
99 MCL. It considers age, performance status, LDH level and leukocyte count as prognostic
100 factors [10]. In MCL, **gastrointestinal** tract involvement has not been identified so far as an
101 adverse prognostic factor [11]. Our patient **presented** with synchronous thyroid and gastric
102 MCL justifying systemic treatment with chemo immunotherapy. The poorest 5-year survival
103 of all the non-Hodgkin's lymphoma subtypes in the NHL classification project was observed
104 with MCL and it is considered to be incurable with standard therapies [12]. CHOP plus
105 rituximab (R) is associated with high response rates but the progression-free survival (PFS) is
106 disappointingly short (median 16–20 months) [13, 14, 15]. A benefit for selected patients
107 using autologous stem cell transplantation (ASCT) consolidation in first remission has been
108 suggested in some phase II studies and registry studies [16–17]. However, many patients are
109 not eligible for autograft and randomized clinical trial did not demonstrate the prolongation in
110 overall survival with this strategy [18]. A better outcome with a regimen consisting of R-
111 hyper CVAD (fractionated cyclophosphamide, vincristine, doxorubicin, dexamethasone plus
112 rituximab) alternating with rituximab plus methotrexate and cytarabine (R-Mtx/AraC) has
113 been reported [19]. But, this regimen can be toxic for patients over the age of 65 and younger
114 patients with co-morbid illness. Since the median age for newly diagnosed mantle cell
115 lymphoma patients is 64, approaches that do not include stem cell transplantation or involve
116 highly aggressive chemotherapy regimens need to be developed. Two large studies show a
117 better PFS for untreated MCL by the application of maintenance rituximab for 2 years
118 following the completion of a moderately aggressive chemo immunotherapy regimen [20-
119 21]. Our patient had an excellent response with R-CHOP, although this regimen is no more
120 considered the first line therapy in MCL. **Two large studies show that induction with**
121 **rituximab and cytarabine-based regimens [22] and the addition of lenalidomide to rituximab-**
122 **bendamustine (R-B) [23] as first-line treatment to elderly MCL patients had been associated**
123 **with a high rate of CR and molecular remission.** In our case, a further follow up is necessary
124 to detect a relapse.

125

126 **Conclusion:**

127 In conclusion, the double localization (thyroid and gastric) and the histological type MCL of
128 the lesion make our patient's case really remarkable.

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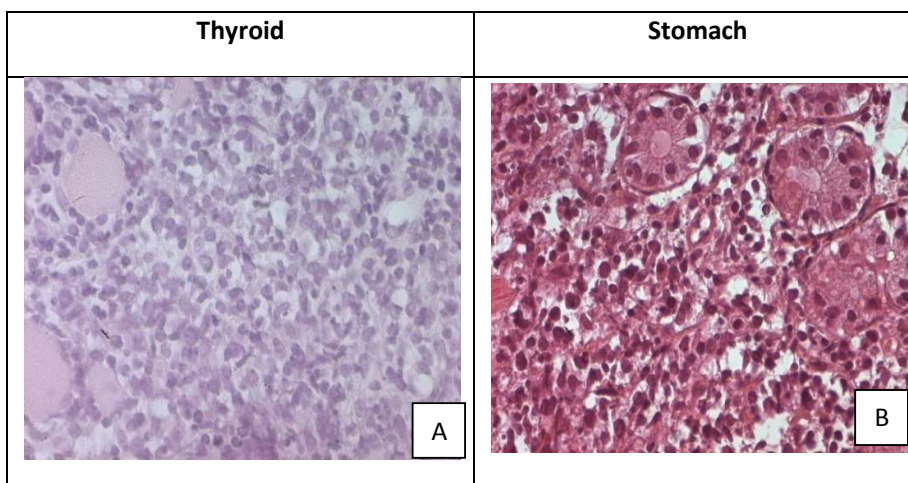
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Figure 1: The mass of the thyroid gland.

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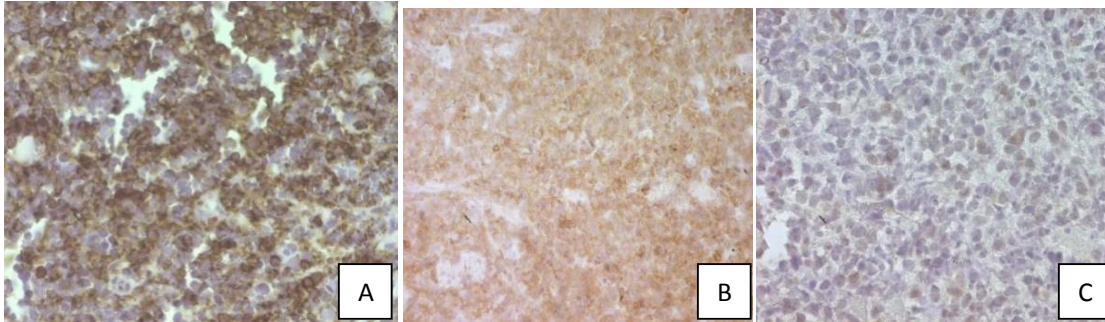


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228 **Figure 2:** (A)-Extensive lymphoid infiltrate destroys the thyroid tissue (hematoxylin-eosin,
229 400X).

230 (B)- The infiltration of the gastric mucosa by the same lymphoid cell proliferation
231 (hematoxylin–eosin, 400X).

232



233

234 **Figure 3:** The tumor cells were positive for CD20 (A), CD5 (B) and cyclin D1 antigen (C)
235 (Original magnification 400X).

236