Immunohistochemistry contribution in the diagnosis of splenic marginal zone lymphoma

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Dear Editor,

Splenic marginal zone B-cell lymphoma is a rare type of non-Hodgkin B-cell lymphoma, without preference for sex, which affects individuals mainly over 50 years. In many cases, the disease is asymptomatic, with absent splenomegaly and lymphadenomegaly. The changes in the peripheral blood, when present, are lymphocytosis, anemia, and thrombocytopenia; cytopenias are mostly due to hypersplenism. The presence of B symptoms and increased LDH is rare. The immunophenotype pattern is characterized by pan-B cell expression with positive CD19, CD20, CD22, CD79a, IgM, and IgD. CD5 is found only in 15%, and CD23 in 30% of cases. It presents as the most common cytogenetic alteration the heterogeneous expressions of chromosome 7, the (9:14) (p13: q32), which overlaps the IgH and PAX-5 genes. The marrow infiltration patterns are highly variable and consist of nodular, paratrabecular, and interstitial. The finding of the intrasinusoidal pattern is more easily observed by CD20 or CD34 immunohistochemistry. Spleen histology shows micronodular infiltration of white pulp with biphasic pattern and a variable degree of red pulp involvement. Differential diagnosis should be made with other chronic B7 cell lymphoproliferative diseases.

The clinical course is mostly indolent with a median global survival of 10 years. There is no standard therapy; so, splenectomy is the most used
A therapeutic approach since it helps in diagnosis as well as in the improvement of symptoms related to cytopenias. Other therapies using chlorambucil or cyclophosphamide, alone or in combination with other agents (CVP, CHOP), have limited results. Immunotherapy with an anti-CD20 antibody is a molecular target for the treatment of non-Hodgkin B cell lymphoma with CD20 positive protein. Results of anti-CD20 monotherapy demonstrated a 17-month duration of response and global survival of 80% up to 6 years and progression-free survival of 60% up to 4 years.

We report the case of a patient with splenic marginal zone lymphoma with positive markers for B-lymphocytes, who presented asymptomatic and with splenomegaly, and review some aspects related to this disease.

A 50-year-old male, businessman, from Fortaleza (CE), presented, in 2019, at a hematology service unit asymptomatic with splenomegaly and blood count results within normal limits. A bone marrow biopsy presented normal results for hematoxylin-eosin (HE), but immunohistochemistry was positive for CD20 in sinusoid, diagnosed with marginal zone lymphoma (Figure 1). A significant white and red pulp labeling for CD20 was observed, while the markers CD5, CD3, cyclin D1, BCL-6, and CD10 were not significant. CD23 marks dendritic and Ki67 follicular cells in 30% of the cells; normal cytogenetics. Splenectomy was performed with the anatomopathological study of the spleen (Figures 2), and the diagnosis was confirmed as marginal splenic zone lymphoma. The first recovery cycle was started and has shown stability and no toxicity until now.

Splenic marginal zone lymphoma infiltrated into the bone marrow, usually used with multiple patterns and usually forming lymphoid nodules with germinal centers; the cellular pattern is similar to the initial site. The sinusoidal pattern is strongly associated with marginal zone lymphoma; however, the most recent reports have shown that this pattern is very nonspecific, rarely happens as the only infiltration pattern and that immunohistochemistry is necessary to identify the infiltrations.

We presented a case that reinforces the importance of anatomopathological examination of the bone marrow, whenever possible, complementary to other laboratory tests (peripheral blood, cytogenetics, immunophenotyping, and molecular), as well as the study of immunohistochemistry and interpretation in

**FIGURE 1. INTRASINUSOIDAL MEDIAL INFILTRATION BY SPLENIC MARGINAL ZONE NHL-B (CD20 +) (200X IMMUNOHISTOCHEMICAL STUDY)**

**FIGURE 2. A: MARGINAL ZONE SPLENIC NON-HODGKIN'S LYMPHOMA, NODULAR ASPECT (HE 200X). B: SPLENIC LNH-B, FROM THE MARGINAL ZONE WITH STRONG EXPRESSION OF CD20 (200X IMMUNOHISTOCHEMICAL STUDY)**
the clinical context. Anatomopathological examination of the spleen was also important to define the diagnosis of splenic marginal zone lymphoma.

Conflict of interest
The authors have no conflict of interest to declare.

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