Simultaneous occurrence of follicular lymphoma and mixed-cellularity Hodgkin’s lymphoma: lymph node and extranodal involvement

Ocorrência simultânea de linfoma folicular e linfoma de Hodgkin celularidade mista: envolvimento nodal e extranodal

Maria do Patrocínio F. Grangeiro; Silvia Maria M. Magalhães; Francisco Valdeci A. Ferreira; Francisco Dário Rocha Filho

Abstract

An unusual and well-characterised case of composite lymphoma in the spleen and lymph node is presented. The simultaneous occurrence of mixed-cellularity Hodgkin’s lymphoma (HL) and follicular non-Hodgkin’s lymphoma (NHL) was demonstrated in a 66-year-old man admitted in our Service with anaemia, hepatosplenomegaly and multiple abdominal lymph nodes. The morphological study of the spleen and lymph node of the splenic hilum showed an infiltrate composed of two distinct neoplasias. The liver was involved by NHL infiltrate and the peripancreatic lymph node exhibited HL. The Reed-Sternberg (RS) cells expressed CD 15 and CD 30, whereas the NHL cells presented standard immunohistochemical features of follicular lymphoma. To our knowledge, this is the fifth case report of concurrent spleen involvement by composite lymphoma. The incidence, clinicopathological and immunohistochemical features of this rare association are discussed.

Keywords

Composite lymphoma
Spleen
Lymph node
Follicular lymphoma
Hodgkin’s lymphoma

resumo

Os autores apresentam um caso raro e bem caracterizado de linfoma composto no baço e linfonodos. A ocorrência simultânea de um linfoma de Hodgkin e um linfoma não-Hodgkin foi demonstrada em um homem de 66 anos admitido com anemia, hepatosplenomegaly e múltiplos linfonodos abdominais. O estudo morfológico do baço e linfonodo do hilo esplênico mostrou infiltração por duas neoplasias distintas. O fígado estava envolvido por linfoma não-Hodgkin e o linfonodo peripancreático mostrava um linfoma de Hodgkin. As células de Reed-Sternberg (RS) expressaram CD 15 e CD 30, enquanto as células do linfoma não-Hodgkin apresentaram as aspectos imunoistoquímicos clássicos do linfoma folicular. Na literatura médica, este é o quinto caso descrito de envolvimento do baço por linfoma composto. A incidência, aspectos clínicos e imunoistoquímicos desta rara associação são discutidos.

Keywords

Linfoma composto
Baço
Linfonodo
Linfoma folicular
Linfoma de Hodgkin

1. Professor of the Department of Pathology and Legal Medicine of Universidade Federal do Ceará (UFC)/Hemoce.
2. Professor of the Department of Clinical Medicine of UFC/Hemoce.
Introduction

The simultaneous occurrence of two different subtypes of lymphoma within the same or in more than one anatomical site is defined as composite lymphoma. Composite lymphoma is rare and its incidence varies among the different types of lymphomas and depends on the number of anatomic sites studied at the time of diagnosis or during the course of the illness (5, 13). In some cases Hodgkin’s lymphoma (HL) is associated with non-Hodgkin’s lymphoma (NHL). The association of non-classical HL and NHL is more common. This phenomenon occurs more frequently than it would be expected by chance, supporting the clonal relationship between the two tumours (6, 7). In most cases, however, the composite lymphoma is a combination of NHL of different histological types (3, 5, 7). In these cases the immunophenotypic and genotypic analysis have indicated that the two histologic types are clonally related (3, 7). The most common anatomic site involved is the lymph node, although cases arising in extranodal sites have also been reported (11).

Case report

A 66-year-old man was admitted to the University Hospital because of asthenia, progressive paleness and weight loss. There was no history of fever, night sweat or any other systemic complaints. Physical examination revealed paleness and moderate hepatosplenomegaly. No peripheral lymphadenopathy was evident. Peripheral cell counts showed a red blood cell count of 3.2 x 10^12/L, haemoglobin level of 8.4g/dl, hematocrit 25%, white blood cell count of 10.9 x 10^9/L and platelet count 297 x 10^9/L. Alkaline phosphatase level was 460U/L (normal range: 50 to 250U/L) and lactate dehydrogenase level was 563U/L (normal range: 150 to 360U/L). An abdominal ultrasonographic study showed moderate liver enlargement and multiple retroperitoneal nodular images suggestive of lymph nodes. The patient underwent diagnostic laparotomy and splenectomy was performed along with biopsy of the liver, omentum and peripancreatic lymph nodes. The macroscopic examination revealed an enlarged spleen (19 x 16 x 8cm) which weighed 900g. Sections of the spleen revealed a prominent white pulp and whitish well-delimited nodule next to the capsule, measuring 2cm in diameter (Figure 1). Peripancreatic and splenic hilum lymph nodes were compact and whitish.

The histological analysis showed composite lymphoma in the spleen and lymph node of splenic hilum. The spleen exhibited a white pulp infiltrated by lymphoid neoplasia composed of a monotonous population of small cells consistent with follicular NHL (Figure 2A). The nodular pericapsular area showed occasional typical Reed-Sternberg (RS) cells against a background of lymphocytes, eosinophils, histiocytes, plasma cells; diagnosis of classical HL was made. The HL infiltration represented 10% of the spleen and was clearly distinct from the NHL component (Figure 2B). The lymph node of splenic hilum exhibited both diseases and the NHL and HL components were also in distinct areas from each other. The HL area infiltrated 40% of the lymph node.

Microscopic examination of the peripancreatic lymph node revealed a diffuse effacement of the architecture by HL. Extensive areas of fibrosis, foci of necrosis and numerous RS cells and variants, admixed with lymphocytes, eosinophils, histiocytes and plasma cells were seen. In the liver, infiltration of the portal space by small cell lymphoma, similar to that seen in the white pulp of the spleen, was evident (Figure 3A).

Immunohistochemical analysis, performed on formalin-fixed paraffin-embedded sections by the PAP method, revealed that the small lymphoid-cells of the liver, spleen and lymph node of the splenic hilum expressed CD 45, CD 20 and Bcl 2 (Figure 3B). The RS cells of spleen and lymph node exhibited expression of CD 15, CD 30 and LMP1/EBV (Table and Figure 4).

After a short-term follow-up, the disease evolved into sepsicaemia and the patient died 17 days after surgery, without having started chemotherapy.
In the present report the authors describe a well-characterised composite lymphoma involving the spleen and lymph node composed of a low-grade NHL and a classical HL. The histological features and the immunological profile were consistent with the diagnosis of follicular non-Hodgkin’s lymphoma and mixed-cellularity Hodgkin’s lymphoma.

Composite lymphoma is rare and is usually described in lymph nodes (1, 2, 4, 11). The concurrent involvement of the spleen has been rarely reported. To our knowledge, this is the fifth report in the English language literature of a composite lymphoma involving the spleen and presenting prominent splenomegaly. In the previous reported cases the simultaneous occurrence of lymphocyte-predominance HL/large-cell NHL and lymphoplasmacytoid/large-cell NHL was presented (8, 10, 11, 12).

Table

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<th>Summary of immunophenotypic features</th>
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HL: Hodgkin’s lymphoma; RS: Reed-Sternberg.
Recent studies using different techniques have focused on identifying a common clonal origin for both disorders\(^1, 11\). Molecular investigations strongly suggest that a common B-cell precursor located in the germinal center gave rise to both HL and NHL. As Hodgkin and Reed-Sternberg cells are clonal and originated from germinal center B-cell precursor in most instances\(^9\), it is reasonable to suppose that the occurrence of B-cell lymphoma concurrent with HL may represent divergent differentiation of the same oncogenic process.

In this paper the authors showed an unusual case of composite lymphoma involving the spleen and lymph node, confirmed by immunohistochemistry. The number of cases reported is too small for a meaningful analysis and conclusion about prognosis. However, there is an emerging consensus that the composite lymphoma has the same prognosis of the more aggressive component and that treatment must point to the aggressive component. In our case sepsicaemia and the consequent patient’s premature death prevented any chemotherapeutic approach.

References