Clinical Notes

Post transplantation human herpes virus-8 unrelated primary effusion lymphoma of the peritoneal cavity in a HIV-negative female

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Summary

Primary effusion lymphoma (PEL) is a recently individualized form of non-Hodgkin lymphoma (WHO classification), developing mainly in HIV-infected males, more frequently homosexual, in advanced stages of the disease (total CD4+ lymphocyte count below 100-200/µl). Occasionally, it appears in other immunosuppressive states (such as solid organs transplantation period) and even, although very rarely, in immunocompetent patients. From a pathogenic point of view, PEL has been related to Kaposi's sarcoma associated herpes virus (also named human herpesvirus 8, HHV 8) and to clinical antecedents of Kaposi's sarcoma. The relatively low frequency of this disease, the absence of a wide casuisticsts allowing a better characterization, and its unfavourable outcome, support the need of a deeper knowledge. We present here the clinico-biological findings of a HIV-negative patient, who was diagnosed of peritoneal PEL, of T cell origin, and not HHV 8-associated, five years after renal transplantation.

Key words: Primary effusion lymphoma (PEL). Non-Hodgkin lymphoma.

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Introduction

Primary effusion lymphoma (PEL) has recently been identified as a distinct subtype of non-Hodgkin lymphoma associated with infection of the neoplastic lymphoid cells by the Kaposi sarcoma-associated herpesvirus/human herpesvirus 8 (HHV-8)1, 2. Primary effusion lymphoma has characteristic clinicopathologic features, including initial presentation as a lymphomatous effusion usually in the absence of a detectable tumor mass, occurs mostly in human immunodeficiency virus (HIV)-positive men, and has a morphologic structure that bridges large cell immunoblastic and anaplastic large cell lymphoma (ALCL)1, 2. Neoplastic lymphoid cells are B cells with a peculiar phenotype. They usually lack surface immunoglobulin and B-cell-associated antigens such as CD19 and CD20 and express CD45, CD30, and antigens associated with late stages of B-cell differentiation such as CD1381, 2. Genotypic analysis of PEL has revealed clonal immunoglobulin gene rearrangements in all cases1, 2. We describe here a unique case among PELs arising in a HIV seronegative female, not associated with HHV 8 infection, after renal transplantation.

Case report

We report a case of a 27-year-old woman who was admitted to our hospital because of dizziness, headache and malaise. Five years earlier the patient had a renal transplantation due to chronic renal failure. From that period the patient was under immunosuppression. She was HIV negative. On physical examination ascites was found but no peripheral lymphadenopathy or hepatosplenomegaly. Laboratory data on admission was as follows: WBC=6.7x10³/ul, hemoglobin=12.5gm/dl, platelets=409x10³/ul, LDH=144U/I, total protein=3g/dl, albumin=1.2g/dl, cholesterol=217mg/dl, liver function tests, amylase and lipase were normal. HIV screen was negative. Peritoneal fluid showed albumin <1g/dl, LDH=144U/I, total protein=3g/dl, albumin=1.2g/dl, cholesterol=217mg/dl, liver function tests, amylase and lipase were normal. The ascitic fluid could not interpret patients’ symptoms, an exploratory laparotomy was performed. During the operation 3lt ascitic fluid were removed but no other abnormalities were found except from a slightly thickened omentum and peritoneum from which biopsy samples were obtained. Cytological examination of the fluid, histologic interpretation of the excised specimens, immunophenotypic and molecular workup was indicative of primary T-cell effusion lymphoma. The patient received chemotherapy with etoposide, farmitrubinc, oncovin and prezocon but three weeks later died due to postoperative complications.

Results

The direct smears from the ascitic fluid showed noncohesive large to very large lymphoid cells with...
abundant basophilic cytoplasm. The cells exhibited features that appeared to bridge large-cell immunoblastic and anaplastic large cell lymphoma. They contained large, pleomorphic, round to multilobated or kidney-shaped nuclei that often disclosed prominent nucleoli. A few cells showed eccentric nuclei, surrounded by a prominent clear perinuclear Golgi zone. Mitotic figures were abundant. Apoptotic neoplastic cells were noted. Histology revealed a diffuse infiltration of the fatty tissue by large lymphoid cells with abnormal nucleus, 1 to 2 prominent nucleoli and amphophilic cytoplasm. Mitotic figures were numerous. Immunohistochemically tumor cells were negative for cytokeratin (KL1, MNF116), and EMA. Most neoplastic lymphoid cells strongly positive for CD30, CD8, and CD3 (Figura 1), and showed weak staining for CD45, whereas they were negative for CD19, CD20, and CD79a. The syndecan (CD138) was also absent except on a few mature plasma cells. Tumor cells did not express CD2, CD5, TIA1, or ALK1. The PCR analysis of T-cell receptor γ chain gene rearrangement showed the presence of a predominant T-cell clone within an oligoclonal T-cell expansion. No clonal rearrangement of IgH chain gene was found. The search for EBV infection was negative both by immunohistochemistry and PCR analysis. No HHV-8 DNA sequences were detected. Patient’s serum did not contain anti-HHV-8 antibodies. So, the diagnosis of primary effusion lymphoma with T-cell immunophenotype was established.

Discussion

PEL selectively involves the serous body cavities, occurs predominantly in immunodeficient patients and is frequently infected by human herpesvirus type-8 and Epstein-Barr virus. Deep immunosuppression promotes the emergence of lymphoproliferative disorders in patients undergoing solid organ transplantation. As with other high-grade lymphomas, prognosis is very poor with median survival of only a few months. Death is usually due to lymphoma.

In the present case, phenotypic and genotypic findings disclosed the T-cell origin of lymphoma cells. Indeed, neoplastic cells were negative for CD19, CD20, CD79a, CD138, and did not exhibit clonal IgH rearrangement by PCR analysis. By contrast, they strongly expressed CD3, and a clonal rearrangement of T-cell receptor γ chain gene was found. Primary effusion lymphoma exhibiting a T-cell phenotype was reported only once, in an HIV-seropositive male patient (3). In the latter case, neoplastic lymphoid cells expressed various T-cell-specific antigens, including CD2, CD3, CD5, and CD7, and no B-cell markers, and both T-cell receptor and immunoglobulin gene rearrangements3. In our case, several features could be suggestive of ALCL. However, ALCLs commonly present with systemic disease, and isolated peritoneal effusion is uncommon. In addition, ALCLs are usually associated with a t(2;5)(p23;q35) translocation that leads to an abnormal expression of ALK, a feature that was not observed in our case4. Moreover TIA1, a cytotoxic marker frequently detected in ALCL was not present4.

With a presentation as a peritoneal effusion and pleomorphic lymphoma cells identified in the fluid, the main diagnoses are anaplastic large cell lymphoma (ALCL) and PEL. A distinction between them is important because of the much better prognosis of the former. Similarities between ALCL and PEL may include similar morphologic features, CD30 expression and apparently null-cell immunophenotype5.

Features that favor PEL include known HIV infection, CD138 and MUM1 expression, and positivity for HHV8 and EBV5. Demonstration of a T-cell phenotype, absence of HHV8 and EBV from the tumor cells and ALK expression would favor an ALCL5.
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