Concurrent Hairy Cell Leukemia and Chronic Lymphocytic Leukemia: Diagnostic and Therapeutic Implications

Sir,

Coexistence of two separate monoclonal hematological cancers poses challenges both from the diagnostic as well as therapeutic perspectives. We describe the concurrent hairy cell leukemia (HCL) and chronic lymphocytic leukemia (CLL) in a previously healthy person.

A 68-year-old male presented with bicytopenia and 7 cm splenomegaly. There were no B-symptoms, jaundice, or lymphadenopathy. Complete blood count showed a hemoglobin of 6 g/dl, total WBC count 8100/mm³, neutrophils 9%, lymphocytes 90%, and platelets 28000/mm³. Direct Coombs test was negative. Peripheral smear examination confirmed lymphocytosis with numerous small mature lymphocytes with clumped chromatin, many smudge cells and a few medium-sized lymphocytes with indistinct nucleoli, weakly basophilic cytoplasm, and irregular “hairy” margins [Figure 1a and b]. Immunophenotyping revealed lymphocytosis with 8% of cells strongly expressing CD25, CD11c, CD103, and CD123, characteristic of HCL. However, 70% of lymphocytes were positive for CD5, CD23, CD79b, SM1g and negative for FMC-7, typical of CLL [Figure 1c]. Trephine biopsy showed hypercellular marrow with lymphoid infiltration comprising of widely spaced mononuclear cells surrounded by clear spaces (“fried egg” appearance) consistent with HCL [Figure 1d].

In the absence of leukocytosis and diffuse marrow infiltration, his CLL was considered to be of Binet stage A and hence not warranting treatment. Bicytopenia and splenomegaly were attributed to HCL, which was treated with cladribine 0.14 mg/kg/day infusion over 2 h for 5 consecutive days. Three months later, the patient was asymptomatic with no splenomegaly, hemoglobin 12 g/dl, WBC 6800/mm³, neutrophils 19%, lymphocytes 74%, monocytes 2%, and platelets 202000/mm³. Peripheral smear [Figure 1e] and flow cytometry [Figure 1f] confirmed total disappearance of HCL clone and persistence of CLL. Patient continues to have no indication for treatment of CLL and will be monitored regularly.

CLL is the most common hematological cancer and can be found incidentally in conjunction with other clonal malignancies. Coexistence of CLL with mantle cell lymphoma and melanoma in a lymph node has previously been reported.[1] In a large retrospective review of 270 patients with CLL, Giné et al. had identified 3 cases of concurrent HCL.[2] Optimal evaluation of peripheral smear morphology can help to suspect such a dual diagnosis. The additional investigations such as flow cytometry, bone marrow biopsy, and immunoglobulin chain rearrangement studies should be used appropriately to confirm the same. In most such patients, CLL is not the cause for the clinical features, and hence, priority should be to treat the coexistent HCL.

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Conflicts of interest

There are no conflicts of interest.
Letters to Editor

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References


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