Primary extra-nodal indolent B-cell lymphoma (small lymphocytic lymphoma/chronic lymphocytic leukemia) of retroperitoneum

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ABSTRACT

Primary retroperitoneal malignant lymphoma is rare, and primary retroperitoneal indolent B-cell lymphoma has not been reported. A 64-year-old woman presented anuria. Imaging techniques showed a large retroperitoneal mass, and biopsies from it revealed non-Hodgkin indolent B-cell lymphoma (small lymphocytic lymphoma/chronic lymphocytic lymphoma). The tumor appeared retroperitoneal and extra-nodal in locations.

Key Words: Retroperitoneum, Non-Hodgkin lymphoma, Indolent B-cell neoplasm, Small lymphocytic lymphoma/chronic lymphocytic lymphoma

1. INTRODUCTION

Retroperitoneum is a soil of many mesenchymal tumors, but primary retroperitoneal tumors are rare. According to Chakrabarti et al.,[1] who examined 71 cases of retroperitoneal masses by fine needle aspiration, showed that out of the 71 cases, 48 cases (68%) were neoplasms which consisted of 35 malignant tumors and 13 benign ones. The 35 malignant lesions were as follows: pancreatic cancer (3 cases), adrenal carcinoma (3 cases), renal cell carcinoma (3 cases), pleomorphic sarcoma (3 cases), malignant peripheral nerve cell tumor (1 case), rhabdomyosarcoma (1 case), non-Hodgkin lymphoma (11 cases), metastatic adenocarcinoma (6 cases), seminoma (3 cases), Yolk sac tumor (1 case), and Wilms’ tumor (2 cases). All the 11 cases of non-Hodgkin lymphoma were nodal non-Hodgkin lymphomas; therefore extra-nodal malignant lymphoma primary in the retroperitoneum is very rare. A literature search could not find cases of extra-nodal indolent B-cell lymphoma primary in the retroperitoneum, although a case of non-Hodgkin lymphoma of undetermined histology in peritoneum successfully treated by chemotherapy was reported.[2] Also, several cases of diffuse large B-cell lymphoma have been reported.[3, 4] Non-Hodgkin lymphoma is now classified according to WHO classification in which lymphomas are roughly classified into B, T, NK, null, lymphoblastic, and unclassifiable lymphomas.[5–13] Primary indolent B-cell lymphoma is a category of B-cell lymphoma comprising of follicular lymphoma, mantle cell lymphoma, lympho-plasmacytic lymphoma, mucosa-associated lymphoid tissue (MALT) lymphoma, and small lymphocytic lymphoma/chronic lymphocytic lymphoma.[5–13]
2. Case Report

A 64-year-old woman presented anuria. The patient did not show evidence for Epstein-Barr (EB) virus infection. Imaging techniques showed a large retroperitoneal mass (see Figure 1). Since retroperitoneal uninvolved lymph nodes were seen, ML was unlikely. No other tumors were seen in the body. No lymphoadenopathy was seen. Needle biopsies of the retroperitoneal tumor were obtained and were subjected to histological and Immunohistochemical studies. Histologically, the biopsy showed a dense proliferation of small lymphoid cells with mild atypia (see Figure 2A-B). The small lymphoid cells showed monotonous proliferation, but very small number of immunoblastic cells were seen in some areas. Plasma cell differentiation was not prominent. No monocytoid cells were seen. Immunophenotypes of the tumor were as follows: cytokeratin (CK) AE1/3 -, CKCAM5.2 -, vimentin ++, CD45 +++, CD20 +++ (see Figure 2C), CD23 ++ (see Figure 2D). CD5 +, CD79a +++, CD138 +/-, bcl-6 -, light chain restriction + (λ > κ) (see Figure 2E), bcl-2 ++ (see Figure 2F), CD45RO -, CD3 -, CD56 -, cyclin D1 -, CD30 -, CD15 -, CD10 -, TdT -, p53 +/-, Ki67 + (see Figure 2G) (labeling index = 8%, KIT -, NSE -, synaptophysin -. EBER for EB-virus was not carried out. A pathological diagnosis of primary retroperitoneal non-Hodgkin lymphoma, indolent B-cell lymphoma, small lymphocytic lymphoma/chronic lymphocytic lymphoma was made. Post-biopsy soluble IL-2 receptor of blood was high (2,294 U/ml). The patient was treated by chemotherapy (R-CHOP and cisplatin) and radiation, and now the tumor was reduced in size with serum IL-2 receptor of 885 U/ml. The patient was discharged and is now followed up during the period of 12 weeks after the chemotherapy. The R-CHOP therapy was done in 6 courses, one course with R followed CHOP in a week. A total of 50 gray of local radiation was carried out with radiation of 2 gray in a day.

3. Discussion

The present retroperitoneal tumor is apparently a lymphoid tumor histologically and immunohistochemically. The histological features are those of indolent B-cell lymphoma. The immunohistology indicated that the tumor was composed of B cells. Positive light chain restriction shows monoclonality of tumor cells. Indolent B-cell lymphoma consists of the following five: small lymphocytic lymphoma/chronic lymphocytic lymphoma, MALT (marginal zone) lymphoma, lympho-plasmacytic lymphoma, follicular lymphoma, and mantle cell lymphoma. The histological and IHC features are those of small lymphocytic lymphoma/chronic lymphocytic lymphoma. The pattern of CD5, CD20, CD23, CD79a, CD138, bcl-2, cyclinD1, bcl-6 etc, is compatible with that of small lymphocytic lymphoma/chronic lymphocytic lymphoma. The low Ki67 labeling index shows low risk of the tumor.

Figure 1. Pelvic CT shows a large retroperitoneal tumor (arrows)
Figure 2. Histological (A,B) and immunohistochemical (C,D,E,F,G) features of the retroperitoneal tumor

A: Low power view shows intense proliferation of small lymphoid cells. HE, ×40. B: High power view shows monotonous proliferation of small lymphoid cells with mild hyperchromasia and without obvious nucleoli. HE, ×200. The tumor cells are positive for CD20 (C), CD23 (D), lambda light chain (E), bcl-2 (F), and Ki67 (G: labeling index 8%). D-G, ×200.
The present small lymphocytic lymphoma/chronic lymphocytic lymphoma appears primary in Ret, because imaging founded no other tumor and no lymphoadenopathy was noted. The present tumor seemed extra-nodal small lymphocytic lymphoma/chronic lymphocytic lymphoma because no lymphoadenopathy was found and non-involved lymph nodes were seen near the retroperitoneal tumor.

4. CONCLUSION
A very rare case of primary retroperitoneal extra-nodal indolent B-cell lymphoma (small lymphocytic lymphoma/chronic lymphocytic lymphoma) is reported.

CONFLICTS OF INTEREST DISCLOSURE
The author declares no conflict of interest.

REFERENCES


