A Unique Case of Sclerosing Mediastinal B Cell Lymphoma

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ABSTRACT

Primary mediastinal B cell lymphoma is a rare subtype of diffuse large B cell Lymphoma (DLBCL) arising from putative thymic B cell origin. It accounts for 2-4 % of NHL. The present case was a 55 years old male presented with sudden onset of cough. Chest x ray and CT thorax showed presence of mildly enhancing solid mass in anterior mediastinum. Fine needle aspiration cytology (FNAC) from the mediastinal mass showed presence of poorly differentiated malignant neoplasm. Trucut biopsy from the same showed presence of tumour with foci of hyalinization. On immunohistochemistry tumour cells expressed CD20, CD 30 and CD 23. Hence the diagnosis of NHL of mediastinal sclerosing B cell type was made.

The present case was one of the rare cases of Non Hodgkin Lymphoma (NHL) with an unusual presentation. Primary mediastinal B cell Lymphoma (PMBCL) with sclerosis is a distinct subtype of NHL with unique clinicopathological aspects and aggressive behavior. Prompt recognition and aggressive treatment may lead to relatively longer survival of patients.

Keywords: NHL, Mediastinum, Sclerosing

Introduction

Primary mediastinal B cell lymphoma is a rare subtype of DLBCL arising from putative thymic B cell origin.[1] It accounts for 2-4 % of NHL, occurring predominantly in young adults with female predominance.[2] It usually present as a localized mass in anterior mediastinum. Bone marrow and lymph node involvement (other than supraclavicular and cervical) is absent by definition to rule out systemic DLBCL with secondary mediastinal involvement.

Case Report

A 55 years old male presented with sudden onset of cough with frequent recurrences since 2 months. On investigations, his hemogram was normal. Chest x ray and CT thorax showed presence of mildly enhancing solid mass in anterior mediastinum of size 110x80x86 mm with presence of necrotic areas which suggested possibility of Lymphoma. There was presence of bilateral pleural effusion. His bone marrow biopsy and bone marrow aspirate showed normocellular marrow. FNAC from the mediastinal mass showed presence of poorly differentiated malignant neoplasm.

Trucut biopsy from the same showed presence of tumor with foci of hyalinization (Fig 1). Tumor cells were medium to large sized with abundant pale cytoplasm and round to ovoid nuclei. Periodic acid Schiff stain showed absence of glycogen. Reticulin stain showed presence of compartmentalising fibrosis (Fig 2a). Hence, a possibility of thymic carcinoma (clear cell type) was given.

On immunohistochemical studies by DAKO-Envision HRP detection system, tumour cells expressed CD20 (Fig 2b), CD 30 and CD 23 while immunonegative for CD3, CK and Vimentin. Hence, diagnosis of NHL of mediastinal sclerosing B cell type was made.

Patient was given treatment following protocol DA-R-EPOCH + G-CSF. It stands for dose adjusted R-EPOCH that is dose adjusted Rituximab- Etoposide, doxorubicin, cyclophosphamide, vincristine and prednisolone + Granulocytic colony stimulating factors (G-CSF). This therapy obviates the need for radiotherapy in these patients and resulted in event free increased survival rate.[1] The present case responded to treatment and then was discharged with continuous follow up.

Fig. 1-H/E STAIN (40X) showing tumour with foci of hyalinisation.
Discussion

PMBCL was first described by Lichenstein et al in 1980. Thereafter many studies have been done regarding its pathogenesis and treatment and was included in WHO classification.

Primary mediastinal B cell lymphoma is a rare subtype of DLBCL arising from putative thymic B cell origin. It accounts for 2-4% of NHL, occurring predominantly in young adults with female predominance. It usually presents as a localized mass in anterior mediastinum. Bone marrow and lymph node involvement (other than supraclavicular and cervical) is absent by definition to rule out systemic DLBCL with secondary mediastinal involvement.

It has a fairly unique clinicopathological presentation. Present case was an adult male which is very unusual presentation. According to Barth et al, elderly patients are more affected by DLBCL as compared to PMBCL which shows involvement of young individuals in 3rd or 4th decade. It is bulky and frequently involves adjacent structures and can involve supraclavicular and cervical lymph node. Bone marrow infiltration is very rare at the time of presentation, as seen in the present case. A study done by Shemmari et al also showed that only 10% of patients had bone marrow involvement.

PMBCL show a wide range of morphology and cytological features depending on individual case. They are usually associated with compartmentalising alveolar fibrosis. The present case also showed sudden onset of cough due to local effects of tumour.

In immunohistochemical studies, tumour cells show positivity for B cell antigens (CD 19, 20, 22, 79a) but as a rule lack Ig. CD 30, CD 15, MUM -1 and CD 23 are also frequently positive. Recent studies show that most cases of PMBCL also express BOB 1, MAL, CD 54, CD 95 and TRAF 1.

In cytogenetic studies, they show gain in chromosome 9p24 and 2p15. It has unique transcriptional signature, NF kb and JAK-STAT signaling pathways. The discordant expression of B cell receptor (CD 79a +, sig -) is characteristic of PMBCL.

PMBCL has an equivalent or slightly better prognosis than typical DLBCL. PMBCL should be differentiated from Hodgkin's lymphoma. PMBCL are CD 30 positive but are usually weak and heterogenous compared to HL. Hoeller et al suggest that expression of BOB 1 favours PMBCL and expression of Cyclin E favours CHL.

Some cases show overlap in immunomarkers expression between PMBCL and HL and are considered to be in grey zone. CD15 is not typically identified in PMBCL and if present should raise the possibility of an overlap diagnosis.

Another important differential diagnosis includes thymic carcinoma in which epithelial component is highlighted by cytokeratin. It should also be differentiated from germinoma as clear cells of PMBCL mimic the germinoma cells. However, absence of glycogen and PAS negativity favours the diagnosis of Lymphoma.

Response to chemotherapy with or without radiotherapy is usually good. Extension into adjacent structures, pleural effusion and male sex has poor prognostic implication as seen in our case. On the other hand, presence of sclerosis denotes good prognosis.
DA-R-EPOCH treatment protocol is usually given. Some studies show use of MACOP-B (Methotrexate + leucovorin rescue, doxorubicin, cyclophosphamide, vincristine, prednisone and bleomycin) chemotherapy in these patients with favourable outcome.[13]

**Conclusion**
The present case is one of the rare cases of NHL with an unusual presentation. PMBCL with sclerosis is a distinct subtype of NHL with unique clinic pathological aspects and aggressive behavior. Prompt recognition and aggressive treatment may lead to relatively longer survival of patients.

**References**
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