Non-Hodgkin lymphomas – T cell type: Histomorphology and immunohistochemistry

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Abstract
Introduction: Non-Hodgkin lymphomas (NHL) comprise a wide variety of neoplasms and are subdivided into two main categories of low grade and high grade lymphomas. The major subtypes among T cell lymphomas are Peripheral T cell lymphomas – Not otherwise specified, Anaplastic Large cell lymphoma, ALK positive, Angioimmunoblastic T-cell lymphoma, T lymphoblastic leukemia/lymphoma and Extranodal NK/T cell lymphoma, nasal type. In this study, the histomorphological diagnostic features and the adjunct immunohistochemical expression of various markers are studied in a tertiary care centre in South India.

Materials and Methods: Patients with lymphadenopathy over a period of two years were included in the study. Lymph nodes excised from these patients were routinely processed and examined. After initial examination with H&E stain, appropriate special stains like Reticulin stain, Periodic Acid Schiff stain and Ziehl-Neelsen stains are performed along with the specific Immunohistochemical markers.

Results: A total of 30 cases were classified as Non-Hodgkin Lymphoma, high grade, T cell type. Peripheral T cell lymphomas-NOS is found to be the most common tumors accounting for 40% of all high grade T cell NHLs. The next common tumor was Anaplastic Large Cell Lymphoma, ALK positive with 36% of cases. T lymphoblastic leukemia/lymphoma accounted for 10% of cases and the rest of the cases were reported as Angioimmunoblastic T-cell lymphoma (7%) and Extranodal NK/T cell lymphoma, nasal type (7%).

Conclusion: Appropriate use of immunohistochemistry aids in the sub-classification of lymphomas according to the recommendations of WHO 2008. But, further molecular studies should be performed for confirmation of the subtyping done by immunohistochemistry.

Keywords: High grade lymphoma, Anaplastic large cell lymphoma, Peripheral T cell lymphoma.
chromogen Diamino benzidine (DAB) and the results were interpreted as either positive or negative and specified as cytoplasmic, membranous or nuclear staining in the tumor cells. After a morphological diagnosis was made, the following panel of markers were used – CD 3, CD4, CD 8, CD 20, CD 10, CD 15, CD 30, TdT, EMA, ALK protein, EBV LMP, BCL 2 and Ki67 (MIB 1) for the differential diagnosis and sub-categorisation of lymphomas.

Results and Observations

Epidemiology of T cell Lymphomas: The total number of lymph node biopsies received in the Department of Pathology during the study period was 378. The various diagnoses given in these 378 lymph nodes are as shown in the Chart 1.

Out of the total 140 cases diagnosed as Non-Hodgkin Lymphoma, 72 cases were diagnosed as high grade (51.4%). This study includes 59 cases from the 72 cases and the 13 cases excluded from the study are due to the unavailability of the tissues for the application of immunohistochemical markers. Among these 59 cases, 30 cases were subclassified as high grade T cell Non Hodgkin Lymphomas (51%). The subclassification of T cell lymphomas is given in Chart 2.

Similarly ALCL’s Fig. 2 are subclassified into two distinct entities with clinical and prognostic significance as ALK protein Positive which accounts for 23% of the total T lymphomas and Negative accounting for 13% of the total.

The majority of the T cell high grade lymphomas were either Peripheral T cell lymphomas-Not otherwise specified (PTCL-NOS) (40%) or anaplastic large cell lymphomas (ALCL) (36%) among all the T cell lymphomas observed in the study.

Among the PTCLs two common categories were observed namely the PTCL-NOS Fig. 1 accounted for 23% of the total T cell lymphomas and the PTCL with Lennert’s morphology accounting for 17% of the total T cell lymphomas.

The other types of lymphomas seen in this study are the T lymphoblastic leukemia/lymphoma accounting for 10% and the rest by angioimmunoblastic T-cell lymphoma Fig. 3 (7%) and Extranodal NK/T-cell lymphoma, nasal type. Fig. 4 (7%).
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Fig. 3: High power view of Angio-immunoblastic variant of PTCL

Fig. 4: High power view of NK/T cell lymphoma, Nasal Type; Arrow – highlights blood vessel wall infiltrated with atypical lymphoid cells

Age & Gender Distribution: The male to female ratio in T high grade NHL’s is 1.6:1 with a slight male preponderance. The age range was from 2-71 with a median age of 45 years. The various age incidences of the lymphomas are shown in the Chart 3.

Chart 3: The median age of the various subsets of high Grade T NHL

Duration of Symptoms and Tumor Bulk: The high grade T cell lymphomas presented with a mean duration of symptoms for 3.8 months, with a range of the shortest of 10 days to two years. The average size of the mass/lymphnodes at presentation was 2.6 x 2 cm.

Staging of Lymphomas: Clinical staging is done for these lymphomas according to the Ann-Arbor Staging system.1 and it is found that B symptoms are less commonly seen in Non-Hodgkin lymphomas with only few cases presenting with fever and weight loss being the most common symptoms observed. The stages of presentation of the different T cell high grade NHLs are shown in Charts 4&5.

Chart 4: Distribution of PTCL in different stages

Chart 5: Distribution of ALCL in different stages

The other subtypes of T cell lymphomas presented predominantly in the earlier stages. Out of the three T lymphoblastic leukemia/lymphomas, two of them presented in Stage I disease and one presented in Stage II.

The two Extranodal NK/T-cell lymphoma, nasal type presented in Stage I disease.

Out of the 2 Angioimmunoblastic T-cell lymphomas, one of them presented with Stage I disease and the other presented with bonemarrow involvement (Stage IV).

Immunohistochemical Profile of T Cell NHL’S: The immunohistochemical panel used for the subcategorisation of the T cell lymphomas and their percentage expression is as shown in Table 1
CD 3, the lineage marker of the T cells is expressed in all (100%) of the PTCLs, AITLs, T-ALLs and Extralodal NK/T cell lymphoma, nasal type. In ALCls this marker expression is seen only in five out of the 11 cases (45%) thus categorising them as ALCls- T cell phenotype. The rest six cases did not express this marker in the tumor population and only background reactive population showed positivity, thus subcategorising them as ALCL- Null cell phenotype.

CD 30 expression is seen in all the 11 cases of ALC (100%) and also found expressed in five cases of the PTCL (42%) thus categorising the PTCLs into a separate subtype based on immunohistochemical expression as the CD 30 positive PTCLs.

EMA is positive in all cases of ALC (100%) and not in other subtypes of T cell lymphomas.

CD 15 is positive in only one of the ALC cases (9%) and was confirmed with the ALK positivity and hence ruling out Hodgkin lymphoma.

ALK protein expression is seen in seven cases of the ALCls thus defining two distinct subcategories among this group into the ALK positive and ALK negative subtypes. ALK expression is not seen in the other subtypes of the high grade NHLs.

CD 10 is found to be positive in all cases of AITLs (100%) but not in other subtypes. EBV-LMP is expressed in one of the two angioimmunoblastic cases (50%). This marker is expressed in both cases of Extralodal NK/T-cell lymphoma, nasal type (100%).

The other markers used are natural killer cell markers, CD 16 and CD 56 which were negative in both the cases of the Extralodal NK/T-cell lymphoma, nasal type.

### Discussion

NHLs arising from the mature T cells constitute the most heterogeneous group of neoplasms namely the Peripheral T cell lymphomas. The recent WHO 2008 classification has three distinct entities in this category namely the Peripheral T-Cell Lymphoma-Not Otherwise Specified (PTCL-NOS), Angioimmunoblastic T cell lymphoma (AITL) and Anaplastic large cell lymphoma (ALCL).

This group represents 10-15% of the NHLs worldwide with slightly higher incidence in Asian countries. These lymphomas generally pursue an aggressive clinical course with both nodal and extra-nodal manifestations and adverse outcome.

PTCL-NOS is a diagnosis of exclusion. The median age of presentation of this group is in the sixth or seventh decade with 65% of the patients having stage IV disease and a poor five year survival rate of 20-30.

Several morphological variants have been described in PTCL-NOS. Variants described in the recent WHO 2008 classification are PTCL-lymphoepithelioid variant (Lennert’s lymphoma), the interfollicular variant and the T zone variants.

ALCL is a rare lymphoma worldwide with an incidence of <5% but represents the most common large cell lymphoma occurring in the pediatric population accounting for 10%-15% of the pediatric/adolescent NHL’s compared to the 2% incidence in adults. Usually, ALCL presents with advanced stage of III/IV with frequent bone marrow and extra-nodal involvement, skin and bone being the most common sites of involvement of systemic ALCL.

Jeffrey et al in their study on ALCL have found several morphological variants in ALCL, with the classical entity (70%) characterized by the neoplastic “hallmark” cells and a background polymorphous infiltrate of lymphocytes, plasma cells and eosinophils. The other variants described are the lympho-histiocytic (5-10%), small cell (5-10%) and sarcomatoid variant (<1%). Another morphological variant included in the WHO 2008 is the Hodgkin like ALCL.

In their study, they have also expressed the significance of IHC in ALK positive ALCL (50-70% of ALCL), in that it correlates with the molecular signature of the t (2,5) by having a specific Golgi region of distribution of ALK protein in these cases.

Summers et al in their study of the specific variant small cell ALCL have found that, this variant has a distinctly aggressive course compared to the other variants with prominent lymphadenopathy, leukemic presentation and disseminated disease with frequent skin involvement.

Jhala et al in their study have reported a rare morphologic entity of neutrophil rich ALCL in two HIV positive patients, concluded that this entity should
be considered in neutrophil rich specimens with otherwise the typical morphology of ALCL and absence of ALK expression.\(^\text{29}\)

Jhala et al in their study on 66 patients of ALK positive and ALK negative ALCLs, have found that BCL2 expression has a significant difference and hence prognosis in these two groups with BCL2 being more frequently expressed in the ALK negative group (57\%) compared to the complete absence of its expression in ALK positive ALCL(0\%).\(^\text{30}\)

Several prognostic parameters characterize this lymphoma with the expression or the non-expression of the ALK protein uniquely characterized by the t (2,5) with fusion of NPM –ALK genes, playing the most important role. Five year failure free survival studies have shown a better prognosis for those with expression of ALK protein (ALK + ALCL), younger age and a worse prognosis for the ALK – ALCL and those expressing the epithelial membrane Antigen (EMA).\(^\text{31} \)\(^\text{32}\)

Berge et al evaluated the role of Epithelial Membrane Antigen (EMA) in 79 cases of ALCLs, HLLs, DLBCLs and PTCLs and have concluded that there is no statistical significant difference in the expression of EMA in these tumors. But this marker is seen consistently to be expressed in ALK positive ALCLs compared to the low or nil expression in ALK negative ALCLs and HLLs.\(^\text{33}\)

The distinction of ALK positive and negative lymphomas with that of the Peripheral T cell lymphoma (PTCL) has been made possible with the appropriate use of IHC markers. Savage et al in their study on 181 patients collected from 22 institutions across the world have emphasized the need for the distinction of ALK negative ALCL from the PTCL – NOS type, since they have found that immunohistochemically as well as in survival studies, the ALK negative group have significant differences.\(^\text{34}\) These express the CD30 antigen strongly and have better five year survival rates (36\%) which were superior compared to the PTCL group (20\%).

In more recent studies, it has been proposed that the diagnosis of ALCL should be revised when EBV markers are positive. But in recent case reports, they have opined that EBV could be expressed in ALCLs and that FISH detection of other EBV markers than EBER could be helpful in actual detection of the viral status in these groups of lymphomas.

Yu zeng and Andrew L Feldman in their study on the genetics of ALCL have studied the role of ALK protein mutations.\(^\text{35}\) They have observed that recurrent chromosomal rearrangements with the ALK protein and its partners namely NPM1 as in t (2,5) contribute to the pathogenesis of ALCL. The other partner genes identified are TPM3, TPM4 and many others. This NPM-ALK fusion protein activates many signal transduction pathways like the JAK/STAT, mTOR, PI3K/AKT pathway which are important signaling pathways in cell proliferation and survival.

They have also observed that ALK negative tumors consistently express two recurrent genetic abnormalities, one involving the DUSP22 (an inhibitor of T cell receptor signaling) and the other being TP53 mutations.

**Conclusion**

Appropriate use of immunohistochemistry aids in the sub-classification of lymphomas according to the recommendations of WHO 2008. But, further molecular studies should be performed for confirmation of the subtyping done by immunohistochemistry.

**References**


