

Lymphoma: Neoplastic proliferation of lymphoid cells





B-Cells in the bone marrow









Normal Lymph Node

- Normal compartments of the lymph node:
- 1- cortex / follicles
- 2- paracortex
- 3- medullary cords
- 4- sinuses





Diagnostic Methods

- Gold standard method: Histomorphologic evaluation of the lymphoid tissues.
- Sections of fixed tissue -embedded in paraffin- for routine H&E staining





Diagnostic Methods

Ancillary studies:

- Immunophenotyping:

- 1- Immunohistochemistry (IHC) and/or
- 2- Flow cytometry

Lymphoid Markers

Lymphoid markers commonly used in immunophenotypic studies:

Pan-hematopoietic: CD45, CD43 B-cell markers: CD19, CD20, CD22, CD79a, Pax5, MUM1 T-cell markers: CD2, CD3, CD5, CD7,CD4, CD8 Germinal center cells: CD10, Bcl-6, HGAL Follicular dendritic cells: CD21, CD23, CD35 Plasma cells: CD138, CD38, MUM1 Immature lymphoid cells: TdT, CD10, CD34, CD1a Activated lymphoid cells: CD30, CD23 NK-cells: CD56, CD57

Diagnostic Methods

• Flow cytometry (<u>multi-parametric</u>) analysis is the method of choice to demonstrate the <u>intensity</u> of antigen expression and simultaneous expression of multiple antigens..

Cell suspensions and flow cytometer













Diagnostic Methods, cont.

Ancillary studies:

- Cytogenetic studies: Conventional Karyotype analysis, FISH analysis, SKY analysis
- Molecular analysis: Southern blot and PCR









DNA microarray platforms: to demonstrate gene expression patterns/profiles; important methodology for the diagnosis and classification of hematological malignancies in the future





LYMPHOID DISORDERS

- 1- Benign (reactive) Lymphadenopathies
- 2- Malignant Lymphoproliferative Disorders/Lymphomas
- 3- Atypical (borderline/grey zone) Lymphoproliferative Disorders

LYMPHOMAS

- Non-Hodgkin Lymphoma (NHL):
 1- B-cell lymphomas (most common)
 2- T-cell and NK cell lymphomas
- Hodgkin Lymphoma (disease):
 - 1- Nodular Lymphocyte Predominance
 - 2- Classical Hodgkin Lymphoma

• **Definition:**

Malignant neoplastic proliferation of lymphoid cells derived from a single transformed cell (monoclonal proliferation).

Non-Hodgkin Lymphoma

- Epidemiology:
- More common in the developed countries
- Frequency of different types varies around the world:

B- vs NK/T-cell..

Burkitt lymphoma..

adult T-cell leukemia/lymphoma..

- Epidemiology, cont.:
- Individuals with immunodeficiencies/immune disorders are at higher risk :
- 1- Congenital (primary) immunodeficiency
- 2- Acquired immunodeficiencies (HIV-related, post-therapy)
- 3- Autoimmune diseases..

Non-Hodgkin Lymphoma

- Etiology:
- Infectious agents and development of lymphoma:
- HTLV-1 infection and T-cell leukemia/lymphoma..
- Epstein-Barr virus (EBV) and Burkitt's lymphoma...
- Human herpesvirus-8 (HHV-8) and primary effusion lymphoma
- HCV and marginal zone lymphoma
- H. Pylori and gastric MALT lymphoma

- Etiology, cont.:
- Genetic alterations and development of lymphoma..
 - t(14;18) translocation and follicular lymphoma..
 - t(8;14) translocation and Burkitt's lymphoma..
 - t(11;14) translocation and mantle cell lymphoma..



- Classification of NHLs:
- * Rappaport Classification:
- Based on morphology; growth pattern (diffuse vs. nodular) and cytomorphology..
- * Working Formulation (1984):
- Based on morphology; divides NHLs into prognostic groups (useful to clinicians) and can be applied in the absence of immunophenotypic studies..

- Classification of NHLs, cont.:
- * Revised European American Lymphoma (REAL) Classification (1994):
- This classification system recognizes specific (real) disease entities based on immunophenotypic, cytogenetic, and molecular features, as well as conventional morphology.

Non-Hodgkin Lymphoma

WHO Classification (2001):

- An updated version of the REAL classification..
- Expanded the principles of the REAL classification to the classification of the myeloid and histiocytic neoplasms..
- Replaced existing classifications..
- The first international consensus..

WHO Classification

 Precursor (immature) B- and T-cell Lymphoid Neoplasms (leukemia/lymphoma):

Precursor lymphblastic leukemia/lymphoma

 Mature (peripheral) B- and T-cell Lymphoid Neoplasms:

Mature B-cell lymphomas Mature NK/T-cell lymphomas



WHO Classification

- Immunodeficiency Associated Lymphoproliferative Disorders:
- 1. Lymphoproliferative diseases associated with primary immune disorders
- 2. Lymphomas associated with infection by HIV
- 3. Post-transplant lymphoproliferative disorders
- 4. Methotraxate-associated lymphoproliferative disorders

Non-Hodgkin Lymphoma

- Clinical manifestations of NHL:
- 1- Lymphadenopathy..
- 2- Organomegaly (infiltration of organs)..
- **3- Systemic/B symptoms** (fever, night sweats, weight loss)..

 Examples of NHL:
 Diffuse Large B-cell Lymphoma (DLBCL), most common lymphoma in the western hemisphere
 Follicular Lymphoma (FL)
 Burkitt Lymphoma (BL)

Non-Hodgkin Lymphoma

Diffuse Large B-cell Lymphoma A diffuse proliferation of large neoplastic B lymphoid cells with a nuclear size equal to or exceeding normal macrophage nuclei or more than twice the size of a normal lymphocyte





- Recent studies using gene expression profiling (GEP) in DLBCL have identified patterns of gene expression, as well as individual genes that appear to have important prognostic significance, related to underlying tumor biology.
- At least two types:
- 1- DLBCL of germinal center cell origin CD10+ and/or BCl-6+
- 2- DLBCL activated B-cell like CD10- and MUM1+

Follicular Lymphoma

A neoplasm of follicle center B cells (a mixture of centrocytes/centroblasts), which has at least a partially follicular/nodular pattern of growth











Burkitt Lymphoma

A highly aggressive lymphoma often presenting at an extra-nodal site or in a leukemic phase, composed of monomorphic medium-sized B-cells with basophilic/vacuolated cytoplasm and numerous mitotic figures









- Therapy of NHL:
- **1- Chemotherapy**
- 2- Radiotherapy
- 3- Combined chemo- and radiotherapy
- 4- Immune therapy: antibodies raised against B-cell antigens (anti- CD20/anti-CD22 antibodies)

Hodgkin Lymphoma

• **Definition:**

Lymphoid neoplasm with a minority of neoplastic lymphoid cells (*Hodgkin cells*) in a predominant inflammatory/reactive background which may mask the tumor cell population

- Epidemiology & Etiology :
- accounts approximately for 30% of all malignant lymphomas
- bimodal age distribution
- EBV has been postulated to play a role in the pathogenesis of classical HL

Hodgkin Lymphoma

- WHO Classification of HL:
- 1- Nodular Lymphocyte Predominance
- 2- Classical Hodgkin Lymphoma

• Nodular Lymphocyte Predominance:

- uncommon variant (6% of HL)
- characteristic nodular pattern of growth
- characteristic variant of Hodgkin cells (known as "popcorn" cells), which are CD20 positive B-cells of follicle center cell origin (CD10+, Bcl-6+)





Classical Hodgkin Lymphoma:

Nodular sclerosis (NSHL) Mixed cellularity (MCHL) Lymphocyte depletion (LDHL) Lymphocyte-rich classical Hodgkin lymphoma (LRCHL)

Classical Hodgkin Lymphoma:

Hodgkin cells: derived from mature B-cells at the germinal center stage of differentiation

- typical Reed-Sternberg cells (binucleated cells)
- variants (mono- and multinucleated forms and lacunar cells)
 Immunophenotype: CD30+, CD15+, Pax5+,

and CD45 negative

Hodgkin Lymphoma

Classical Hodgkin Lymphoma: Nodular sclerosis (NSHL):

 characterized by Reed-Sternberg cells and *lacunar cells* within a polymorphic reactive background, and a characteristic *fibrosis* (bands of collagen surrounding nodules)













Classical Hodgkin Lymphoma: <u>Mixed cellularity (MCHL):</u>

 classical Reed-sternberg cells in a mixed, polymorphic bachground; no lacunar cells or collagen bands

Lymphocyte depletion (LDHL):

- uncommon subtype
- rich in Hodgkin's cells

Hodgkin Lymphoma

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- Therapy of HL:
- 1- Chemotherapy
- 2- Radiotherapy