

# Lymphoma and Myeloma

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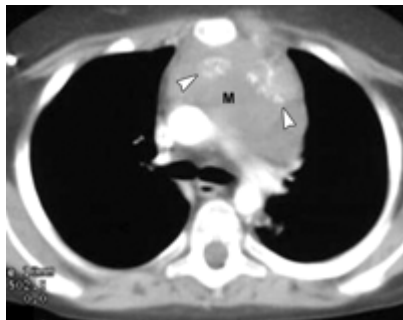
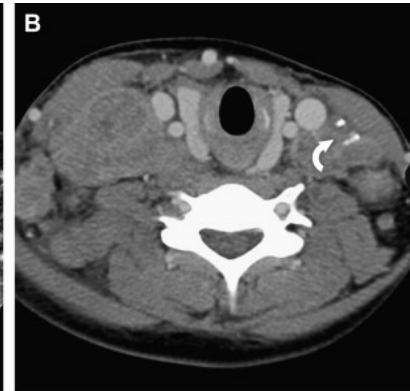
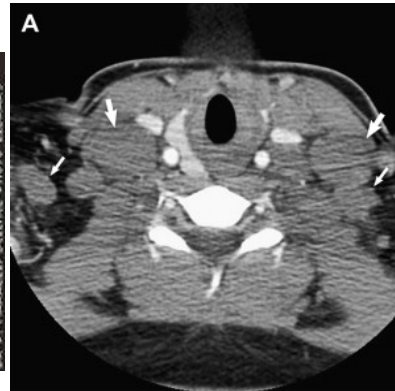


# NHL/ HD(Lymphoma): common features

- 1- Lymph node enlargement: painless, decides stage
- 2- May be associated with B-symptoms: fever, night sweats, weight loss
- 3- Compression symptoms may occur
- 4- Extra-nodal involvement
- 5- Needs LN BX for diagnosis
- 6- Each has different histology types
- 7- Both have similar staging system

# Case 11: NHL

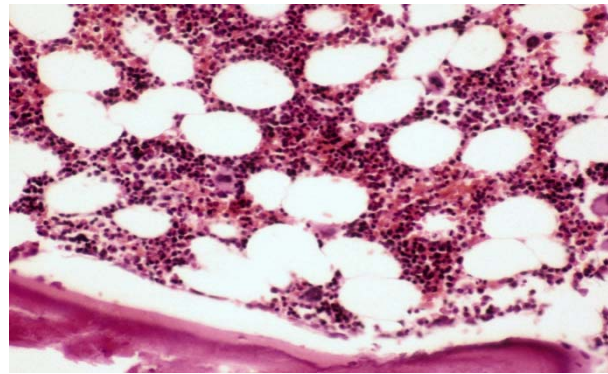
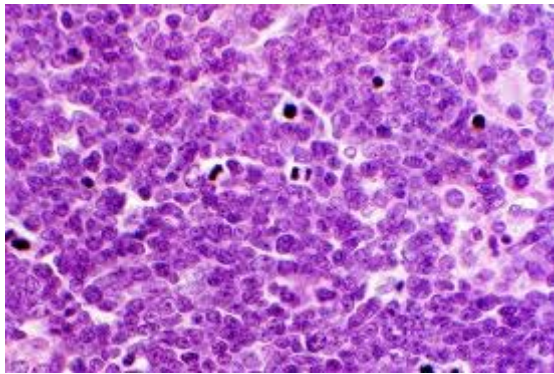
27 yr old male, presented with weight loss, low grade fever and profuse sweating for the last 6 wks. P/E, he had generalized lymphadenopathy, hepato-splenomegaly and enlarged tonsils.



## Case 11

Hb 10g/dl, MCV 100, WBC 14k, Normal diff, Plt 196k, LDH 820, S.uric .a 7.5, Creat, Ca, PO4 NI,

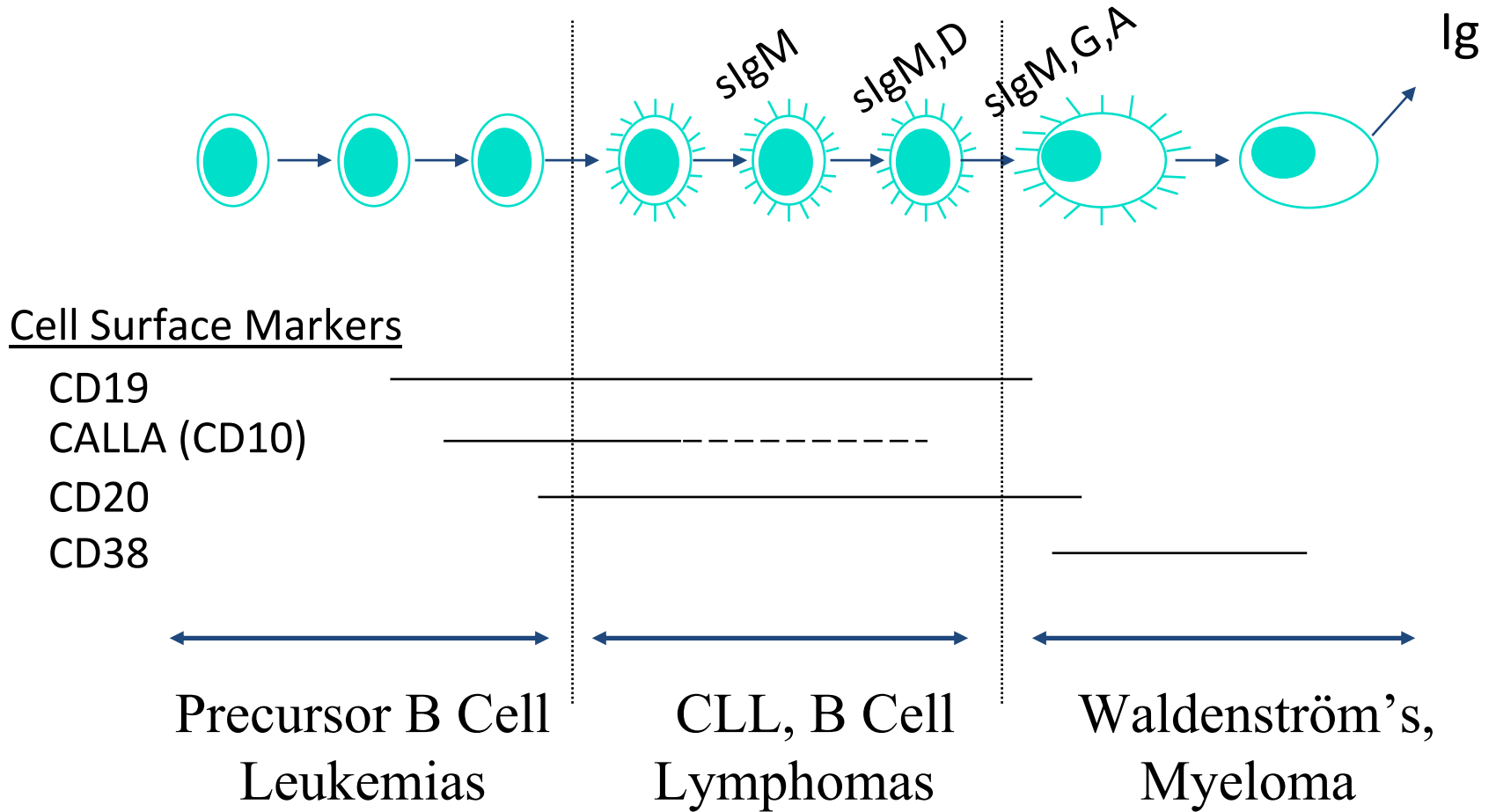
LN.Bx



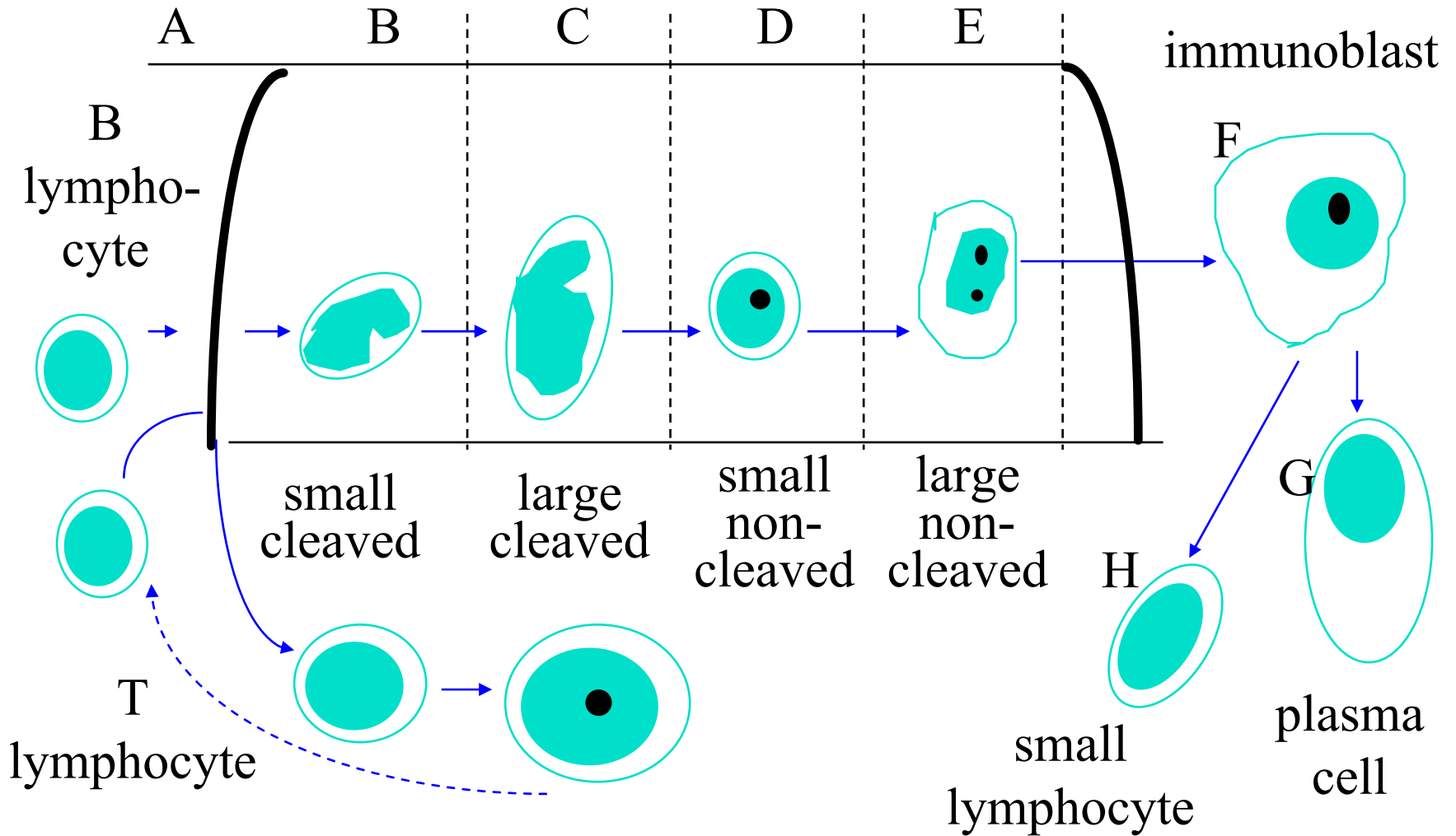
# NHL

- The types of non- Hodgkin's lymphoma reflect the developmental stages of lymphocytes.
- Each type of lymphoma can be viewed as a lymphocyte arrested at a certain stage of development and transformed into a malignant cell.
- 85% B cell origin, the rest T or null cell.

# B CELL DIFFERENTIATION



# MATURATION IN LYMPHOID FOLLICLE



# Etiology of NHL

- Idiopathic
- Immune suppression
  - congenital (Wiskott-Aldrich)
  - organ transplant (cyclosporine)
  - AIDS
  - increasing age
- DNA repair defects
  - ataxia telangiectasia
  - xeroderma pigmentosum



# Etiology of NHL

- Chronic inflammation and antigenic stimulation
  - *Helicobacter pylori* inflammation, stomach
  - *Chlamydia psittaci* inflammation, ocular adnexal tissues
  - Sjögren's syndrome
- Viral causes
  - EBV and Burkitt's lymphoma
  - HTLV-I and T cell leukemia-lymphoma
  - HTLV-V and cutaneous T cell lymphoma
  - Hepatitis C

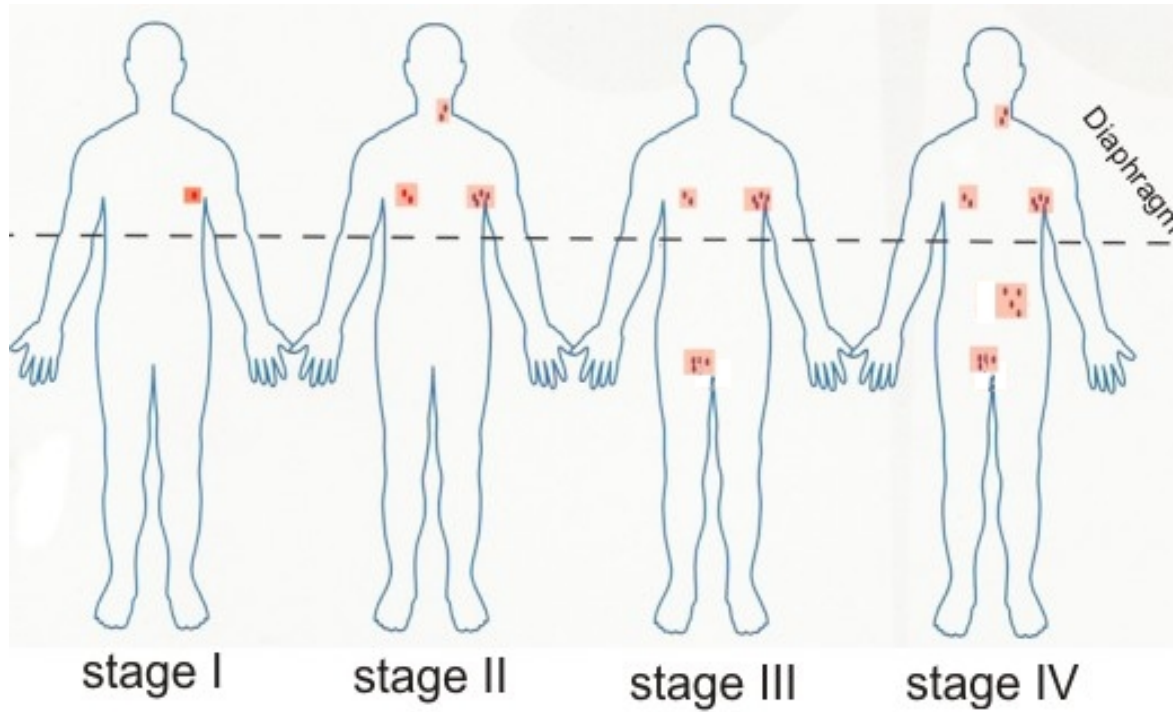
# Diagnosis of NHL

- Chromosome changes
  - 14;18 translocation in follicular lymphoma
    - *bcl-2* oncogene
  - t(8;14), t(2;8), t(8;22) in Burkitt's lymphoma
    - *c-myc* oncogene
  - t(11;14) in mantle cell lymphoma
    - *cyclin D1* gene

# Staging: Ann Arbor

- I. 1 lymph node region or structure
- II. >1 lymph node region or structure, same side of diaphragm
- III. Both sides of diaphragm
- IV. Extranodal sites diffuse, beyond “E” designation

## Staging System for Lymphomas (ANN ARBOR CLASSIFICATION)



# Revised European-American Lymphoma (REAL) Classification: B-Cell Neoplasms

Indolent	Aggressive	Very Aggressive
<ul style="list-style-type: none"> <li>• CLL/SLL</li> <li>• Lymphoplasmacytic/IMC/WM</li> <li>• HCL</li> <li>• Splenic marginal zone lymphoma</li> <li>• Marginal zone lymphoma               <ul style="list-style-type: none"> <li>– Extranodal (MALT)</li> <li>– Nodal</li> </ul> </li> <li>• Follicle center lymphoma, follicular, grade I-II</li> </ul>	<ul style="list-style-type: none"> <li>• PLL</li> <li>• Plasmacytoma/Multiple myeloma</li> <li>• MCL</li> <li>• Follicle center lymphoma, follicular, grade III</li> <li>• DLCL</li> <li>• Primary mediastinal large B-cell lymphoma</li> <li>• High-grade B-cell lymphoma/Burkitt's-like</li> </ul>	<ul style="list-style-type: none"> <li>• Precursor B-lymphoblastic lymphoma/Leukemia</li> <li>• Burkitt's lymphoma/B-cell acute leukemia</li> <li>• Plasma cell leukemia</li> </ul>

## ***Prognostic factors in non-Hodgkin's lymphoma***

- ***Adverse factors:***
- ***Age > 60 years***
- ***Stage III or IV, i.e. advanced disease***
- ***High serum lactate dehydrogenase level***
- ***Performance status (ECOG 2 or more)***
- ***More than one extranodal site involved***

# Treatment Options in Advanced Indolent Lymphomas

- *Observation only.*
- *Radiotherapy to site of problem.*
- *Systemic chemotherapy*
  - *oral agents: chlorambucil and prednisone*
  - *IV agents: CHOP, COP-R, FC-R*
- *Antibody against CD20: rituximab*
- *Stem cell or bone marrow transplant.*
- *New monoclonal antibodies*

# Reasons to Treat in Advanced Indolent Lymphomas

- Constitutional symptoms
- Anatomic obstruction
- Organ dysfunction
- Cosmetic considerations
- Painful lymph nodes
- Cytopenias



## Treatment Options: Aggressive Lymphomas

### ***Aggressive***

- Diffuse large cell lymphoma, large cell anaplastic lymphoma, peripheral T cell lymphoma.

### ***Very Aggressive***

- Burkitt's lymphoma and lymphoblastic lymphoma.

# Treatment Options for Aggressive Lymphomas

- \*potentially curable

- \*disseminates through bloodstream early

- \*must use systemic chemotherapy

  - ❖CHOP-R x ?8 cycles

  - ❖CHOP-R x 3 cycles followed by radiotherapy

- \*Bone marrow transplantation for some cases

- \*New monoclonal antibodies

# Standard Treatment for Aggressive Lymphomas

- Systemic chemotherapy

  - CHOP-R

- ± Intrathecal chemotherapy

  - AIDS patients and CNS involvement

- ± Radiotherapy

  - Spinal cord compression, bulky disease

  - ??BMT

## Burkitt's Lymphoma

- Treated with multidrug regimen similar to pediatric leukemia/lymphoma regimens.
- BMT

# **Hodgkin's Disease/Lymphoma Treatment**

**With appropriate treatment about 85% of patients with Hodgkin's disease are curable**

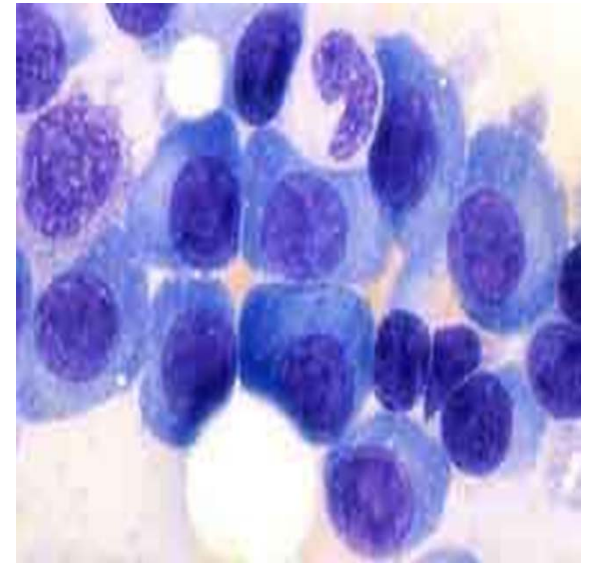
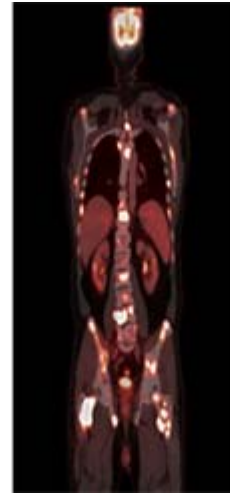
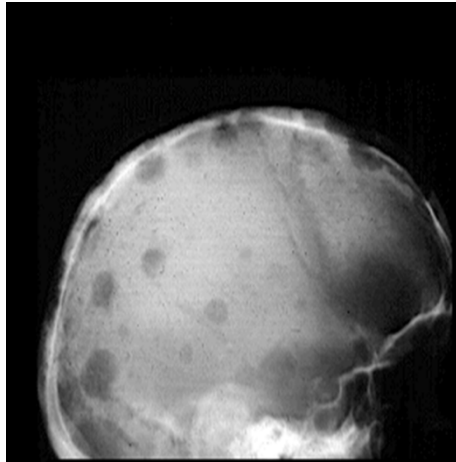
- I A,B                                      chemo.Radiation  
Therapy in selected cases**
- II A                                        Chemo.XRT in selected  
cases**
- IIB; IIIA,B; IVA,B                  Chemo  
(+/- radiotherapy in selected cases)**

# Chemotherapy Regimens for Hodgkins disease

- MOPP
  - **M**echlorethamine, **O**ncovin, **P**rocarbazine, **P**rednisone
- ABVD
  - **A**driamycin, **B**leomycin, **V**inblastine, **D**acarbazine
- BEACOPP
- BMT for relapse or resistance

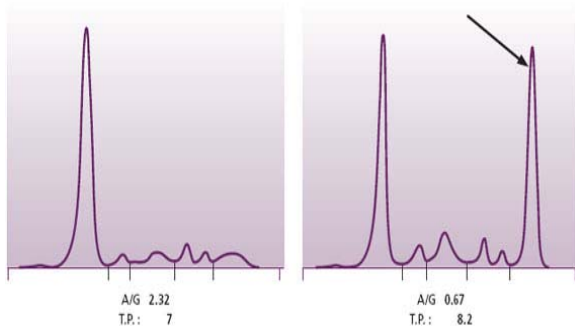
## Case 11 B: Multiple Myeloma

67 yr old male has been complaining of back pain for several months. He recently noticed exertional dyspnea. He was admitted because of severe pain in his arm.



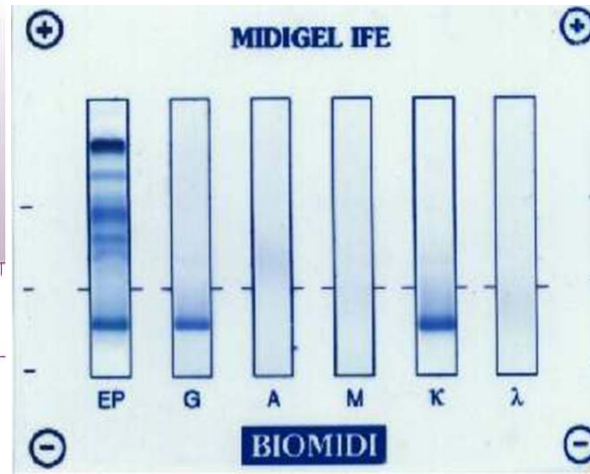
# Case 11 B Investigation and Diagnosis

- 1- Hb 8 g/dl, normocytic normochromic, WBC 8K, Plt 180K, ESR 112mm/1<sup>st</sup> hr. Bld film. B2-microglobulin 6mg/l, serum albumin 3 g/dl, P/E 1gG 15 g/dl and IF/ igGK
- 2- Multiple myeloma igG/k stage III (C).



Fractions	%	Ref. %	g/dl	Ref. g/dl
Albumin	69.9	52.9-66.9	4.9	3.7-4.9
Alpha	3.5	3.0-5.8	0.2	0.2-0.4
Alpha 2	8.8	7.5-13.4	0.6	0.5-1.0
Beta	9.3	8.5-13.7	0.7	0.6-1.0
Gamma	8.5	8.8-19.2	0.6	0.6-1.4

Fractions	%	Ref. %	g/dl	Ref. g/dl
Albumin	40.2	52.9-66.9	3.3 L	3.7-4.9
Alpha	4.8	3.0-5.8	0.4	0.2-0.4
Alpha 2	11.0	7.5-13.4	0.9	0.5-0.9
Beta	7.8	8.5-13.7	0.6	0.6-1.0
Gamma	36.2	8.8-19.2	3.0 H	0.6-1.4





# Clinical Features

- Symptoms related to BM infiltration: bone pain, osteolytic lesions and fractures, anemia, hypercalcemia
- Secretion of abnormal proteins: renal and neurological or visceral manifestations
- Hyperviscosity syndrome
- Recurrent infections
- Amyloidosis

## Related organ or tissue impairment

B<sub>uy</sub> – Lytic bone lesions – visible on x-ray in **85%** of patients. **Hint** – Osteoclasts activated, not osteoblasts.

C – Hypercalcemia (Ca > 11 mg/dL)

A – Anemia (Hb < 10)

V – Hyperviscosity – especially common in the rare  
IgM secreting myeloma

I – Bacterial infections (>2)

A – Amyloidosis

R – Renal (Crt > 1.96 mg/dL): **HINT** – occurs 50% of the time because most often the light chains are toxic to the tubules.

These are the end organ manifestations of myeloma

# Clinical features

- common tetrad of multiple myeloma is ***CRAB***
  - **C = Calcium (elevated)**
  - **R = Renal failure**
  - **A = Anemia**
  - **B = Bone lesions**

## **Active Multiple Myeloma**

Both criteria must be met:

- 1- Clonal bone marrow plasma cells 10% or biopsy-proven bony or extramedullary plasmacytoma
- 2- Any one or more of the CRAB features

## **Smouldering MM**

Both criteria must be met:

Serum monoclonal protein (IgG or IgA) 3 g/dL, or urinary monoclonal protein 500 mg per 24 h and/or clonal bone marrow plasma cells 10%–60%

Absence of myeloma defining events or amyloidosis Mu

# Diagnosis and Staging Workup

- Bone marrow biopsy and aspirate
- Serum protein electrophoresis and immunofixation
- Skeletal survey/ MRI/PET Scan
  - Plain x-rays are better than bone scan.
  - Lytic lesions do not show up well on bone scan.
- Quantitative immunoglobulins

# Prognostic Factors

## **International staging system**

I (good prognosis)

Serum albumin > 3.5 g/dl

Serum  $\beta$ 2 microglobulin < 3.5 mg/dl

II

Not I or III

III

$\beta$ 2 microglobulin: >5.5 mg/dl

# Categories with Potential Prognostic Significance

<b>Factor</b>	<b>Abnormality</b>	<b>Median Survival</b>
	plasmablastic morphology	5 – 23 mos
<b>Surface Markers</b>	CD38+/CD45- Peripheral blood	>10 cells/uL 37 mos
<b>Kinetics</b>	S phase 1- 3% S phase > 3%	22 mos 12 mos
<b>Conventional cytogenetics</b>	Deletion 13	15 mos
<b>FISH</b>	t(4;14)	29 mos

Gertz, Morie A. , (2007) 'Relevant prognostic features of multiple myeloma and the new International Staging System', Leukemia and Lymphoma, 48:3, 458 - 468

# Treatment of Multiple Myeloma

## Standard Chemotherapy

- Dexamethasone and Thalidomide
- Dexamethasone and Bortezomib (Velcade) / Lenalidomide
- Melphalan and prednisone
- High Dose Chemotherapy  
with Bone marrow transplant (auto)