



PATHOLOGY AIIMS PYQ

Medsynapse by Dr. Nikita



The phenomenon where subsequent generations are at the risk of earlier and more severe disease is known as:

future anticipate.

- a. Mosaicism
 - b. Imprinting
 - c. Pleiotropy
 - d. **Anticipation**
- silencing → Methylation mutes
- future
- Trinucleotide repeat → Huntington's



FAST

Genetic terms

TERM	DEFINITION	EXAMPLE
Codominance *	Both alleles contribute to the phenotype of the heterozygote.	Blood groups A, B, AB; α_1 -antitrypsin deficiency; HLA groups. - AB
Variable expressivity <i>in 2 diff phs</i>	Patients with the same genotype have varying phenotypes.	Two patients with neurofibromatosis type 1 (NF1) may have varying disease severity. *
Incomplete penetrance <i>multiple</i>	Not all individuals with a disease show the disease. % penetrance \times probability of inheriting genotype = risk of expressing phenotype.	BRCA1 gene mutations do not always result in breast or ovarian cancer.
Pleiotropy	One gene contributes to multiple phenotypic effects.	Untreated phenylketonuria (PKU) manifests with light skin, intellectual disability, musty body odor.
Anticipation Q	Increased severity or earlier onset of disease in succeeding generations.	Trinucleotide repeat diseases (eg, Huntington disease).
Loss of heterozygosity <i>above sibling</i>	If a patient inherits or develops a mutation in a tumor suppressor gene, the wild type allele must be deleted/mutated/eliminated before cancer develops. This is not true of oncogenes.	Retinoblastoma and the "two-hit hypothesis," Lynch syndrome (HNPCC), Li-Fraumeni syndrome.
Epistasis Q	The allele of one gene affects the phenotypic expression of alleles in another gene.	Albinism, alopecia.
Aneuploidy	An abnormal number of chromosomes; due to chromosomal nondisjunction during mitosis or meiosis.	Down syndrome, Turner syndrome, oncogenesis.

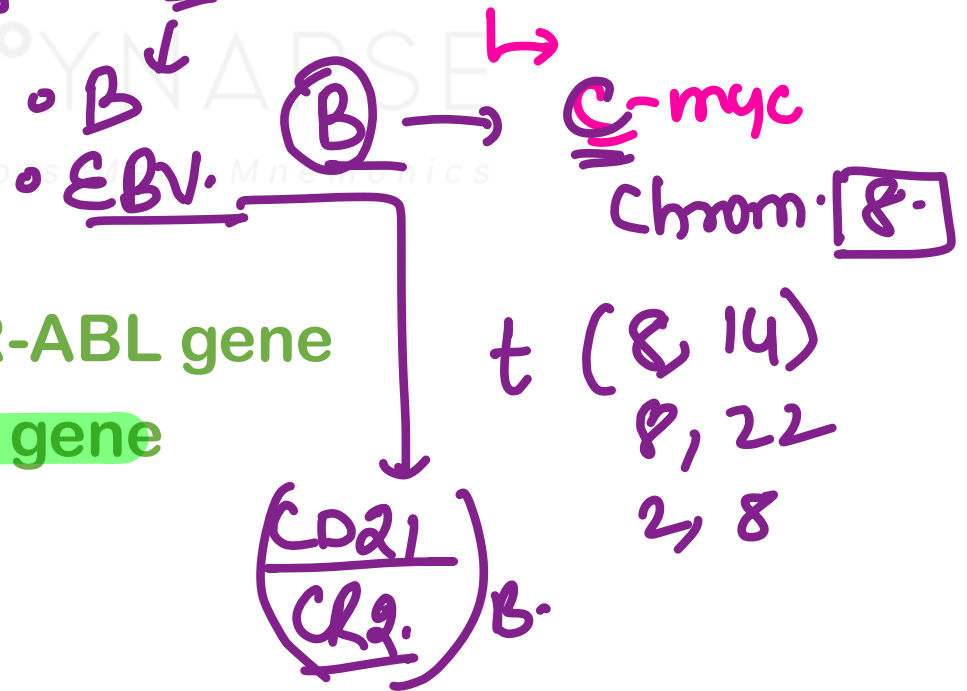
- * Pleiotropy
- * Epistasis
- * variable exp
- * Incomplete penetrance



A 10-year old boy was presented with a mass in abdomen. On imaging, the para-aortic lymph nodes were enlarged. On biopsy, **starry sky appearance** was seen. What is the **underlying abnormality**?

Africa → jaw swelling → Burk Burk stars

- a) p53 gene mutation
- b) Rb gene mutation
- c) Translocation involving BCR-ABL gene
- d) Translocation involving **Myc gene**





After an incised wound, new collagen fibrils are seen along with a thick layer of growing epithelium. The approximate age of the wound is:

a) 4–5 days

b) About 1 week

c) 12–24 hours

d) 24–72 hours



MEDSYNAPSE
Where Concepts Meet Mnemonics

after 3 days



FAST

PHASE OF WOUND HEALING	EFFECTOR CELLS	CHARACTERISTICS
① Inflammatory (up to <u>3 days</u> after wound)	Platelets, neutrophils, macrophages	Clot formation, ↑ vessel permeability and <u>neutrophil</u> migration into tissue; <u>macrophages</u> clear debris 2 days later
② Proliferative (day <u>3</u> –weeks after wound)	Fibroblasts, myofibroblasts, endothelial cells, keratinocytes, macrophages	Deposition of <u>granulation tissue</u> and type <u>III collagen</u> , <u>angiogenesis</u> , <u>epithelial cell proliferation</u> , <u>dissolution of clot</u> , and <u>wound contraction</u> (mediated by myofibroblasts) Delayed second phase of wound healing in <u>vitamin C</u> and <u>copper</u> deficiency
③ Remodeling (1 week–6+ months after wound)	Fibroblasts	Type III collagen replaced by <u>type I collagen</u> , ↑ <u>tensile strength</u> of tissue Collagenases (require <u>zinc</u> to function) break down type III collagen <u>Zinc deficiency</u> → <u>delayed wound healing</u>

Tree-Thr

clot

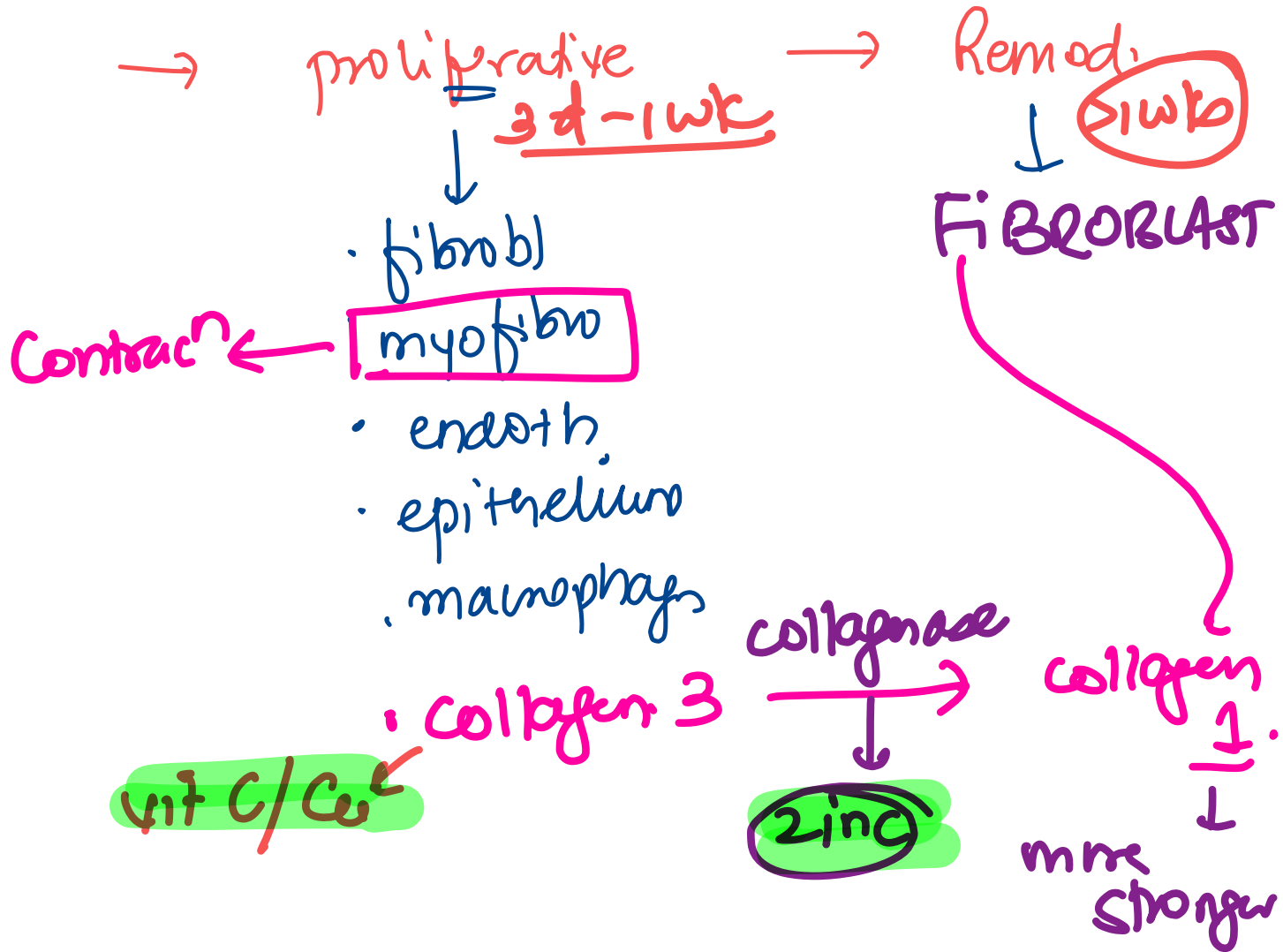
epithelium contraction

fibrosis

collagen: 80%

first - fibroblasts fibrosis

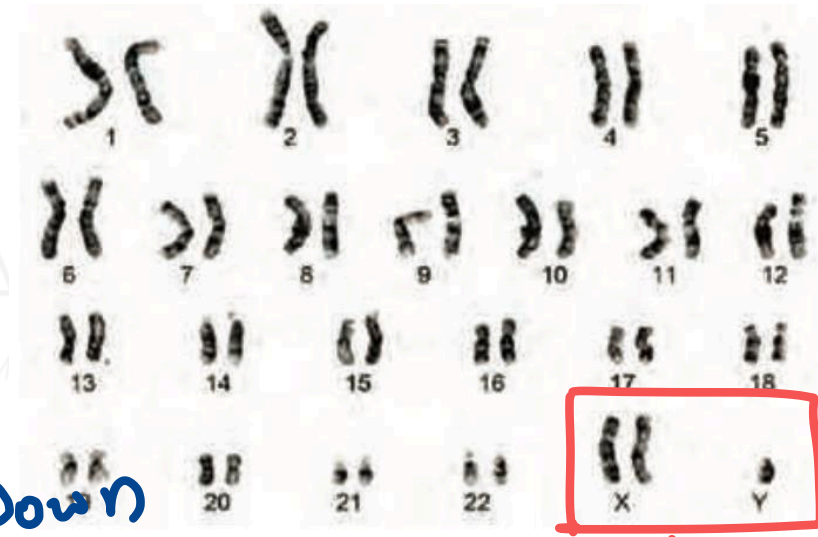
Inflammatory up to 2d.
↓
neutrophils
macrophages.



vit C/Cu²⁺



Which of the following abnormality is seen in the given karyotype?



a) High pitched cry → Cri du chat

b) Round face with protruding tongue

c) Short stature with webbed neck

↳ Down
↳ Turner

d) Gynecomastia with long thin limbs

↳ XXY
Klinefelter

↑ breast CA

Klinefelter → ↑ FSH ↑ LH
Hypogonadotropic



Klinefelter syndrome

Male, 47,XXY.

Small, firm testes; infertility (azoospermia); tall stature with eunuchoid proportions (delayed epiphyseal closure → ↑ long bone length);

gynecomastia; female hair distribution.

May present with developmental delay.

Presence of inactivated X chromosome (Barr body). Common cause of hypogonadism seen in infertility workup

↑ risk of breast cancer.

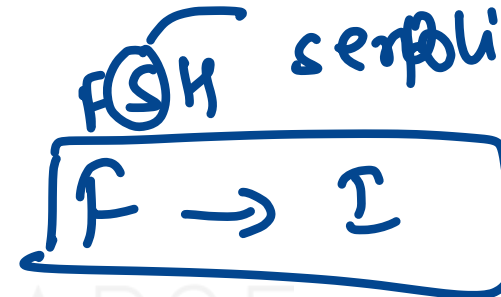
XXY

Dysgenesis of seminiferous tubules

→ ↓ inhibin B → ↑ FSH.

Abnormal Leydig cell function → ↓ testosterone

→ ↑ LH.



LH → Leydig



MEDSYNAPSE
Where Concepts Meet Mnemonics



Inheritance of ABO blood group is:

INI 23

° AB

° myotonic dystrophic

↓
AD

chrom. 19.

° CTG

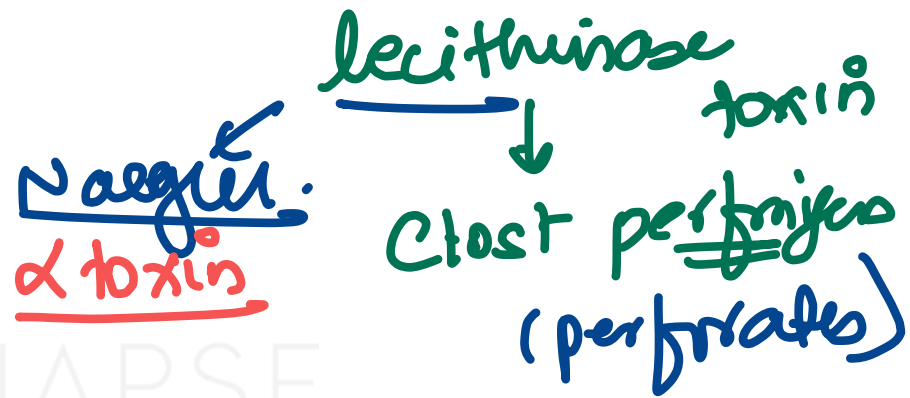


- a) X-linked inheritance
- b) Autosomal Recessive inheritance
- c) Mitochondrial inheritance

d) Codominance → eg. ABO, HLA · AB



Most common nephropathy associated with malignancy is:



- a. Focal segmental glomerulosclerosis (FSGS)
- b. Minimal change disease
- c. IgA nephropathy
- d. Membranous glomerulonephritis → hepC, malign
↳ phospholipase A₂ Ab ⊕

Membranous nephropathy

Also called membranous glomerulonephritis. Can be 1° (eg, antibodies to phospholipase A₂ receptor) or 2° to drugs (eg, NSAIDs, penicillamine, gold), infections (eg, HBV, HCV, syphilis), SLE, or solid tumors. ↑ risk of thromboembolism (eg, DVT, renal vein thrombosis).

Diffuse capillary and GBM thickening

Granular due to immune complex (IC) deposition

“Spike and dome” appearance of subepithelial deposits

↑ cellularity

lines → good pasture (Ab)

↑ WBC

PSCGN

HBV/HCV

→ MPCGN / membranous

Main videos - Patho-nephritic / nephrotic



Oil red 'O' stain is used for:

fat →

① oil red → frozen.
 ② Sudan Black → fixed

* (PTAH → muscle.
 ↳ ATP ↗

* luxol's fast blue
 ↓
 myelin (fat word)

- a) Glutaraldehyde fixed specimen
- b) Alcohol fixed specimen
- c) Formalin fixed specimen
- d) Frozen specimen ✓

* Verhoeff → elastic fibres.

* Alzheimer → Bielschowsky silver



Which of the following is responsible for adhesion of platelets to the vessel wall?

7- heven extrinsic

VWD

Main - Bleeding disorder

• ↑BT + ↑aPTT &

↓
↓vWF
plt adhesion

↓
↓fVIII
↓
intrinsic

8-12 except
10 → comm

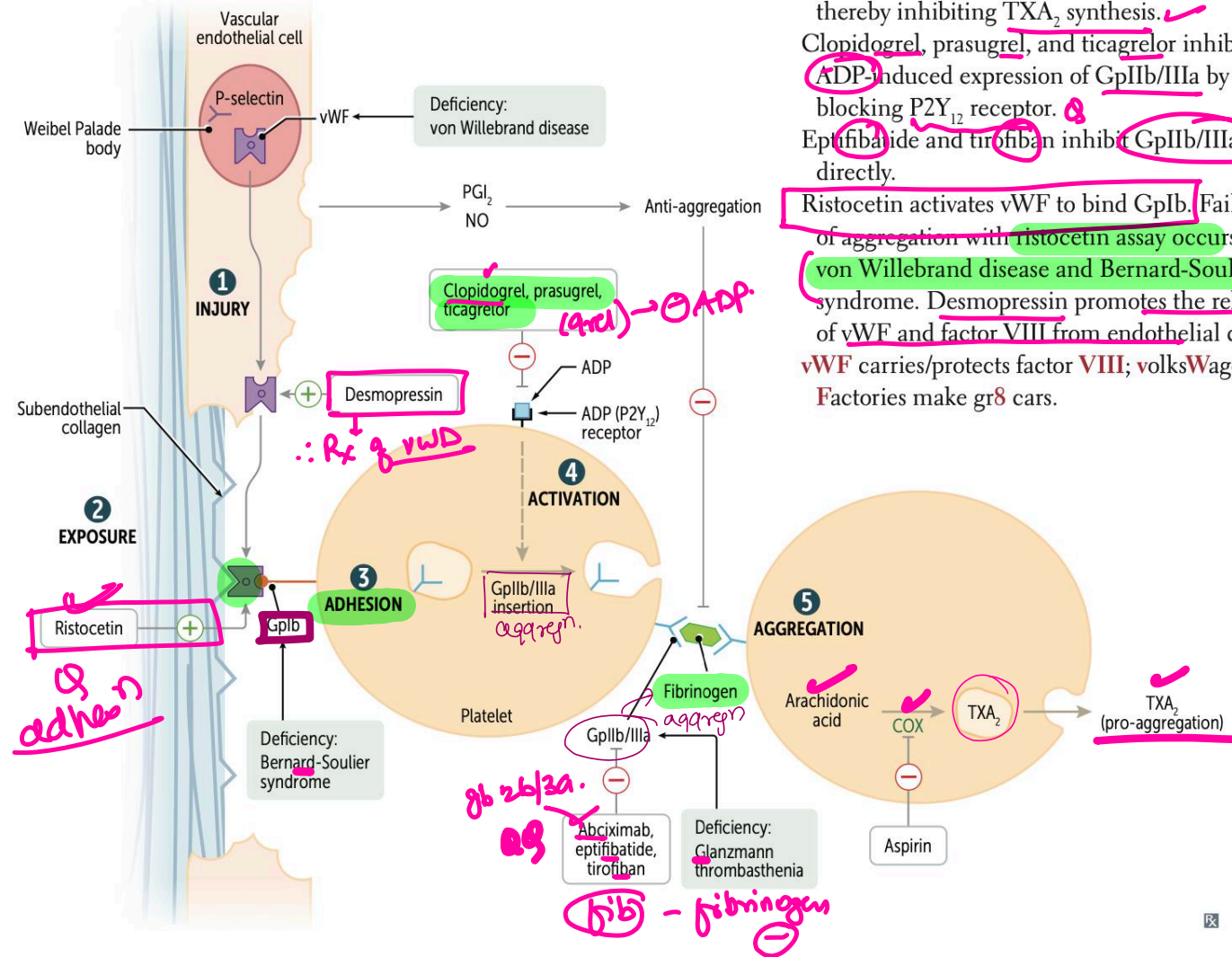
Happi

- a) Factor IX
- b) Von Willebrand factor
- c) Fibrinogen
- d) Fibronectin



MEDSYNAPSE
Where Concepts Meet Mnemonics

Thrombogenesis



Formation of insoluble fibrin mesh.
 Aspirin irreversibly inhibits cyclooxygenase, thereby inhibiting TXA₂ synthesis.
 Clopidogrel, prasugrel, and ticagrelor inhibit ADP-induced expression of GpIIb/IIIa by blocking P2Y₁₂ receptor.
 Eptifibatide and tirofiban inhibit GpIIb/IIIa directly.
 Ristocetin activates vWF to bind GpIb. Failure of aggregation with ristocetin assay occurs in von Willebrand disease and Bernard-Soulier syndrome. Desmopressin promotes the release of vWF and factor VIII from endothelial cells. vWF carries/protects factor VIII; volksWagen Factories make gr8 cars.

FAST



adhesion

∴ Rx of vWD

g6 26/30. ag

(fibrinogen)

- plt adh defen



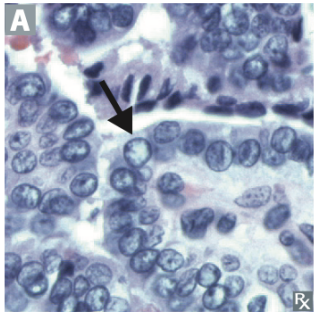
Lymphatic spread is most commonly seen in which type of thyroid malignancy:

→ papillary
• prognosis good
• radn

- a) Papillary carcinoma → lateral aberrant thyroid
- b) Follicular carcinoma → hematog
- c) Medullary carcinoma
- d) Anaplastic carcinoma



Papillary carcinoma



Most common. Empty-appearing nuclei with central clearing (“Orphan Annie” eyes) **A**, psamMoma bodies, nuclear grooves (Papi and Moma adopted Orphan Annie). ↑ risk with RET/PTC rearrangements and BRAF mutations, childhood irradiation. **08**
Papillary carcinoma: most prevalent palpable lymph nodes. Good prognosis. ✓

◦ Radⁿ ✓
◦ lymph nodes ✓
◦ Orphan annie ✓

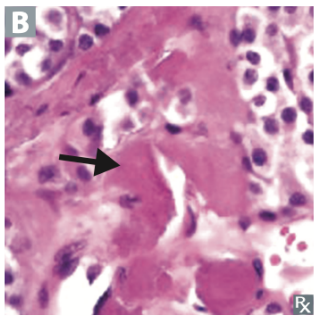
→ note on FNAC ✓

Follicular carcinoma



Good prognosis. Invades thyroid capsule and vasculature (unlike follicular adenoma), uniform follicles; hematogenous spread is common. Associated with RAS mutation and PAX8-PPAR-γ translocations. Fine needle aspiration cytology may not be able to distinguish between follicular adenoma and carcinoma. ✓

Medullary carcinoma



From parafollicular “C cells”; produces calcitonin, sheets of polygonal cells in an amyloid stroma **B** (stains with Congo red). Associated with MEN 2A and 2B (RET mutations). ✓



Undifferentiated/ anaplastic carcinoma

R53

Older patients; presents with rapidly enlarging neck mass → compressive symptoms (eg, dyspnea, dysphagia, hoarseness); very poor prognosis. Associated with TP53 mutation.



This image was taken by attaching the camera to the microscope. What is the requirement for such a microscope?

Fluorescence microscopy

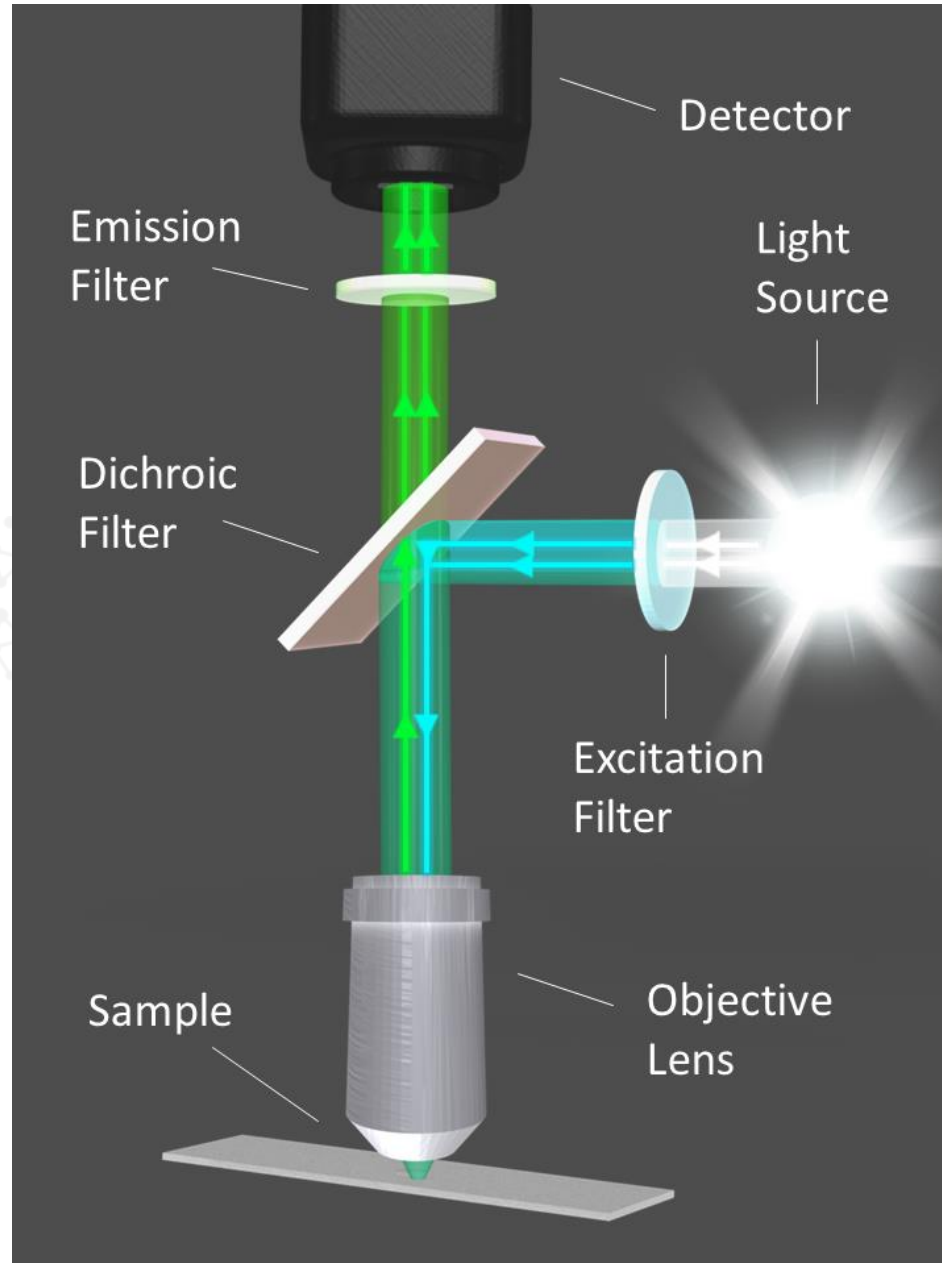
a) Dark field condenser

b) Phase shifter

c) Dichroic mirror

d) Cathode ray tube

→ reflects low wavelength
transmits high wavelength.





In **Langerhans Cell Histiocytosis**, the characteristic abnormality seen is:

N → ⊕ in skin → APC

• CD1a, S100
207.

- a) Foamy macrophages
- b) Giant cell
- c) Plasma cell
- d) Birbeck's granules**

Q



→ Birbeck

LCH

Tennis

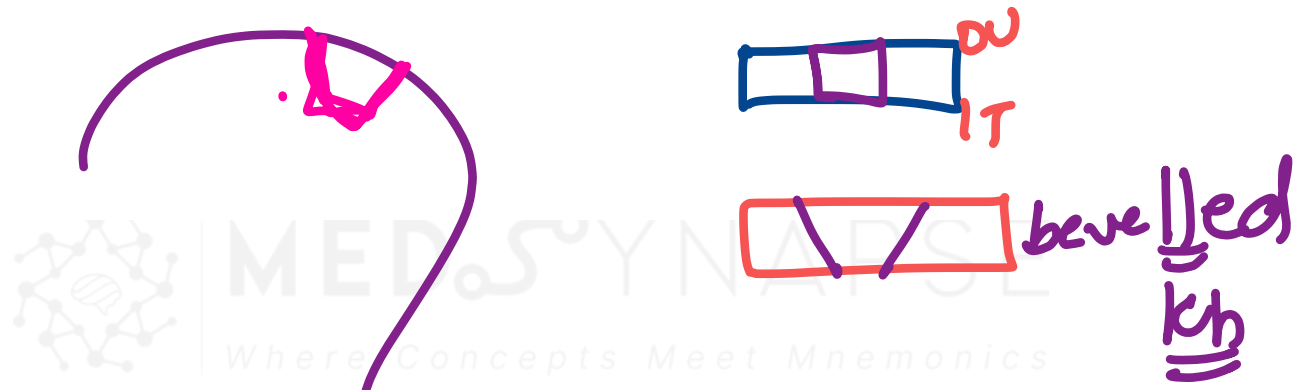
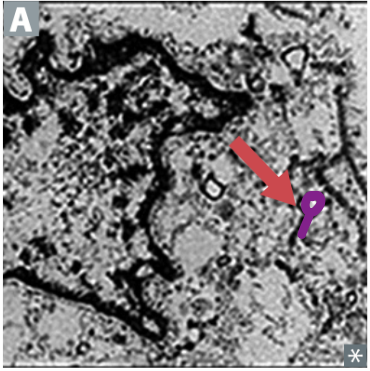
EM

EDSYNAPSE
Where Concepts Meet Mnemonics



Langerhans cell histiocytosis

Collective group of proliferative disorders of Langerhans cells (antigen-presenting cells normally found in the skin). Presents in a child as lytic bone lesions and skin rash or as recurrent otitis media with a mass involving the mastoid bone. Cells are functionally immature and do not effectively stimulate primary T cells via antigen presentation. Cells express S-100 and CD1a. Birbeck granules ("tennis rackets" or rod shaped on EM) are characteristic A.



Hand Schuller Christian dis →
(CDE)
o dI
o cavernial lytic
o exophthalmos



Type of multifocal, unisystem Langerhans cell histiocytosis, as demonstrated by HPE.

Triad of:

- Lytic bone lesions (often in skull)
- Exophthalmos
- Diabetes insipidus (due to pituitary stalk involvement)

[Seen in only one-third of patients]