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Pathology

CELL ADAPTATIONS

Cell adaptation

00:03:25

Cell organelles affected during cell injury :

- mitochondria.
- Endoplasmic reticulum.
- Nucleus.
- Plasma membrane.

Causes of cell injury :

- **Hypoxia** : Decreased Oxygen supply to a tissue.
most common cause of cell injury : **Hypoxia**
most common cause of hypoxia : **Ischemia**
Ischemia refers to decreased blood supply.
Ischemia is a **more severe** form of cell injury than hypoxia, as when blood supply decreases, supply of all nutrients including O_2 is diminished.
Cells most sensitive to hypoxia : **Neurons**.
Cells least sensitive to hypoxia : **Fibroblasts, skeletal muscle**.
- Physical agents e.g. , Radiation.
- Chemical agents e.g. , Carcinogens.
- Infectious agents. e.g. , Bacteria, viruses.
- Genetic abnormalities e.g. , Mutation.
- Immunologic agents.
- Nutritional imbalances : Deficiency (PEM), excess (obesity).

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Manifestation of cell injury

00:11:00

Cell injury may lead to :

- **Adaptation** : Hypertrophy, hyperplasia, metaplasia, atrophy.
- **Reversible cell injury** : On removing the injurious stimuli, the cell will revert to normal state.

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- Irreversible cell injury (cell death) : Necrosis, apoptosis, pyroptosis, ferroptosis, necroptosis.
- Intracellular accumulation.
- Pathologic calcification.

Hypertrophy

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Hypertrophy : Increased cell size but no increase in number of cells.

Mechanism : Occurs due to increase synthesis of cellular proteins.

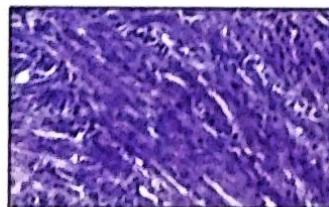
- GATA4
 - NFAT
 - MEFA
- } 3 proteins that are responsible for hypertrophy

Hypertrophy usually occurs in permanent/ non-dividing cells.

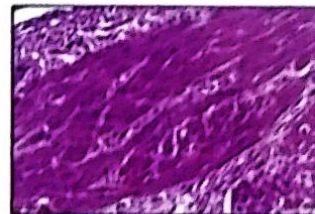
Type of cells based on division ability :

- Permanent/non-dividing cells : Cannot divide at all e.g., cardiac muscle, skeletal muscle.
- Stable cells : Liver or pancreas.
- Labile cells : Rapidly dividing cells e.g., bone marrow, skin epidermis.

Physiological hypertrophy	Pathological hypertrophy
<ul style="list-style-type: none"> • Uterus during pregnancy. • Breast during lactation. • Skeletal muscle in body builders. 	<ul style="list-style-type: none"> • Left ventricular hypertrophy. • In case of bladder outlet obstruction due to stone, area proximal to stone will undergo hypertrophy.



Normal smooth muscles cells of uterus



Hypertrophied smooth muscle cells

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Hyperplasia

00:24:19

Increase in the number of cells which will lead to increase in size of the organ.

mechanism: Growth factor induced proliferation of mature cells.

usually occurs in the **dividing cells**.

Dividing cells can undergo both hyperplasia and hypertrophy.

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Physiological hyperplasia		Pathological hyperplasia
Hormonal: <ul style="list-style-type: none"> Breast during pregnancy. Breast during puberty. 	Compensatory: Liver after partial hepatectomy.	Occurs due to hormonal excess: <ul style="list-style-type: none"> Increased androgen leads to benign prostatic hyperplasia. Increased estrogen leads to endometrial hyperplasia.

Hyperplastic proliferations are a fertile soil in which cancers can develop.

For e.g., endometrial hyperplasia can lead to endometrial cancer.

Examples of both hypertrophy and hyperplasia:

- Breast during puberty/pregnancy
- uterus during pregnancy.

Atrophy

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Atrophy: Decreased cell size and number leading to decreased organ size.

mechanism:

- Decreased protein synthesis.
- Increased protein degradation.
- Autophagy.

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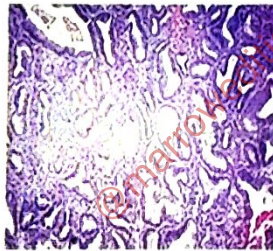
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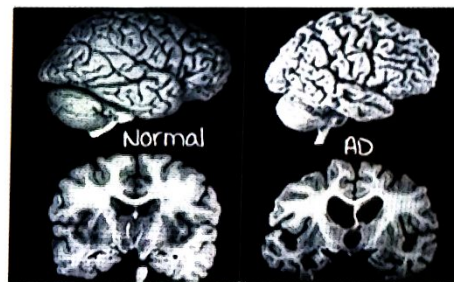
Physiologic atrophy	Pathologic atrophy
<ul style="list-style-type: none"> • The disappearance of notochord. • Disappearance of thyroglossal duct at puberty. • Involution of uterus after parturition. 	<ul style="list-style-type: none"> • Senile atrophy. • Ischemic atrophy (decreased blood supply leading to atrophy). • Denervation atrophy (loss of nerve supply). • Pressure atrophy (in case of a tumour, surrounding structures get atrophied due to pressure). • Disuse atrophy. • Nutritional atrophy.



→ Normal endometrial gland



→ Hyperplasia gland



Normal brain

Atrophied brain

Metaplasia

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It is a reversible change in which one **differentiated cell type/ mature cell type** is converted to another type.

epithelial metaplasia : Epithelium gets converted to another type of epithelium.

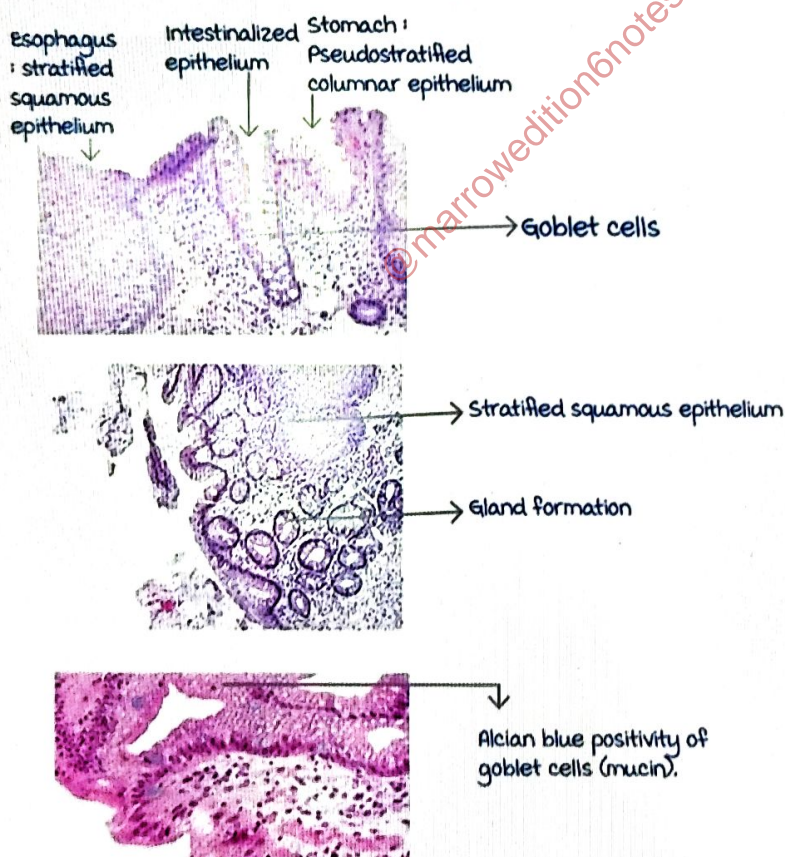
mesenchymal metaplasia : mesenchyme gets converted to another type of mesenchyme.

mechanism : Reprogramming of stem cells.

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Examples :

- **most common** metaplasia : In smokers pseudostratified ciliated columnar epithelium of respiratory tract, converts into stratified squamous epithelium (**squamous metaplasia**). Squamous metaplasia is reversible after cessation of smoking.
- **Barrett's esophagus/ Columnar lined oesophagus (CLO)** : Stratified squamous epithelium of esophagus is converted to columnar epithelium on exposure to GERD.
- On HPE of Barrett's esophagus : **Intestinal metaplasia** and **goblet cells** are seen.
- Special stain for Barrett's esophagus : **Alcian blue**.
- Goblet cells produce mucin which stains with Alcian blue.
- Barrett's esophagus is a risk factor for **adenocarcinoma of esophagus**.



Vitamin A deficiency can lead to metaplasia.

Example of connective tissue metaplasia : **myositis ossificans**.

MCQs

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Q. A 57 year old man comes to the physician for a follow up evaluation of chronic, retrosternal chest pain. The pain is worse at night and after heavy meals. He has taken oral pantoprazole for several months without any relief of his symptoms. Upper endoscopy shows ulcerations in the distal esophagus and a proximally dislocated Z-line. A biopsy of the distal esophagus shows mature columnar epithelium with goblet cells. Which of the following microscopic findings underlies the same patho mechanism as the cellular changes seen in this patient?

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- A. Pseudostratified columnar epithelium in bronchi.
- B. Squamous epithelium in bladder.
- C. Paneth cells in duodenum.
- D. Simple columnar epithelium in endocervix.

The given scenario is that of Barrett's esophagus which shows metaplasia.

Other options are examples of epithelium normally present in those tissues.

Q. Which of the following statements is false :

- A. Atrophy is a type of reversible cell injury.
- B. Plasma membrane blebbing occurs in reversible cell injury.
- C. On removal of stimulus, necrosis can be reversed.
- D. Chronic sublethal stimulus can cause cellular aging.

Necrosis is irreversible cell injury.

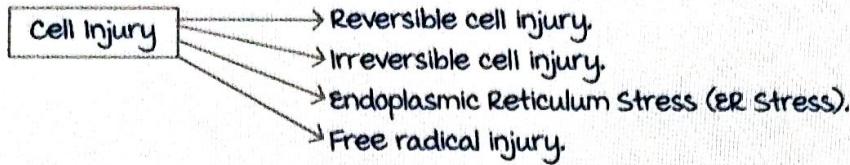
Q. A 20 year old woman had Goodpasture syndrome which progressed to chronic renal failure. She is 165 cm tall and weighs 55 kg. She now has blood pressure measurements in the range of 150/90 to 180/110 mmHg, but does not regularly take medications. Laboratory studies show her blood urea nitrogen is over 100 mg/dL and she requires chronic dialysis. A chest x-ray shows an enlarged heart. The size of her heart is most likely to be the result of which of the following processes involving the myocardial fibres?

- A. Hypertrophy.
- B. Fatty infiltration.
- C. Hyperplasia.
- D. Fatty degeneration.
- E. Edema.

CELL INJURY

Cell injury

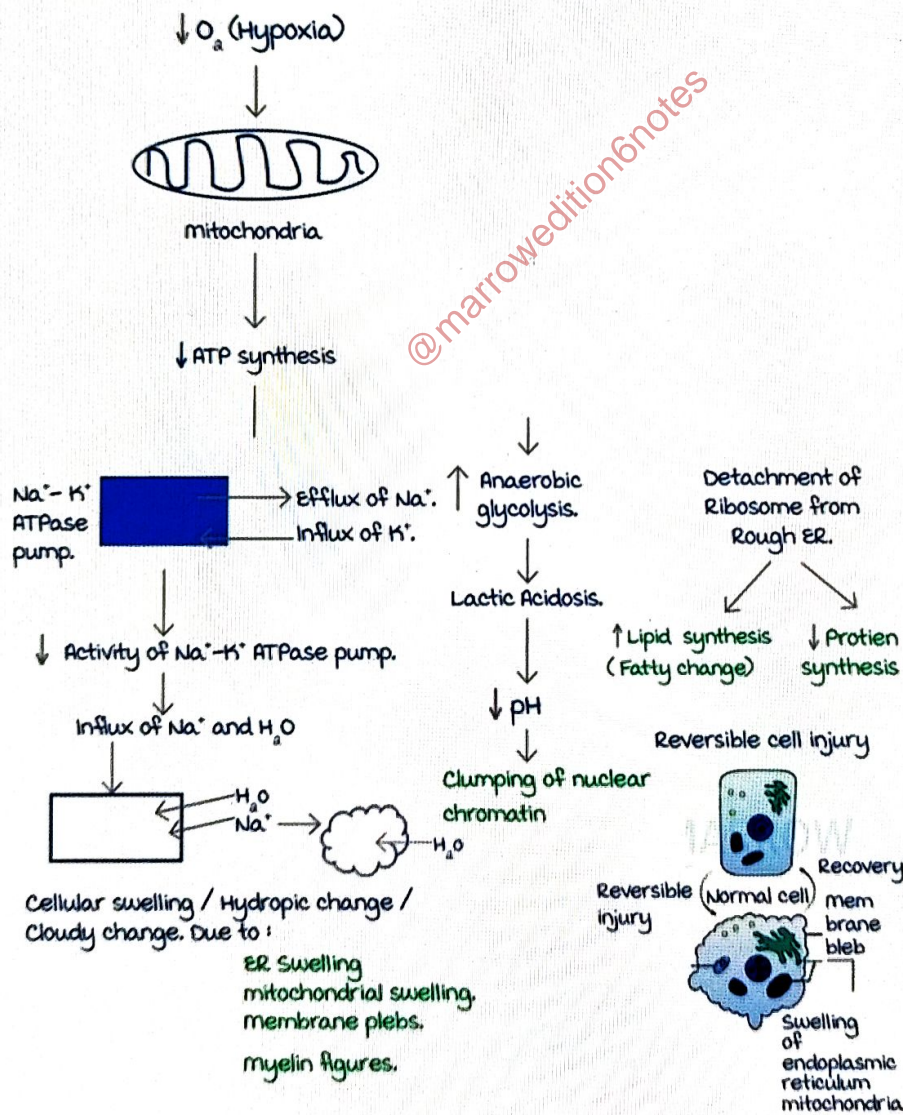
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Reversible Cell Injury :

If injurious stimuli are removed, cell can go back to its normal state.


Mechanism :



myelin figures : Due to damage to phospholipid bilayer, Composed of phospholipid Ca²⁺.

m/c organelle affected in cell injury : mitochondria.
 most important morphological feature of reversible cell injury : Cell swelling / Hydropic changes.

myelin figures :

- Seen in both reversible and irreversible cell injury.
- Composed of Ca^{2+} and phospholipids.
- Looks like myelin. 

Irreversible cell injury

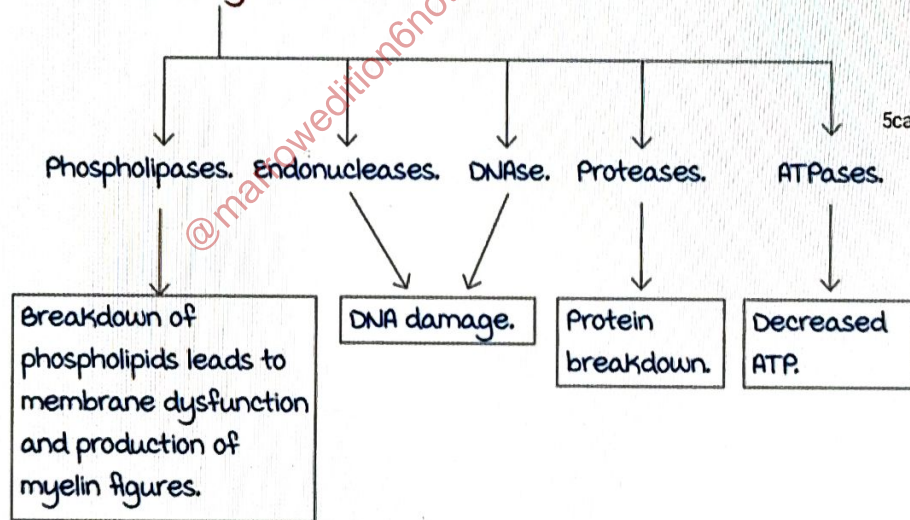
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2 things characterize irreversibility :

- mitochondrial dysfunction.
- membrane dysfunction.

membrane dysfunction :

Persistent injurious stimuli → Loose its selective permeability
 → Increased cytosolic Ca^{2+} .



Clinicopathological correlation :

In disorders like MI or liver disease : Enzymes can be measured in blood because :

- There is membrane dysfunction.
- They leak out of the cell.

Mitochondrial dysfunction

00:

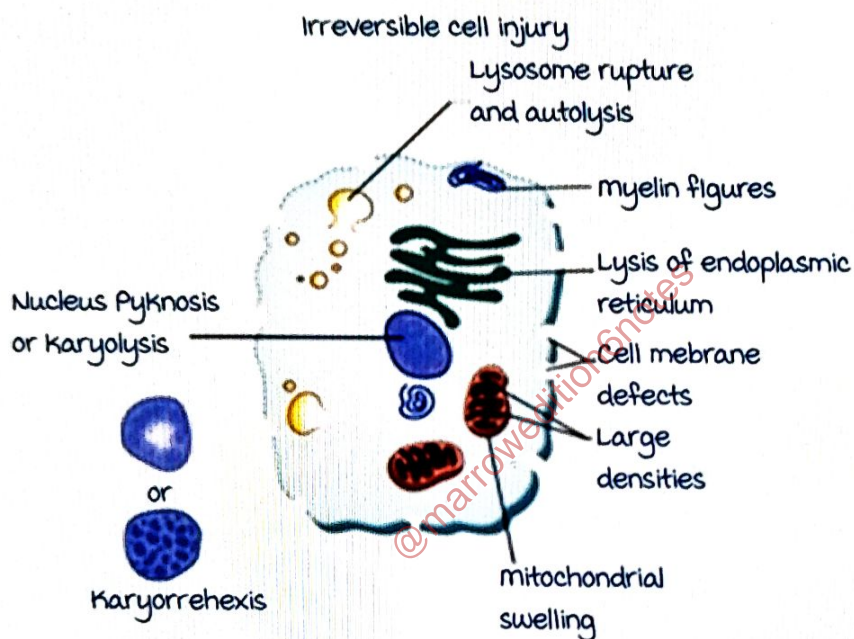
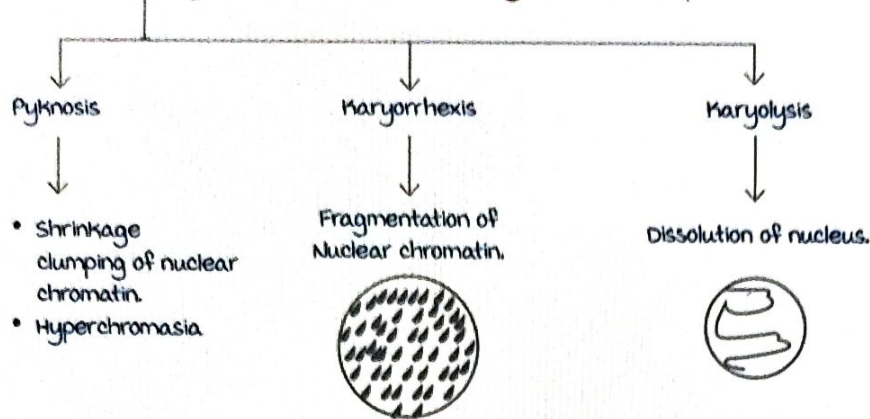
Large, flocculent, amorphous densities : Characteristic feature of irreversible injury.

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Only been seen on Electron microscopy.

Nuclear changes : most important light microscopic feature.



Free radical injury

00:27:33

Definition : molecule with one or more unpaired electrons in their outermost orbit.

Eg : O_a^- , H_aO_a , OH^- , $OOONO^-$.

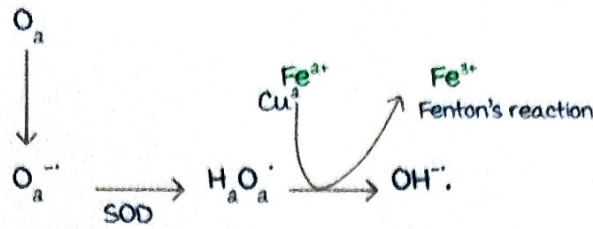
most potent free radical : OH^- .

Excess free radicals \rightarrow Oxidative stress

1. Ageing.
2. Cancers.
3. Neuro degenerative disorders : AD
Alzheimer's.
4. Reperfusion Injury.

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Production of free radicals :



Fe and Cu proteins are transferrin, ferritin and ceruloplasmin. Hence always bound and cannot always produce free radicals.

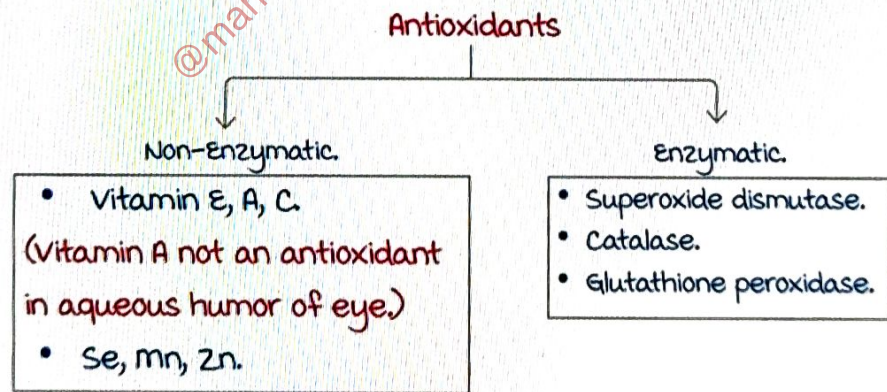
Enzymes leading to free radicals production

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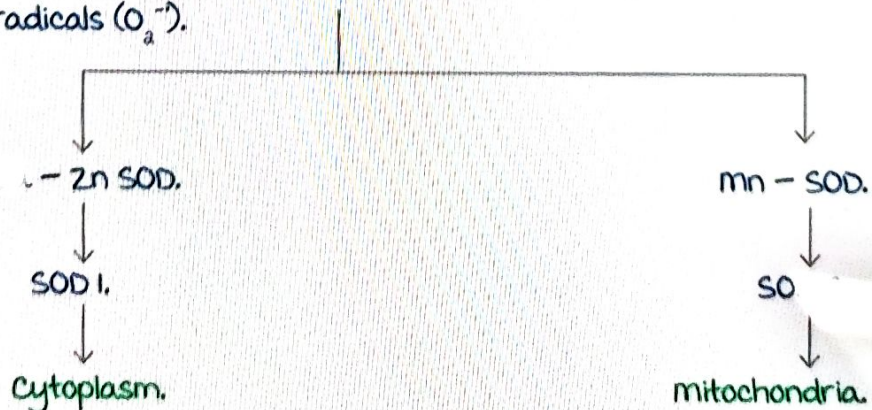
- NADPH oxidase.
- Xanthine oxidase.
- Superoxide Dismutase.

They cause injury by :

1. Oxidative modification of proteins.
2. Damage to DNA.
3. Lipid peroxidation of membranes.



Superoxide dismutase (SOD) : Inactivates superoxide free radicals (O_2^-).



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Brain is protected from free radical injury by SOD 1.

Clinicopathological correlation :

mutation of SOD 1 → Amyotrophic lateral sclerosis.

Catalase → Inactivates H_2O_2 and present in Peroxisomes.

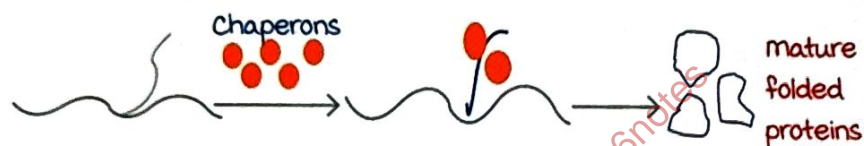
Glutathione Peroxidase : Inactivates both H_2O_2 and OH^- and present in both cytoplasm and mitochondria.

Intracytoplasmic ratio of oxidized glutathione to reduced glutathione : Important indicator of oxidative state of a cell.

Endoplasmic reticulum stress

00:44:50

ER : site of protein synthesis.



Chaperons helps in Proper folding of proteins.

When there is excess protein misfolding → Transported to cytoplasm → Degraded by the ubiquitin proteasome pathway.

Excess protein misfolding leads to misfolded protein disease.

- Familial hypercholesterolemia → LDL receptor.
- Tay sach's Disease → Hexosaminidase α Subunit.
- I AT Deficiency → α I AT.
- CJD → Prion proteins.
- Alzheimer's disease → AB amyloid.
- Cystic fibrosis → CFTR.

MCQs

Q. A 65 year old male patient presents to the emergency with substernal chest pain radiating to the left shoulder. The level of troponin I and CK-MB enzyme was done and it came out to be high. A diagnosis of myocardial infarction was made. Few hours later, the person died. What is the most likely reason for the enzyme leak?

A. Clumping of nuclear chromatin.

- B. Swelling of mitochondria.
- C. Defects in cell membrane.
- D. Autophagy by lysosomes.

Q. In an experiment, a tissue preparation is subjected to oxidant stress. There are increased numbers of free radicals generated within the cells. Generation of which of the following enzymes within these cells is the most likely protective mechanism to reduce the number of free radicals?

- A. Glutathione peroxidase.
- B. Catalase.
- C. Hydrogen peroxide.
- D. NADPH.
- E. Myeloperoxidase.

Q. A 53-year-old man suffers a cardiac arrest and his wife calls emergency services. The paramedics arrive a few minutes later and begin life support measures. A regular heart rate is established after 40 minutes of resuscitative efforts as he is being transported to the hospital. A thrombolytic agent (tPA) is administered. Which of the following cellular processes is most likely to occur in his myocardium following administration of the tPA?

- A. Apoptosis.
- B. Free radical injury.
- C. Heterophagocytosis.
- D. Squamous metaplasia.
- E. Accumulation of cytokeratins.

This is case of ischemia-reperfusion injury.

CELL DEATH

Irreversible cell injury : Cell death.

Mechanisms of cell death

00:00:32

1. Necrosis.
2. Apoptosis.
3. Necroptosis.
4. Pyroptosis.
5. Ferroptosis.
6. Autophagy.

Necrosis :

- It is a form of pathological cell death.
- mechanism :
 - I. Denaturation of proteins.
 - II. Enzymatic digestion of cells.
- Leads to damaged plasma membrane → Contents leak out of cell → Inflammatory reaction (to clear debris) → **Accidental cell death.**

Types of necrosis :

1. Coagulative.
2. Liquefactive.
3. Caseous.
4. Fat.
5. Fibrinoid.
6. Gangrenous.

Coagulative necrosis

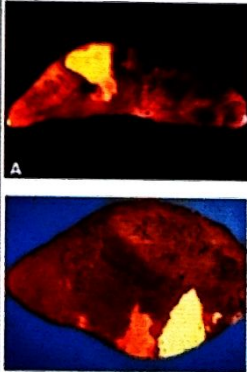
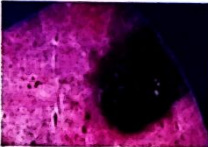
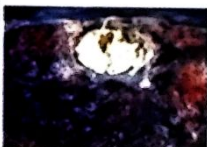
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most common form of necrosis.

morphological features of coagulative necrosis :

- **Densely eosinophilic appearance** (Loss of cytoplasmic RNA).
- Appear **glassy** (No glycogen).
- **moth eaten appearance** (Organelles digestion by lysosomal enzymes).


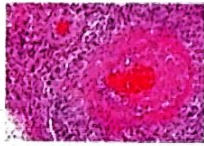

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Type of necrosis	Coagulative necrosis	Liquefactive necrosis	Caseous necrosis
Notes.	<p>Infarct is a localised area of coagulative necrosis.</p> <p>Type of dry gangrene.</p>	<p>A.K.A Colliquitive necrosis : Due to enzymatic digestion of cells.</p> <p>Type of wet gangrene.</p>	<p>Cheese like appearance.</p> <p>Combination of coagulative + liquefactive necrosis.</p>
Occurrence	<p>Occurs in all solid organs, except brain.</p> <ul style="list-style-type: none"> • Kidney. • Spleen. • Heart (MC). • Liver etc. 	<ul style="list-style-type: none"> • Brain. • Abscess. • Fungal infections. 	
microscopy	<p>Cell outlines preserved.</p> <p>Densely eosinophilic glassy, moth eaten appearance.</p>	<p>Cell outlines not preferred.</p>	
Examples	<p>1. Burns.</p> <p>2. Dry gangrene.</p> <p>3. Zenker's degeneration (Seen in typhoid : Affects rectus abdominis or skeletal muscle).</p>	<p>Brain.</p> <p>Abscess.</p> <p>Fungal infections.</p>	<p>Tuberculosis (Caseating granulomas) : mycolic acid decomposition.</p> <p>Fungal infections : Histoplasmosis. Coccidiomycosis</p>
			

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Other necrosis

00:16:38

Type of necrosis	Fat necrosis	Fibrinoid necrosis	Gangrenous necrosis
Notes	Enzymatic or traumatic	Immune complexes → Deposited in vessel wall → Fibrin like appearance → Fibrinoid necrosis.	Gangrene : Dry or wet.
Occurance	<ul style="list-style-type: none"> Traumatic : Breast. Enzymatic : Omentum, pancreas, mesentery. 	Seen in type 2 or 3 hypersensitivity reaction.	Seen in lower limbs.
Examples	Pancreatitis → Lipase → Release of free fatty acids → Contact with Calcium → Saponification → Chalky white deposits.	Aschoff nodules (Rheumatic heart disease). malignant hypertension. Polyarteritis nodosa (vasculitis).	
			

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Apoptosis

00:27:25

- Genetically programmed cell death.
- Apoptosis : "Falling off".
- most studies on apoptosis done on a nematode : *Caenorhabditis elegans*.
- Single cell death.
- mechanism of cell to eliminate unwanted cells.
- Both physiological and pathological death.

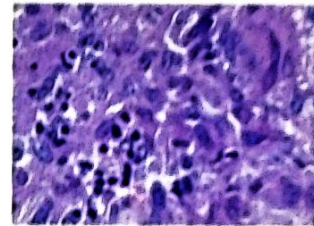
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- I. Physiological apoptosis :
 1. Organogenesis / embryogenesis.
 2. Involution of hormone dependent tissue up hormone withdrawal.
 3. Endometrial shedding during menstruation.
 4. Death of self-reactive lymphocytes.
 5. Cells which have completed their purpose (neutrophils after inflammation).
- II. Pathological apoptosis :
 1. DNA damage.
 2. misfolded protein diseases (Cystic fibrosis, alpha 1 anti trypsin deficiency).
 3. Diseases with councilman bodies (Hepatitis B).

Morphological features of apoptosis

00:34:42

- **Cell size shrinkage** : Earliest morphological feature.
- **Plasma membrane intact** → No inflammation.
- **Peripheral chromatin condensation** : most characteristic morphological feature.
- Cell membrane blebs, which will disintegrate → **Apoptotic bodies** : membrane bound with organelles.



Mechanisms of apoptosis

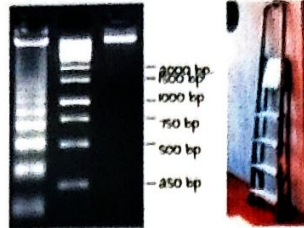
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- Three phases :
 1. Initiation phase.
 2. Execution phase.
 3. Removal of apoptotic bodies.
- 2 enzymes : Important in apoptosis :
 1. **Caspases** :
 - Cleaves near the aspartic acid residues.
 - 2 types :
 - Initiator caspases (Cas 8, 9, 10).
 - Executional caspases (Cas 3, 6, 7).

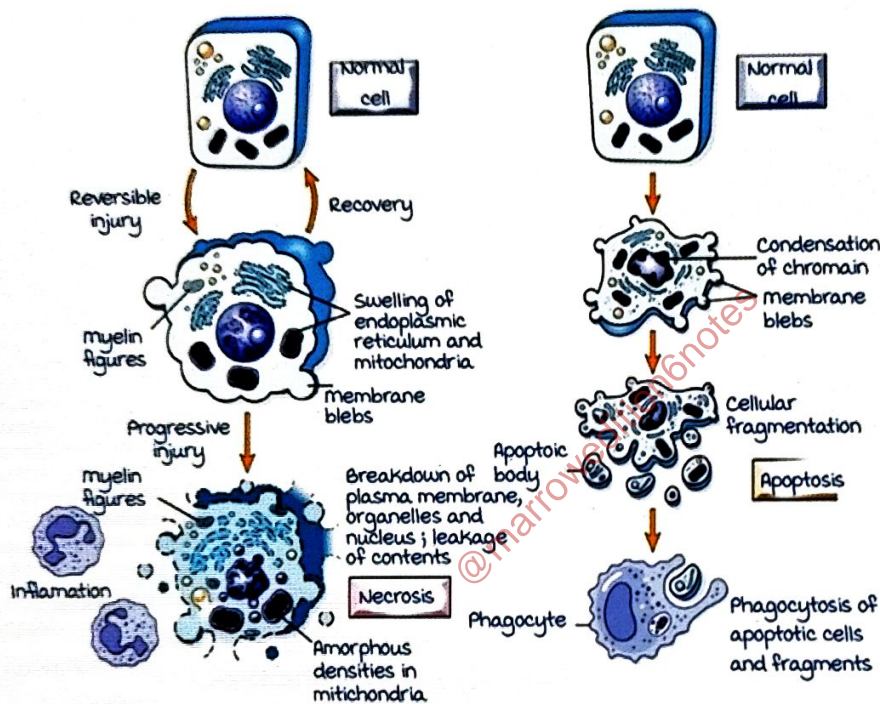
2. Endonucleases :

- Breakdown of DNA.
- When apoptotic cells subject to DNA electrophoresis → DNA fragments (which are in various base pairs) → Appear like a ladder → Stepladder pattern.

In a necrotic cell, plain pattern → Smear pattern (as there are no endonucleases)



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Feature	Necrosis	Apoptosis
Definition	Enzymatic or ischemic process	Genetically programmed cell death
Mechanism	Passive	Active
Cell	Group of cells	Single cell
Type of death	Always pathological	Both physiological and pathological
Cell size	Increases	Decreases
Cell membrane	Affected	Intact
Inflammation	Present	Absent
marker	No	Annexin V, CD 95
PAGE	Smear	Step ladder

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Regulators of apoptosis

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3 basic regulators :

1. Pro-apoptotic factors : Initiate apoptosis.

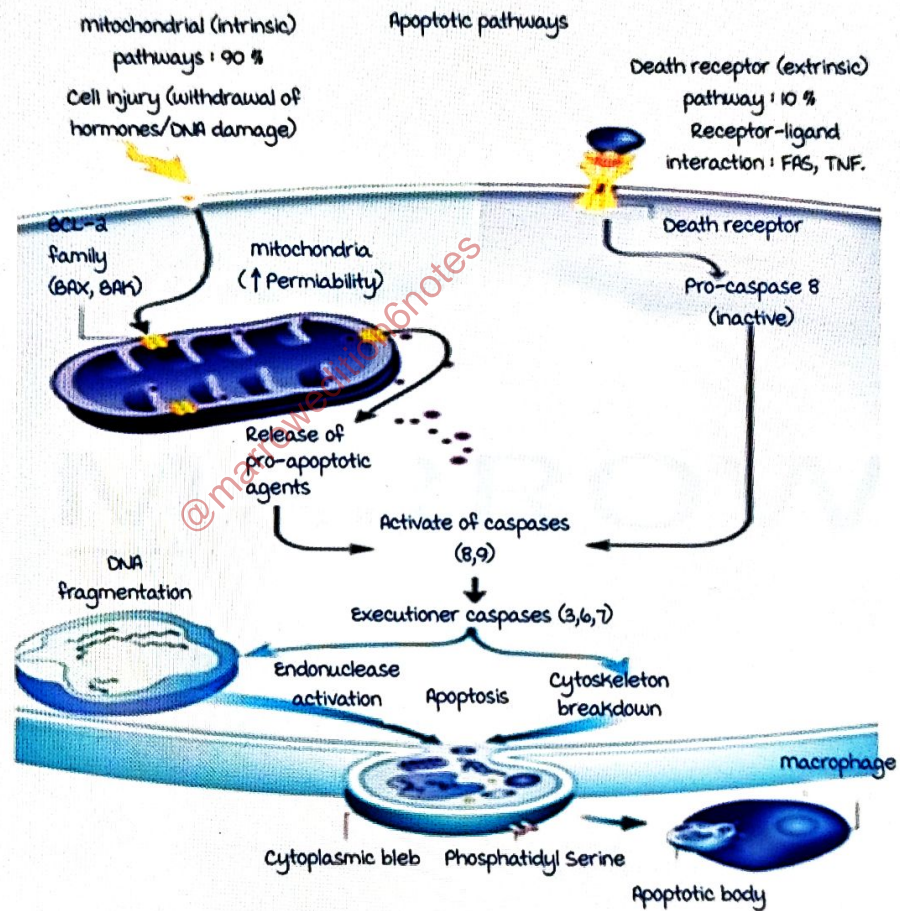
- Bax and Bak.

2. Anti-apoptotic : Inhibit apoptosis.

- BCL-2, BCL-XL, MCL1.

3. Stress sensors : Regulated Initiators of apoptosis.

- bim, bid, bad, PUMA, NOXA.



mechanism :

Initiation phase : 2 pathways :

1. Intrinsic.
2. Extrinsic

Intrinsic : mitochondrial pathway.

- 90% cases.
- MC used
- MC organelle affected in apoptosis : mitochondria.

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- Normal cell → Normal mitochondria → Permeability maintained by **BCL 2** family proteins (Anti-apoptotic) → Cytochrome c inside the mitochondria.
- Signal/any kind of trigger → Activation of **stress sensors** (Bim, bid, bad, NOXA, PUMA) → Activation of **pro-apoptotic factors** (Bax, bak) → Bax, bak channel formed between inner and outer membrane of mitochondria → Release **cytochrome c** → Leakage out of the cells → Combines with **apoptosis activating factor 1 (Apaf 1)** → Forms **apoptosome** → Activates **caspase 9** (Initiator caspases) → Activates **caspases 3, 6 and 7** (Executor caspases) → Apoptosis.
- Apoptosis :
 - Inhibitors of intrinsic pathway : **IAP**.
 - **SMAC and DIABLO** inhibit IAP (Pro-apoptotic).

2. Extrinsic : Death receptor mediated pathway.

10% cases.

- **FAS Ligand (FAS-L)** of T-lymphocyte engage with **FAS** on another cell → 4 death domains on the cell membrane combines with each other → Forms **FADD (FAS Associated Death Domain)** → Converts **pro-caspase 8** → **Caspase 8** → Activates **caspase 3, 6, 7** → Execute apoptosis
- Inhibitor of extrinsic pathway : **FLIP**.

Removal of death cells

01:02:10

- Normal cell → **Phosphatidyl serine** on inner membrane → macrophage can't recognise this molecule.
- Apoptotic cells → **Phosphatidyl serine** moves to outer membrane (**phosphatidyl serine flip**) → macrophages recognize apoptotic cells through **Annexin 5** → Binds to them → Produces a color to them → Macrophages eat them up.

marker of apoptotic cell : **Annexin 5**.

efferocytosis : Phagocytosis of apoptotic cells.

Necroptosis

01:08:44

- Necrosis + apoptosis.
- Cell starts as **apoptosis (mechanism)**, end as **necrosis (morphological features)**.
- mechanism : **Caspase independent**.
- **Programmed necrosis**.
- mechanism :
TNF + TNF-RI → Activates RIP K1 and RIP K3 → Phosphorylation of MLKL → **MLKL enters plasma membrane** → Damage of Plasma membrane + inflammation + free radical injury (morphologically similar to necrosis).
- Seen in :
Development of mammalian growth plate (Physiological).
Pathological conditions:
 1. Acute pancreatitis.
 2. Acute steatohepatitis.
 3. Neurodegenerative disorders.

Pyroptosis

01:13:30

- Cell death associated with **fever inducing cytokine (IL1)**.
- **microbial toxin** → Enters cell → Recognized by **NOD like receptors** → Activates **inflammasome** → Activates caspase 1 → Activates **IL 1** → Fever + inflammation.

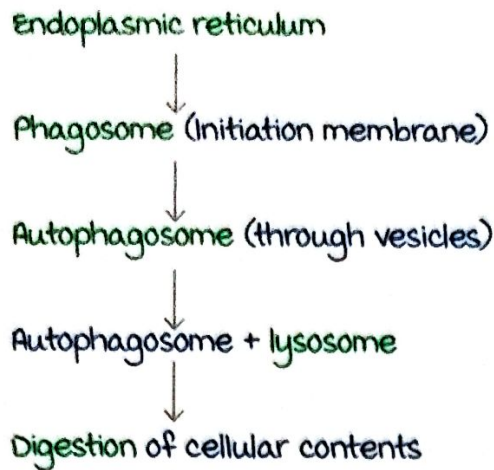
Ferroptosis :

- Cell death caused by excess iron.
- Discovered in 2012.
- **Excess intracellular iron** → Lipid peroxidation of membrane → Free radical injury → Plasma membrane damage → Cell death.

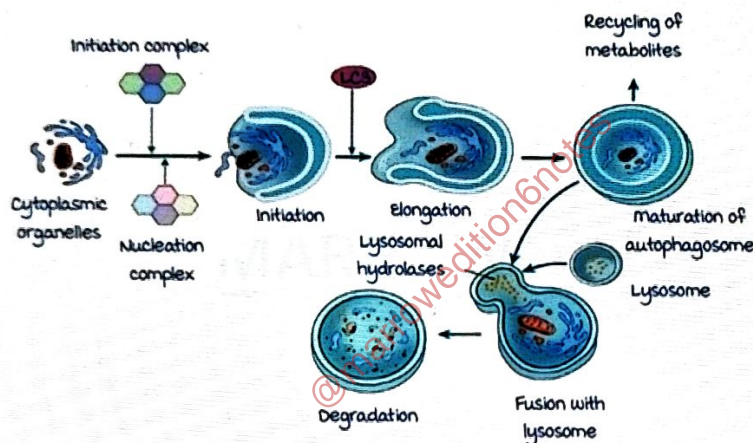
Autophagy

01:17:45

- Cell eats its own contents.
- Survival mechanism by cell during nutrient deprivation.
- mechanism :



- Genes required for formation of autophagosome.
LC 3 : marker for autophagy.
ATG 16 L1 : Seen in Crohn's disease.



Cellular stresses like nutrient deprivation activate an autophagy pathway.

This proceeds through several phases such as initiation, nucleation, elongation of isolation membrane, and eventually creates a double membrane bound vacuoles known as autophagosomes.

Cytoplasmic materials and cellular organelles are sequestered in autophagosomes, and are degraded after fusion with lysosomes.

In the final stage, digested materials are released for recycling of metabolites.

Q. A 35 year old man who works at a facility processing highly radioactive substances accidentally receives a high, whole-body dose of ionizing radiation estimated to be 1500

rads (15 gray). He dies 1 week later. At autopsy, histologic examination of the skin shows scattered, individual epidermal cells with shrunken, markedly eosinophilic cytoplasm and pyknotic, fragmented nuclei. These morphologic changes most likely indicate which of the following processes?

- A. Apoptosis.
- B. Coagulation necrosis.
- C. Liquefaction necrosis.
- D. Mutagenesis.
- E. Tumor initiation.

Q. Which of the following is not true for necroptosis:

- A. morphological features of necrosis.
- B. Programmed cell death.
- C. Caspase mediated.
- D. mediated by RIP1 and RIP 3.
- E. Seen in neurodegenerative disorders.

Q. Which of the following types of cell death is induced by lipid peroxidation?

- A. Pyroptosis.
- B. Necroptosis.
- C. Ferroptosis.
- D. Programmed cell necrosis.

Q. A 40 year old woman has the sudden onset of severe abdominal pain. On physical examination she has diffuse tenderness in all abdominal quadrants, with marked guarding and muscular rigidity. She has laboratory findings that include serum AST of 43 U/L, ALT of 30 U/L, LDH 630 U/L, and lipase 415 U/L. An abdominal CT scan reveals peritoneal fluid collections and decreased attenuation along with enlargement of the pancreas. Which of the following cellular changes is most likely to accompany these findings?

- A. Coagulative necrosis.
- B. Dry gangrene.
- C. Fat necrosis.
- D. Apoptosis.
- E. Liquefactive necrosis.

INTRACELLULAR ACCUMULATIONS

Intra = Inside Cellular = Cell

These are accumulations that get deposited inside the cell following cell injury.

Depositions :

- Proteins.
- Lipids.
- Glycogen.
- Water.
- Hyaline.
- Calcium.
- Pigments.

Glycogen

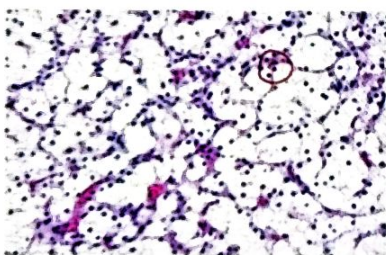
00:02:01

Can be deposited in glycogen storage disorders & in severe diabetic nephropathies (glycogen accumulation known as **Armani Ebstein lesions** seen in PCT of kidneys).

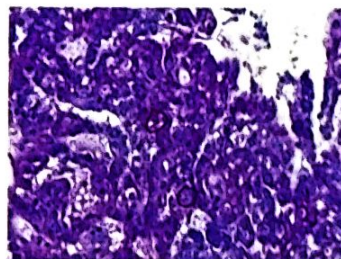
HPE : **Clear vacuoles** as it dissolves in aqueous fixative.

Special stain : PAS (Periodic Acid Schiff) → **Pink/magenta**.

Other PAS + : Lymphoblasts, basement membrane and fungi.



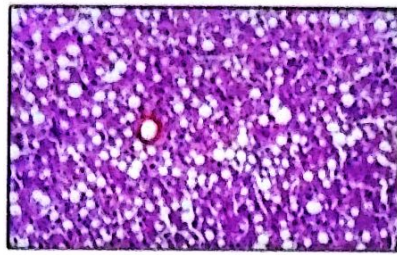
HPE of Clear cell RCC



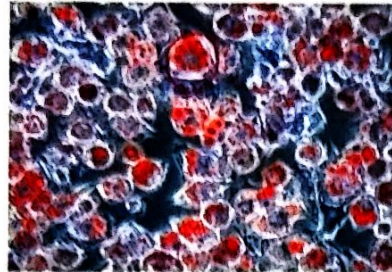
Special stain → PAS

Lipids/Fat :

- Triglycerides → Fatty liver/Steatosis.
- Cholesterol and cholesterol esters → Atherosclerosis, xanthomas, cholesterosis.



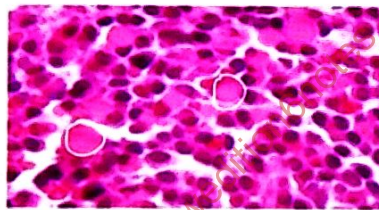
Liver Biopsy : Fat Steatosis.
 ↳ micronodular. ↳ macronodular.



Special Stain :
 Oil red - O.
 Sudan black.

Proteins :

Russel body : Intracytoplasmic inclusion.
 Dutcher body : Intranuclear inclusion. } → multiple myeloma



Russel bodies in multiple myeloma.

Reabsorption droplets in proteinuria in renal tubules.

HPE : Eosinophilic, granular appearance.

Hyaline :

2 forms → Intracellular & extracellular.

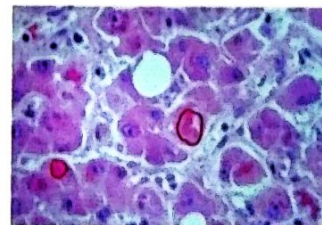
HPE : Pink/eosinophilic, smooth appearance.

Mallory hyaline body : E.g. of intracellular hyaline deposition.
 Commonly seen in alcoholic liver disease.

Other conditions where mallory hyaline bodies are seen :

(mnemonic : New Indian WATCH).

- NASH.
- Indian Childhood cirrhosis.
- Wilson's disease.
- Alcoholic Liver disease.
- Tumors like HCC.
- Cirrhosis like Primary biliary cirrhosis.
- Focal nodular Hyperplasia.



Mallory hyaline bodies are composed of intermediate filaments like CK 8 and CK 18.

Active space

Calcification

00:18:56

Deposition of calcium with small amounts of other minerals.
2 types → Dystrophic and metastatic.

Dystrophic calcification :

- Dead tissues.
- No abnormality in calcium metabolism.
- Serum Ca^{2+} → Normal.

Eg : mnemonic → **RAT**.

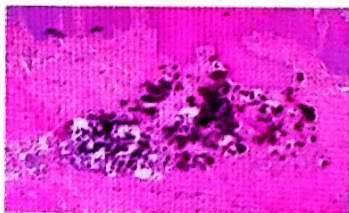
- **R**heumatic vegetations.
- **A**theromatic plaques.
- **T**B lymph nodes.
- Necrosis.
- Dead parasites.
- Monckeberg's medial calcific sclerosis (calcification in tunica media of blood vessels).
- **P**sammoma bodies → Foci of dystrophic calcification.

Seen in

1. Papillary carcinoma of thyroid.
2. Papillary renal cell cancer.
3. Meningioma.
4. Prolactinoma.
5. Serous Cystadenocarcinoma of ovary.
6. Psammoma bodies appear as concentric lamellations.

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HPE : Densely basophilic, gritty.

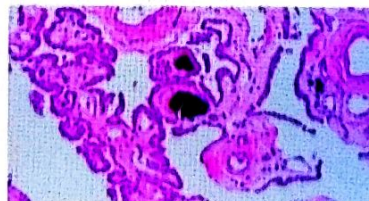


Calcifications
Squamous Epithelium



Monckeberg's medial
calcific sclerosis

Psammoma bodies with
concentric lamellations.



Active space

metastatic calcifications :

- In living tissues.
- Abnormality of calcium metabolism.
- Serum Ca^{2+} → High.

Examples :

1. Vitamin D related disorders.
2. Bone diseases like multiple myeloma and Paget's disease.
3. Parathyroid diseases.
4. RCC and breast carcinoma.
5. Sarcoidosis.
6. Milk alkali syndrome.

Calcification begins in **mitochondria** (except kidney → Begins in basement membrane of renal tubules).

m/c organ affected with calcification →

Lung alveoli > Gastric mucosa.

Special stain for Ca^{2+} :

- **Von Kossa** : Black color.
- **Alizarin red S** : Red color.

(Can even pick up small quantities).



Test for bone mineralization → Tetracycline Labelling Index.

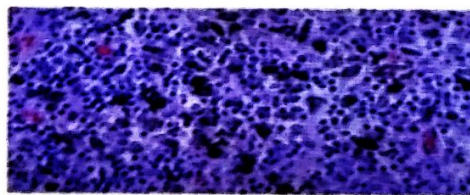
Pigments

00:36:13

Coloured substance deposited in various tissues and organs of the body.

2 types : Exogenous (external substance) and endogenous.

Exogenous : Tattoos, Anthracosis (deposition of carbon or black pigments in lungs).



Endogenous : Lipofuscin, hemosiderin, melanin.

Lipofuscin :

Pigment is derived from lipid peroxidation of membranes.

Telltale sign of free radical injury.

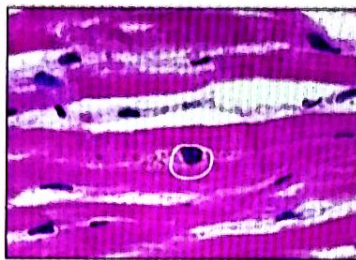
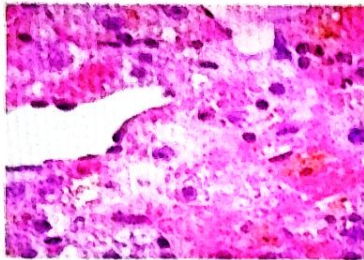
bnvssprasanth7@gmail.com

AKA Ageing pigment/wear and tear pigment.

On ageing, atrophy of organs occur, causing free radical injury. Lipofuscin gets deposited on this, hence it is responsible for brown atrophy of liver and heart.

HPE : Perinuclear brown pigment.

Special stain : Oil red - O.



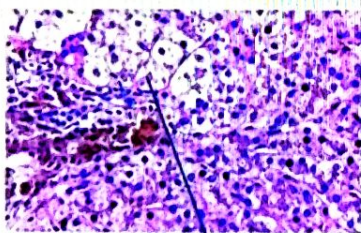
Hemosiderin :

Deposited in conditions of Iron overload (eg, blood transfusion, bruise, hemorrhage).

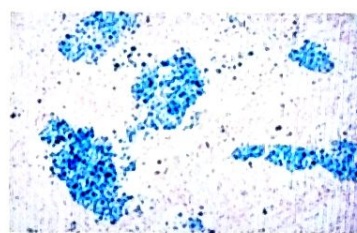
Color of hemosiderin on light microscopy : Golden yellow/brown and refractile.

Special stain : Prussian blue stain → Perl's reaction.

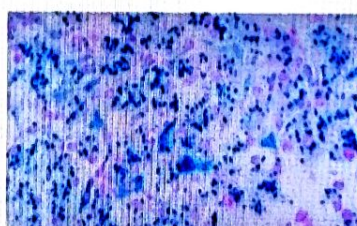
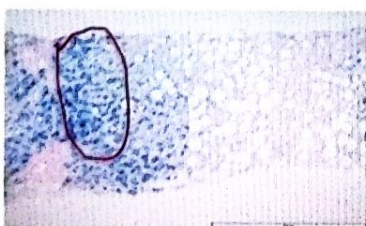
Principle : Potassium ferrocyanide → Ferric ferrocyanide.



Hemosiderin



Special stain → Prussian blue : Perl's reaction



Active space

Melanin

00:49:18

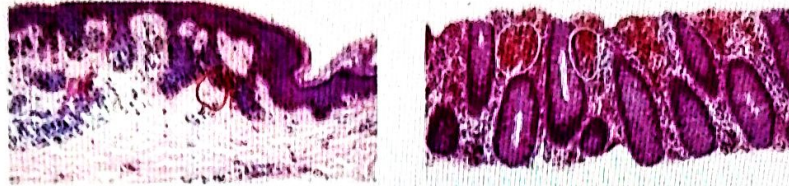
Present everywhere in our body. bnvssprasanth7@gmail.com

In Brain → Substantia Nigra.

Pale Substantia Nigra → Parkinson's disease.

Black colored pigment.

Derived from tyrosine.



Special stain : masson Fontana (mf).

a massons in pathology :

masson Fontana stain

masson's trichrome for collagen.

Dopa reaction (most specific), Schmorl's test.

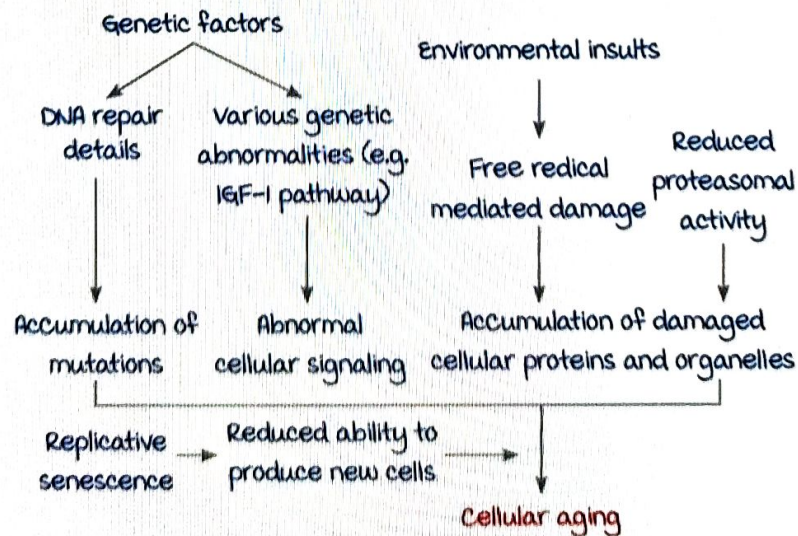
markers for malignant melanoma :

- HMB - 45
- S - 100
- melan - A.

Hemochromatosis → Bronze like pigmentation of skin is due to melanin.

Cellular ageing

00:55:38



Active space

Concepts in ageing :

m/c theory of cellular ageing → Free radical mediated damage.

- DNA damage :

Werner syndrome → Syndrome of premature ageing.

Premature ageing due to **defect in DNA helicase**.

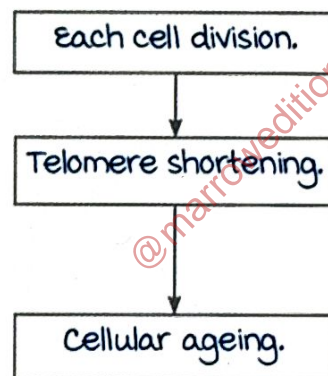
- Cellular senescence :

Hayflick limit → Cells divide only approximately 60 -70 times in their entire lifespan.

Telomeres → Short repeated sequence of nucleotides (TTAGGG) at the ends of chromosomes.

Telomeres are present as a protective mechanism that prevents the chromosome from breaking or fusion.

Telomere attrition :



Telomerase :

Enzyme which synthesizes telomeres. 5ca2793ec88d500486113130

Prevents cellular ageing.

AKA Immortality gene.

Cells with high telomerase activity → Germ cells, Stem cells.

Nil telomerase activity → Somatic cells.

Cancer cells usually have high telomerase activity.

- Dysregulated nutrient sensing :

It is a mechanism of cellular ageing.

Calorie restriction has shown to increase lifespan.

Sirtuins :

They are **NAD dependent protein deacetylases.**

They inhibit **cellular ageing** by reducing free radical injury, increasing insulin sensitivity, increasing DNA repair.

Sirtuins levels can be increased by :

- Calorie restriction.
- Wine consumption.

Sirtuins have a role in ageing, DM and cancer.

Cell/Condition	Stain
m/c stain in Histopathology	Hematoxylin and Eosin.
m/c in Hematology	Romanowsky like Leishman Geimsa.
Reticulocyte	Supravital (Brilliant cresyl blue), New methyl blue.
Lymphoblast	PAS.
myeloblast	NSE, SBB, Oil Red-O.
monoblast	NSE.
Hairy cell	TRAP.
Lipid	Oil red O, sudan Black.
Iron	Prussian Blue.
Calcium	Von Kossa, alizarin red S.
Glycogen	PAS.
Copper	Rhodamine, rubeanic acid.
mast cell	Toluidine blue.
mucin	Alcian blue.
Reticulin Fibres	Silver.
Elastin fibres	Van geison, VVG.
Collagen	masson trichrome.
melanin	Masson Fontana.
H pylori	Warthin starry silver.
Cryptococcus	Indian ink.
Fungi	Silver methenamine, PAS, GMS.
Amyloid.	Congo red.

Active space

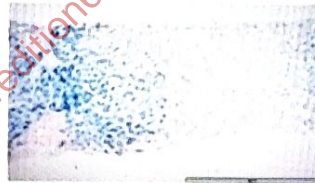
MCQs :

Q. A 54 year old man with a chronic cough has a squamous cell carcinoma diagnosed in his right lung. While performing a pneumonectomy, the thoracic surgeon notes that the hilar lymph nodes are small, 0.5 to 1.0 cm in size, and jet black in colour throughout. Which of the following is the most likely cause for this appearance to the hilar nodes?

- A. Anthracotic pigment.
- B. Lipochrome deposits.
- C. Melanin accumulation.
- D. Hemosiderosis.
- E. Metastatic carcinoma.

Q. The figure below shows the liver biopsy of a 45 year old patient who presents with raised blood sugar levels, pedal edema, and brownish skin. Investigations show reduced total iron binding capacity. What is the special stain used and the thing stained?

- A. Masson's trichrome and collagen.
- B. Prussian blue and iron.
- C. Reticulin stain and fibrosis.
- D. Brilliant cresyl blue and reticulocytes.



Q. While in a home improvement center warehouse buying paint, a 35 year old man hears 'Look out below!' and is then struck on the leg by a falling pallet rack, which strikes him on his left leg in the region of his thigh. The skin is not broken. Within 2 days there is a 5 x 7 cm purple colour to the site of injury. Which of the following substances has most likely accumulated at the site of injury to produce a yellow-brown colour at the site of injury 16 days later?

- A. Lipofuscin.
- B. Bilirubin.
- C. Melanin.
- D. Hemosiderin.
- E. Glycogen.

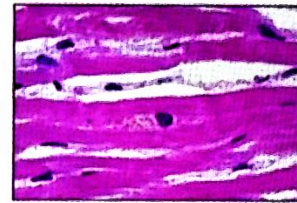
Active space

Q. Which of the following is not true about sirtuins?

- A. NAD dependent protein deacetylases.
- B. Levels increased by calorie restriction.
- C. Have a role in aging, cancer.
- D. Decrease life span.

Q. An 84 year old man dies from complications of Alzheimer disease. At autopsy, his heart is small (250 gm) and dark brown on sectioning. Microscopically, the section is given below. Which of the following substances is most likely increased in the myocardial fibers to produce this appearance of his heart?

- A. Hemosiderin from iron overload.
- B. Lipochrome from wear and tear.
- C. Glycogen from a storage disease.
- D. Cholesterol from atherosclerosis.
- E. Calcium deposition following necrosis.



6. A 49 year old man with a history of alcohol abuse has increasing abdominal girth. On examination his liver edge is firm. A liver biopsy shows cirrhosis, and individual hepatocytes contain red, globular inclusions positive for cytokeratin with immunohistochemical staining. Which of the following structural elements are these intracellular globules most likely to contain?

- A. Actin and myosin.
- B. Cholesterol esters.
- C. Fatty acids.
- D. Fibronectin.
- E. Intermediate filaments.
- F. microtubules

Alcohol liver disease → Mallory hyaline bodies made of intermediate filaments.

ACUTE INFLAMMATION

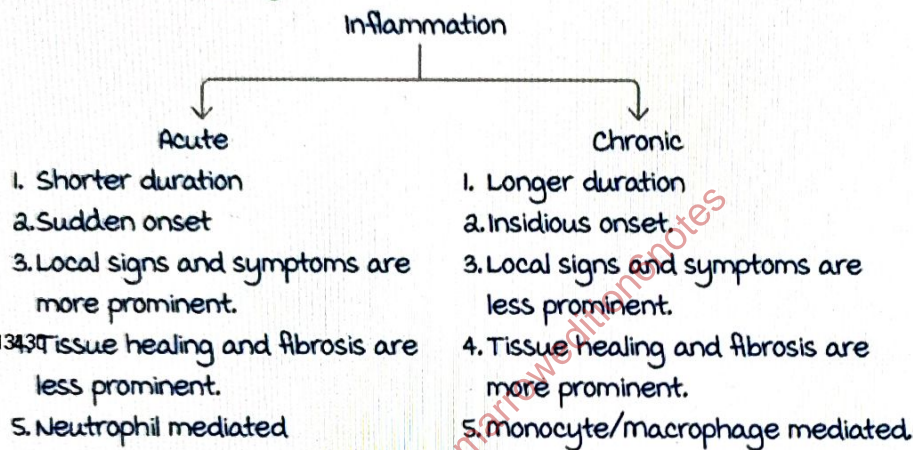
Inflammation

00:01:02

Inflammation is the response of **vascularized connective tissue** to injurious stimuli.

Injurious stimuli / precursors can be :

- Infection.
- Immune reactions.
- Foreign bodies.
- Tissue injury.



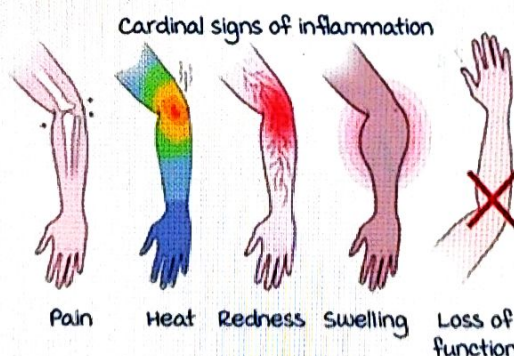
Signs of inflammation

00:05:52

4 cardinal signs of inflammation as given by Roman scholar

Celsus :

- Rubor/redness.
- Tumor/swelling.
- Calor/heat.
- Dolor/pain.



Virchow : Father of modern pathology.

He added the **5th sign** of inflammation : **Functio laesa**/loss of function.

Active space

Mechanism of acute inflammation

00:10:16

All blood vessels are lined by **endothelial cells**.

CD-34 is the endothelial marker.

In a blood vessel the leukocytes occupy the center, and flow in a **laminar fashion**, surrounded by RBCs and plasma proteins or fluid which occupy the outer parts.

In acute inflammation, leukocytes in the middle must cross the **endothelium** and basement membrane to reach **site of injury** to kill the **microbe**.

Steps of acute inflammation :

- **vascular events :**

1. Early transient vasoconstriction : Lasts only a few seconds.
2. Vasodilation.
3. Increased **vascular permeability**.
4. Stasis.

- **Cellular events :**

1. Margination.
2. Rolling.
3. Adhesion.
4. Transmigration.
5. Chemotaxis.
6. Opsonization.
7. Phagocytosis.

Vasodilation

00:16:17

Post the early transient vasoconstriction → **Arteriolar dilation**.

Histamine is the usual mediator.

Increase in blood flow causes redness/**rubor** and heat/**calor**.

Increased vascular permeability is seen in the **venules**.

Histamine is the usual mediator.

This is the hallmark of **acute inflammation**.

Swelling / tumor due to **leakage** of protein rich fluid/**exudate** outside the blood vessel.

Difference between exudate and transudate :

Exudate	Transudate
Specific gravity > 1.020	Specific gravity < 1.020
Inflammatory edema.	Non-inflammatory edema.
Rich in proteins and cells.	Poor in proteins and cells.
Increased LDH.	Decreased LDH.

mechanisms of increased vascular permeability :

1. Endothelial cell contraction / retraction / formation of endothelial gaps :
 - usually affects post capillary venules.
 - usually mediated by histamine and leukotrienes.
 - Responsible for immediate transient response.
2. Direct endothelial injury :

mild	Severe
As in burns.	As in sepsis.
Responsible for delayed prolonged response.	Responsible for immediate sustained response.

3. Leucocyte mediated endothelial injury.
4. Increased transcytosis.

Stasis

00:27:42

Stasis is the slowing of blood flow due to leakage of fluid outside causing the RBCs to accumulate inside leading to hyper viscosity of blood

Margination / pavementing

00:30:00

The process of redistribution of leukocytes from the centre to the margins of the blood vessels.

Active space

Rolling

00:32:23

The leukocytes begin to form loose attachments over the endothelium i.e., roll over the endothelium.

Rolling is mediated by certain molecules called **Selectins**.

Selectins are of three types, namely :

- E-selectin : Present on the endothelium.
- P-selectin : Present on the platelets and endothelium.
- L-selectin : Present on the leukocytes.

GlyCAM1, CD-34 are receptors present on the endothelium for L-selectin.

Sialyl Lewis X modified glycoproteins are receptors present on leukocyte for E and P selectin.

The expression of selectins is induced by IL-1 and TNF.

Redistribution of P-selectins present in Weibel-Palade bodies is mediated by histamine and thrombin.

Adhesion

00:40:49

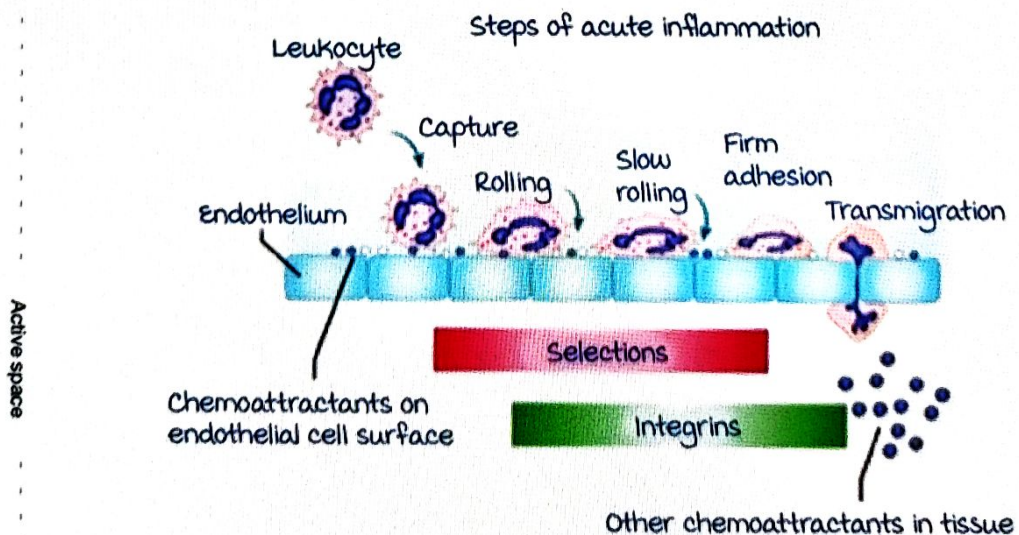
Firm adhesion of the leukocyte to the endothelium is called as adhesion.

Adhesion is mediated by certain molecules called **Integrins**.

Integrins are of two types, namely :

- $\beta 1$ integrin / **VLA4**.
- $\beta 2$ integrin / **LFA1** / **MAC1**.

Both of which are present on the leukocytes.



VCAM1 receptor present on endothelium for $\beta 1$ integrin / VLA4.
ICAM1 receptor present on endothelium for $\beta 2$ integrin/LFA1/
MAC1.

Transmigration / Diapedesis :

movement of leukocyte across the endothelium or basement
membrane (BM).

mediated by PECAM1 / CD31.

The neutrophils synthesize collagenases / matrix
metalloproteinases (MMP's) which dissolve the BM and
facilitate crossing over the BM.

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Chemotaxis

00:47:29

The movement of leukocyte in the direction of a chemical
stimuli towards the site of injury.

It is unidirectional and targeted movement.

Chemotactic mediators are broadly :

- Exogenous mediators : Secreted outside the cell, e.g.,
bacterial cell wall products like N-Formyl methionine.
- Endogenous mediators : Secreted by the leukocyte itself,
e.g., LTB_4 , IL-8, $C5a$ [mnemonic : LIC].

mechanism of chemotaxis :

Ligand binds to 7-transmembrane G protein coupled receptor

→ Increase in cytosolic calcium → Polymerization of actin

→ Chemotaxis.

Opsonization

00:52:53

It is the coating of microbes so that they are easily
phagocytosed.

Opsonins used are :

- Fc fragment of IgG (Best opsonin).
- Complement products $C3b$, $C4b$, $C5b$.
- Serum proteins like fibrinogen, C-reactive protein.

Phagocytosis :

It is the process of killing the microbe.

Discovered by **Elie metchnikoff**.

3 steps of phagocytosis :

- Recognition and attachment :

By various receptors like **scavenger receptors**, **mannose binding receptors** and **receptors for opsonins**.

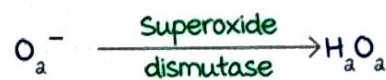
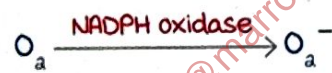
- Engulfment :

The neutrophil enters the site of injury, where they form **pseudopods** which facilitates cup formation around the bacteria. → The cup then detaches to form a **vesicle** hosting the bacteria, this vesicle is called **phagosome**. → The phagosome then fuses with the lysosome to form **phagolysosome** containing **lysosomal enzymes**.

- Killing :

It may take place via an **oxygen dependent mechanism** or an **oxygen independent mechanism**.

In oxygen dependent mechanism,



This HOCl kills the bacteria.

$H_a O_a^-$ - halide is the **most effective** bacterial killing system.

Another minor oxygen dependent killing mechanism involves **reactive nitrogen species** which will lead to the formation of **peroxynitrite** which can kill the bacteria.

In **oxygen independent mechanism**, which is a minor killing system is mediated by **enzymes** like **lysozyme**, **lactoferrin**, **major basic protein** which is present in eosinophils and has got an **anti-parasitic effect**.

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Active space

Frustrated phagocytosis :

It occurs when cell encounters materials that cannot be phagocytosed.

e.g., Immune complexes bound to basement membrane.

This is accompanied by the increased release of lysosomal enzymes.

Leukocyte function defects

01:08:40

Leukocyte function defects :

- Leukocyte adhesion deficiency Type 1 (LAD 1).
- Leukocyte adhesion deficiency Type 2 (LAD 2).

LAD 1	LAD 2
Both are autosomal recessive.	
Patients will present with recurrent infection.	
Pathogenesis : Defect in synthesis of β_2 integrin CD11 / CD18.	Pathogenesis : Defect in synthesis of sialyl Lewis x modified glycoprotein.
Delayed separation of umbilical stump.	Sometimes associated with Bombay blood group.

Chronic Granulomatous Disease (CGD) :

75% cases are X linked recessive.

25% cases are autosomal recessive.

∴ more common in males.

Pathogenesis : Defect in NADPH oxidase which results in defective oxygen dependent killing.

Clinically, patient may present with increased risk of infections with catalase positive organisms.

5ca2793ec88d500486113120 Nitro blue tetrazolium test / NBT is the screening test for CGD.

Dihydrorhodamine test / DHR is the confirmatory test done by flow cytometry.

Active space

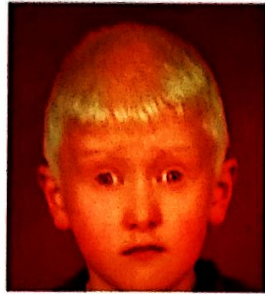
Chediak Higashi syndrome :

Autosomal recessive mode of inheritance.

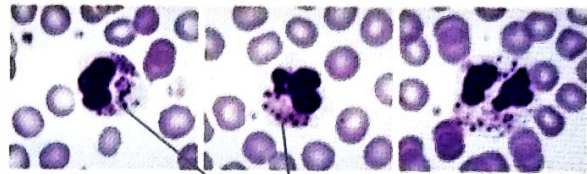
Pathogenesis : Defect in **LYST**/lysosomal trafficking regulator protein required for phagolysosome fusion.

Clinically, in addition to fever and recurrent infections patient can have oculocutaneous albinism, nerve defects or deafness and thrombocytopenia.

Peripheral smear can show **giant granules** in neutrophils.



Chediak Higashi syndrome (Albinism).



Giant granules in neutrophils.

NET / Neutrophil extracellular traps

01:19:08

NETs are extracellular fibrillar network.

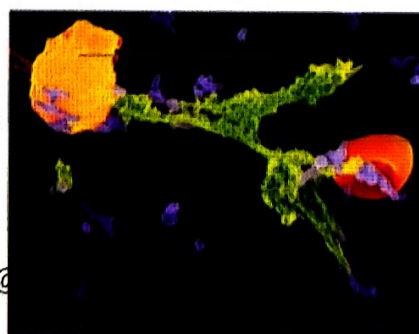
They are produced by neutrophils in response to severe infections.

It helps in limiting the spread of infection.

It produces a lot of antimicrobial substances.

Arginine is the amino acid which help in NET formation.

Some studies show that an increase production of these may cause an increased risk of autoimmune diseases like SLE.



→ Neutrophil

→ Fibrillar network

NET/Neutrophil extracellular traps

Active space

bnvssprasanth7@

Emperipolesis :

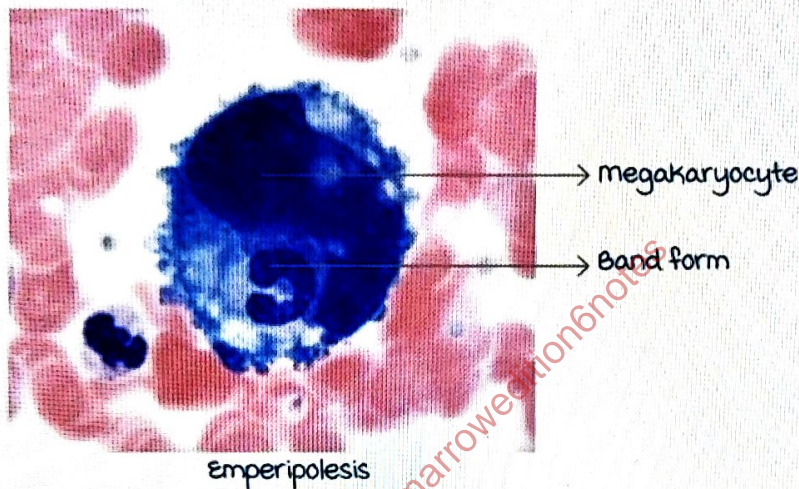
Cell within a cell appearance.

most important differential diagnosis is phagocytosis.

The cell inside can come out with no abnormality which is not the case in phagocytosis.

Seen in :

1. Rosai Dorfman syndrome.
2. Chronic lymphocytic leukaemia.
3. Haematolymphoid disorders.
4. myelodysplastic syndrome.



MCQs

Q. A 3 year old child has a history of recurrent infections with pyrogenic bacteria, Staph aureus. Neutrophilic leucocytosis is also present. Microscopic examination of biopsy obtained from that area shows microbial organisms but few neutrophils. An analysis of neutrophil function shows a defect in rolling. The child's increased susceptibility to infection is most likely caused by a defect in which of the following molecules :

- A. Selectins.
- B. NADPH oxidase.
- C. LTB₄.
- D. Integrins.

Q. A 2 year old boy presents with recurrent infections involving multiple organ systems. Extensive investigation results in diagnosis of chronic granulomatous disease. Which

of the following most closely characterizes the abnormality in his phagocytic cells?

- A. Decreased killing of microorganisms because of enhanced production of hydrogen peroxide.
- B. Deficiency of NADPH oxidase.
- C. Impaired chemotaxis and migration.
- D. Inability to kill streptococci.

Q. Which statement is true regarding NETS?

- A. Produced by neutrophils in response to infectious pathogens and inflammatory mediators.
- B. Present the spread of microbes by trapping them in their fibrils.
- C. Provide a high concentration of antimicrobial substances at the sites of infection.
- D. All of the above.

Q. In an experiment, Enterobacter cloacae organisms are added to a solution containing leukocytes and blood plasma. Engulfment and phagocytosis of the microbes is observed to occur. Next a substance is added which enhances engulfment, and more bacteria are destroyed. Which of the following substances in the plasma is most likely to produce this effect?

- A. Complement C3b.
- B. Glutathione peroxidase.
- C. Ig m.
- D. P-Selectin.
- E. NADPH oxidase.

Q. Sequence of events in acute inflammation :

- A. Vasodilation → Stasis → Transient vasoconstriction → Increased permeability.
- B. Transient vasoconstriction → Stasis → vasodilation → Increased permeability.
- C. Transient vasoconstriction → vasodilation → Stasis → Increased permeability.
- D. Transient vasoconstriction → vasodilation → Increased permeability → Stasis.

Active space

CHRONIC INFLAMMATION

Chronic inflammation : Longer duration and insidious onset.

most important cell : monocyte/macrophage.

In chronic inflammation,

1. Infiltration of tissue with mononuclear cells like lymphocytes, plasma cells or monocytes/macrophages.
2. Tissue destruction and tissue injury (hallmark of chronic inflammation).
3. Attempts at healing or repair.

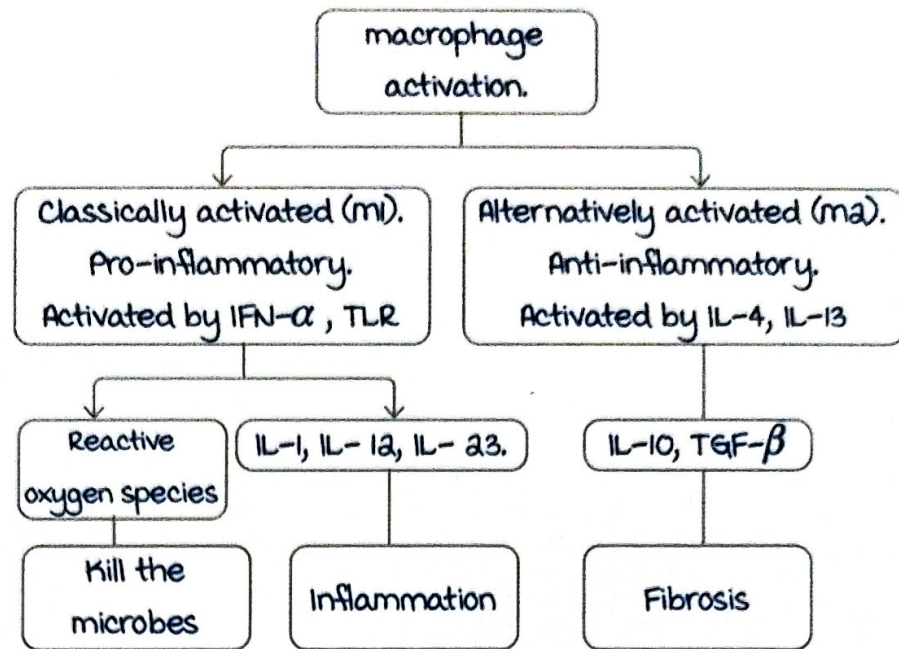
Cells of chronic inflammation : monocyte/macrophage.

monocyte : Cell with horseshoe shaped nucleus which is produced from hematopoietic stem cell. Present in blood.

Tissue specific monocytes are called macrophages.

monocytes take about 1 day to go to tissue from blood.

Tissue	Name of macrophage
Brain	microglia
Lymph nodes	Sinus histiocytes
Bone	Osteoclast
Lung	Pulmonary alveolar macrophages (dust cells)
Liver	Kupffer cells
Spleen	Littoral cells
Placenta	Haußbauer cells
Kidney	mesangial cells



Other cells of chronic inflammation : CD4+ T lymphocytes, plasma cells.

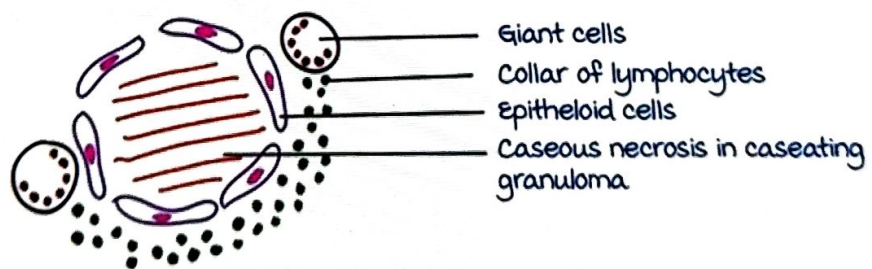
Granuloma

00:10:29

Collection of modified/activated macrophages : **epithelioid cells.**

Epithelioid cells are surrounded by a collar of lymphocytes. Giant cells may be seen.

Caseous necrosis in caseating granuloma.

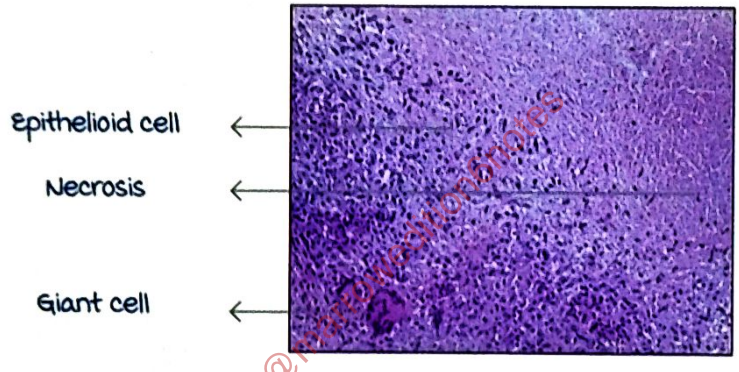
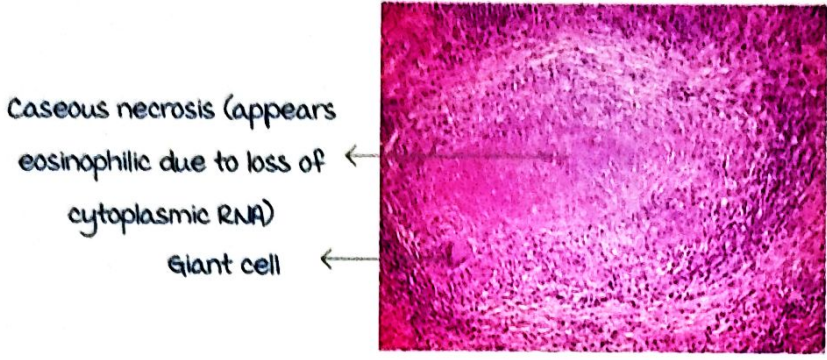
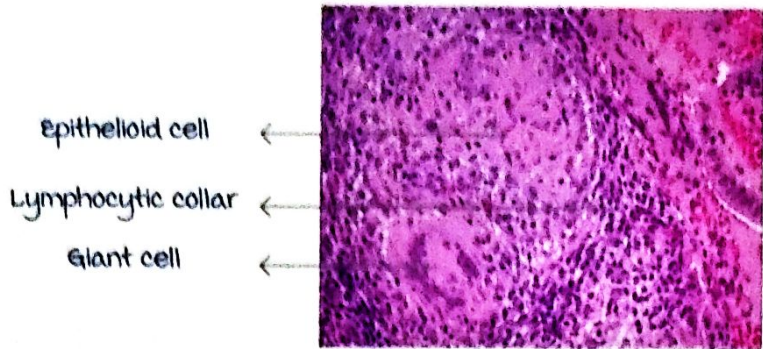


Epithelioid cells : Epithelium like appearance with slipper shaped nucleus.

most important cell in a granuloma.

Giant cell is formed by the fusion of a large number of epithelioid cells.

Active space

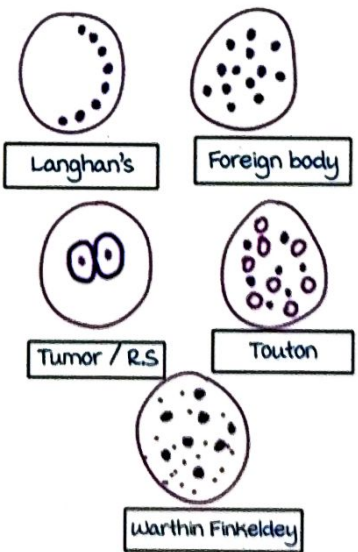


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Types of giant cells

00:16:28

- Langhans giant cell :
Seen in TB.
Horseshoe/necklace arrangement of nuclei.
- Foreign body giant cell :
Haphazard arrangement of nuclei
Foreign bodies like talc, sutures.
- Tumor giant cells :
Seen in giant cell tumor of bones.
Reed-Sternberg cells seen in Hodgkin's lymphoma.
Nuclei in RS cell have owl's eye appearance.

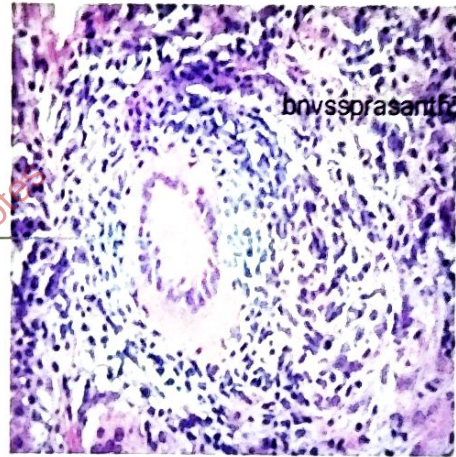


Active space

- Touton giant cells :
Seen in **xanthomas** (lipid tumor).
Foamy cytoplasm or vacuolated cytoplasm.
- Aschoff giant cell :
Seen in rheumatic fever, rheumatic heart disease.
Aschoff body : Aschoff giant cell, lymphocytes, plasma cells, caterpillar cells.
- Warthin Finkeldey cells :
Seen in **measles**.
Intracytoplasmic or intranuclear inclusions seen.

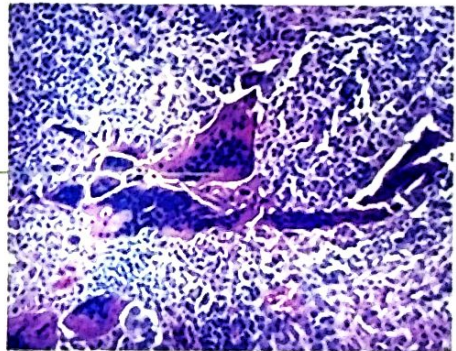
Physiological giant cells : megakaryocytes, osteoclast.

Langhan's giant cell (TB)

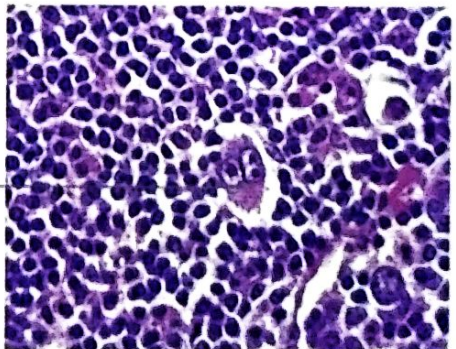


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Foreign body giant cell.

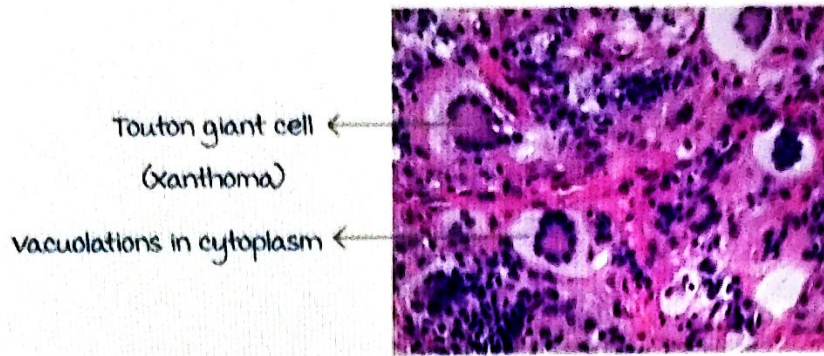


Reed-Sternberg cell.

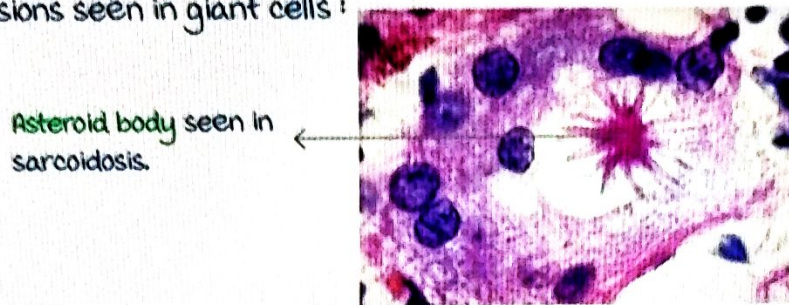


Owl's eye appearance :
Nucleus of Reed-Sternberg cell.
CMV inclusions.

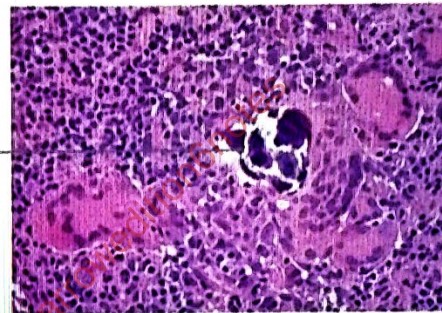
Active space



Inclusions seen in giant cells :



Schaumann body seen in Sarcoidosis :
Composed of calcium, so densely basophilic.



Pathogenesis of granuloma formation

00:23:38

Granuloma formation : Type IV hypersensitivity.

most important cytokine : $IFN-\gamma$.

Antigen presenting cell combines with $CD4+$ TH-1 lymphocyte to produce $IFN-\gamma$.

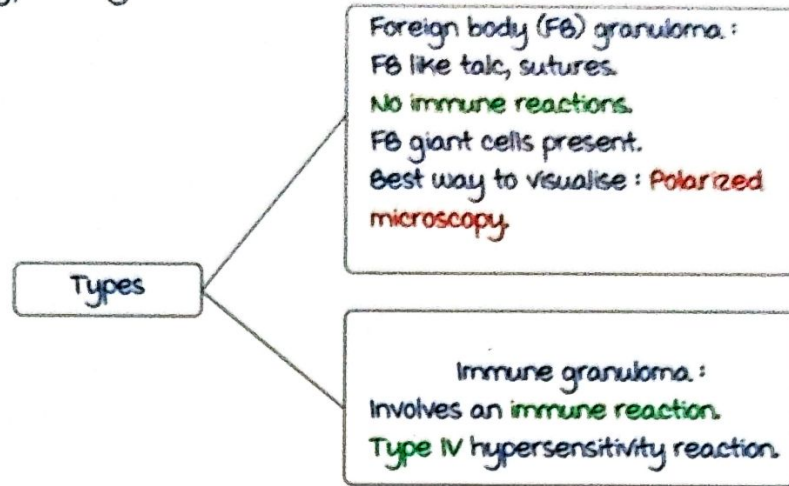
$IFN-\gamma$ → Acts on macrophage → Activated macrophage → Epithelioid cells (hallmark of granuloma formation).

Epithelioid cells are fused to form giant cells and gradually forms a granuloma.

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Active space

Types of granulomas :



Granulomatous disorders

00:28:16

Infective causes :

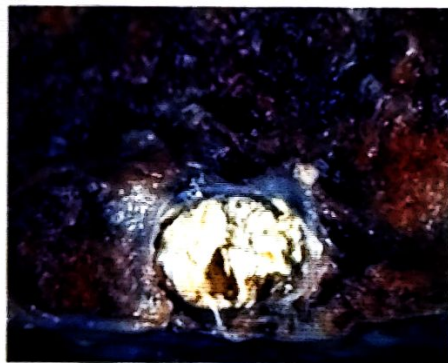
- TB : Both caseating granuloma (m/c) and non-caseating granuloma.

Gross specimen shows yellowish cheesy granuloma :

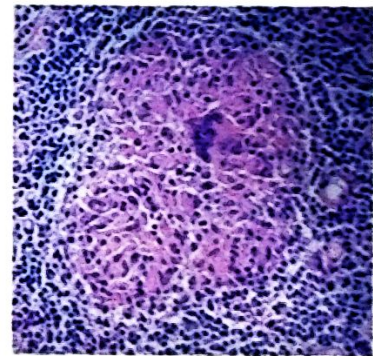
Caseating granuloma

microscopically : Epithelioid cells, Langhan's giant cell with pink caseating necrosis.

Ziehl-Nielsen stain is done to visualize the acid-fast bacilli.



Caseous necrosis : Gross appearance



Microscopic appearance

- Leprosy : Tuberculoid leprosy (granuloma formation due to intact immunity as compared to lepromatous).

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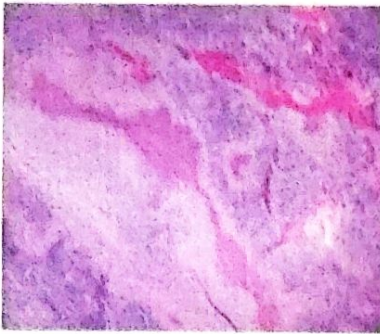
Syphilis : Tertiary syphilis/gumma

Accompanied by lot of plasma cells.

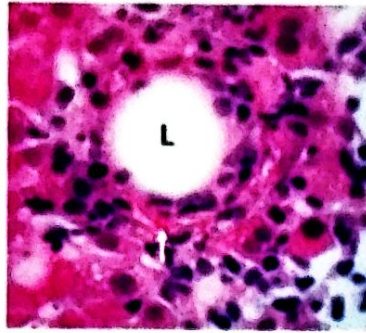
- Durk's granuloma in malaria.
 - Cat-scratch disease.
 - Lymphogranuloma venerum.
- } Stellate granuloma.

Active space

- Q fever : Doughnut granuloma/fibrin ring granuloma.
Drug causing Q fever: Allopurinol.



Stellate granuloma.

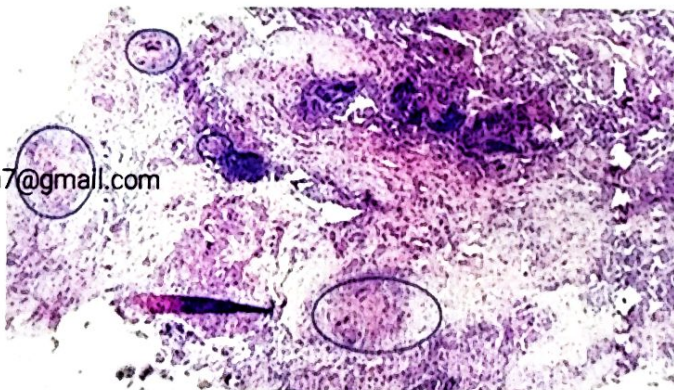


Doughnut ring granuloma.

Non-infective causes of granulomatous diseases

00:35:36

- **Sarcoidosis** :
Presence of a non-caseating granuloma (naked granuloma).
Caseating granuloma can also be seen.
Asteroid body and Schaumann body seen (giant cell inclusions).
metastatic calcification.
Absence of lymphocytic collar : Naked granuloma.
- Crohn's disease.
- **Giant cell arteritis (granulomatous arteritis)**.
- Churg-Strauss syndrome : Eosinophilic granuloma.
- Berylliosis.



Naked granuloma.

Active space

Systemic effects of inflammation :

- Fever : Cytokines involved are IL-1, TNF- α , IL-6.
- Acute phase reactants :

Positive (increased during inflammation)	Negative (decreased during inflammation)
CRP	Transferrin
Fibrinogen	Albumin
IL-6	Transcortin
Hepcidin	
Ferritin	
Haptoglobin	
Ceruloplasmin	
Factor VIII	
vWF	

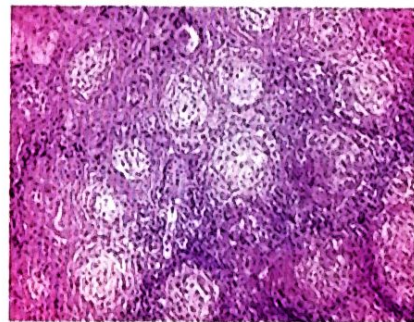
Q. Which of the following is not a feature of sarcoid granuloma?

- Non caseating.
- Giant cells have cytoplasmic inclusions.
- Fibroblastic proliferation at the periphery of granuloma.
- Peripheral mantle of lymphocytes.

Ans : Naked granuloma is seen.

Q. The figure below is from a hilar lymph node from a 54 year old man who sought medical care for low grade fever, anorexia, fatigue, night sweats

and persistent cough with hemoptysis. A chest X ray revealed a right apical infiltrate with cavitation while sputum examination revealed acid fast bacilli. This condition is typified



by a form of inflammation that invariably includes which of the following?

- Caseous necrosis.
- Multinucleated giant cells.
- Clusters of epithelioid cells.
- Prominent granulation tissue.

Ans : A granuloma is not formed without a cluster of epithelioid cells.

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Active space

Q. A 40-year-old woman had laparoscopic surgery 3 months ago. Now she has a small 0.5 cm nodule beneath the skin at the incision site that was sutured. Which of the following cell types is most likely to be most characteristic of the inflammatory response in this situation?

- A. mast cell
- B. Eosinophil
- C. Giant cell.
- D. Neutrophil
- E. Plasma cell

Ans : Likely to be a foreign body granuloma.

Q. Macrophages play an important role in phagocytosis and chronic inflammation. Which of the following cells perform the same function in liver?

- A. Merkel cell
- B. Sinusoidal cell
- C. Kupfer cell
- D. Hepatocytes

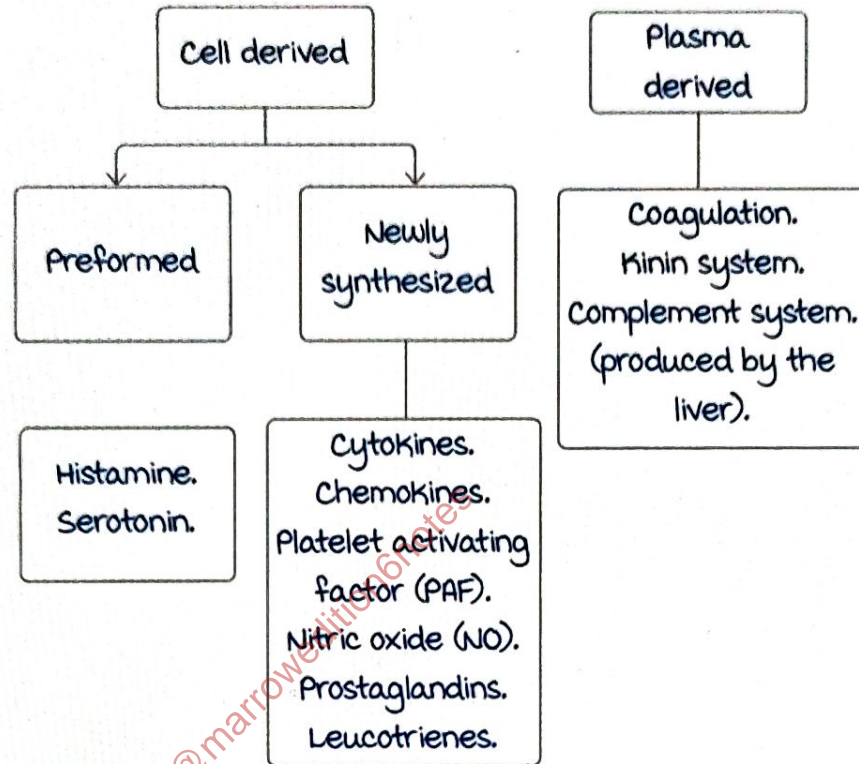
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MEDIATORS OF INFLAMMATION

Mediators of inflammation

00:00:59

mediators :



Role of mediators :

	Histamine	Serotonin
Sources	<ul style="list-style-type: none"> • mast cell (richest source of histamine). • Basophil. • Platelets. Stain for mast cell → Toluidine Blue.	<ul style="list-style-type: none"> • enterochromaffin cells of gastrointestinal tract (richest source). • Platelets.
Precursor	Histidine.	Tryptophan.
Functions common to both	1. Vasodilation. 2. Increased vascular permeability. 3. Bronchoconstriction.	1. Vasodilation. 2. Increased vascular permeability. 3. Bronchoconstriction.

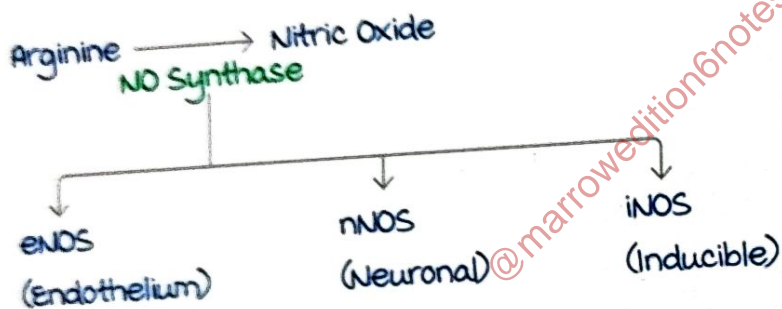
Active space

Platelet activating factor (PAF)

- Produced by:
 1. Platelets.
 2. Endothelial cells.
 3. Neutrophils.
- 1000 times more potent than histamine.
- Functions:
 1. Activates platelets (platelet aggregation).
 2. Vasoconstriction.
 3. Bronchoconstriction.

Nitric oxide (NO):

- Colorless and odorless gas.
- Also called **EDRF** (Endothelium Derived Relaxation Factor).
- Produced from **Arginine** (amino acid).



Functions:

1. Anti-microbial.
2. Smooth muscle relaxation.
3. Vasodilation.
4. Reduces platelet aggregation.

Cytokines

00:15:04

- Soluble polypeptides.
- Cytokines



Interleukins:

1. Acute infection: **IL-1, IL-6.**

2. Chronic infection : IL-12, IL-17.
3. most common cytokine in fever : IL-1.
4. most common cytokine in acute phase reactant : IL-6.

TNF alpha :

- MC Cytokine in cancer cachexia.
- Reduces appetite by mobilizing lipid and protein.
- Cachexia : Lean and thin (emaciation).

Interferon alpha :

- Antimicrobial action.

Interferon gamma :

- Granuloma formation (MC cytokine : chronic inflammation).

Function	Cytokine
Pro inflammatory	IL 1, IL2, IL 4, IL6, IL8, IFN gamma, TNF alpha
Anti inflammatory	IL 4, IL 6, IL 10, TGF beta
Both pro and anti	IL 4, IL 6
Cancer cachexia	TNF alpha
Granuloma formation	IFN gamma
Fibrosis	TGF beta, PDGF
Angiogenesis	VEGF
Fever	IL 1
Eosinophil activation	IL 5

Chemokines

00:23:48

- Small molecules which act as chemo attractants (chemotaxis) for specific cell types.
- 3 types :
 1. CXC (α) : Cysteine-X-Cysteine (X : Any amino acid other than cysteine).
Chemoattracted to neutrophils, Ex : (IL 8).
 2. CC (β) : Cysteine-Cysteine together.
Specific for : Eosinophils (Eotaxin), macrophages

Active space

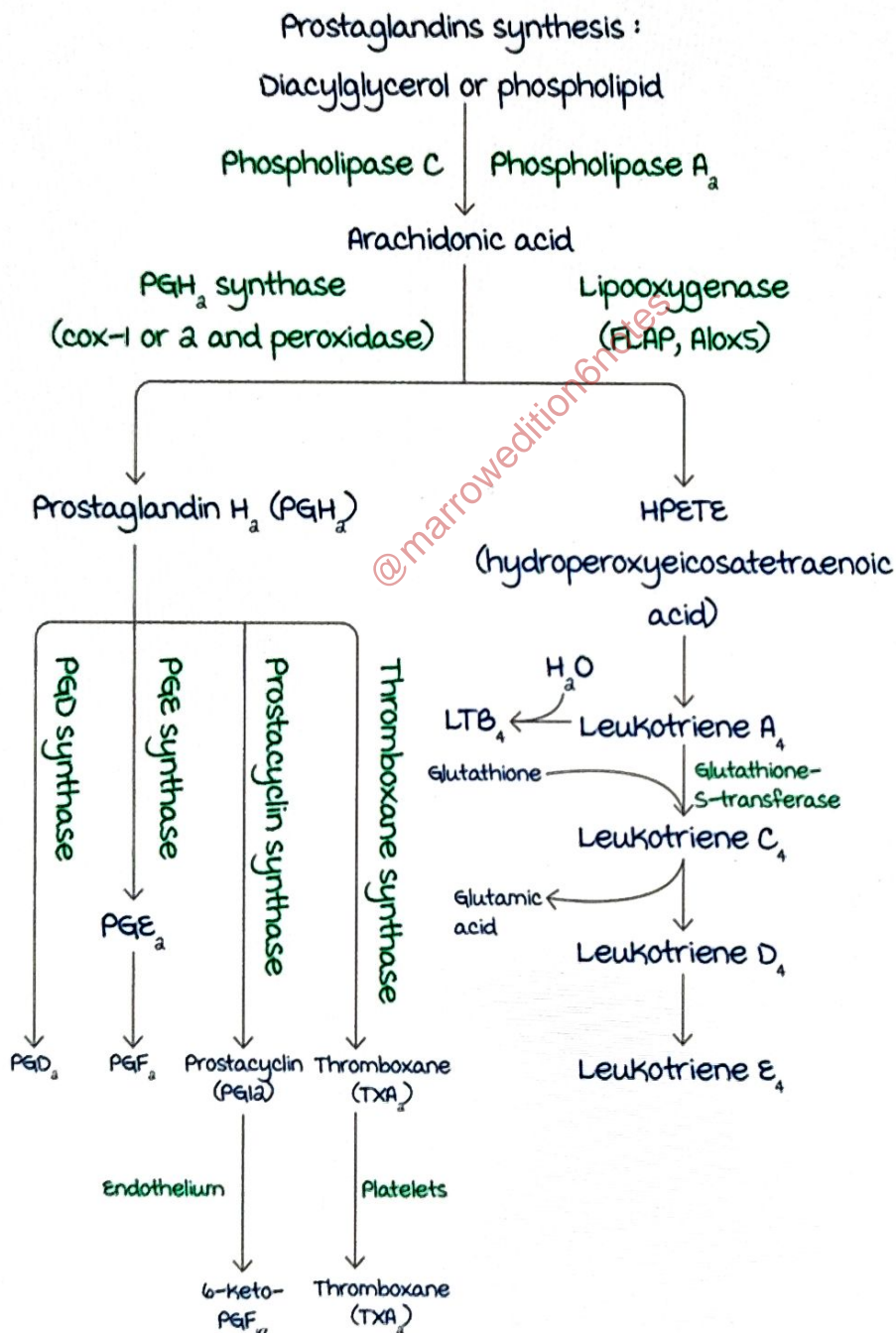
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- (MIP-1 α), monocytes (MCP-1).
- 3. Chemokine (γ) : Cysteine.
Specific for : Lymphocytes (Lymphotoxin).
- 4. CX3C chemokine (δ) :
Specific for : monocytes (Fractalkine), T-cells.

Arachidonic acid metabolites

00:28:38

- Prostaglandins and Leukotrienes :
- 20 carbon poly unsaturated fatty acid (PUFA).



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Active space

Functions of prostaglandins

00:33:08

1. Prostacyclins (PGI₂):
 - Vasodilation.
 - Decreases platelet aggregation.
 - Present on endothelial cells.

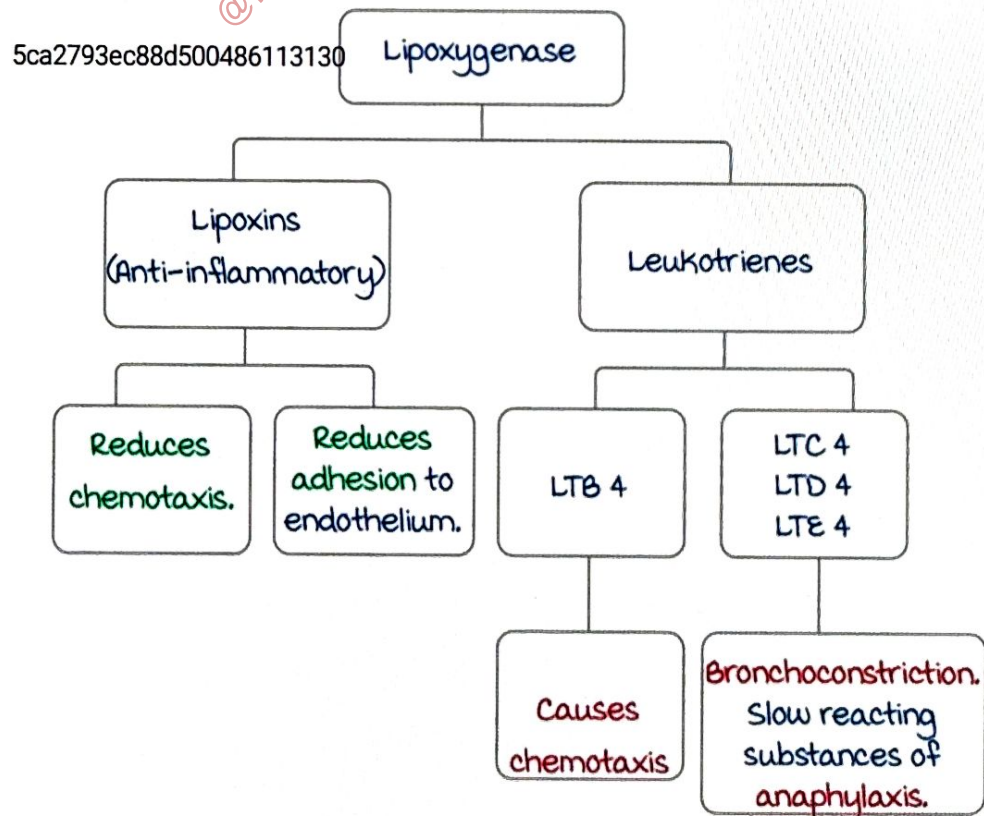
2. Thromboxanes (TXA₂):
 - Vasoconstriction.
 - Increases platelet aggregation.

3. PGD₂: Neutrophil chemotaxis.
 PGE₂: Fever, Pain.
 PGF₂α: uterine and bronchial smooth muscle contractions.

Cyclooxygenase pathway is caused by 2 enzymes:

- COX 1 → most tissues.
- COX 2 → Inducible in inflammation.

Lipoxygenase pathway:



Active space

Pharmacological applications of AA pathway 00:40:05

1. Steroids :

- Broad spectrum anti-inflammatory drugs.
- It inhibits **Phospholipase A₂** → AA not formed → No inflammation.

2. Aspirin, Ibuprofen :

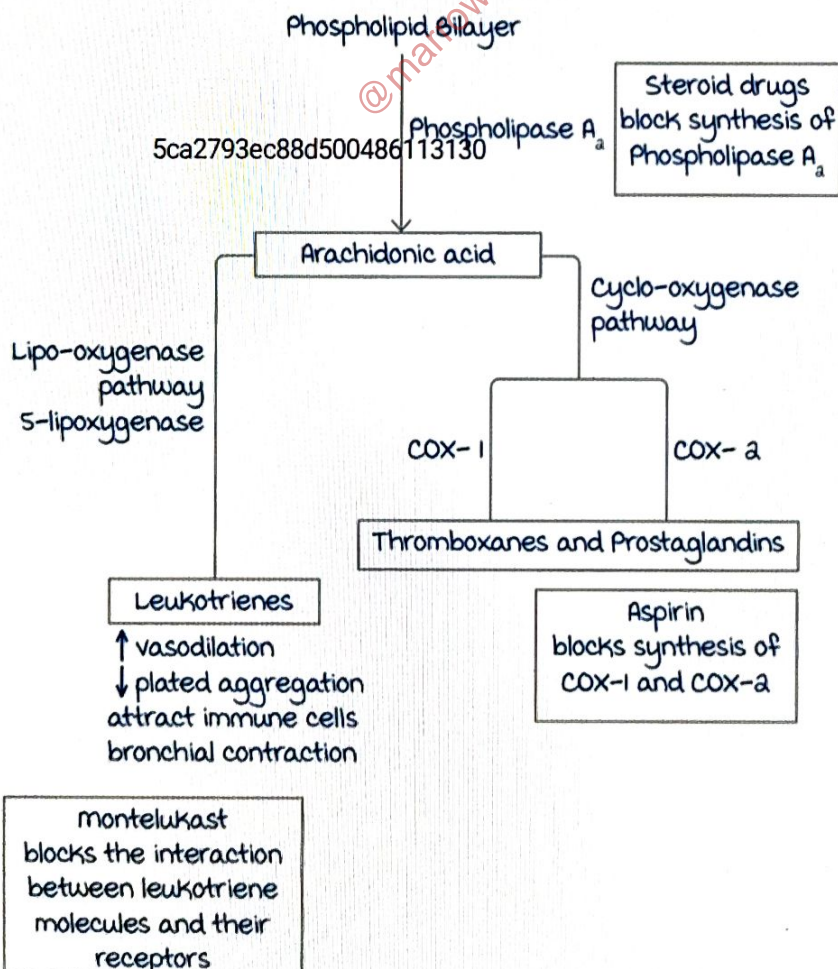
- Cyclooxygenase inhibitors (**COX-1 & COX-2 inhibitors**) → No prostaglandins → No inflammation → No pain.
- Aspirin causes **gastric ulcers** (COX-1 : Gastro protective).

3. Selective COX-2 inhibitors → more potent and less side effects.

4. Leukotriene receptor antagonists :

- Leukotrienes → Bronchoconstriction → Inhibition can lead to Bronchodilation → used for treatment of **bronchial asthma**.
- Drugs : **montelukast, Zafirlukast**.

Drugs inhibiting COX and LOX pathway
Eicosanoids Derived from Arachidonic Acid : Two Pathways



Action	mediator
Vasodilation	PGI ₂ , PGD ₂ , PGE ₂
Vasoconstriction	TXA ₂ , LTC ₄ , D ₄ , E ₄
Increased vascular permeability	Leukotrienes
Chemotaxis	C ₄ , D ₄ , E ₄
Bronchoconstriction	LT C ₄ , D ₄ , E ₄

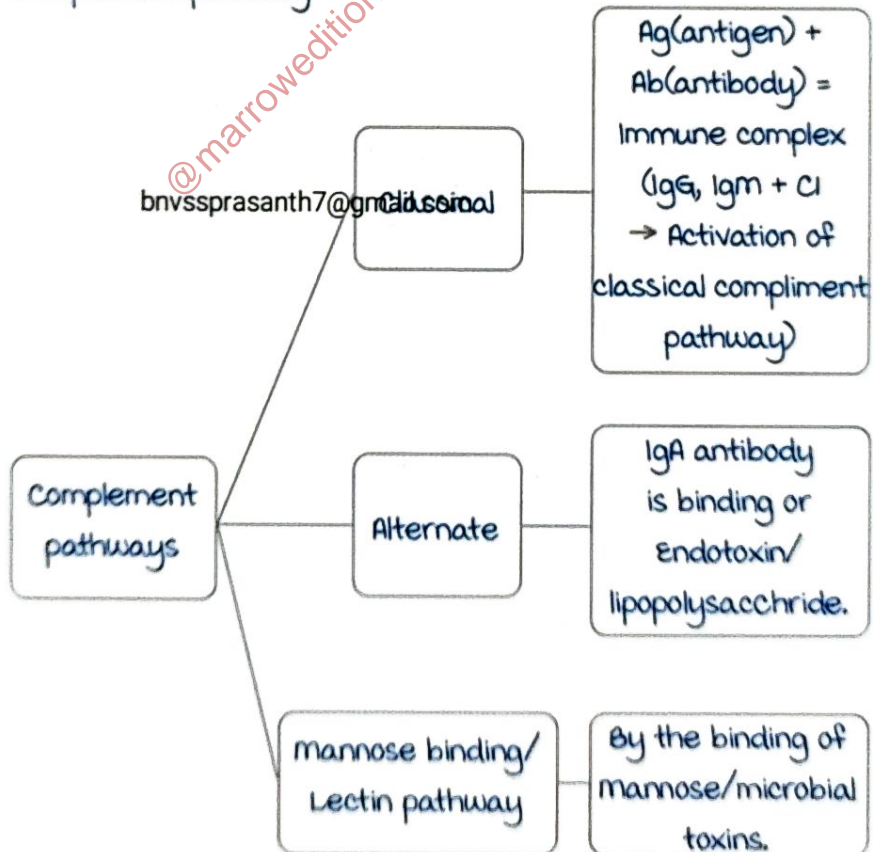
Plasma derived mediators

00:47:13

1. Complement system :

- It is a series of 20 complement proteins in plasma, secreted by liver.
- Inactive precursors $\xrightarrow{\text{Proteolysis}}$ Active precursors.

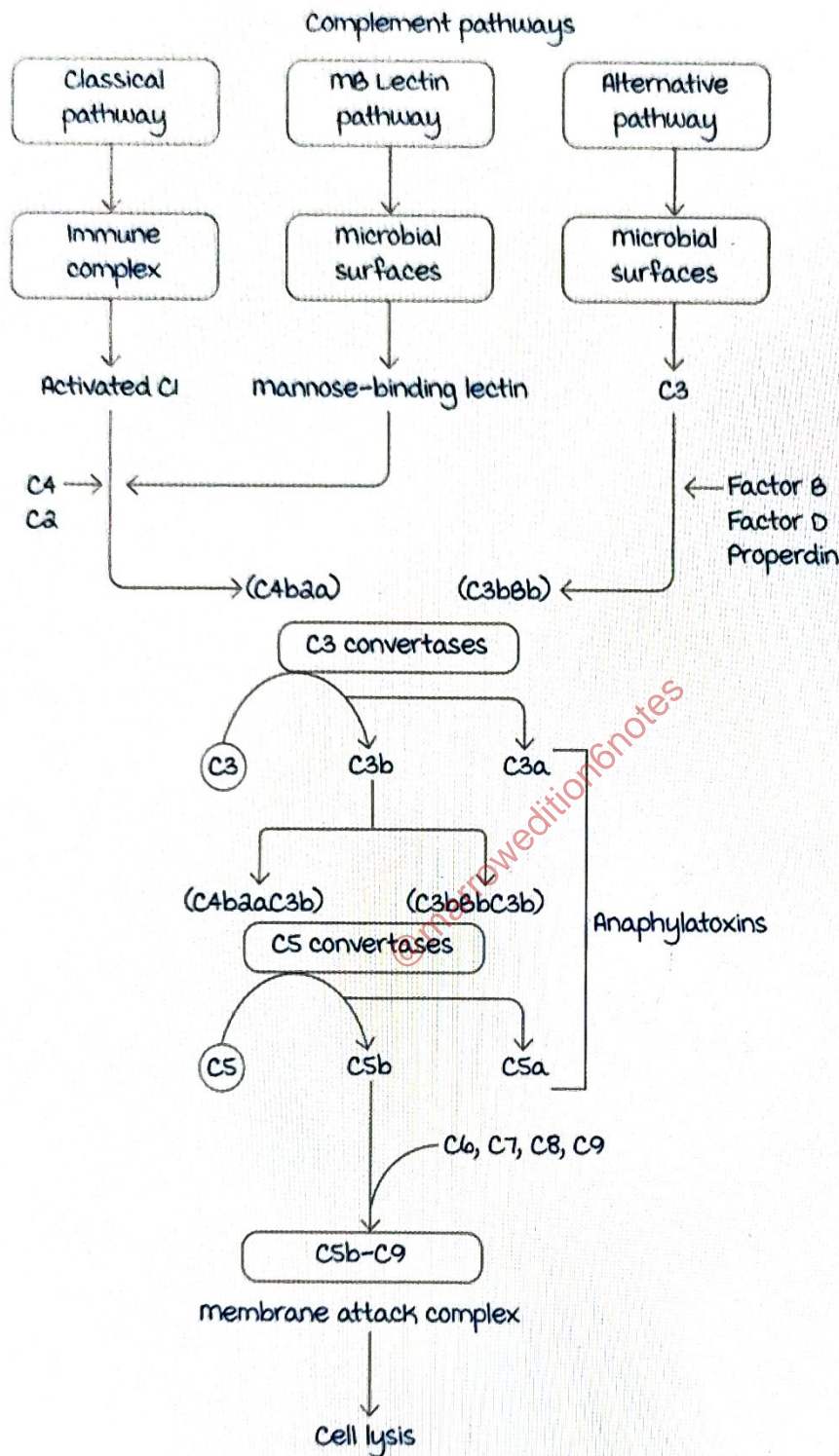
Complement pathway :



- most critical/important step in complement cascade : Activation of C₃.

Active space

- Final step in Compliment activation : Formation of MAC pool (membrane attack complex) or C5b-9



Functions of compliment pathways

00:52:46

1. Anaphylatoxins → C3a & C5a.
2. Chemotactic → C5a.
3. Opsonin → C3b.
4. MAC → C5b-9.

Active space

Clinical application :

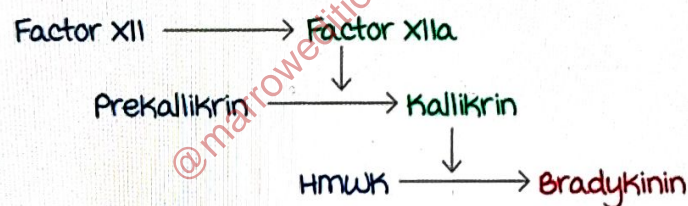
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Compliment factor deficiencies :

1. C1 inhibitor → Hereditary angioneurotic edema.
 - Swelling of lips, eyes, larynx in patients.
2. C2 (early component) → Increased risk of autoimmune disorders like SLE.
 - most common compliment factor deficiency.
3. Deficiency of MAC → Increased risk of Neisseria infections.
4. Deficiency of CD55 (DAF)/CD59 (MIRL) → Increased risk of PNH (Paroxysmal Nocturnal Hemoglobinuria).
 - Decay-accelerating factor (DAF).
 - membrane inhibitor of reactive lysis (MIRL)
5. Deficiency of factor H, I, CD 46, Properdin → Atypical Hemolytic uremic Syndrome (HUS).

Kinin system

00:58:25



Bradykinin causes pain & vasodilatation.

MCQ

Q1. Which one of the following substances is produced by the action of lipoxygenase on arachidonic acid, is a potent chemotactic agent and causes aggregation and adhesion of leucocytes?

- A. C5a.
- B. TXA2.
- C. LTB4.
- D. IL 8.

Q2. A 22 year old man develops marked right lower quadrant abdominal pain over the past day. On physical examination there is rebound tenderness on palpation over the right lower quadrant. Laparoscopic surgery is performed, and

the appendix is swollen, erythematous, and partly covered by a yellowish exudate. It is removed, and a microscopic section shows infiltration with numerous neutrophils. The pain experienced by this patient is predominantly the result of which of the following two chemical mediators?

- A. Complement C3b and IgG.
- B. Interleukin-1 and tumor necrosis factor.
- C. Histamine and serotonin.
- D. Prostaglandin and bradykinin.
- E. Leukotriene and HPETE.

Q3. A 45 year old woman has had a chronic, non-productive cough for 3 months, along with intermittent fever. She has a chest radiograph that reveals multiple small parenchymal nodules along with hilar and cervical lymphadenopathy. A cervical lymph node biopsy is performed. Microscopic examination of the biopsy shows noncaseating granulomatous inflammation. Cultures for bacterial, fungal, and mycobacterial organisms are negative. Which of the following chemical mediators is most important in the development of her inflammatory response?

- A. Interferon gamma.
- B. Bradykinin.
- C. Complement C5a.
- D. Histamine.
- E. Prostaglandin E₂.

Q4. In a lab exercise for medical students, an unknown compound is studied. The students are informed that the compound has been isolated from endothelial cells and its synthesis can be inhibited by aspirin. In the lab, the students demonstrate that the compound is a potent vasodilator and platelet anti aggregant. The substance is most likely which of the following mediators?

- A. LTCA.
- B. LXA₄.
- C. TXA₂.
- D. PGI₂.

Q5. An episode of marked chest pain lasting 4 hours brings a 51 year old man to the emergency room. He is found to have an elevated serum creatine kinase. An angiogram reveals a complete blockage of the left circumflex artery 2 cm from its origin. Which of the following substances would you most expect to be elaborated around the region of tissue damage in the next 3 days as an initial response to promote healing?

- A. Histamine.
- B. Immunoglobulin G.
- C. Complement component C3b.
- D. Leukotriene B₄.
- E. vascular endothelial growth factor.

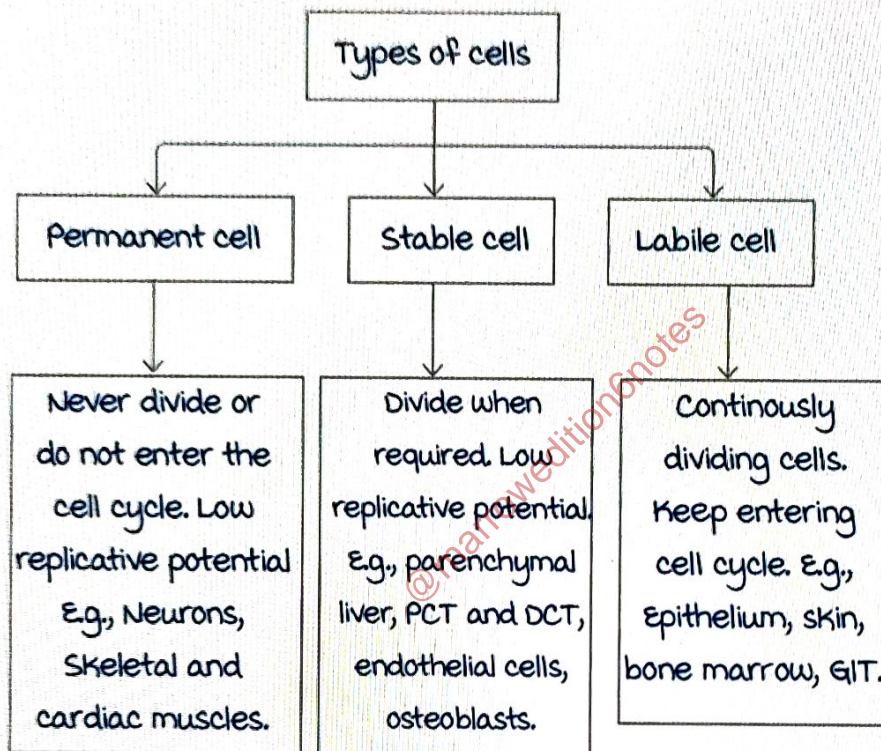
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WOUND HEALING AND TISSUE REPAIR

Regeneration : Dead cells are replaced by **same parenchymal cells**.

E.g., Following resection of a lobe of liver for tumour, rest of the liver regenerates via hyperplasia.

Repair : Dead cells are replaced by **fibrous connective tissue**.

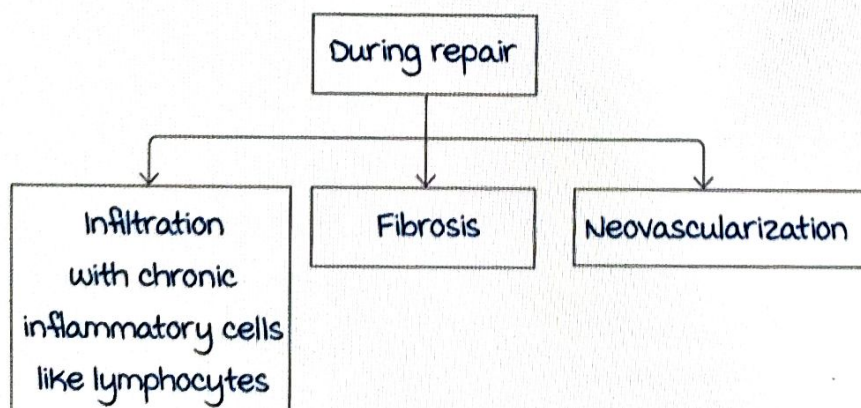


Following a brain injury, repair can happen only by neuroglia, not neurons.

Skeletal and cardiac muscle can adapt only by hypertrophy.

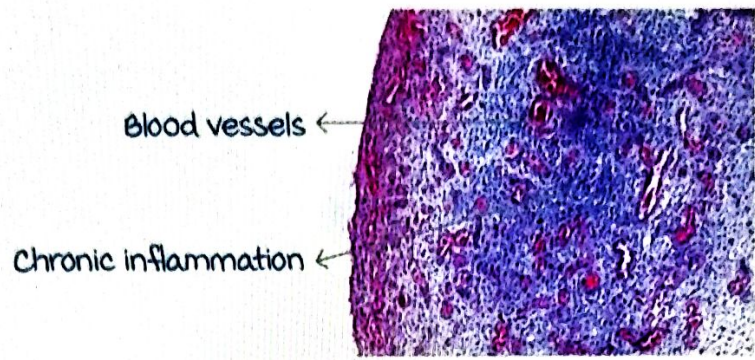
Repair

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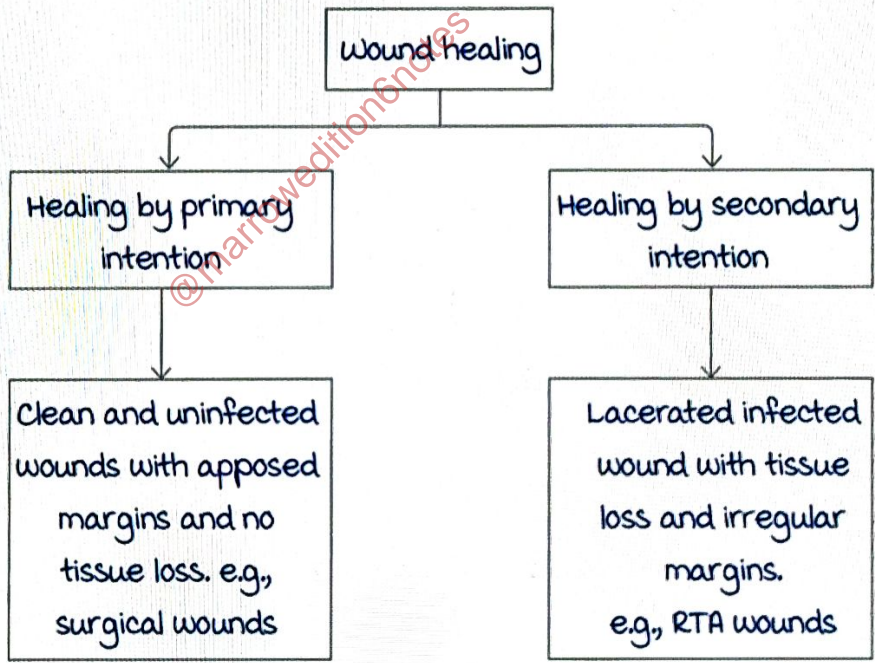
Active space

Hallmark of repair : Granulation tissue formation.
 Hallmark of granulation tissue : New blood vessel formation.
 Cytokine helping in neovascularization : VEGF.



Granulation tissue appears red because of new blood vessels formed, and edematous as these vessels are leaky.

Types of wound healing :



Healing by primary intention



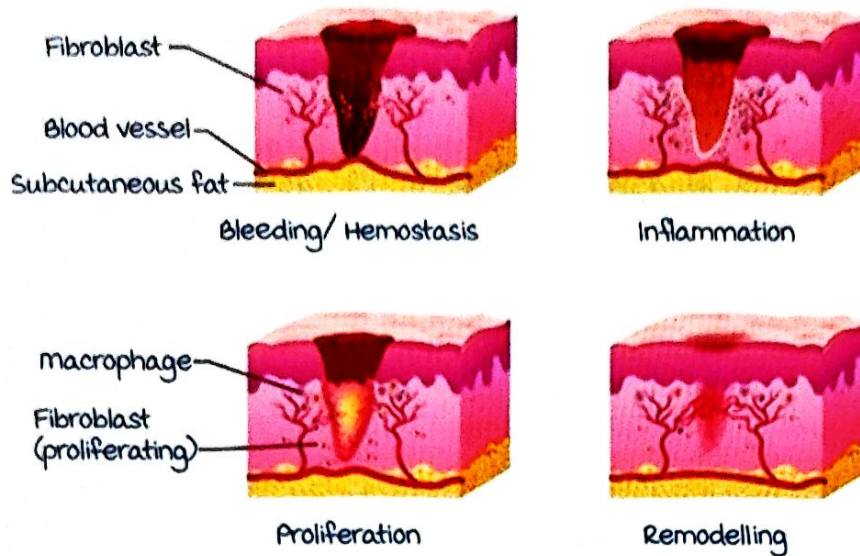
Healing by secondary intention

Active space

Steps in wound healing

00:13:32

Stages of wound healing



Stages :

- 1) Hemostasis : Stoppage of bleeding.
- 2) Inflammation : Acute inflammatory cells (neutrophils) arrive at the site of injury. This is followed by proliferation of monocytes, lymphocytes.
- 3) Proliferative phase : Repair occurs via proliferation of macrophages and fibroblasts.
- 4) Remodelling : Scar formation.

At 0 hours (immediate) :

- Hemostasis.
- with the help of fibrin.

Within 24 hours :

- Neutrophils from margins start coming towards the clot.
- mitosis begins in the basal layer of epidermis.

After 24-48 hours :

- Dense neutrophilic infiltrate.
- Thin continuous epithelial layer is formed.

On Day 3 :

- Neutrophils are replaced by macrophages.
- Early granulation tissue.
- Collagen fibres are evident at the margins of wound.

Active space

On Day 5 :

- maximum granulation tissue.
- maximum neovascularization.
- Collagen fibres bridge the incision.

3rd week :

- Decreased inflammation.
- Decreased edema.
- Decreased neovascularization.
- Increased fibroblastic proliferation.
- maximum collagen.

On Day 28 :

- Scar formation.

In secondary intention :

- more inflammatory cells.
- Bigger clot.
- more granulation tissues.
- Bigger scar.
- Wound contraction mediated by myofibroblasts occurs here, not seen in primary intention.

After 1 week : wound regains 10% of its normal strength.

After 3 months (approx 12 weeks) : 70-80% of its strength.

Wound never regains its original strength.

Initial collagen formed is type III collagen.

Type III is replaced by Type I collagen.

Type I collagen :

more abundant,

Strong and has the highest tensile strength.

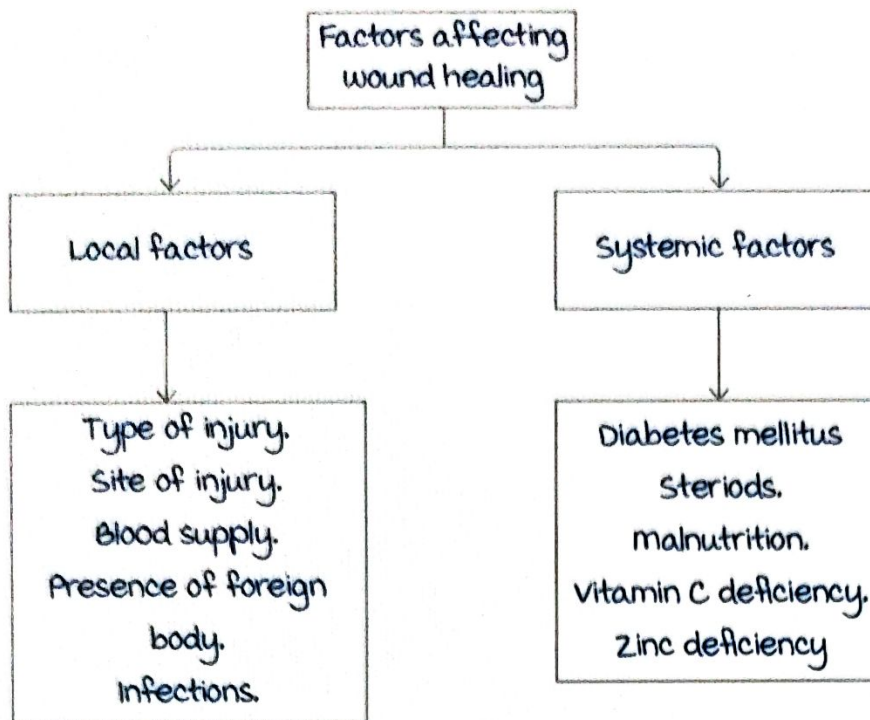
At the end the ratio of Type I collagen : Type III collagen : 4:1.

Active space

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Factors affecting wound healing

00:26:42



most common cause of delayed wound healing : Infections.

Collagens :

Triple helical structure.

Vitamin C is required for hydroxylation and crosslinking of collagen.

4 types :

Type I : most abundant, maximum tensile strength. Seen in skin, bones and tendons.

Type II : Present in vitreous humour and cartilage.

Type III : Present in keloid, uterus and granulation tissue.

Type IV : Basement membrane.

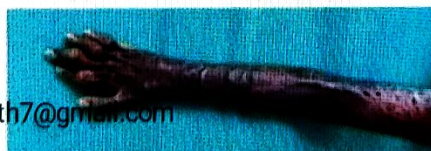
Disorders due to defective wound healing

00:31:58

Keloid

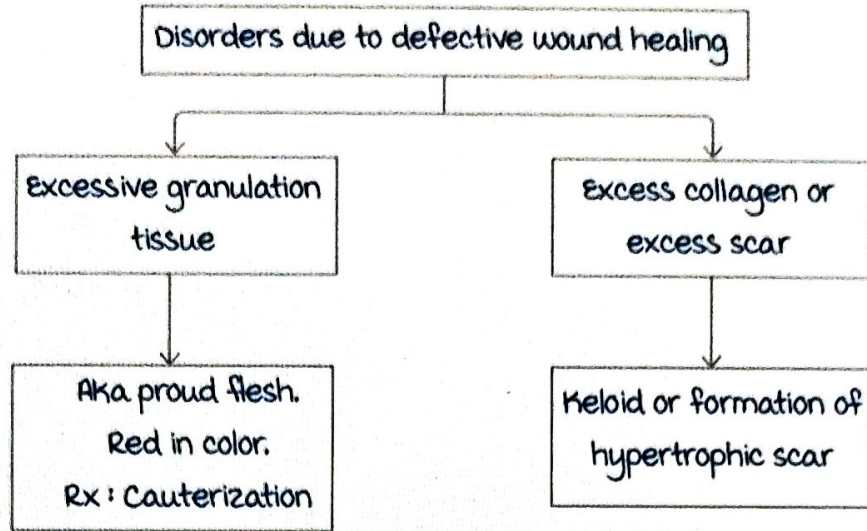


Hypertrophic scar



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Active space



Keloid	Hypertrophic scar
Scar crosses wound margins	Scar raised just above the surface
Donot regresses spontaneously	Spontaneous regression
Thick, haphazard collagen bundles	Thin orderly arrangement of collagen bundles
masson's trichrome stain is used for demonstation	-

Keloid induction for cosmetic purposes :



Desmoid : Excessive proliferation of fibroblasts.

Q. A 19 year old truck driver is involved in a collision. He incurs blunt force abdominal trauma. In response to this injury, cells in tissues of the abdomen are stimulated to enter the G1 phase of the cell cycle from the G0 phase. Which of the following cell types is most likely to remain in G0 following this injury?

- a. Smooth muscle.
- b. Endothelium.
- c. **Skeletal muscle.**
- d. Fibroblast.
- e. Hepatocyte.

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Q. A 36 year old woman has a laparotomy performed for removal of an ovarian cyst. She recovers uneventfully, with no complications. At the time of surgery, a 10 cm long midline abdominal incision was made. The tensile strength in the surgical scar will increase so her normal activities can be resumed, most of the tensile strength will likely be achieved in which of the following time periods?

- a. One week.
- b. One month.
- c. Three months.
- d. Six months.
- e. One year.

Q. A 22 year-old man incurs a stab wound to the chest. The wound is treated in the emergency room. Two months later there is a firm, 3x2 cm nodular mass with intact overlying epithelium in the region of the wound. On examination the scar is firm, but not tender, with no erythema. This mass is excised and microscopically shows fibroblasts with abundant collagen. Which of the following mechanisms has most likely produced this series of events?

- a. Keloid formation.
- b. Development of a fibrosarcoma.
- c. Poor wound healing from diabetes mellitus.
- d. Foreign body response from suturing.
- e. Staphylococcal wound infection.

most common site of keloid formation : Chest or sternal region.

Treatment of keloid :

Intralesional injections like Triamcilon.

Hypertrophic scar regresses spontaneously.

No treatment required.

Exuberant granulation tissue treated by cautery.

Active space

HEMODYNAMIC DISORDERS

Hyperaemia & congestion

00:02:00

Increased blood volume in the dilated vessels.

Hyperaemia	Congestion
Active	Passive
Red in colour (due to increased Oxygenated blood)	Blue, red in color (due to deoxygenated blood)
Arteriolar dilatation	Impaired venous outflow
Example : Inflammation	Example : Right heart failure

Examples of congestion :

Acute pulmonary congestion :

Characterized by focal hemorrhages, alveolar and septal edema.

Chronic pulmonary congestion :

Heart failure cells.

In CHF : Hemosiderin laden macrophages in lungs are seen.

Stain : Prussian blue stain/pearl's stain.

Acute hepatic congestion : Centrilobular necrosis, fatty change.

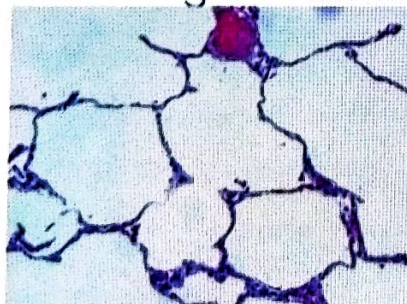
Chronic hepatic congestion : Nutmeg liver.

Chronic venous congestion in spleen : Gamma gandy body.

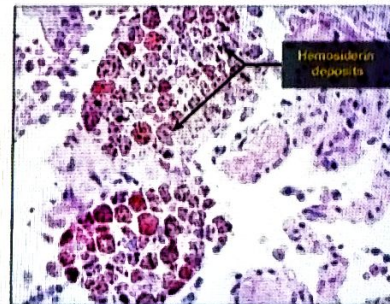


Fibrosis, calcification, hemorrhage and hemosiderin.

Normal lung alveolae

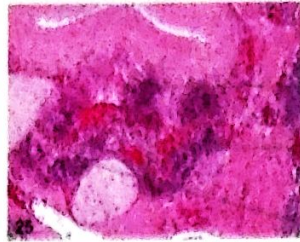
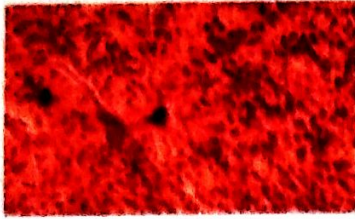


Heart failure cells



Active space

Nutmeg liver



→ Hemosiderin
→ Hemorrhage
→ Calcification

Thrombosis

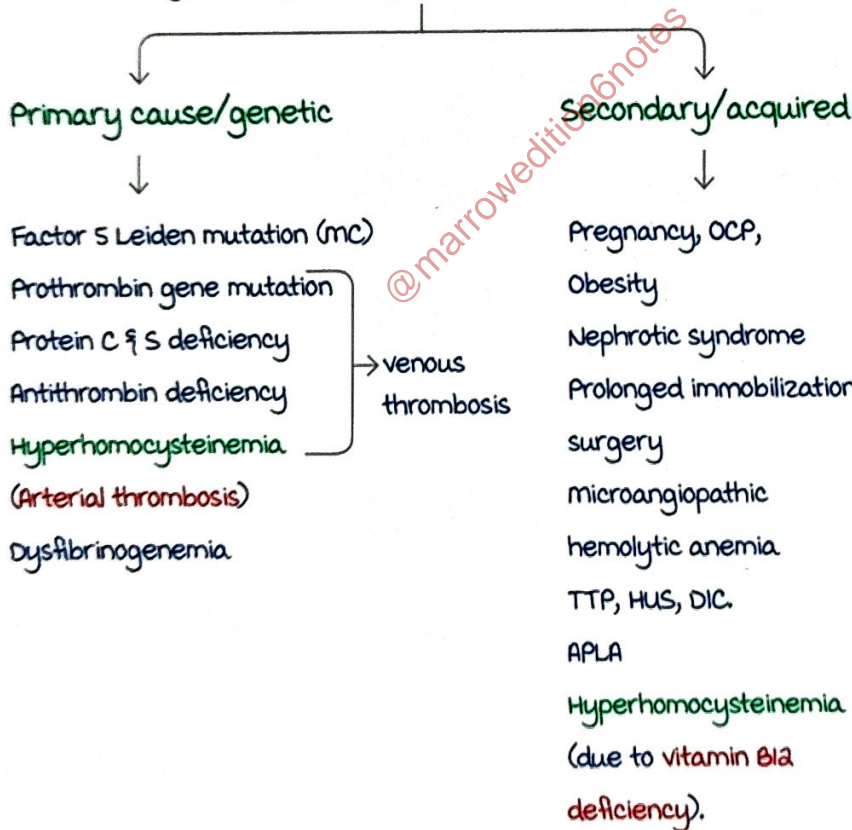
00:10:35

Thrombi is an aggregate or a mass of platelets.

Causes of thrombosis : **Virchow triad** :

1. Endothelial injury.
2. Alterations in normal blood flow (stasis or turbulence).
3. Hypercoagulability of blood.

Causes of Hypercoagulability of blood :



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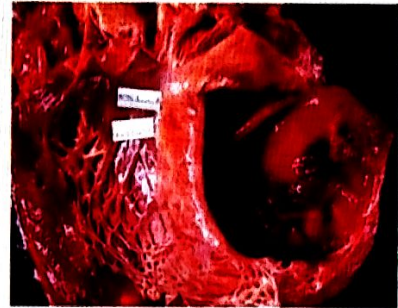
Active space

Types of thrombus

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mural thrombi	Arterial thrombi	venous thrombus
Originates in heart / aorta & attached to the wall	Occurs in rapidly flowing arterial blood	Occurs in slow moving blood of veins
	Turbulence	Stasis
	Propagates retrograde to the point of attachment	Develops along the direction of blood flow
	Grossly white in color as more platelets present.	Red in color, more RBCs present.

mural thrombi



Lines of Zahn :

Alternate light (due to platelets) and dark (due to RBC) areas.

Seen in arterial & venous thrombosis.

To differentiate,

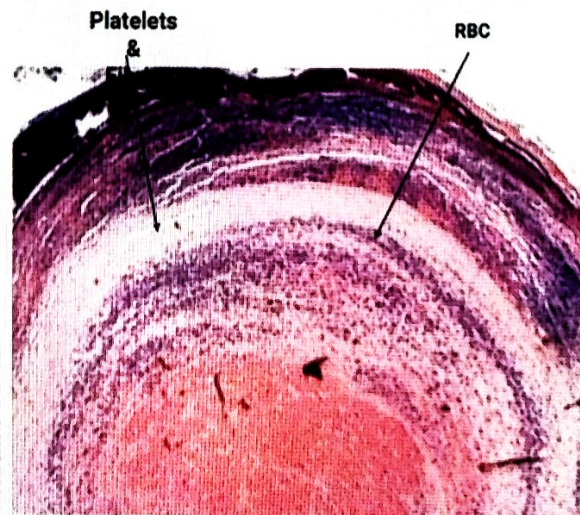
Antemortem clot : Lines of Zahn present.

Post-mortem clot : Lines of Zahn absent.

It is gelatinous.

Lower portion is red as RBC settle down by gravity.

Upper portion is yellowish because of plasma accumulation.



Active space

Fate of thrombus : (mnemonic : **DOPE**)

- Dissolution
- Organisation
- Propagation
- Embolism

Emboli

00:22:54

embolus is a **detached** intravascular solid, liquid/gaseous mass carried by the blood stream to a **site distant** from its original site.

Pulmonary emboli [PE]

mcc : **DVT** [deep vein thrombosis].

usually, asymptomatic due to the dual blood supply.

Two types PE :

Saddle embolus :

Present at the **bifurcation** of the pulmonary vasculature.

It can lead to **sudden cardiac death**.

Paradoxical emboli :

It can pass through inter arteriolar/interventricular defect and reach the systemic circulation.



Saddle
Pulmonary
Embolus

Active space

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Fat embolism :

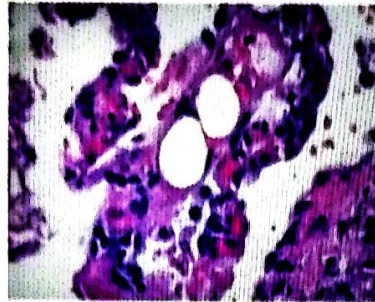
mcc : Fracture of a long bone.

Develops 1 to 3 days after a fracture

C/F ; Dyspnoea, delirium, decreased platelet count.

microscopically : Fat globules in urine.

On chest x ray : Bat wing appearance.



Air embolism and Amniotic fluid embolism

00:28:56

Air embolism :

A/K/A Caisson's disease, Decompression sickness.

Seen in deep sea divers :

Rapid change in atmospheric pressure



Increased production of nitrogen bubbles

>100 ml air is required to produce air embolism.

C/F : Bends (gas bubbles in the joint produce pain).

Chokes (gas bubbles in lung).

Amniotic fluid embolism :

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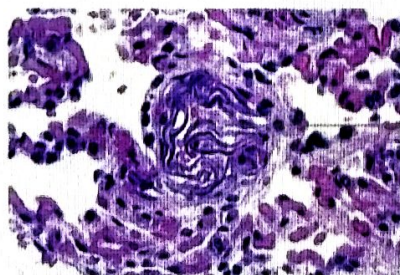
It is seen as a pregnancy / labour complication.

It is seen in post-partum period / during labour.

Tear in placental membrane : The fetal tissue enters maternal circulation.

microscopically : Squamous epithelial cells, lanugo hair, fat

from vernix caseosa.



→ Squamous cells of fetal skin

Active space

Infarct

00:33:28

An infarct is a localised area of **coagulative necrosis**.
It is **wedge shaped**.

Two types :

Haemorrhagic/red infarct	Pale/white infarct
Seen in organs with dual blood supply . e.g., GIT, lungs.	Seen in organs with end arterial circulation . e.g., heart, kidney, spleen.
Seen in loose tissues - ovaries.	Seen in solid organs .
Ill-defined margins	Well defined margins

Liver shows **both** white & red infarcts.

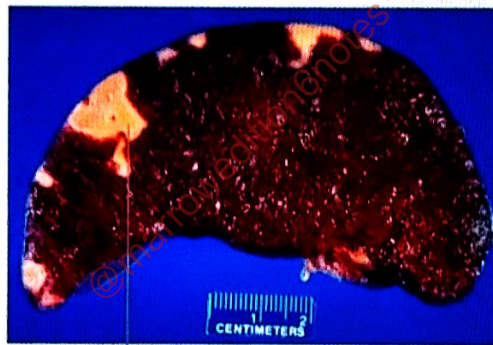
Lung : Red infarct

Wedge shaped

spleen : white infarct



Red infarct



White infarct

MCQs :

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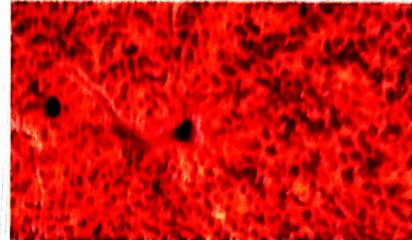
Q. A 23 year old man undergoes surgery for fracture of pelvis and left femur resulting from a motor vehicle accident. The next day he develops dyspnea, speech difficulty and a petechial skin rash. Which of the following types of embolism is the likely cause of these findings?

- A. Air
- B. Amniotic fluid
- C. Fat
- D. Paradoxical

Active space

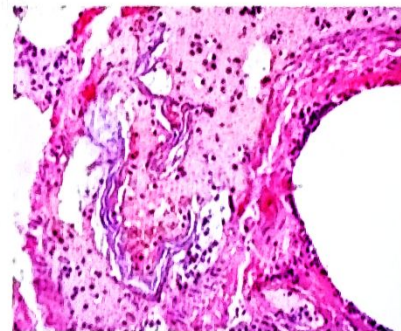
Q. What does the image below indicate :

- A. Sago spleen - Amyloidosis.
- B. Nutmeg liver - red areas are viable pericentral, white areas are periportal necrotic.
- C. Red areas are necrotic near central vein while white areas are viable, fibrotic periportal areas.
- D. Lardaceous spleen.



Q. An autopsy from the lung revealed presence of laminated swirls of squamous cells in pulmonary arteriole as shown below. What is the most likely pathogenesis?

- A. Pulmonary embolism.
- B. Air embolism.
- C. Marrow embolism.
- D. Amniotic fluid embolism.



@marroweducation6notes

NEOPLASIA BASICS

Basics of neoplasia :

- Neo means **New**.
- Plasia means **Growth**.

Neoplasm : Any new growth independent of growth factors.

Divided into two types:

- Benign tumours.
- malignant tumours.

Desmoplasia :

- **Extreme fibrosis** leading to hard tumours.
- more likely to be malignant.

Terminologies

00:05:12

1. **Benign tumours :**

- usually ends in suffix : **oma**.
- Epithelial origin : **Papilloma**.
- mesenchymal origin :

Lipoma.

Osteoma.

Chondroma.

Fibroma.

a. **malignant tumours :**

- Epithelial origin :
Called as carcinoma.
Eg : Squamous cell carcinoma.
Transitional cell carcinoma.
Adenocarcinoma.
- mesenchymal origin :
Called as sarcoma.
Eg : Osteosarcoma.
Chondrosarcoma.

Active space

Fibrosarcoma.

Leiomyosarcoma.

Exceptions :

malignant tumours ending with suffix oma :

- melanoma.
- Chloroma :

Soft tissue involvement of AML.

most common AML resulting in chloroma : AML M₂.

most common site of chloroma : Orbit.

most common presentation : Proptosis.

Also known as granulocytic sarcoma.

- Seminoma.
- Lymphoma.
- Teratoma.

3. mixed tumours :

Different germ layer derivative.

Teratoma :

Derivative of 2 or 3 germ layers.

- Benign : mature teratoma.
- malignant : Immature teratoma.
- monodermal teratoma :

Single germ layer.

Struma ovarii.

Pleomorphic adenoma :

- usually affects salivary gland.
- most commonly affected : Parotid gland.
- Biphasic tumour
- Shows two components :
 - Epithelial component : Glands.
 - mesenchymal component : Chondromixoid tissue.

Wilm's tumour :

- Triphasic tumour.
- Three components :

Epithelial component.

mesenchymal component.

Blastemal component.

4. Choriostoma :

- Ectopic rest of normal tissue.
- Normal tissue in abnormal location.
- Eg : Pancreatic tissue in the stomach.

5. Hamartoma :

- Haphazard/abnormal/disorganized proliferation of tissues indigenous to the site of origin.
- Eg : Pulmonary hamartoma :
Now considered as a benign tumour.
Some rearrangement in chromosome 12 have been identified.

Properties of a tumour

00:17:04

1. Anaplasia :

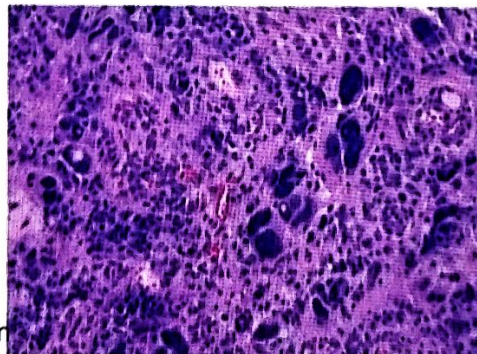
Lack of differentiation.

Differentiation : Defined as the structural and functional similarity to original cell.

- Pleomorphism : Variation in size and shape of cell.
- High Nucleus : Cytoplasmic (N/C) ratio :
Normal N/C ratio : 1:4 to 1:6.
malignant cells N/C ratio : 1:1.
- Hyperchromatic nuclei.
- Prominent nucleoli.
- Loss of polarity.
- Abnormal mitosis :

Bipolar mitosis.

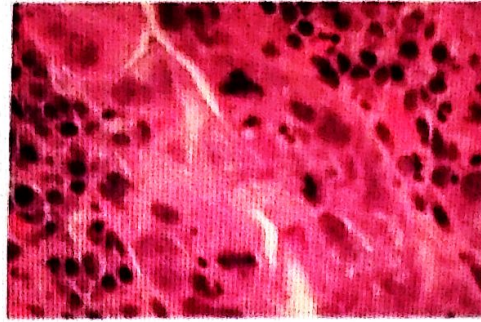
Anaplasia



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Active space

Tripolar atypical mitosis : mercedes Benz sign.



2. Rate of growth :

- Benign tumor : Slowly growing.
- malignant tumor : Rapidly growing.
- minimum tumour weight which can be clinically detected : $1\text{ g}/10^9$ cells.
- maximum tumour weight usually compatible with life : $1\text{ kg}/10^{12}$ cells.

3. Local invasion

Benign tumours :

- usually encapsulated
- No local invasion.

malignant tumours :

- Non encapsulated.
- Local invasion present.
- Eg : Carcinoma lung invading trachea, oesophagus.

4. metastasis :

- Distant spread of tumour.
- most important point to differentiate benign from malignant tumour : metastasis > local invasion.
- Hallmark of malignancy : Anaplasia.

	Benign	malignant
Anaplasia	Absent	Present
Growth	Slow	Rapid
Local invasion	Absent	Present
metastasis	Absent	present

Active space

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Two malignancies which do not metastasize :

- Basal cell carcinoma (Rodent ulcer).
- Glioma.

Routes of metastasis

00:32:25

1. Lymphatic route :

most carcinomas metastasize by this route

Exceptions :

- Follicular carcinoma thyroid.
- Choriocarcinoma.
- Hepato cellular carcinoma.
- Renal cell carcinoma.

2. Haematogenous :

most sarcomas metastasize by this route.

Exceptions :

- Synovial sarcoma.
- Rhabdomyosarcoma.

Invasive vein more than artery (Due to thinner walls of vein).

3. Direct seeding of body cavities :

- mucinous carcinoma ovary/appendix spread through peritoneum : Pseudomyxoma peritonei.

4. Transcoelomic spread :

- Krukenberg tumour :

Spread of carcinoma stomach to other organs via peritoneum, like ovary.

Terms

00:37:33

1. metaplasia :

- Reversible change.
- One differentiated cell type to another differentiated cell type.
- Examples :
Ciliated epithelium to squamous epithelium in smokers.

Barrett's oesophagus.
vitamin A deficiency.
myositis ossificans.

1. **Desmoplasia :**

- Abundant fibrosis/collagen laid down by a tissue.
- This makes the tissues hard.

2. **Anaplasia :**

- Lack of differentiation.
- Irreversible.
- Breach of basement membrane.

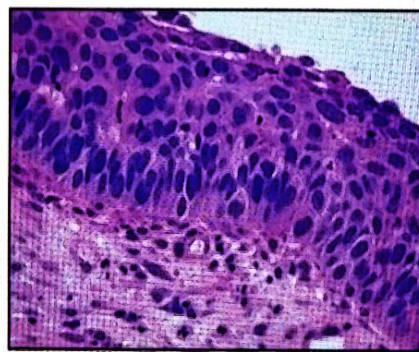
3. **Dysplasia :**

- Disordered growth/proliferation.
- Show anaplasia.
- Partially reversible.
- Example : Carcinoma in situ.

Basement membrane is intact.

malignant cells are limited by basement membrane.

PAS stain used for identification of carcinoma in situ
(stains basement membrane: magenta colored).



Cell cycle

00:44:42

Phases of cycle :

1. **G0 Phase :**

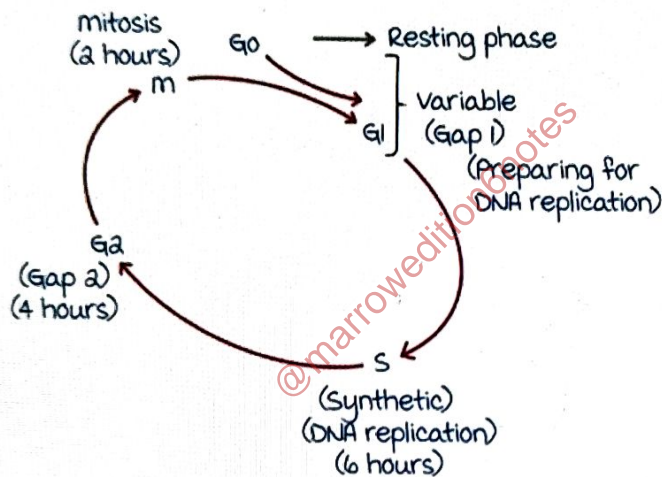
- Resting phase.

2. **G1 Phase :**

- Gap I phase.

Active space

- Prepares itself for DNA replication.
3. S Phase :
- Phase of **DNA replication**.
 - Approximately 6 hours duration.
 - Phase of **no return**.
4. G₂ Phase :
- Gap 2 phase.
 - Prepares itself for mitosis.
 - Approximately 4 hours duration.
5. m Phase :
- mitosis.



Longest phase : **G₀ or G₁ phase** (variable).

most radiosensitive phase of cell cycle : **G₂m phase**

($m > G_2$).

most radio resistant phase of cell cycle : **S phase**.

most radio sensitive cell in the body : **Lymphocyte**.

most radio resistant cell in the body : **Platelet**.

most radio sensitive tumour in the body : **Ewing's sarcoma**.

Cell cycle regulation :

1. Cyclins and CDK (cyclin dependent kinases) :

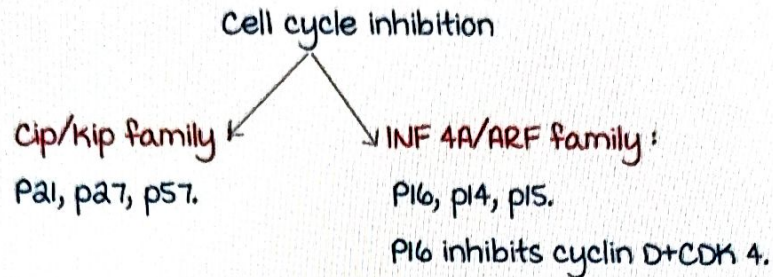
Cyclins combine with CDK and phosphorylates it, thus moving to next phase of cell cycle.

Cyclins		CDK
D	+	4, 6
E	+	2
A	+	2
B	+	1

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Cell cycle inhibition

00:52:08



Cyclin D associated with mantle cell lymphoma.

Cyclin E associated with breast cancer.

Questions :

1. An experiment is conducted in which proliferating cells are subjected to ionizing radiation. The ionizing radiation leads to arrest in a checkpoint that monitors completion of DNA replication. It is observed that there are increased numbers of chromosomal abnormalities in these cells. Which of the following is the checkpoint affected by the ionizing radiation?
 - A. G₀/G₁.
 - B. G₁/S.
 - C. S/G₂.
 - D. G₂/M (most radiosensitive phase).
 - E. M/G₀.

2. A study is performed to analyse characteristics of malignant neoplasms in biopsy specimens. The biopsies were performed on patients who had palpable mass lesions on digital rectal examination. Of the following microscopic findings which is most likely to indicate that

the neoplasm is malignant?

- A. Pleomorphism.
- B. Atypia.
- C. Invasion.
- D. Increased N: C ratio.
- E. Necrosis.

3. A 60 year old man who has a 90 pack year history of cigarette smoking has had a chronic cough for the past ten years. He has begun to lose weight (3kg) during the past year. No abnormal findings are noted on physical examination. He has a chest radiograph that reveals a right hilar mass. A sputum cytology shows atypical, hyperchromatic squamous cells. What is the most common initial pathway for metastases from this lesion?

- A. Bloodstream.
- B. Pleural cavity.
- C. Contiguous spread to chest walls
- D. Lymphatics (since epithelial malignancy).
- E. Bronchi.

4. A 62 year old man has complained of pain on urination for the past week. He is afebrile. On cystoscopy, a slightly erythematous 1cm diameter area is seen on the bladder mucosa. This area is biopsied and on microscopic examination shows cells with marked hyperchromatism and increased nuclear/cytoplasmic ratio involving the full thickness of the epithelium. However, these changes are confined to the epithelium above the basement membrane. Which of the following terms best describe these biopsy findings?

- A. metaplasia
- B. minimal dysplasia.
- C. microinvasion.
- D. Hyperplasia.
- E. Carcinoma in situ.

5. A 53 year old woman feels a lump in her right breast. Her nurse practitioner palpates an irregular 3cm mass that is not movable because it appears fixed to the overlying skin, which is retracted. A mastectomy is performed and the pathologist on sectioning the breast finds a 3 X 3.5cm ovoid mass that does not have discrete borders, but appears to infiltrate into the surrounding fibrofatty breast stroma. The mass is firm, white, and has a fibrous consistency. Which of the following features is most likely demonstrated by the gross appearance of this mass?

- A. Anaplasia
- B. Aplasia
- C. Desmoplasia
- D. Dysplasia
- E. metaplasia

@marroweditionsnotes

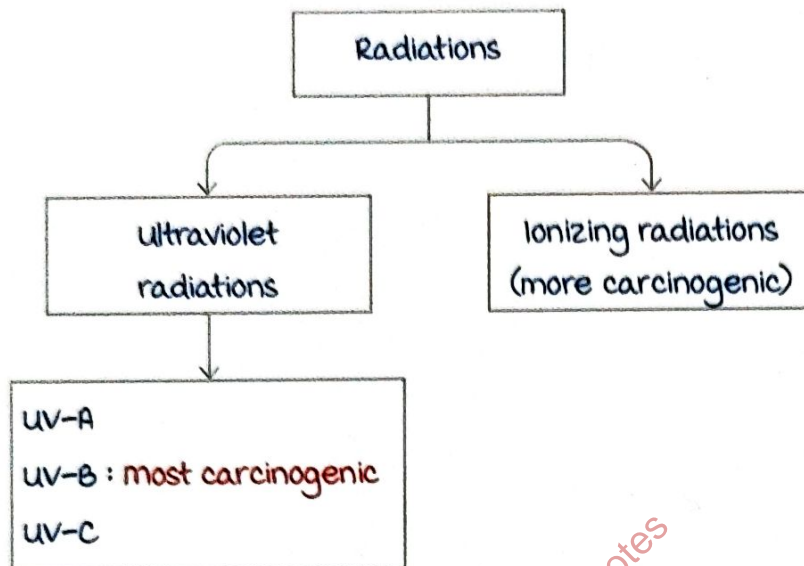
Active space

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TYPES OF CARCINOGENESIS

Radiation carcinogenesis

00:00:57



UV-B radiation causes skin cancers like basal cell carcinoma and malignant melanoma.

Theoretically, UV-C is most carcinogenic but does not reach down due to ozone layer.

most common cancer caused by ionizing radiation → acute myeloid leukemia.

Leukemia which is never caused by radiation → CLL.

Other cancers associated with radiation :

Papillary carcinoma of thyroid.

- Risk factor for follicular carcinoma thyroid is long standing goiter/iodine deficiency.

Carcinoma breast.

Carcinoma lung.

Organs most resist to radiation : Bone, gut.

Active space

Chemical carcinogenesis

00:05:19

Chemical	Cancer
Polycyclic aromatic hydrocarbons (PAH)	Lung cancer
Arsenic	Skin cancer, hepatic angiosarcoma
Asbestos	Lung adenocarcinoma → most common malignancy caused. malignant mesothelioma → most specific malignancy caused
Aflatoxin	HCC
Beta naphthylamine/ azo dyes. People working in dry cleaning industry. Those who store wool in naphthalene balls.	Bladder carcinoma
Benzene	Leukemia, AML
Diethylstilbesterol (DES)	Clear cell carcinoma vagina
Polyvinylchloride (PVC)	Hepatic angiosarcoma
Cadmium	Prostate cancer

most common lesion caused by asbestos → pleural plaque.
HPE of asbestosis : Ferruginous bodies/asbestos bodies → asbestos fibers coated with iron.

- Stained with prussian blue.
- Dumb-bell shaped/beaded/fusiform rod like structures.

marker for mesothelioma : Calretinin.

Active space

Amphibole asbestos fibers are more carcinogenic.

mothers exposed to DES → Clear cell carcinoma of vagina or cervix in daughter.

Chemical carcinogens are of two types :

Directly acting carcinogens	Indirectly acting carcinogens
Do not require any conversion	Requires conversion to active metabolites by cytochrome p450 enzymes
Less potent	more potent
e.g. : Chemotherapeutic agents, alkylating agents	e.g. : PAH

Chemical carcinogenesis takes place in 2 phases :

Initiation phase → Irreversible.

Promotion phase.

Microbial carcinogenesis

00:13:31

Parasites	Viruses	Bacteria	Fungi
Schistosoma haematobium → bladder cancer (SCC).	Hepatitis B, C → HCC. HTLV-1 → adult T-cell leukemia.	H. pylori	Aspergillus ↓ Aflatoxin ↓ HCC.
Clonorchis, Opisthorchis → cholangio carcinoma.	HHV-8. EBV. HPV.		

Schistosoma converts transitional epithelium to squamous epithelium by metaplasia → SCC.

Helicobacter pylori :

Causes the following cancers :

- Gastric adenocarcinoma → most common malignancy caused

Active space

- **MALToMa** → most **specific** malignancy caused

Gram negative bacilli.

Pathogenesis :

Two toxins produced are → **cag A** and **vac A** → causes cancer.

mostly affects the pyloric antrum of the stomach → antral biopsy.

Does not penetrate the stomach mucosa.

- Seen floating over the mucosa.

Special stains used are :

Warthin's starry silver stain → small black coloured bacilli seen floating over the mucosa.

modified Giemsa stain.

Steiner stain.

Viral carcinogenesis

00:19:43

Adult T-cell leukemia.

Caused by HTLV-1 (human T-cell lymphoma virus).

HPE : **Clover leaf cells**.

Pathogenic factor : **Tax gene**.

Diseases caused by HHV- 8 :

Primary effusion lymphoma.

Kaposi's sarcoma.

- Borderline blood vessel tumor (vascular tumor).
- usually seen in HIV positive patients or those with immunodeficiency.
- On microscopy : **Spindle shaped cells** with slit like spaces.

multicentric Castleman's disease.

Ebstein Barr virus (EBV) :

Binds to CD21 receptor on B-cell.

Diseases caused by EBV :

- Infectious mononucleosis/kissing disease.

Downey cells → ballerina skirt appearance.

- Hodgkin's lymphoma.
Nodular lymphocyte predominant > nodular sclerosis variant : Not associated with EBV.
- B- cell lymphomas.
- Burkitt's lymphoma.
Translocation t(8: 14) → amplification of c-myc.
marker : bcl- 6.
microscopic appearance : Starry sky appearance.
- Leiomyosarcoma.
- Post-transplant lymphoproliferative disorder.
- Nasopharyngeal carcinoma.

Pathogenesis of EBV :

LMP-1 : Latent membrane Protein- 1.

Increased activation of NF- κβ pathway → increased growth signaling → increased cell proliferation.

EBNA-2.

IL-10.

Human papilloma virus (HPV) :

Strains of HPV are :

Low risk	High risk
HPV 1, 2, 4, 6, 11	HPV 16, 18
Genital warts. CIN I.	CIN II, III. Cervical cancer. Penile cancer. Laryngeal cancer.

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Pathogenesis of HPV :

Produces two proteins E6 and E7.

E6 combines with p53 (tumor suppressor gene) and degrades it.

E7 combines with Rb and causes its degradation.

Both causes increased cell proliferation.

Active space

HPE :

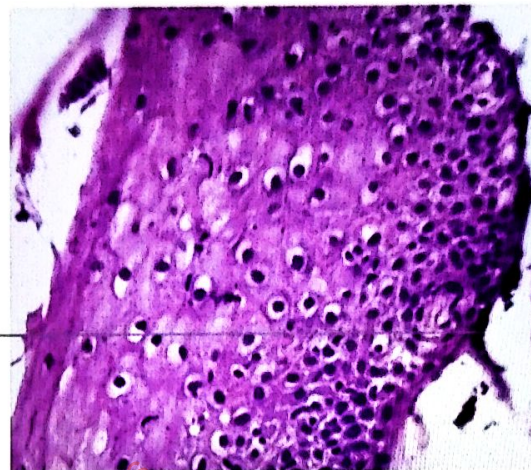
Koilocytes → large cell with thick membrane and raisinoid nucleus, with perinuclear halo.

Produced by E4 protein.



Thick membrane
Raisinoid nucleus
Peri- nuclear halo

Koilocyte



Koilocyte ←

microbe	Cancer
H. Pylori	Gastric adenocarcinoma. MALTOMA.
HPV	Cervical cancer. Anogenital cancer. Laryngeal cancer.
HTLV I	Adult T cell leukemia
HBV	Hepatocellular carcinoma
HHV 8	Kaposi's sarcoma. Primary effusion lymphoma multicentric Castleman disease..
EBV	HL. NHL. Burkitt's lymphoma. Nasopharyngeal carcinoma. PTLD.

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Active space

Q. A 51 year old man has worked for 10 years in a factory producing plastic pipe but not following safety standards. He has noted weight loss, nausea, and vomiting worsening over the past 5 months. On examination he is afebrile. There is generalized muscle wasting. Laboratory studies show the serum alkaline phosphatase is 405 U/L with AST 47 U/L, ALT 35 U/L, and total bilirubin 1.2 mg/dL. An abdominal CT scan reveals a 12 cm right liver lobe mass. Liver biopsy reveals a neoplasm composed of spindle cells forming irregular vascular channels. With immunohistochemical staining the cells demonstrate vimentin positivity and cytokeratin negativity. Exposure to which of the following substances most likely led to development of this neoplasm?

- A. Benzene.
- B. Radon.
- C. Cyclophosphamide.
- D. Asbestos.
- E. Vinyl chloride.

Q. Pathogenic mechanism of HPV in cervical cancer is :

- A. Degradation of cyclin D1.
- B. upregulation of BCL2.
- C. Instability of E6 and E7.
- D. Down regulation of p16INK4a.

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Q. A clinical study is performed of oncogenesis in human neoplasms. It is observed that some neoplasms appear to develop from viral oncogenesis, with serologic confirmation of past viral infection. Which of the following neoplasms is most likely to arise in this manner?

- A. Retinoblastoma.
- B. Small cell anaplastic carcinoma.
- C. T-cell leukemia.
- D. Prostatic adenocarcinoma.
- E. Hepatic angiosarcoma.

Active space

Q. HHV 8 is related to all except :

- A. Kaposi's sarcoma.
- B. Primary effusion lymphoma.
- C. Adult T cell lymphoma.
- D. Castleman's disease.

Q. A 56 year old man has had a chronic cough for the past year. He is a non-smoker. He had an episode of hemoptysis 3 days ago. No abnormal findings are noted on physical examination. A chest X-ray demonstrates a 6 cm perihilar mass. A sputum sample is collected, and the sputum cytology report reads, Atypical cells present suggestive of squamous cell carcinoma. Which of the following environmental exposures is most likely to be associated with these findings ?

- A. Asbestos.
- B. Radon.
- C. Silica.
- D. Benzene.
- E. Formaldehyde.

Q. A previously healthy 42 year old man has a skin nodule on his right hand that has become larger and darker with more irregular outlines over the past 3 months. On physical examination this lesion is 1.2 cm diameter, darkly pigmented, and a slightly raised nodule on the dorsum of his right hand. No other skin lesions are noted. Three non tender enlarged lymph nodes are palpable in the right axilla. The lesion is excised and microscopic examination shows a neoplasm composed of darkly pigmented polygonal and spindle cells. Which of the following risk factors is most important for development of this neoplasm ?

- A. Cigarette smoking.
- B. Allergy to latex gloves.
- C. Inheritance of a faulty RB gene.
- D. Prior job-related handling of asbestos.
- E. Chronic exposure to ultraviolet radiation.

HALLMARKS OF NEOPLASIA

Hallmarks of carcinogenesis

00:01:51

1. Self sufficiency in growth signals.
2. Insensitivity to growth inhibitory signals.
3. Limitless replicative potential.
4. Evasion of apoptosis.
5. Sustained angiogenesis.
6. Altered cellular metabolism.
7. Invasion and metastasis.
8. Escape of immune recognition.

Self sufficiency in growth signals

00:03:08

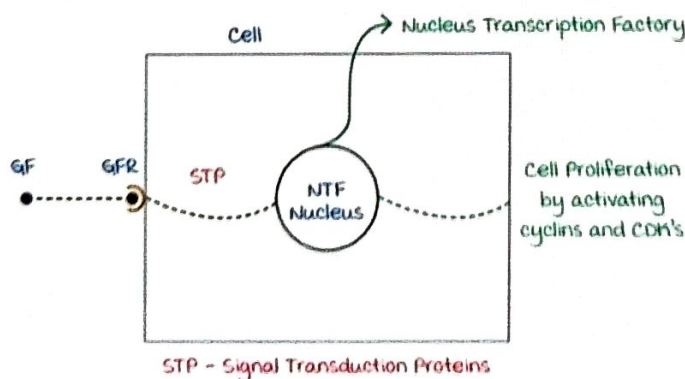
Proliferation without growth factors.

Proto-oncogenes : Normal genes which are required for cell proliferations.

Mutation in these genes converts proto-oncogenes into **oncogenes** leading to production of cancer.

This mutation is called gain of function mutation.

- Every cell requires growth factor for proliferation.
- Growth factor combines with cell through growth factor receptor on the surface of the cell.
- Growth factor after combining with growth factor receptor enters the cell through signal transduction protein (STP).
- STP transmits signals to nucleus.
- Nuclear transcription factors are present inside the cell which leads to cell proliferation by activating cyclins and cyclin dependent kinases.



Active space

Any defect in growth factor/ growth factor receptor/ STP/ NTF/ cyclins → Cancer.

Growth factors	Cancer
Hepatocyte growth factor	Hepatocellular carcinoma.
HST - 1	Osteosarcoma.
PDGF - β / sis	Increased risk of astrocytoma.
Growth factor receptors	Cancer
EGFR - 1 (ERB B 1)	Lung adenocarcinoma.
EGFR - 2 (ERB B 2 / Her 2 neu)	Breast and ovarian cancer.
ALK gene on chromosome 2	Anaplastic large cell lymphoma. Inflammatory myofibroblastic tumor. Adenocarcinoma of lung.
C-Kit	Gastrointestinal stromal tumor and seminoma.
RET on chromosome 10. (Gain of function mutation).	Increased risk of medullary carcinoma of thyroid and MEN II syndrome.

Loss of function mutation in RET : **Hirschsprung disease.**

Signal transduction proteins :

1) RAS : **m/c oncogene** affected in human malignancy.

(**m/c gene** affected in human malignancy : p53).

RAS is divided into 3 types : K-RAS, H-RAS and N-RAS.

K-RAS : Increased risk of **KPL** tumors.

K : Colon cancer.

P : Pancreatic cancers.

L : Lung cancer.

H-RAS : Bladder cancer.

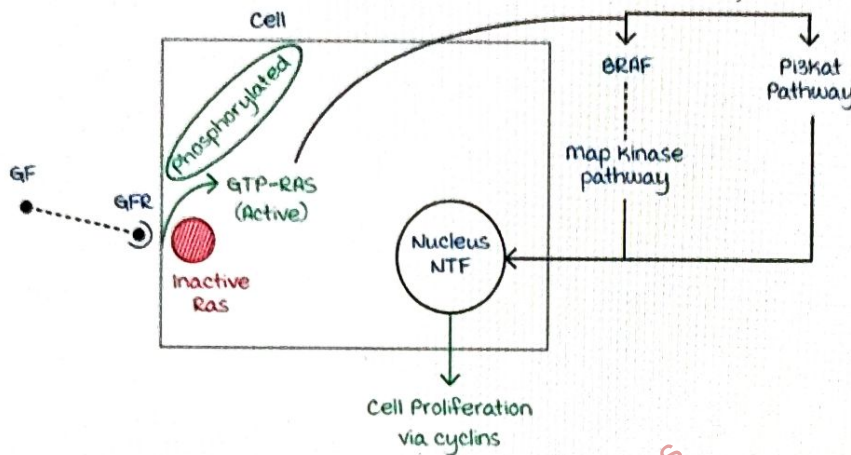
N-RAS : melanoma.

- RAS inside the cell is inactive due to combination of GDP.
- GF (signal) binds with GFR enters the cell & phosphorylates RAS.
- Phosphorylated RAS is a GTP RAS (active), it is a proto oncogene.

Active space

- GTP RAS activates 2 pathways :
 1. BRAF Kinase MAP pathway.
 2. PI3KAT pathway.
- Through these pathway signals are sent to nucleus containing NTF.
- NTF are activated leading to cyclin activation resulting in cell proliferation.

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Any defect in RAS/BRAF/PI3KAT → Cancer.

BRAF is affected in :

- Hairy cell leukemia (HCL)
- Langerhans cell histiocytosis (LCH).
- melanoma.
- Pilocytic astrocytoma.
- Papillary carcinoma of thyroid.
- Colon cancer.

NOTCH gene mutation implicated in T-ALL.

ABL gene mutation : t(9 : 22) : CML.

1. If products are 210 Kda (Kilo dalton) : CML.
2. If products are less than 190 Kda : ALL.

Nuclear transcription factors :

myc is a oncogene.

3 types :

N-myc : Amplification leads to neuroblastoma.

L-myc : Lung cancer (small cell).

C-myc : Amplified in Burkitt's lymphoma.

Cyclins and CDK's :

t(11 : 14) : mantle cell lymphoma.

On chromosome 11 : Cyclin D 1.

On chromosome 14 : Ig H locus.

Translocation causes over expression of cyclin D 1 leading to increased cell proliferation resulting in mantle cell lymphoma.
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Oncogenes in human malignancy :

Gene	Cancer
C KIT	GIST
RET	medullary carcinoma of thyroid and MEN II.
ALK	ALCL, Adenocarcinoma of lung and inflammatory myelofibroblastic tumor.
ABL	CML
K RAS	Colon and pancreatic cancer
H RAS	Bladder cancer
N RAS	melanoma
C myc	Burkitts lymphoma.
L myc	Small cell lung cancer
N myc	Neuroblastoma
NOTCH	ALL-T

Insensitivity to growth inhibitory signals

00:28:46

Tumor suppressor genes :

Normal genes which decreases cell proliferation.

Loss of function mutation leads to increased cell proliferation and cancer.

1) RB gene :

Located on chromosome 13q 14.

Mutation of RB leads to increased risk of retinoblastoma and osteosarcoma.

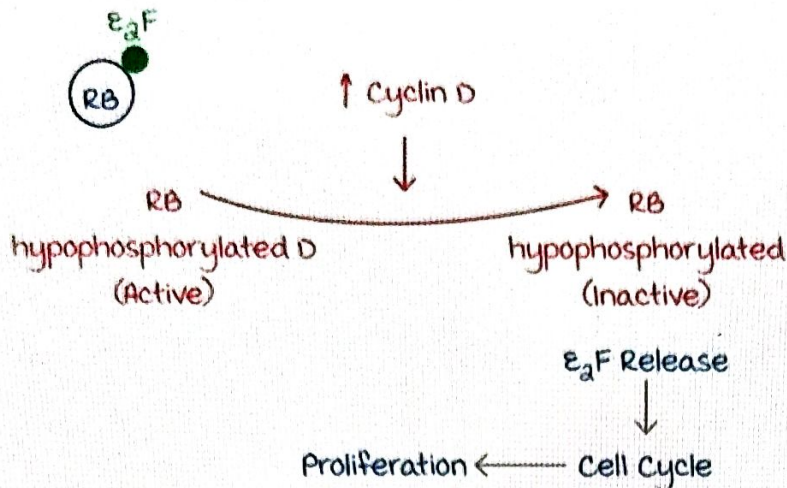
Governor of genome.

RB hypophosphorylated : Active RB.

RB hyperphosphorylated : Inactive RB.

Active space

Role of RB in cell cycle :



RB is active in hypophosphorylated state.

RB has E₂F transcription factor in its pocket.

Increase in cyclin D/CDK 4 inactivates the activated RB.

Due to release of E₂F transcription factor (required by the cell for cell cycle), cell cycle is progressed leading to proliferation.

Role of RB is that it regulates G₁ S checkpoint of cell cycle. Hence called as governor of cell cycle.

Knudson's two hit hypothesis:

First described for retinoblastoma.

Both alleles are defective.

For retinoblastoma to develop, both the alleles have to be mutated.

Loss of heterozygosity :

To develop retinoblastoma, 1st mutation has to occur by birth.

But the disease does not present by birth.

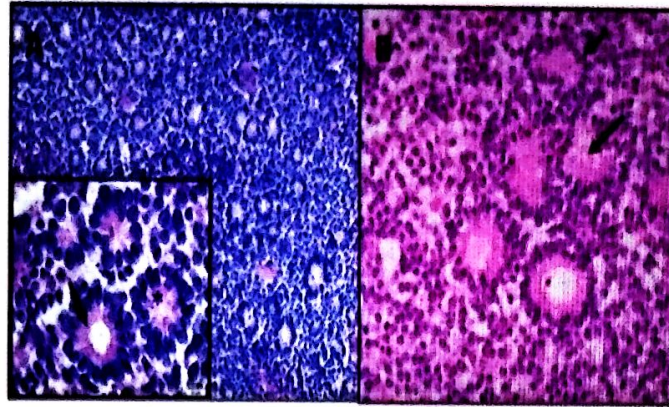
2nd mutation acquire later on and and develop retinoblastoma.

This is known as **loss of heterozygosity**.

HPE of retinoblastoma : Small round blue cells with scanty cytoplasm.

Flexner Wintersteiner rosettes (it is a true rosette : Central space is empty).

Fleurettes is also seen.



P53 gene :

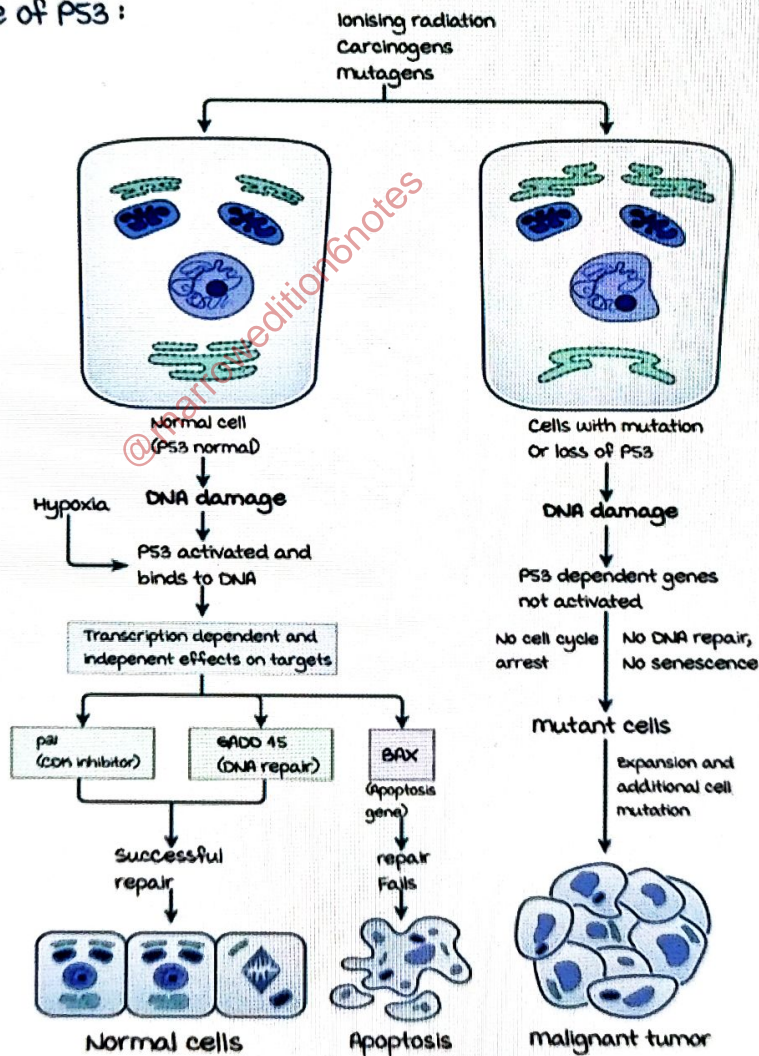
Chromosome 17p.

m/c affected gene in human malignancy (> 50% of cancers).

Guardian of genome or molecular policeman of cells.

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Role of P53 :



Active space

In case of DNA damage :

P53 is activated, which inturn activates p21 causing stoppage of cell cycle.

P53 will try to repair DNA by activating GADD 45.

In stubborn cells where those two mechanisms don't work, p53 activates BAX causing apoptosis.

Loss of p53: mutant cells are produced due to lack of repair, cell cycle arrest or senescence.

In congenital mutation of p53: Li Fraumeni syndrome. Many cancers can develop in this syndrome such as:

4 B's:

Bone cancer.

Breast cancer.

Blood cancer.

Brain cancer.

p63: Helps in squamous cell differentiation.
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TSG's:

Gene	Chromosome	Tumors
RB	13q	Retinoblastoma and osteosarcoma.
p53	17p	Li Fraumeni syndrome
NF1	17	Neurofibroma and meningiomas
NF2	22	Schwannoma
BRCA1	17	Breast cancer and ovarian cancer
BRCA2	13	male breast cancer, female breast cancer and prostate cancer.
WT1	11	Wilms tumor
WT2	11	Wilms tumor
PTEN	10	Endometrial cancer and prostate cancer, Cowden syndrome.
VHL	3	Clear cell RCC, Cerebellar hemangioblastoma.
APC	5	FAP

Limitless replicative potential

00:50:20

Telomerase:

maximum telomerase activity present in cancer cells.

Elaboration of telomerase: No telomere shortening and cells won't die.

Active space

Evasion of apoptosis

00:51:26

Increased synthesis of antiapoptotic factor :

t(14 : 18) : Follicular lymphoma.

Chromosome 18 : Ig H locus.

Chromosome 14 : bcl 2.

Due to translocation : Increased activity of bcl2 (anti apoptotic gene) leading to decrease apoptosis resulting in increased cell proliferation → Follicular lymphoma.

Sustained angiogenesis

00:53:04

Increase secretion of :

Proangiogenic factors and Anti angiogenic factors.

Cancer cells increase pro angiogenic factors (increase blood supply) like :

Vascular endothelial growth factor (VEGF).

Platelet derived growth factor (PDGF).

Fibroblast growth factor (FGF).

Anti angiogenic factors (decrease blood supply) like :

Vasculostatin.

Endostatin.

Angiostatin.

Thrombospondin.

Altered cellular metabolism

00:54:55

Warburg effect :

Sir Otto warburg : Nobel prize for discovering this effect.

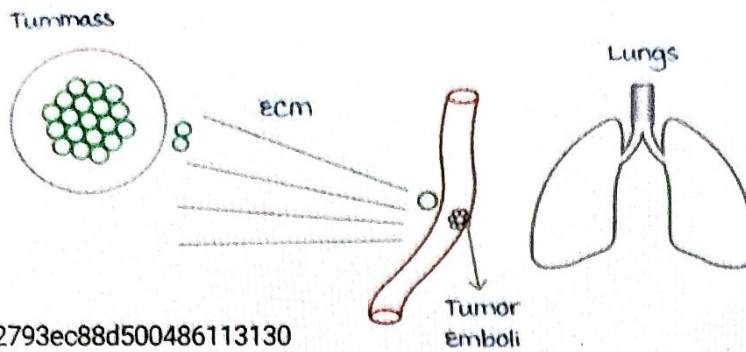
Cancer cells undergoes aerobic glycolysis.

PET scan is based on this warburg effect.

Invasion and metastasis

00:56:06

Tumor has to cross extra cellular matrix to reach the lung :



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All tumor cells are joined by E-cadherin.

1. Detachment of cells by **loss of E-cadherin**.
2. These detached cells attaches to extracellular matrix by **integrin**.
3. Degrades extracellular matrix by : **matrix metallo proteinases (2,9)**.

Can express type IV collagen, cathepsins, urokinase.

Elaborate collagenase and enters blood vessels.

4. Epithelial to mesenchymal transition (EMT).

mediated by **Snail & Twist**.

5. Tumor cells attaches with platelets is called as tumor emboli → either reach lymphatics or organs.

(Hematogenous route common for sarcomas.

Lymphatic route common for carcinomas.

Except HCC & RCC : Hematogenous route.)

Escape of immune recognition

01:04:10

Decreased expression of MHC antigens.

Selective outgrowth of antigen negative variants.

Repair pathway defects

01:05:29

3 pathways :

1. Defect in nucleotide excision repair : Xeroderma pigmentosa.
2. Defect in mismatch repair : HNPCC.
3. Homologous recombination :
 - a. Fanconi anemia.
 - b. Ataxia telangectasia.
 - c. Bloom syndrome.

Q. A change in bowel habits prompts a 53 year old woman to see her physician. On physical examination there are no lesions noted on digital rectal examination, but her stool is positive for occult blood. A colonoscopy is performed and reveals a 6 cm friable exophytic mass in the caecum. A biopsy of this mass is performed and microscopic examination shows a moderately differentiated adenocarcinoma. Which of the following laboratory findings is most likely to be present in this patient?

- A. K-RAS mutation in the neoplastic cells.
- B. Neoplastic cells positive for vimentin.
- C. Stool culture with *Shigella flexneri*.
- D. Presence of HIV-1 RNA.
- E. DNA topoisomerase I autoantibody.

Q. A 52 year old man has had increasing fatigue for the past 6 months. On physical examination he has a palpable spleen tip. Laboratory studies show a WBC count of 189,000/microliter. The peripheral blood smear shows many mature and immature myeloid cells present. Cytogenetic analysis of cells obtained via bone marrow aspiration reveals a t(9:22) translocation. This translocation leads to formation of a hybrid gene that greatly increases tyrosine kinase activity. Which of the following genes is most likely translocated to cause these findings?

- A. p53.
- B. RB.
- C. ABL.
- D. NF-1.
- E. RAS.

Q. Mother of a 4 year old boy notices that his abdomen is enlarged. Physical examination shows an ill defined abdominal mass. An abdominal CT shows a 9 cms mass in the region of right adrenal gland. The mass is removed and microscopic appearance shows small blue cells with Homer wright rosettes. Which of the following genes is most likely to have undergone alterations to produce these findings?

- A. K RAS.

- B. BCL2.
- C. N myc.
- D. P53.

Q. A 64 year old man has noted a 5 kg weight loss along with increasing fatigue over the past year. He has experienced dull abdominal pain for the past week. He has developed abdominal distention with lack of stools in the past two days. On physical examination, bowel sounds are reduced. An abdominal CT scan reveals a mass involving the descending colon. At laparotomy, a partial resection of the left colon is performed, with removal of an encircling mass. Microscopically, the mass is found to be a moderately differentiated adenocarcinoma. Which of the following laboratory test findings is most likely to be present in this man?

- A. microcytic hypochromic anemia.
- B. Positive antinuclear antibody test.
- C. Hyperglycemia.
- D. Elevated alpha-fetoprotein.
- E. Lactate dehydrogenase.

Q. In an experiment, it is observed that chronic, increased exposure to ionizing radiation results in damage to cellular DNA. As a consequence, a protein is now absent that would arrest the cell in the G1 phase of the cell cycle. Subsequent to this, the cell is transformed to acquire the property of unregulated growth. The absent protein is most likely the product of which of the following genes?

- A. RAS.
- B. TP53.
- C. MYC.
- D. ABL.
- E. BCL-2.

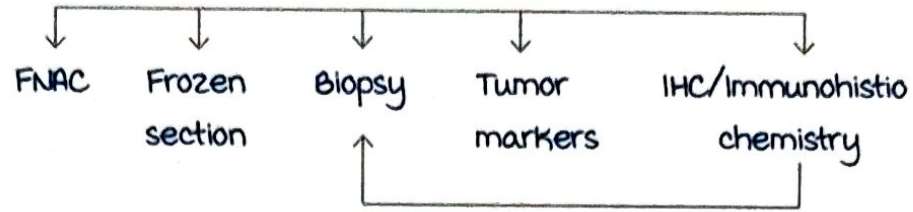
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Active space

LAB DIAGNOSIS OF CANCERS

Investigations

00:00:45



Fine Needle Aspiration Cytology (FNAC)

00:02:45

23-30 G small bore needle.

Technique : The needle is pierced into the swelling and aspirated. The sample is placed on a slide, stained and visualized under a microscope.

used for easily accessible organs : Lymph node, breast, thyroid.

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FNAC of lymph nodes is usually done if TB is suspected.

If the material taken in the syringe can indicate the condition of the patient.

If the material is cheesy, caseous or yellow in colour : AFB stain must be done.

Advantage of the procedure : Less invasive.

Disadvantage of the procedure : Target is missed in the procedure resulting in false negative reports.

Sample of thyroid is contaminated with blood with only few cells left for FNAC, as it vascular organ.

Fine needle non aspiration cytology (FNAC) :

As we do not aspirate, this technique can be used for highly vascular organs like thyroid.

PAP smears / Exfoliative cytology :

Lung cancers : Broncho alveolar lavage specimen.

CIN / cervical intraepithelial neoplasia : PAP smear.

Image guided FNAC :

On ultrasound guided FNAC : lesion is localized hence target is not missed. ↑ Sensitivity of FNAC.

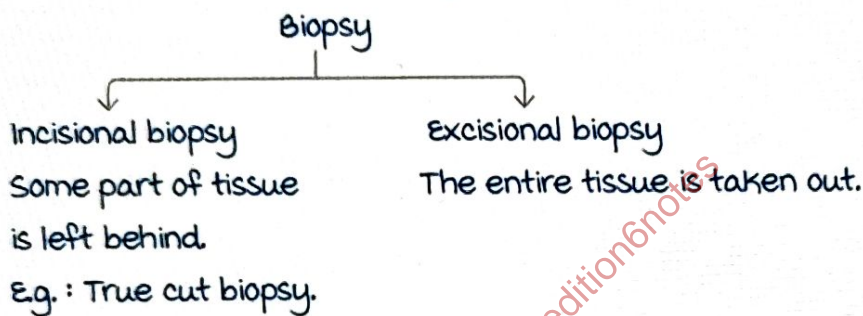
For very small or deeper lesions.

Follicular carcinoma of thyroid cannot be diagnosed by FNAC. Because on FNAC, Follicular Adenoma and follicular carcinoma cannot be differentiated because capsular and vascular invasion is not visualized.

Biopsy

00:09:20

To be done to confirm the diagnosis based on FNAC.



True cut biopsy :

An incision is made (in breast : Around the areola) and true cut biopsy gun is inserted through the incision.

An ultrasound probe can be used for better localization.

Once localized the biopsy gun is triggered to cut and remove the tissue.

Useful for breast cancer and soft tissue lesions

Fixatives :

Biopsy specimen tissue is fixed most commonly 10% neutral buff formalin for histopathology.

For electron microscope, fixative used is 2.5% glutaraldehyde.

For testicular biopsy : Bowin's fluid (sperms are destroyed by formalin).

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Active space

Stains used in Pathology :

Cell/ condition	Stain
most common stain in histopathology	Hematoxylin and eosin
most common in hematology	Romanowsk like Leishman/ Geimsa
Reticulocyte	Supravital
Lymphoblast	PAS
myeloblast	NSE, SBB, OIL RED O
monoblast	Non specific esterase /NSE
Hairy cell	TRAP
Lipid	Oil red O, sudan black
Iron	Prussian blue
calcium	Von Kossa, Alzarine red S
Glycogen	PAS
Copper	Rhoamine, rubeanic acid
mast cell	Toluidine blue
mucin	mucicarmine Alcian blue
Reticulin fibres	Silver
Elastin fibres	Van geison
Collagen	masson trichrome
melanin	masson fontanna
H. pylori	Warthin starry silver
Cryptococcus	India ink
Fungi	Silver methanamine PAS
Amyloid	Congo red

Immunohistochemistry / IHC

00:15:40

On biopsy specimen, antigen antibody reaction is done and visualised.

Active space

Uses of IHC :

- Origin of the tumor.

E.g. : In a poorly differentiated tumor to differentiate epithelial, mesenchymal and vascular origin, all 3 markers are added. The positive marker gives the origin of the tumor.

- In diagnosis of unknown primary.

- Prognostic and therapeutic significance.

E.g. : Especially in breast cancer cases, 3 markers (ER, PR & HER 2 neu) are added :

Estrogen receptor/ER	}	+ : Good prognosis. Treated by Tamoxifen.
Progesterone receptor/PR		

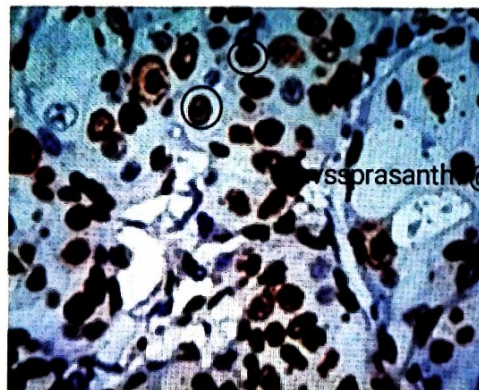
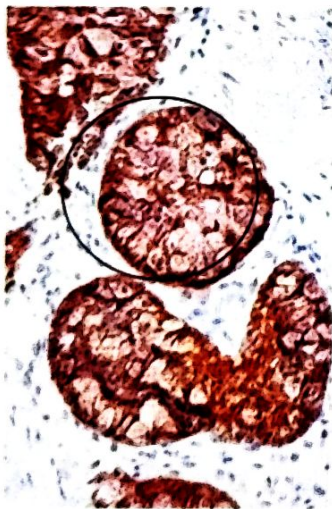
Her2 neu + : Poor prognosis.

Treated by Trastuzumab, Herceptin.

Technique :

Certain cancers have particular cell types, with specific antigens.

E.g. : If epithelial malignancy is suspected, they have Cytokeratin antigen. Cytokeratin antibodies are added, if they react with the antigen, a positive reaction gives a color. No colour is produced in the absence of epithelial malignancy. If IHC is positive, usually gives a brown color.



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Brown color indicates + IHC.

Active space

Important IHC markers applied on Biopsy :

Cell of origin/tumor	marker
Epithelial origin	Cytokeratin
mesenchymal origin (Sarcomas)	Vimentin
Glial	GFAP / Glial fibrillary acid protein
Smooth muscle (e.g. : Leiomyosarcoma)	SMA / Smooth muscle actin
Skeletal muscle / Rhabdomyosarcoma	Desmin, myogenin, myo D1
Vascular e.g : Angiosarcoma.	Vwf, CD31, VEGF, Factor VIII.
Neuroendocrine e.g. : Pheochromocytoma, Neuroblastoma, medullary carcinoma of thyroid, Paraganglioma.	NSE Chromogranin Synaptophysin
Hepatic e.g : Hepatocellular carcinoma	Hep par1, arginase 3 Alpha feto protein / AFP
GIST / Gastro Intestinal Stromal Tumor	DOG1, CD 34, CD117
malignant melanoma	Hmb 45, S 100, melan A
malignant mesothelioma	Calretinin, CK 5/6
Ewings sarcoma	CD 99, MIC 2
Osteosarcoma	Osteopontin, osteonectin, osteocalcin

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Tumor markers are released in the blood :

marker	Condition
PSA / Prostate Specific antigen	Prostate ca
PAP	Prostate ca
Calcitonin	medullary ca thyroid
PSA & PAP are organ specific but not cancer specific.	
CEA	Colon ca, pancreatic ca
HCG	Chorio ca

marker	Condition
AFP/ Alpha fetoprotein	Hepatocellular ca, NSGCT Non seminomatous germ cell tumor like yolk sac tumor, Hepatoblastoma
Immunoglobulins	Multiple myeloma
Carbohydrate Antigen/CA 19-9	Colon ca, pancreatic ca
CA 125	Ovarian ca
CA 15-3	Breast ca
Catecholamines	Pheochromocytoma

Clinical scenario :

A 3 year old child presents with a testicular mass, microscopy showed presence of Schiller Duval bodies. What is the diagnosis and which Tumor marker used is ?

Schiller Duval bodies are seen in Yolk sac tumor (testicular tumor). Tumor marker : Alpha fetoprotein.

markers for unknown primary : CK7/CK20 profile

CK 7+/CK20+ : Bladder Ca, Stomach, pancreas.

CK 7-/CK 20- : Hepatocellular carcinoma.

Renal cell carcinoma.

CK 7+/CK20- : Cancers of female genital tract, breast, cervix, endometrium, lung and thyroid.

CK 7-/CK20+ : Colorectal Cancer.

Frozen section

00:29:28

It is an intra-surgical procedure.

Done during lumpectomy, to check if the margins are involved or not.

The sample is taken and margins are labelled & sent for histopathology lab.

Quick procedure compared to normal processing of the tissue.

Stain used : Oil red O.



Active space

Paraneoplastic syndromes

00:31:25

Are symptom complexes in cancer patients which cannot be explained by the local or indigenous spread of tumor or by the elaboration of the hormones.

Syndrome	Tumor	Substance
SIADH / Syndrome of inappropriate secretion of ADH	Small cell ca lung	ADH
Cushing's syndrome	Small cell ca lung	ACTH
Hypercalcemia	SCC lung Breast Ca	PTHrP/ Parathyroid hormone related peptide.
Polycythemia	RCC	Erythropoietin
migratory thrombophlebitis	Ca pancreas Ca colon	
Hypertrophic pulmonary osteoarthropathy	Small cell ca lung	
Acanthosis nigricans (velvety thickening)	Ca stomach Ca colon	Epidermal growth factor
myasthenia garvis	Thymoma Ca lung	

most common paraneoplastic syndrome : Hypercalcemia.

most common endocrinopathy : Cushing syndrome.

Tumor producing maximum paraneoplastic syndromes :

Small cell carcinoma of lung.

Clinical scenario :

60 year old male smoker presents with a centrally located mass in the lung. He has moon like face and striae on the body. The histopathology image is given. what is the marker to be used?

microscopy showed Small cell carcinoma of the lung.
marker used due to its neuroendocrine origin :
NSE, chromogranin & synaptophysin.

Cancer cachexia :

Loss of body fat, muscle mass, anorexia.
TNF- α is the responsible cytokine.

Tumor lysis syndrome :

Seen in rapidly proliferating tumors (Burkitt lymphoma).
Results in : **Hyperuricemia, hyperkalemia, hypocalcemia.**

Recent updates :

Role of microRNAs in cancer :

- Oncogenic micro RNA 155, 200 : Seen in B cell lymphomas.
- Tumor suppressor micro RNA 15, 16 : Detected in CLL.

DICER mutation : Seen in cancers of female genital tract.

Q. A 62 year old man with an 80 pack year history of smoking experiences an episode of hemoptysis. On physical examination, he has puffiness as well as plethora of the face, pedal edema, bruises of the skin, & a blood pressure of 165/100 mm Hg. A chest radiograph reveals a 5 cm right upper lobe lung mass. A fine needle aspirate of this mass yields cells consistent with small cell anaplastic lung carcinoma. A bone scan shows no metastases. Immunohistochemical staining of the tumor cells is likely to be positive for which of the following hormones?

- Parathormone related peptide.
- Erythropoietin.
- ACTH.
- Insulin.
- Gastrin.

Patient has small cell carcinoma with Cushing's syndrome as a paraneoplastic syndrome, it shows raised ACTH.

Q. Which of the following markers is used for the diagnosis of Rhabdomyosarcoma?

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Active space

- A. Desmin.
- B. Cytokeratin.
- C. Myeloperoxidase.
- D. Synaptophysin.

Q. Which of the following IHC markers is positive in a neuroendocrine tumor?

- A. Cytokeratin.
- B. Calretinin.
- C. GFAP.
- D. Synaptophysin.

Q. A 5 year old child who presented with proptosis of one of the eyes was found to have a desmin positive tumor. What is the probable diagnosis?

- A. Ewings sarcoma.
- B. Embryonal rhabdomyosarcoma.
- C. Leukemia.
- D. Retinoblastoma.

Ewings sarcoma : CD99, mic 2. Retinoblastoma : RB.

Q. A 49 year old man complains of pain in his left thigh for 3 months. On physical examination his thigh is increased in size, compared to the right. A plain film radiograph reveals the presence of a 15 cm solid mass that does not appear to arise from bone, but it does have infiltrative margins. A biopsy of this mass is taken, and on microscopic examination the mass is composed of highly pleomorphic spindle cells. Which of the following immunohistochemical markers is most likely to be demonstrated in the cells of this mass?

- A. Cytokeratin.
- B. Factor VIII.
- C. Alpha fetoprotein.
- D. Lambda light chain.
- E. Vimentin.

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Suspected : Leiomyosarcoma (non bony origin in thigh).

TRICKS TO DIAGNOSE TUMORS

Squamous cell carcinoma :

Identified by :

Desmosomes (in HPF).

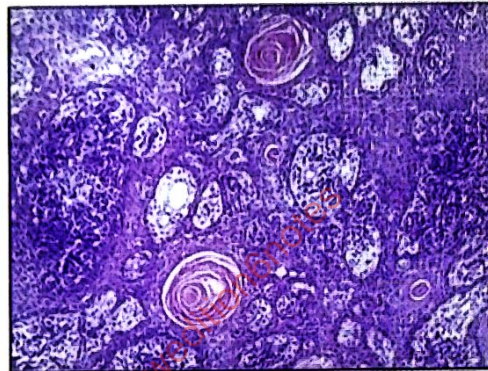
Keratin pearls.

marker : Being an epithelial malignancy.

- Cytokeratin.
- p63.



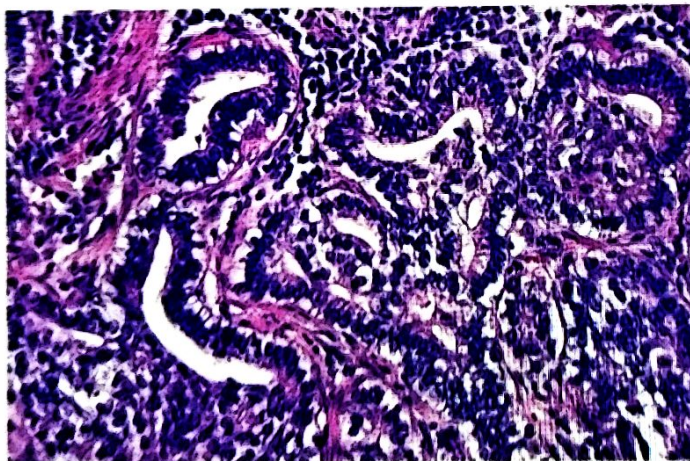
Stratified squamous
epithelium



Keratin pearls

Adenocarcinoma :

Glands with the lumen inside lined by pleomorphic cells.



Papillary tumor :

Identified by :

- Papillae : Finger like projections with fibrovascular core.

Exception :

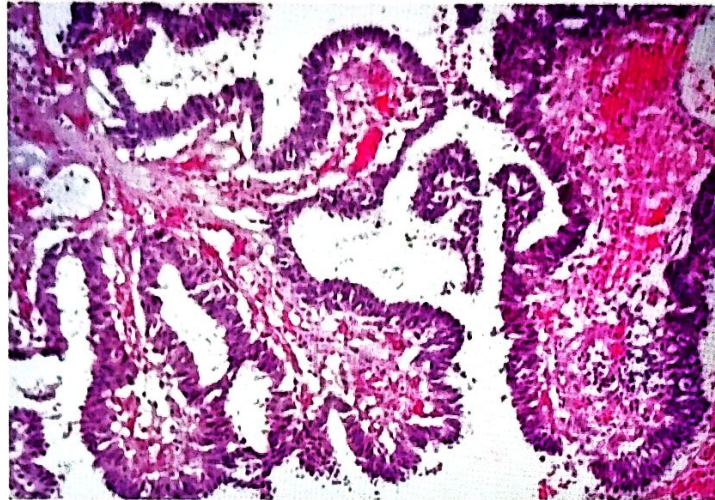
In papillary RCC, no fibrovascular core (foamy histiocytes +nt).

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Active space

In papillary carcinoma, thyroid papillae are lined by Orphan Annie eye nuclei that are optically clear nuclei.

- Psammoma bodies :
Foci of dystrophic calcification.
Basophilic : Dense blue colour.



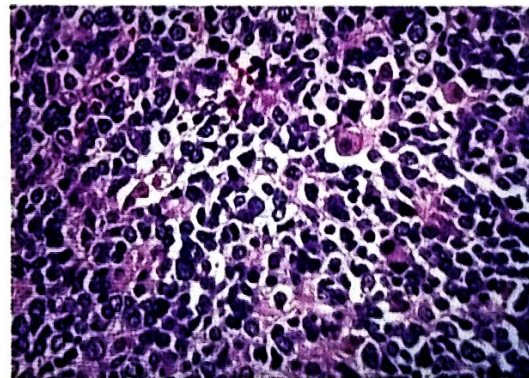
Neuroendocrine tumour :
History of diarrhea, flushing.
Identified by : Cells with salt and pepper chromatin.
Cells are usually arranged in nests.

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- Commonly seen in :
- Small cell carcinoma of lung.
 - Pheochromocytoma.
 - Carotid body tumor.
 - Paraganglioma.
 - Neuroblastoma.

Positive IHC marker :

- NSE.
- Synaptophysin.
- Chromogranin.



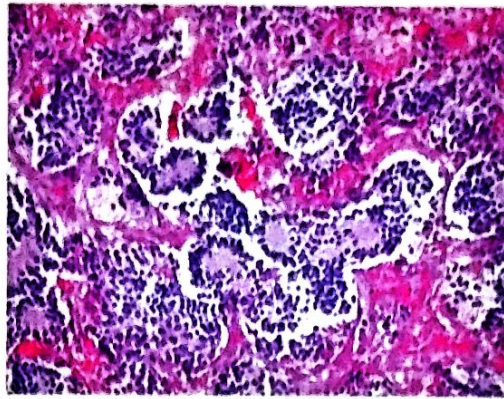
In electron microscopy, shows :
Dense, core Neurosecretory granules.

Rosette :
Purple coloured tumor cells arranged like flower.
Sheets of cells with scanty cytoplasm.
Seen in round small blue cell tumours of childhood.

Active space

E.g.:

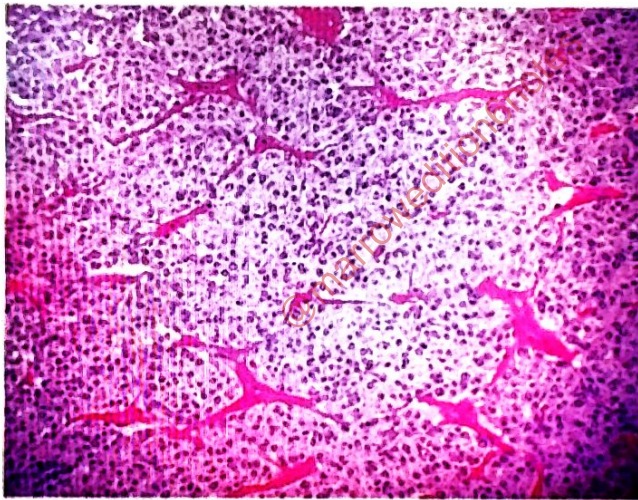
- Neuroblastoma.
- Retinoblastoma.
- Hepatoblastoma.
- Lymphoma.
- medulloblastoma.
- Rhabdomyosarcoma.
- Ewing Sarcoma/PNET.
- Nephroblastoma/ Wilm's Tumor.



Pheochromocytoma :

A 30 year old male with episodic hypertension, palpitations, headache. CT scan shows a mass in adrenal. What is the most likely diagnosis ?

A. Pheochromocytoma



On microscopy shows Zell Ballen pattern (pink colour cells and nests in between are seen).

Identify by history /the salt and pepper chromatin .

Electron microscopy and the markers are same as that of neuroendocrine tumors.

Cribriiform pattern :

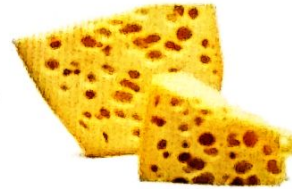
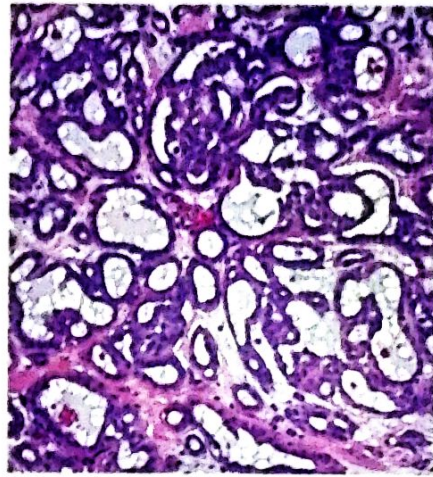
Cookie cutter pattern / Swiss cheese pattern.

Seen in :

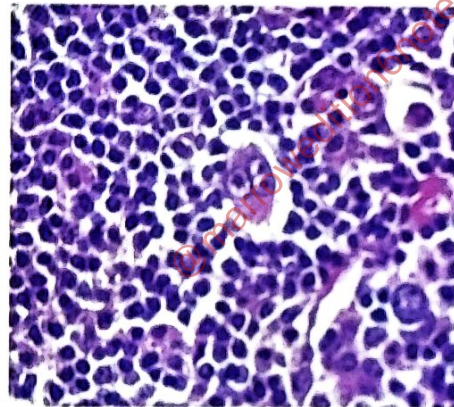
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Cribriiform DCIS (ductal carcinoma in situ).

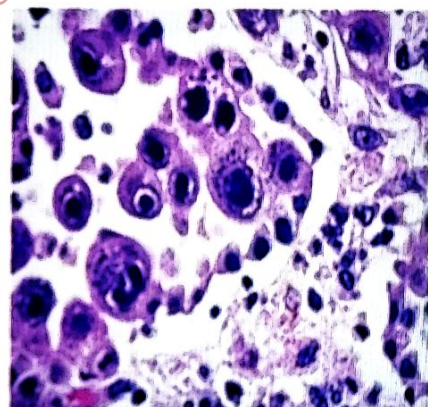
Adenoid cystic carcinoma of salivary gland.



Owl's eye :
Seen in :
Reed-Sternberg cells in
Hodgkin lymphoma.
Cytomegalovirus inclusions.

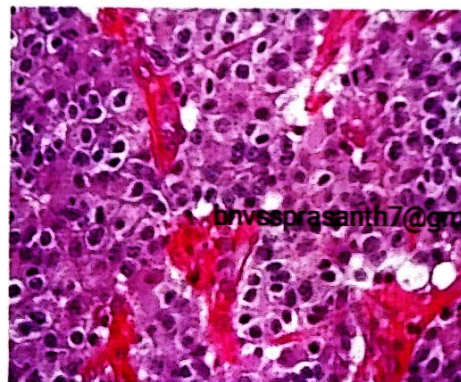


RS cells



CMV inclusions.

Fried eggs appearance :
Seen in :
Oligodendroglioma (grade 2 astrocytoma).
Bone marrow biopsy of hairy cell leukemia.



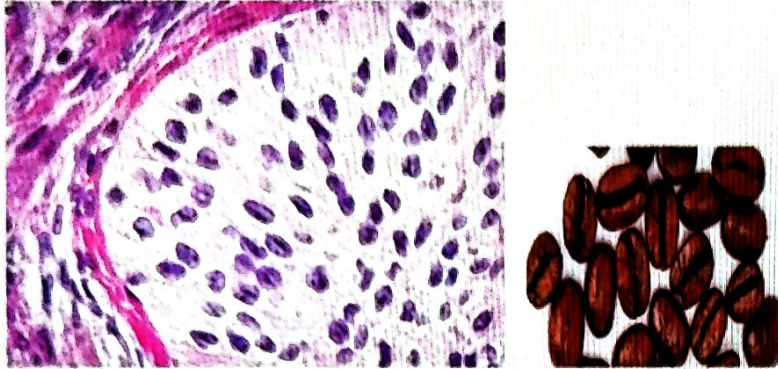
Active space

Coffee bean nuclei :

Longitudinal groove present.

Seen in :

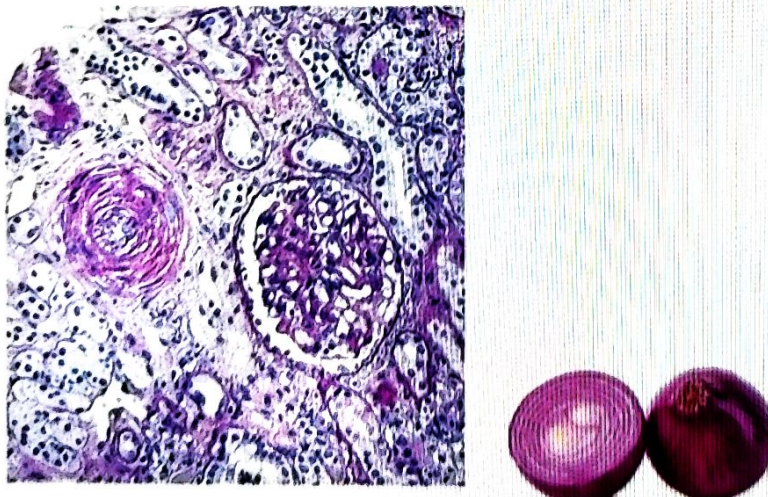
- Papillary carcinoma thyroid.
- Langerhans cell histiocytosis.
- Brenner's tumor.
- Chondroblastoma.
- Granulosa cell tumor.



Onion skin appearance :

Seen in :

- Biopsy of malignant hypertension.
- Biopsy of Chronic inflammatory demyelinating polyneuropathy (CIDP).
- Biopsy of Primary sclerosing cholangitis.
- In gross appearance of spleen in SLE.
- X ray of Ewing's sarcoma.
- In Electron microscopy of Tay sach's disease.



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Active space

Schiller duval body/ Glomeruloid body :

Seen in :

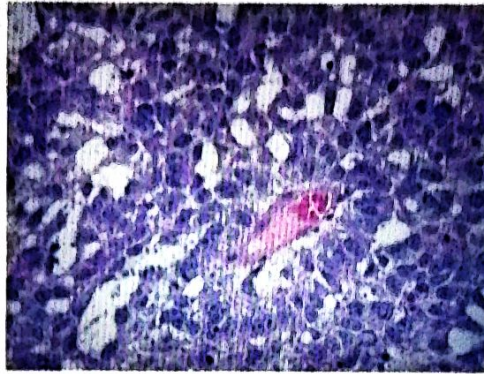
- Yolk sac tumor.

markers :

Alpha fetoprotein.

Alpha 1 antitrypsin .

- Glioblastoma multiforme (grade 4 brain tumor)



Blood vessel with RBCs in the center and another layer of cells outside :Schiller Duvel body .

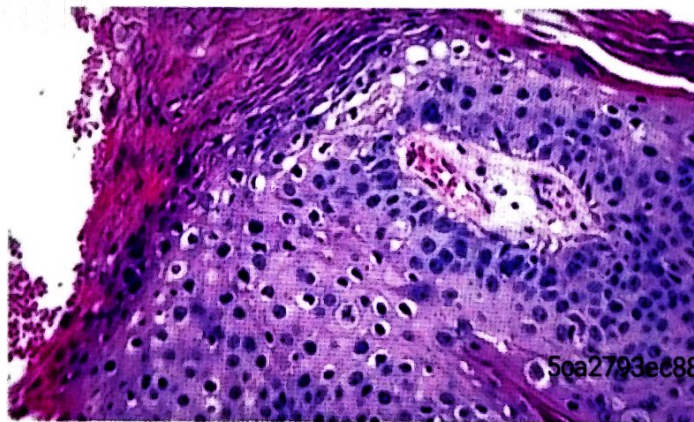
Koilocyte :

History of a genital wart or cervical cancer → Human Papilloma Virus infection.

Identified by : koilocyte

Characteristic feature :

- Thick membrane.
- Resinoid nucleus.
- Perinuclear halo.



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GENETICS - BASIC CONCEPTS AND DIAGNOSIS

Introduction

00:02:00

Genetics is the study of genome.

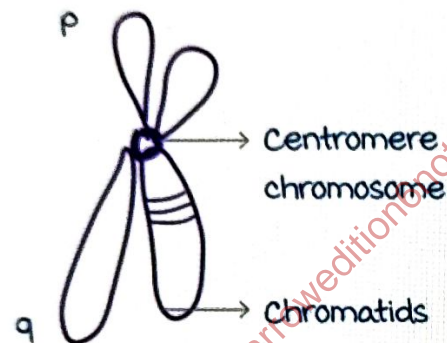
There are approx. 20,000-30,000 genes in our body.

These genes are located on the chromosome.

The short arm of every chromosome is called as P (petite).

The long arm is called as Q.

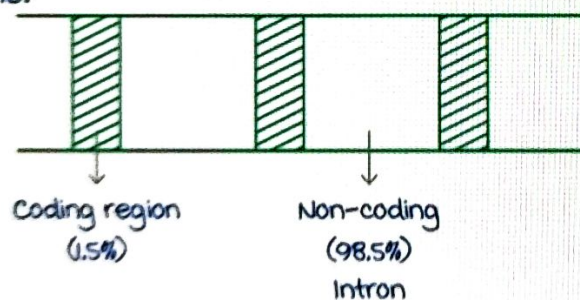
Every gene has got 2 alleles, Out of which one is received from the father and the other from the mother.



Structure of a gene :

Coding region, called the **exon**, consist of 1.5 % of the gene.

Non-coding region called the **intron**, comprises 98.5% approx of the gene.



Terminologies in genetics

00:04:14

1. Homozygous :
When both the alleles are same AA or aa.
2. Heterozygous :
When both alleles are different : Aa.
3. Dominant disorder:
It can manifest in a heterozygous state.

Active space

4. Recessive :

It can only **manifest** in the homozygous state.

The heterozygous state in a recessive disease is known as a **carrier**.

5. Co-dominant : Both the alleles will act dominant & will **simultaneously express**.

Example : ABO blood group, HLA typing.

6. Incomplete penetrance :

This is a property of **autosomal** disorder.

For example : Suppose 100 individuals are affected with marfans, in which only 80 people will show symptoms and the rest will escape symptoms.

Here the penetrance is 80%.

7. Variable expressivity : **Different expression** of clinical features. Despite having the same defective gene is seen in autosomal dominant disorders.

For example: If there are 4 individuals with the defective gene of neurofibromatosis - 1.

Individual 1 : manifests skin lesion.

Individual 2 : manifests neurofibromas.

Individual 3 : manifests with eye lesions.

Individual 4 : manifests with skin and eye lesion.

Here there is variable expression of the clinical features.

8. Pleiotropy :

A single mutant gene can produce multiple end effects.

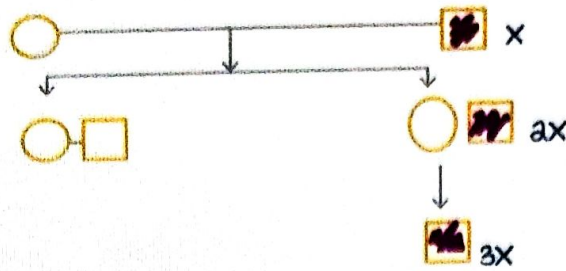
For example : In sickle cell anaemia, glutamic acid is replaced by valine at the 6th position of the beta globin chain.

All the defect is at a single site, it can lead to multiple organ defects like splenomegaly, auto-splenectomy, microvascular occlusions.

9. Anticipation : In some cases the severity of the diseases increased with each successive generation.

For example: If severity of a disease is "x", severity become 2x in next generation & so on.

Seen in trinucleotide repeat mutations like **Fragile X syndrome** (CGG repeats).



10. Polymorphism : Here people differ in genome only by 0.5%. (99% of the genome is almost same).
It is of 2 types : Single nucleotide polymorphism and copy number variations.
11. mutation : It is a permanent heritable gene in DNA.
Can be divided into 2 types ;
Point mutation : If a single loci/nucleotide is affected, it is called point mutation.
It can be classified into :
- Silent mutation : Single nucleotide change but it does not produce a different amino acid or protein.
 - missense mutation : A single nucleotide change but produces a different amino acid and protein.
(Example : Sickle cell anemia)
 - Nonsense mutation : There is a single nucleotide change but it will produce a stop codon (UAA, UAG, UGA). Example : Beta-thalassemia.
 - Frame shift mutation : Insertion or deletion of 1 or 2 nucleotides results in a shift in the reading frame of DNA.
Example : Beta-thalassemia.

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Micro RNA

00:19:21

It is a non-coding RNA.

(DNA leads to RNA and RNA becomes a protein, if it does not become a protein, it is called a non-coding RNA)

It is only 22 nucleotides in length.

It has some role in post transcriptional silencing.

It is of 2 types :

- Tumor suppressing miRNA (Good) : miR 15, 16.
In cases of CLL, there is a deletion of miRNA 15,16.

- Oncogenic miRNA (Bad) : miRNA 155, 200.
- The various B cell lymphomas can be associated with increased expression of miRNA 150, 200.

Epigenetics :

They are hereditary chemical modifications in the DNA/ Histones/ chromatin.

They are reversible.

No change in the nucleotide sequence.

Occur by 2 processes :

1. DNA methylation (more common).
2. Histone deacetylation.

The role of epigenetics :

- Regulation of gene expression.
- X chromosome inactivation.
- Involved in cellular aging .
- Involved in various cancers.

It can be diagnosed in lab by 2 techniques :

Bisulphate sequencing.

Immunoprecipitation assays.

Gentic diagnostic techniques

00:25:53

Cytogenetic analysis :

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- Karyotyping.
- FISH.

The cytogenetic techniques are used for known defects.

They can easily pick the chromosomal disorders.

They are usually for bigger defects.

molecular genetic assays :

- MLPA (multiplex ligation probe analysis).
- PCR (polymerase chain reaction).
- Sequencing.
- Array.

molecular techniques are used for unknown mutations or unknown loci.

Also used in molecular defects.

PCR :

Types :

1. Sanger sequencing :
It is the gold standard for sequence determination.
2. Pyrosequencing : When the specimen is contaminated or the sample is very small.
3. Single base primer extension :
When there is a known genetic defect.
4. Restriction fragment length analysis :
Unknown genetic defects.
5. Real time PCR : Quantitative estimation like in CML, to determine the load of cells with t (9:22).
6. Genome wide association studies :
To see the trend of disease in a population.
7. Amplicon length analysis : Done for repetition of genome like trinucleotide repeat mutation.

Questions :

1. A 63 year old female presented with a breast lump measuring 6 X 5 cm. Truecut biopsy reveals IDC. Tumor cells are sent for genetic testing. The number of tumor cells are very less and the sample had lost fat. Which technique is best suited?
 - a. Sanger sequencing.
 - b. RFLP.
 - c. RT PCR.
 - d. Pyrosequencing.
2. A patient has been diagnosed with CML and is started on Imatinib mesylate. The patient shows good response with it and is taken up for evaluation of Bcr: abl fusion remaining copies. Which of the following is the most suitable technique?
 - a. Sanger sequencing
 - b. RFLP.
 - c. RT PCR.
 - d. GWAS.

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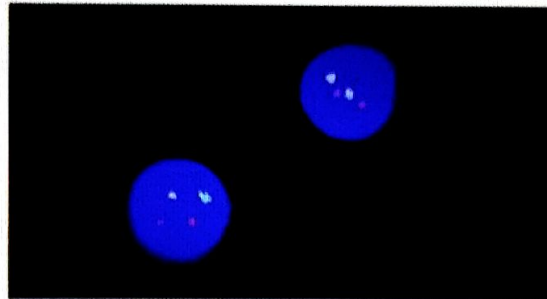
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FISH/ Fluorescent in situ hybridisation.

00:33:56

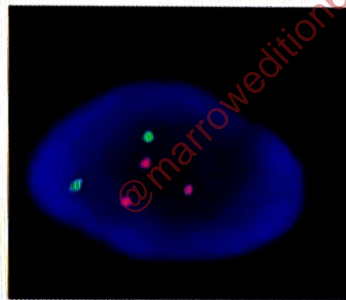
Applications :

- Chromosomal disorders (aneuploidy, deletion, trisomy)
- Translocations.
- Amplifications.

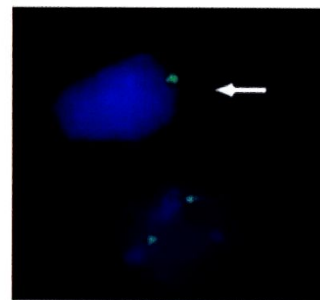


Normal : 2 green and 2 red dots (signals) \rightarrow double chromosomes of each type.

Q. 1) 1 year old child with simian crease. Red is for chromosome 21. Image of FISH shows trisomy 21 (Down's syndrome).



1) Down's syndrome



2) Turner's syndrome

Q. 2) 16 year old girl with primary amenorrhoea and webbed neck. Green signifies X chromosome. 3 green signal. Loss of 1 X, which means patient has XO : Turner's syndrome.

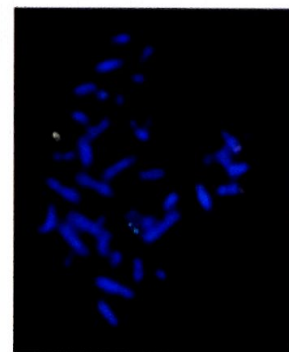
Q. 3) 14 year old male patient from Bihar with massive splenomegaly.

In image, red is chromosome 9 &

green is chromosome 22

Here 1 red and green signal has fused with each other.

Diagnosis is CML.

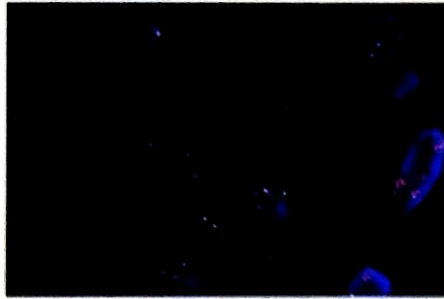


3) CML

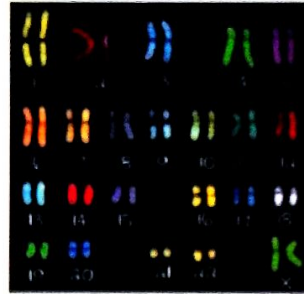
Active space

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Her2 neu amplification : Here multiple red and green signals can be appreciated, which is feature of amplification. useful in a patient for **breast cancer**, if IHC for HER 2 neu come equivocal, FISH is often done.



Her2neu



Spectral karyotyping

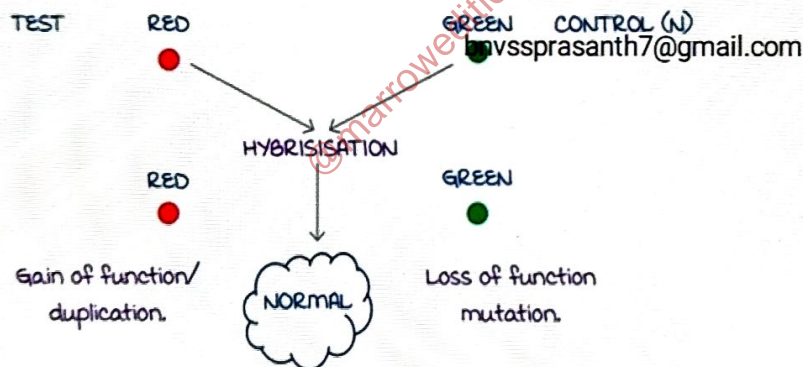
FISH is usually done in **interphase** of cell cycle.

Spectral karyotyping :

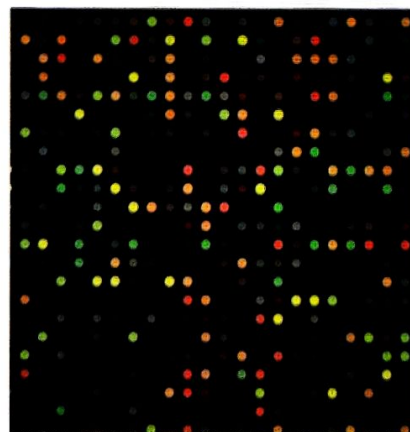
It is a modification of FISH, which is actually a **5 colour FISH**.

Comparative genomic hybridisation

00:42:41



This is called as a **microarray**. multiple people can be tested together.

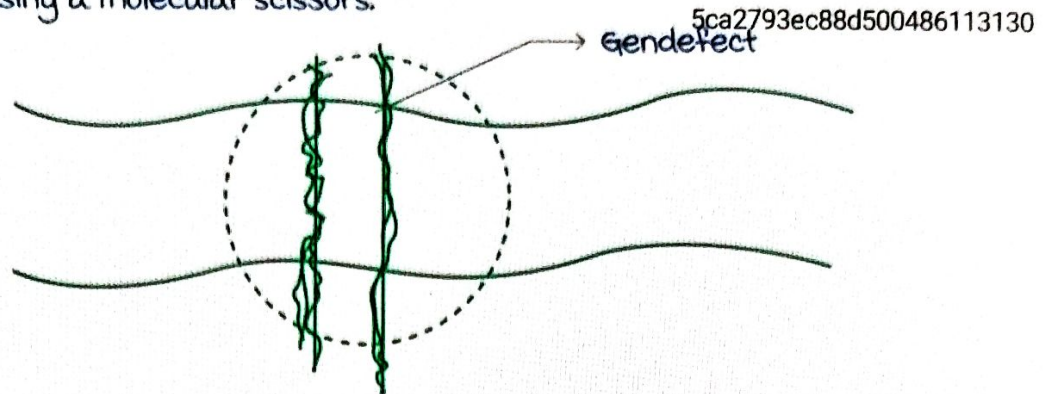


microarray

Active space

GENE (Genome editing with engineered nucleases) :

Here the genome is being edited by engineered nucleases. Here DNA can be deleted, or it can be inserted in to genome using a molecular scissors.



The molecular scissors will create site specific breaks and join the ends by non-homologous end joining.

Nucleases available :

- TALEN.
- Zn-finger endonuclease.
- **CRISPR-CAS9.**

Drawbacks : Can be misused.

Question :

1. Biopsy from a 8 year old child with leg swelling was showing small round blue tumour cells consistent with a diagnosis of Ewing's sarcoma. What is the best method to detect translocation in this malignancy?

- a. Next generation sequencing.
- b. FISH.
- c. PCR.
- d. Conventional karyotyping.

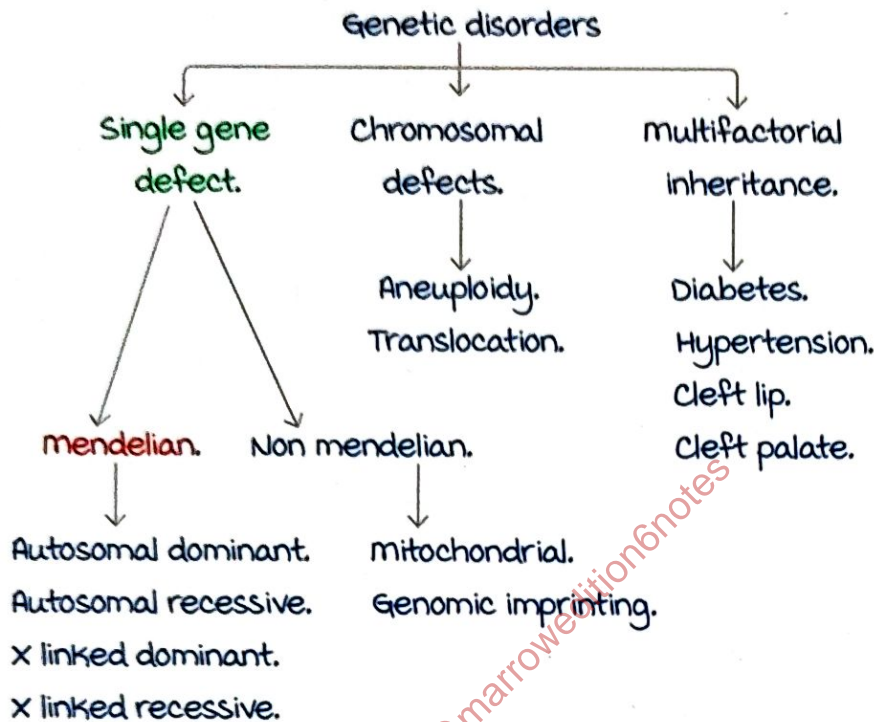
2. A patient who has autosomal dominant gene for type I osteogenesis imperfecta has blue sclera and slightly reduced height while his brother has multiple fractures and deformities. This is an example of :

- a. Polymorphism.
- b. mutation.
- c. **variable expressivity.**
- d. Reduced penetrance

GENETICS - MENDELIAN MODES

Classification of genetic disorders

00:00:43



Autosomal dominant disorders

00:01:33

males = females.

Can be expressed in a **heterozygous state**.

At least one parent of index case is affected.

manifests in **adulthood**.

Due to defects in **structural proteins**.

Better prognosis.

Skip generations are **absent**.

Two properties :

- **Incomplete penetrance** →

Eg: 100 people have defective gene for achondroplasia.

80 people show disease.

Penetrance = 80%.

Active space

- **variable expressivity** → Clinical features vary in individuals with same genotype.

Genotypes in autosomal dominant disorder :

- Only genotypes possible: Aa & aa .
- A → Affected ; a → Normal.

Homozygous (AA) dominant is incompatible with life.

Q. Father is affected (Aa) & mother is normal (aa). what percentage of children will be affected?

Answer :

	a	a
A	Aa	Aa
a	aa	aa

50% will be affected and 50% will be normal.

Examples : mnemonic → **He Has A Very DOMINANT Father.**

- Huntington's disease.
- Hereditary spherocytosis.
- Achondroplasia.
- VWD, VHL.
- Dystrophia myotonica.
- Osteogenesis imperfecta.
- Marfan's syndrome.
- Intermittent porphyria.
- NF-1.
- Adult onset polycystic kidney disease.
- NF-2.
- Tuberous sclerosis.
- Familial adenomatous polyposis, Familial hypercholesterolemia.

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Active space

Marfan syndrome

00:13:33

Defect in **Fibrillin 1 (FBN 1)** gene on **chromosome 15**.

Defect in **Fibrillin 2 (FBN 2)** gene : **Congenital contractural arachnodactyly.**

Pathogenesis :

Fibrillin 1 gene is a helical protein with TGF-beta inside.
mutation of fibrillin → Excess of TGF-beta → Destroys elastin fibers.

Clinical presentation :

Skeletal defects →
most striking feature.

- Tall stature :
 Long bones (span exceeds height).
- Long Spidery fingers & toes.
- Hyperextensible joints :
 most common involved → Thumb.
- Dolicocephalic head.
- High arched palate.
- Pectus excavatum & scoliosis can occur.

**Ocular defects :**

- Ectopia lentis (Supero temporal dislocation of lens).
- myopia.

CVS defects :

- mitral valve prolapse →
 most common cardiovascular defect.
- Aortic dissection →
 most common cause of death.

Life expectancy is halved due to cardiovascular complications.

Diagnosis of marfan's syndrome (Revised Ghent's criteria):

- Family history.
- Clinical signs & symptoms.
- Fibrillin 1 gene mutation.

Biopsy of vessels → Cystic medial degeneration.

Stain for elastin → Verhoeff's Van Gieson (VVG) stain.

Treatment :

TGF-Beta blocker (Losartan).

It is thought that President Abraham Lincoln & Julius Caesar may have had marfan's syndrome.

Neurofibromatosis Type 1 & Type 2 :

Neurofibromatosis Type 1 (NF-1) :

Gene on **chromosome 17** encodes for **neurofibromin**.

Presentation :

- Cafe-au-lait spots (> 6 spots).
- Neurofibromas.
- Lisch nodules (Pigmented Iris hamartomas).

Patient can develop tumors :

- Optic nerve gliomas.
- meningiomas.
- Pheochromocytoma.

Can be associated with Juvenile myelo monocytic leukemia.

multiple neurofibromas



Lisch nodules



Neurofibromatosis type 2:

Gene on **chromosome 22**.

Increased risk of **bilateral acoustic neuroma** or **schwannomas**.

Autosomal recessive disorders

00:26:46

males = Females.

Only expressed in **homozygous states**.

When heterozygous \rightarrow Carrier/ trait.

Show **complete penetrance**.

Usually due to **enzyme deficiency**.

Skip generations **present**.

manifests in childhood/ infancy.

Examples :

mnemonic → ABCDEF~~GH~~I.

- Alpha antitrypsin deficiency, Ataxia telangiectasia, Alkaptonuria.
- Beta thalassemia.
- Cystic fibrosis, Congenital adrenal hypoplasia.
- Deafness.
- Emphysema.
- Fredrich's ataxia.
- Gaucher's disease, Glycogen storage disorders, Galactosemia.
- Hemochromatosis, Homocystinuria.
- Inborn errors of metabolism.

2 lysosomal storage disorders that are not autosomal recessive :

- Fabry's disease.
- Hunter's disease.

Hematological enzyme deficiency that is not autosomal recessive:

- G-6PD deficiency.

Lysosomal storage disorders

00:31:03

Disorders	Enzyme deficiency
Sphingolipidoses	
Gaucher's disease.	Glucocerebrosidase.
Tay sach's disease.	Hexosaminidase A.
Fabry's disease.	Alpha galactosidase A.
Krabbe disease.	Galactocerebrosidase.
Niemann pick disease.	Sphingomyelinase.
mucopolysaccharidoses	
Hurler syndrome.	Alpha L iduronidase.
Hunter syndrome.	Iduronate 2 sulfatase.

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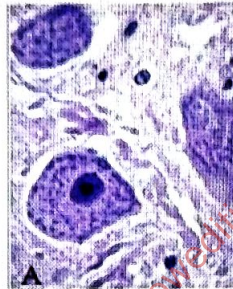
Tay Sachs disease :

Mnemonic → TAYSACHS

- Autosomal recessive.
- Young adults.
- Cherry red Spot.
- Common in Askenazi jews.
- CNS defects.
- Hexosaminidase alpha subunit deficiency leads to accumulation of GM 2 Ganglioside.
- Onion skinning appearance in electron microscopy.

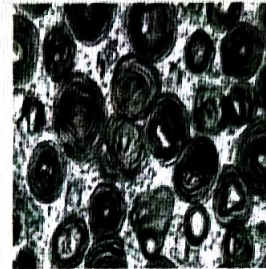
Brain biopsy :

Ballooned neurons.



Electron microscopy :

Onion skin appearance.



Onion skin appearance in medicine :

- Biopsy of malignant hypertension.
- Nerve biopsy of CIPD.
- Biopsy of primary sclerosing cholangitis.
- Gross specimen of spleen in SLE.
- X-ray of ewing's sarcoma.
- Electron microscopy of Tay sach's disease.



Neimann pick disease :

Defect of sphingomyelinase.

Sphingomyelin accumulate in lysosomes.

Electron microscopy : Zebra bodies.



Gaucher's disease :

most common lysosomal storage disorder.

Deficiency of glucocerebrosidase.

Clinically manifests in three forms :

1. Non neuronopathic →

most common form.

Affects adults.

Some glucocerebrosidase activity present.

No CNS symptoms.

Hepatosplenomegaly, fractures, bone pain.

2. Neuronopathic →

Less common.

Absent glucocerebrosidase activity.

Infants.

CNS symptoms present.

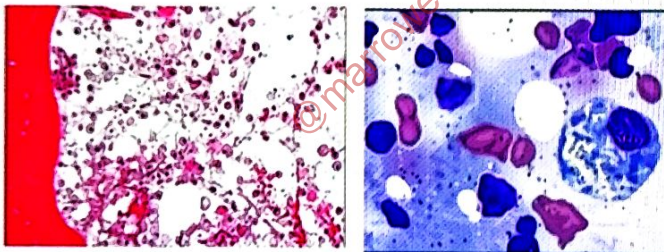
3. Intermediate.

Biopsy → Gaucher's cells.

wrinkled/crumpled tissue paper appearance.

Stains :

- PAS+
- Oil red O +
- Pearl's stain +



Pseudo gaucher's cell → Chronic myeloid leukemia (PAS-).

X linked recessive disorders

00:41:45

males >>> Females.

Females are usually carriers.

Females can be affected when there is random X inactivation during Lyon's hypothesis.

Examples :

mnemonic → Lady Hardinge College Girls Don't Care About Foolish Words.

- Lesch Nyhan syndrome.
- Hemophilia A & B, Hunter's disease.
- Colour blindness.

Active space

- G6PD deficiency.
- Duchenne muscular dystrophy.
- Chronic granulomatous disease.
- Agammaglobunemia.
- Fabry's disease, Fragile X syndrome.
- Wiskott aldrich syndrome.

Father is affected, mother is normal. :

	X	X
X ⁰	XX ⁰	XX ⁰
Y	XY	XY

All daughters are carriers.

All sons are normal.

Father doesn't inherit the disease to sons, all daughters are carriers.

X linked dominant disorders

00::46:15

Rare disorders.

Examples :

Mnemonic →: RAVI.

- Rett's syndrome.
- Alport syndrome.
- Vitamin D resistant rickets.
- Incontinentia pigmenti.

Alport syndrome :

- Inherited by any mode.
- most common mode : X linked dominant inheritance.

Clinical scenarios

00::47:04

Q. Father has achondroplasia, mother is normal. What percentage of children will have achondroplasia?

Answer :

mode of inheritance → Autosomal dominant (Heterozygous state)

	a	a
A	Aa	Aa
a	aa	aa

Active space

50% will be affected and 50% will be normal.

Q. Husband has sickle cell anemia, wife is a carrier of sickle cell anemia. what is the percentage of children affected with sickle cell anemia?

Answer :

mode of inheritance → Autosomal recessive.

	A	a
A	AA	Aa
A	AA	Aa

50% will be affected, 50 % carriers.

Q. Husband has HbA₂ : 4.8%, wife has HbA₂ : 2%. what is the percentage of children affected with thalassemia major?

ANS:

Normal HbA₂ level: 2-3.5%.

Beta Thalassemia trait HbA₂ level: 4-9%.

mode of inheritance: Autosomal recessive.

	A	a
A	AA	Aa
A	AA	Aa

50% kids will be carriers, while 50% will be normal.

Hence no children with thalassemia major.

Q. A 22 year old man is evaluated for mitral regurgitation due to mitral valve prolapse. Examination reveals a tall, slender, young man with long extremities and long tapering fingers. Pupillary dilation reveals bilateral dislocation of lens. This patient is potentially at increased risk for development of which of the following?

- A. Aortic dissection (marfan's syndrome).
- B. Lisch nodules.
- C. Non caseating granuloma.
- D. Rapidly progressive renal failure.

Q. Which of the following dyads are correct?

- A. Marfan syndrome : AR.
- B. PKU : AD.
- C. Vit. D resistant rickets : AD.
- D. Alkaptonuria : AR.
- E. DMD : XLR.

Q. A patient has an autosomally inherited condition. The patient and his grandfather show evidence of disease, but the patient's father is asymptomatic. This is an example of?

- A. Mutation.
- B. Polymorphism.
- C. Variable expressivity.
- D. Reduced penetrance.

@marroweditionsnotes

Active space

GENETICS : NON-MENDELIAN MODES AND PEDIGREE

Mitochondrial inheritance

00:02:44

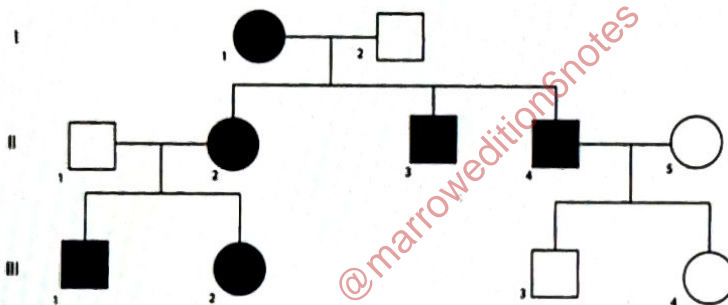
Exclusively maternally inherited.

Pathogenesis :

During fusion of ova and sperm, the ova eliminates all sperm mitochondrial DNA by ubiquitin protease pathway.



Zygote DNA contains only maternal mitochondrial DNA
mother transmits disease to all children, whereas father to none.



Mitochondrial genome

00:07:26

mitochondrial genome : 37 genes, 12 t-RNA, 2 r-RNA.

Heteroplasmy : Presence of normal + mutant mitochondrial DNA in same person.

Organs affected are : CNS, eye, skeletal muscles.

Disorders :

- **MELAS** : mitochondrial Encephalopathy, Lactic acidosis and Stroke like episodes : **mc**.
- Leigh syndrome.
- Leber's hereditary optic neuropathy.
- Pearson syndrome.
- CPEO : Chronic Progressive External Ophthalmoplegia.
- Kearnsare syndrome.
- NARP syndrome.

Active space

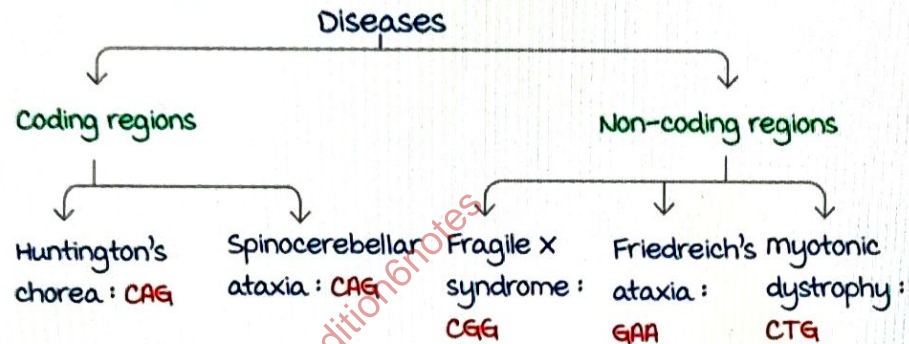
Trinucleotide repeat mutation

00:12:48

mutations which causes increased number of codons that causes disease.

1st generation : CAG 40 : Normal.
 ↓
 and generation : CAG 100 : Premutation. ↑ mutation
 ↓
 3rd generation : CAG 1000 : Full mutation.

Anticipation : The severity of the disease increases with each successive generation.

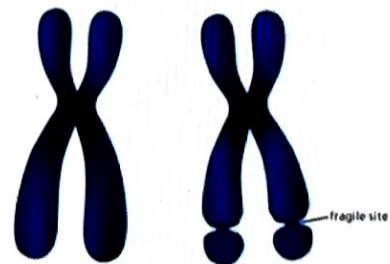


These diseases mostly involve cytosine and guanine nucleotides.

Fragile X syndrome

00:22:53

X linked recessive disorder
 when cells are cultured in folate deficient media, constriction is seen in X chromosome.



Fragile X chromosome

Due to loss of function mutation in FMR 1 gene leading to increased CGG repeats.

Normal : 6 to 55 CGG repeats.

Premutation : 55 to 200 CGG repeats.

Full mutation : 200 to 4000 CGG repeats.

Fragile X syndrome, Huntingtons chorea follow non mendelian mode of inheritance despite being X linked and Autosomal dominant diseases respectively

Active space

Clinical features :

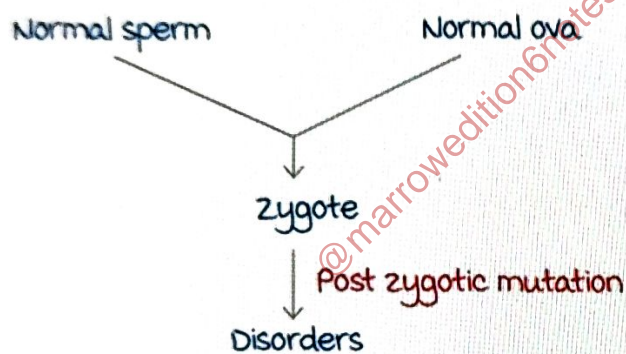
- 2nd MC genetic cause of mental retardation.
- MC **inherited** cause of mental retardation.
- **macroorchidism** (large testis) : most distinctive feature.
- Large head.
- Large everted ears.
- Large jaws/mandible.

Fragile X tremor / ataxia : Gain of function mutation in FMR 1 gene (Premutation).

Fragile X associated ovarian failure : Gain of function mutation in FMR 1 gene (premutation).

Gonadal mosaicism :

Type of autosomal dominant inheritance.



No risk to the sibling.

Examples :

- Tuberous sclerosis.
- Osteogenesis imperfecta.

Genomic imprinting

00:40:16

Gene **silencing** / inactivation.

Physiologically either paternal / maternal allele is inactivated

: Only one

allele is functional.

Happens by the process of **epigenetics** : Lowers expression of genes.

- DNA methylation.

- Histone deacetylation.

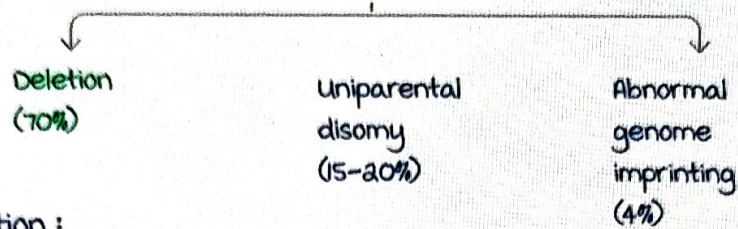
Chromosome 15q11

- Prader willi syndrome
- Angelman syndrome

Prader willi syndrome

00:46:11

Pathogenesis :



Deletion :

maternal allele is silenced and only paternal allele is functioning normally.

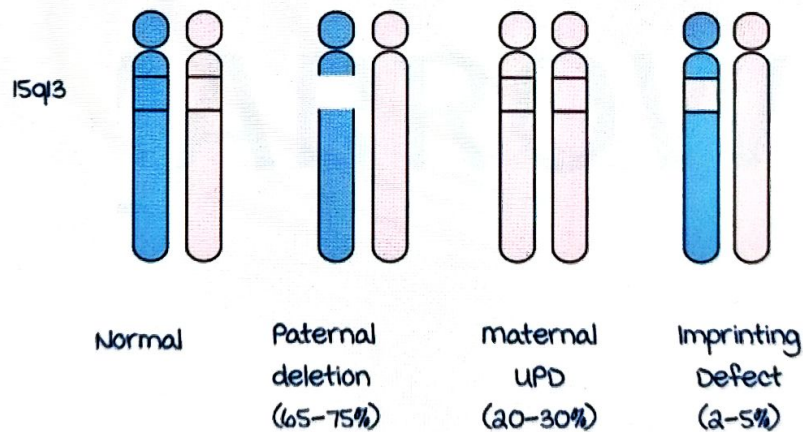
If paternal allele is deleted and maternal allele is silenced :

Prader willi syndrome.

uniparental disomy :

If both the alleles are maternal (maternal disomy) which are non-functional they lead to uniparental disomy.

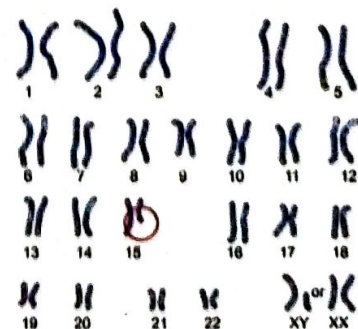
p m p m p m p m



SNORP gene mutation have been observed lately.

Clinical features

- Small bird like head.
- mentally retarded.



Active space

- Respiratory problems.
- Obesity.
- Short lifespan.
- Hyperphagia.
- Hypogonadism.
- Hypotonia.

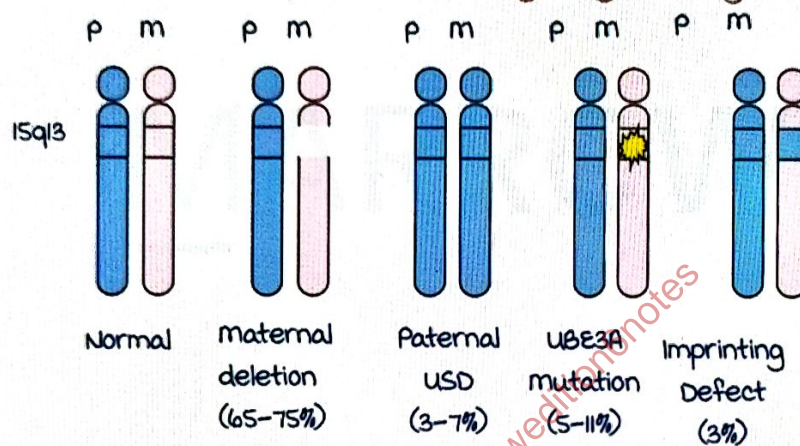


Angelman syndrome

00:53:16

Paternal gene is silenced, maternal gene is functional.

maternal deletion + paternal silencing : Angelman syndrome.



UBE3A gene mutation have been implicated lately.

Clinical features

- Inappropriate laughter : Happy puppet syndrome.
- Stiff / ataxic movements : Hand flapping.
- microcephaly.
- Seizures.
- mental retardation.

Prader willi syndrome	Angelman syndrome
Chromosome 15	Chromosome 15
maternal imprinting	Paternal imprinting
Paternal deletion	maternal deletion
maternal disomy	Paternal disomy
SNORP gene	UBE3A gene
Hyperphagia, obesity, mental retardation	Happy puppet syndrome

Active space

Pedigree analysis

01:00:24

Types :

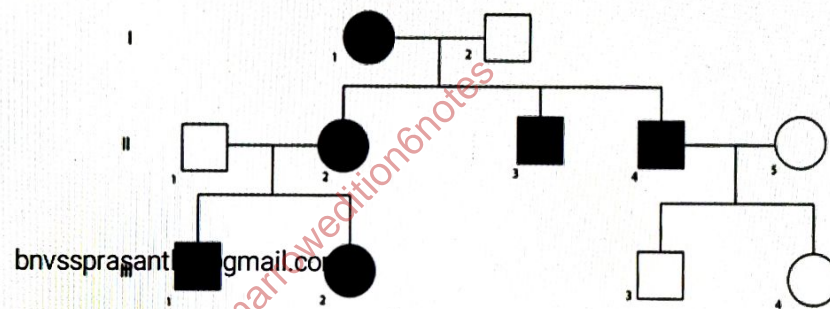
- mitochondrial inheritance
- Autosomal dominant
- Autosomal recessive
- X linked recessive
- X linked dominant
- Gonadal mosaicism

Steps for pedigree analysis :

1. Rule out mitochondrial inheritance

Affected mother transmits disease to all kids

Affected father does not transmit disease to kids



2. Rule out whether dominant/recessive disorders.

Dominant : skip generation absent

Recessive : skip generation present

If recessive,

male = female : Autosomal

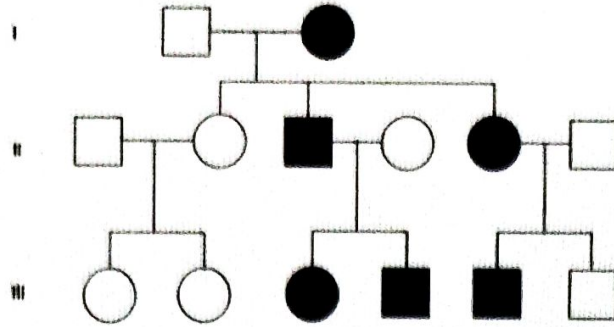
male >>> female : X linked

If dominant,

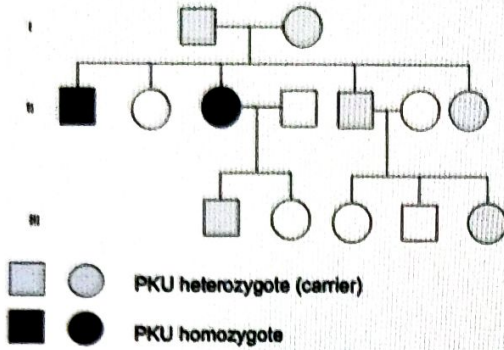
Autosomal : Father to son transmission present.

X linked : Father to son transmission absent and 100% daughters are affected.

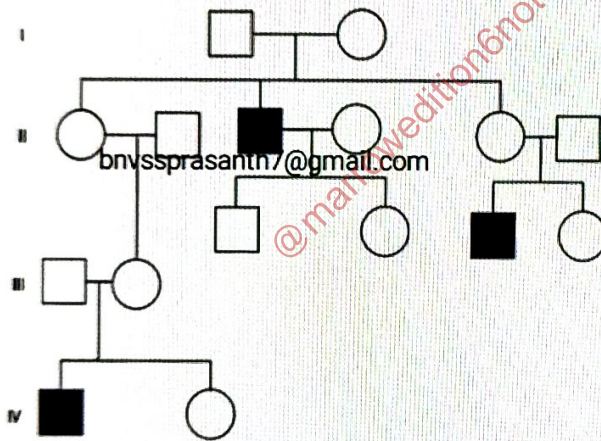
Gonadal mosaicism : One child affected, but none of the parents affected.



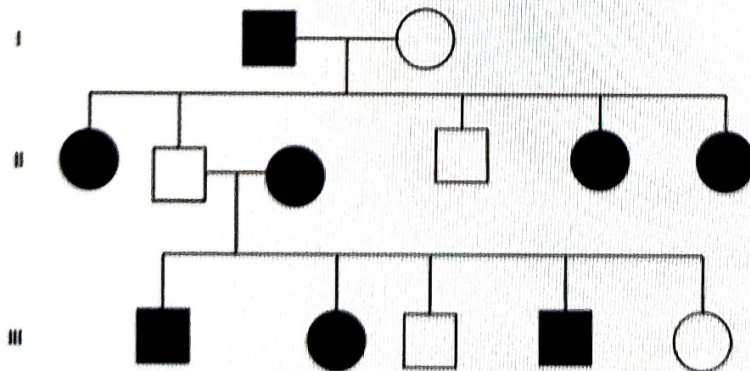
1. Autosomal Dominant



2. Autosomal Recessive



3. X-linked recessive



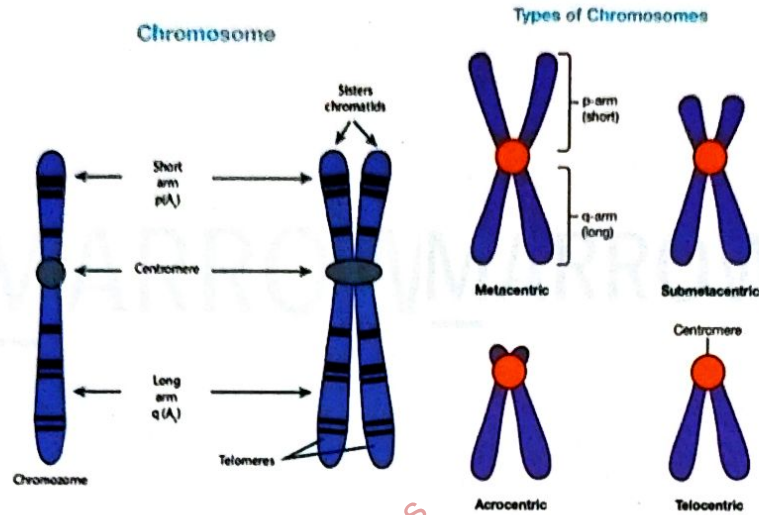
4. X-linked Dominant

Active space

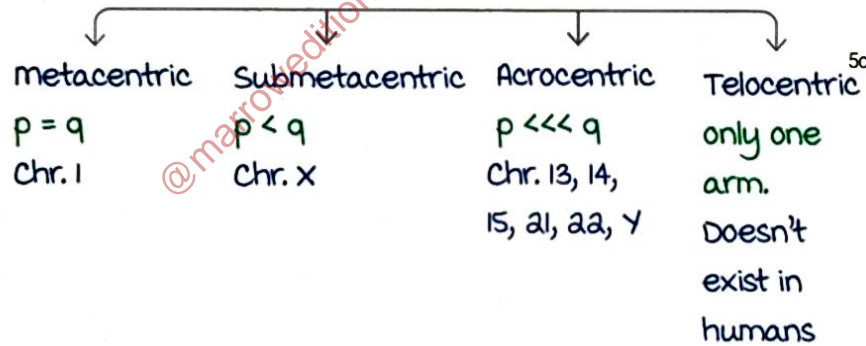
GENETICS- CHROMOSOMAL DISORDERS

Structure of chromosome

00:01:12



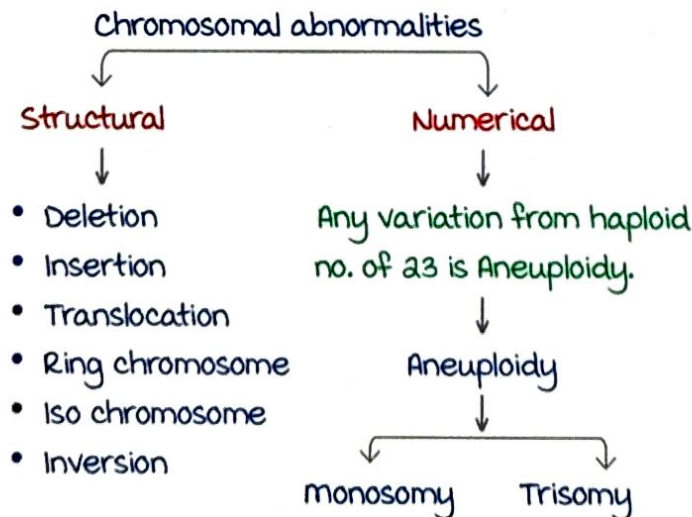
Depending on position of centromere, types of chromosomes :



5ca2793ec88d50048611313

Chromosomal abnormalities

00:07:54



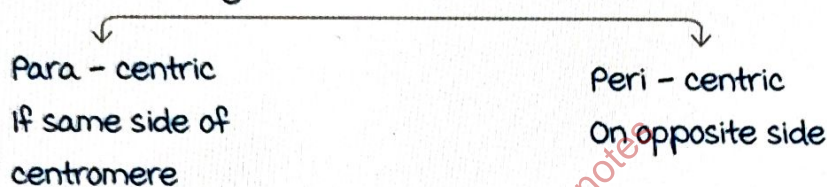
Only monosomy compatible with life : Turner's syndrome.

Active space

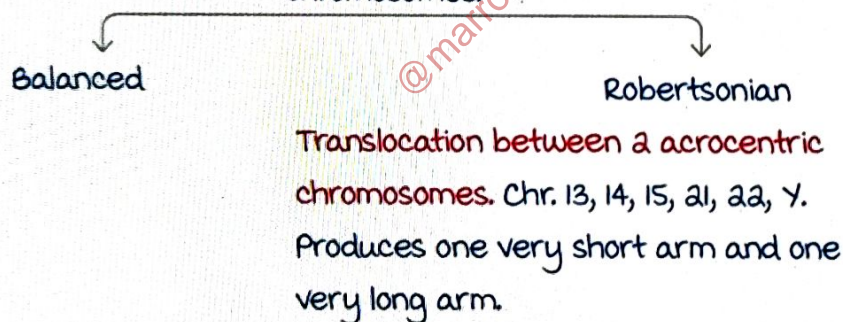
Structural abnormalities

00:11:22

1. **Insertion** : Portion of a chromosome inserted.
2. **Deletion** : Portion of a chromosome deleted.
3. **Ring chromosome** : Break at both ends, fusion of damaged sticky ends. Functionally inactive.
Ring chromosome seen in Turner's syndrome.
4. **Isochromosome** : If division happens horizontally leads to Two short/ two long arms.
MC Isochromosome : iXq
MC Isochromosome in cancer : $i17q$
MC Isochromosome in testicular tumor : $i12p$
5. **Inversion** : Two breaks with attachment of inverted segment.



6. **Translocation** : Exchange of material between two chromosomes.



+ (14:21) Robertsonian translocation seen in 4% cases of Down's syndrome.

Down's syndrome

00:22:26

Trisomy 21.

MC chromosomal disorder.

MC genetic cause of mental retardation.

MC inherited cause of mental retardation : **Fragile X syndrome.**

Pathogenesis :

- meiotic non-dysjunction (95%) : Occurs in oogenesis.
- 47 chromosomes are present. Maternal age is a risk factor.

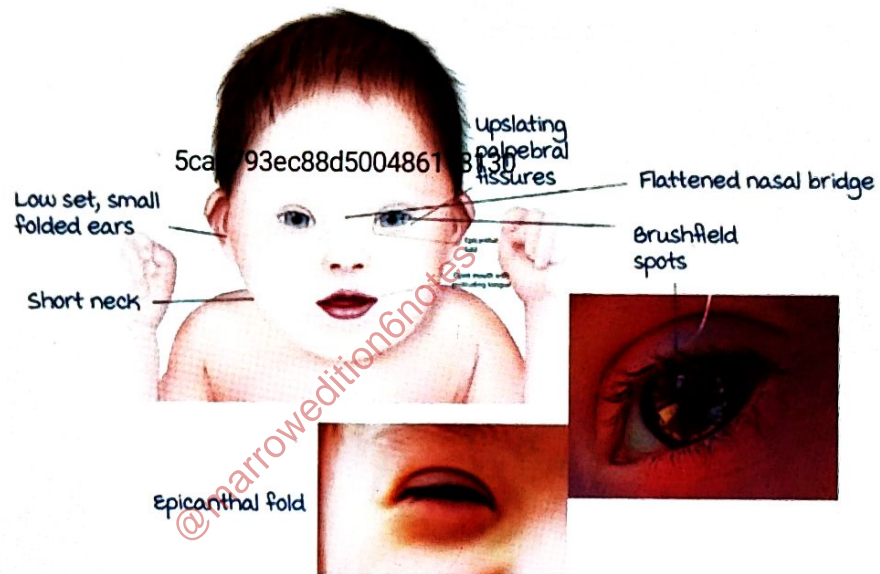
- Robertsonian translocation (4%).
46 chromosomes are present.
maternal age is not a risk factor.
- mosaics (1%).

Triple test : AFP, β - HCG, unconjugated estradiol.

Quadruple test : Triple test + Inhibin.

Clinical features and complications of Down's syndrome

00:30:25



All these facial characteristic features give rise to an appearance called as **mongloid Idiocy**.



Sandle Gap/Saddle toe



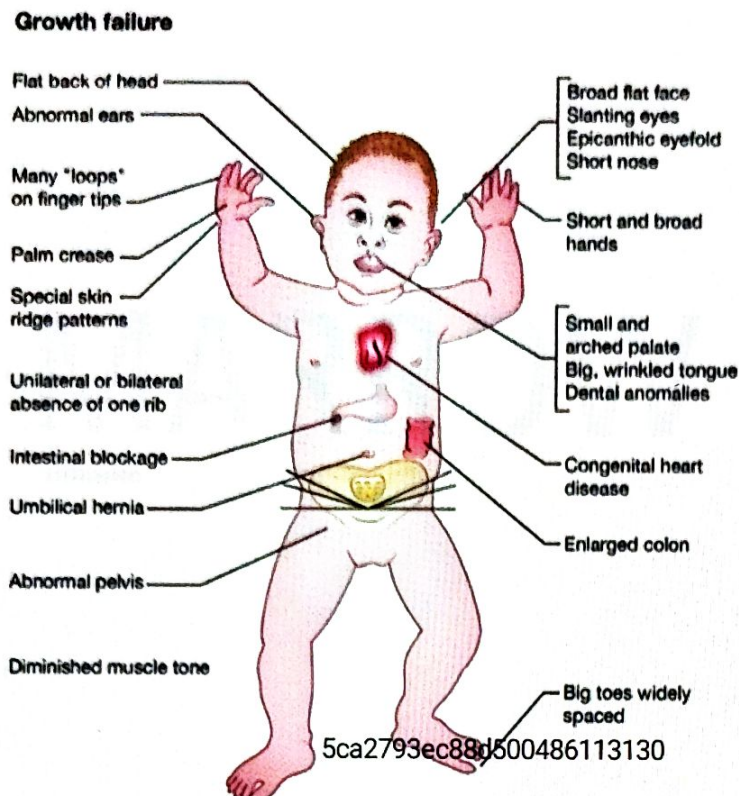
Normal Palm crease



Simian crease

- Simian crease : Single palmer crease.
- Clinodactyly.
- Overlapping fingers.
- Sandle Gap/ Saddle toe : Increased gap between 1st & 2nd toe.

Active space



Complications of Down's syndrome :

1. Cardiovascular defect :
MC Cardiovascular defect : Endocardial cushion defect, VSD.
2. GIT : Annular pancreas, Duodenal atresia, Hirschsprung disease.
3. Acute leukemia :
MC leukemia : ALL
MC leukemia in children < 3 yrs : AML
MC subtype of AML : AML M7
4. Endocrine : Hypothyroidism.
5. CNS : Premature Alzheimer's disease.

Edward and Patau syndrome

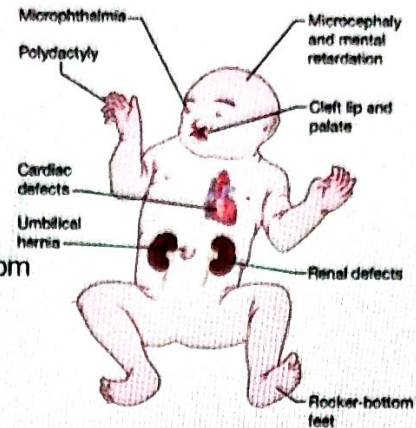
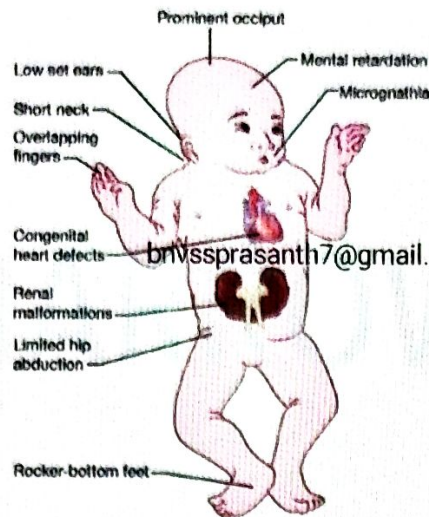
00:39:55

Common features of both syndromes :

- mental retardation.
- CVS defects.
- Rocker bottom feet.
- Prominent occiput.

Active space

Edward syndrome : Trisomy 18 Patau syndrome : Trisomy 13



Incidence: 1 in 15,000 births
 Karyotype:
 Trisomy 13 type : 47,XX,+13
 Translocation type : 46,XX,+13,(t(13;14)(q10;q10))
 Mosaic type : 46,XX/47,XX,+13

del 22 q9 ll.2

A/T/A DiGeorge syndrome or velo cardio facial defects.
 (mnemonic : CATCH 22)

- Cleft lip/ palate
- Abnormal facies
- Thymic hypoplasia
- Cardiac defect
- Hypocalcemia

del 22q11.2

Defect in development of 3rd & 4th pharyngeal pouch.
 Causes thymic & parathyroid hypoplasia

Trisomy 22 (Cat eye syndrome)

multiple colobomas in the eye.

Cat cry syndrome (del 5p)

- Cat like cry.
- Behavioural abnormalities.
- Developmental delay.

CRI-DU-CHAT Syndrome

Sex chromosomal disorders

00:47:10

Lyon's hypothesis :

Only one of the X chromosomes is genetically active.
 Other X, of either paternal or maternal origin becomes

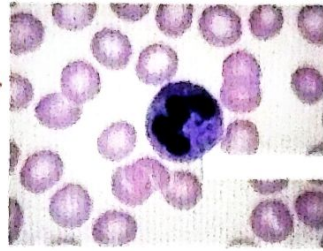
Active space

inactive (occurs randomly).

Inactivated X chromosome : **Barr body**.

Shape : **Drumstick appearance**.

Sample : Buccal mucosa.



Barr body

Clinical applications :

Normal male	XY	No barr body
Normal Female	XX	1 barr body

No. of barr body = No. of X chromosome : 1

Clinical applications of barr body :

Turner's syndrome (XO) : No barr body.

Klinefelter syndrome (XXY) : Extra barr body.

Klinefelter's syndrome

00:53:43

MC cause of male hypogonadism.

more the no. of X chromosome, more is the mental retardation.

Pathogenesis :

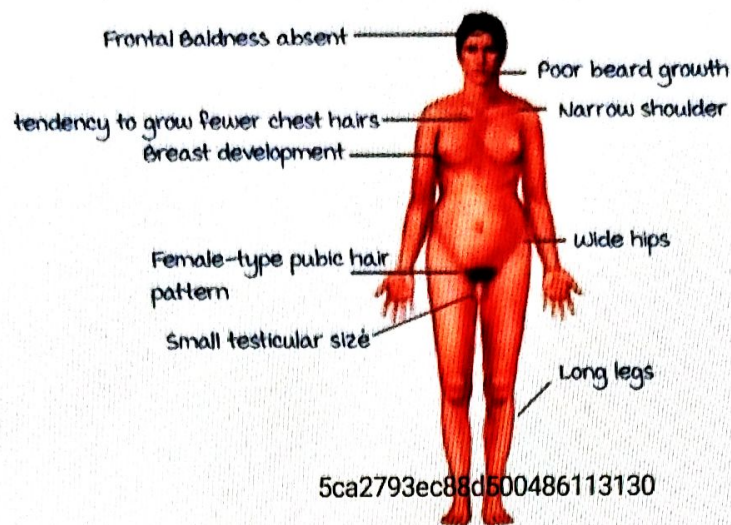
meiotic non disjunction (occurs in both oogenesis & spermatogenesis)

1. Syndromes associated with advanced **maternal age** :
Down's syndrome, XXX syndrome.
2. Syndromes associated with advanced **paternal age** :
 - marfan syndrome.
 - Osteogenesis imperfecta.
 - Achondroplasia.
3. Syndrome associated with both advanced maternal and paternal : Klinefelter.

Clinical features :

- Tall stature.
- Poor muscle tone.
- Reduced secondary sexual characteristics.
- Gynecomastia.
- Eunuchoid body habitus.
- Long extremities.

- Frontal baldness absent.
- Testicular atrophy : Infertility.



Testicular Biopsy : Atrophy of seminiferous lobules.
 Hyalinisation of seminiferous lobules.
 Leydig cell hyperplasia.

Hormonal changes :

Increased FSH, LH and decreased testosterone.

CVS changes : MVP.

Increased risk of developing extragonadal germ cell tumour and Breast CA.

Turner's syndrome

01:02:40

MC cause of female hypogonadism.

Pathogenesis :

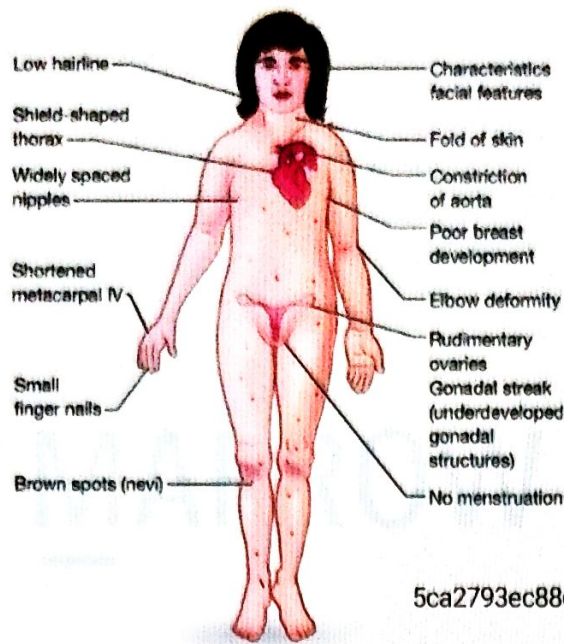
1. meiotic non -dysjunction.
2. Anaphase lag.
3. Ring chromosome.
4. Isochromosome.
5. mosaics.

Webbed neck due to lymphedema.

Elbow deformity : Cubitus valgus.

Ovaries : Streaked ovaries.

Clinical features :



mcc primary amenorrhea : Turner's syndrome.
 mc CVS defect : Bicuspid aortic valve
 mc cause of death : Co-arcetation of aorta

mnemonic : CLOWNS

Cardiac abnormalities, Cubitus valgus, Cystic hygroma.

Lymphedema

Streaked Ovaries

Webbed neck

Normal intelligence, Nipples widely spaced

Short stature, Short 4th metacarpal

Increased risk of developing Gonadoblastoma.

Noonan's syndrome :

Same Clinical features of Turner's syndrome.

Normal karyotype.

mutation in chromosome 12.

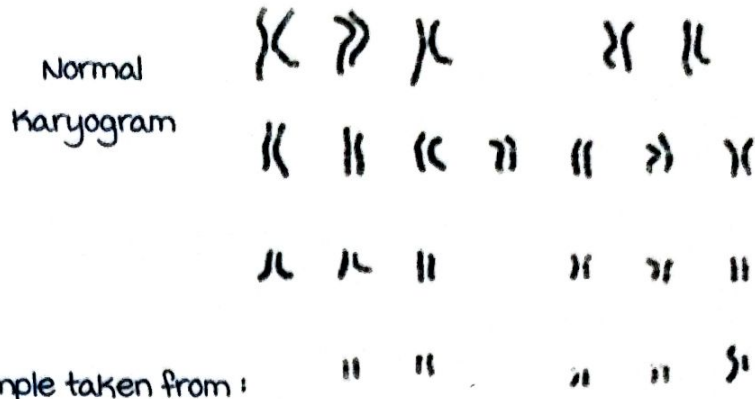
Karyotyping

01:10:59

Karyogram : Arrangement of chromosomes in **descending order** of length followed by sex chromosomes.

Uses : To diagnose structural & numerical abnormalities of chromosome.

Active space



Arrest the cells in metaphase (Using colchicine).

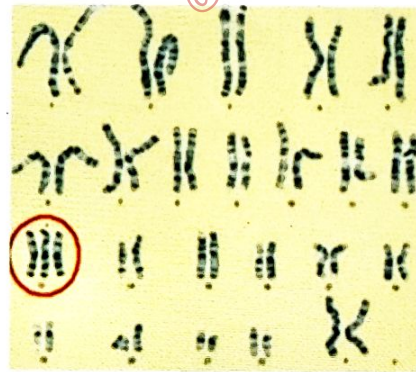
Staining : MC G-banding (Giemsa banding).

Q - banding (quinacrine).

using light microscope,

resolution required for karyotyping : 5 mb.

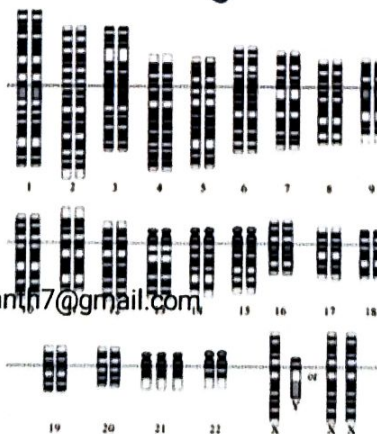
Patau syndrome



Edward syndrome



Down's syndrome



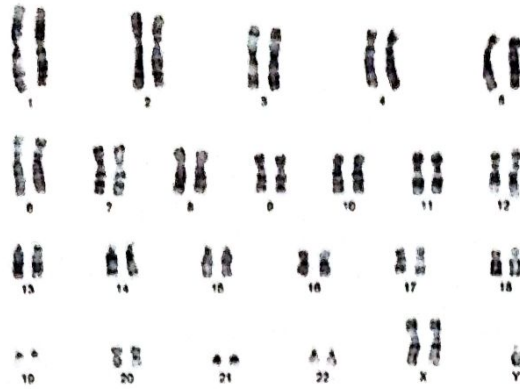
Cri-Du-Chat syndrome



Active space

bnvssprasant7@gmail.com

Klinefelter's syndrome



MCQs :

Q. All of the following are characteristic of Turner's syndrome except:

- A. Webbed neck
- B. Coarctation of aorta
- C. Cubitus valgus
- D. Umbilical hernia

Q. Which chromosome contains the most known genetic disease of any human chromosome?

- A. I
- B. II
- C. 9
- D. 6

Q. A tall man with gynecomastia and testicular atrophy has a testicular biopsy that shows sparse, completely hyalinized seminiferous tubules. Leydig cells are present in large clumps. Which of the following genetic disorders should be suspected?

- A. Trisomy 18
- B. Trisomy 21
- C. 45, XO
- D. 47, XXY

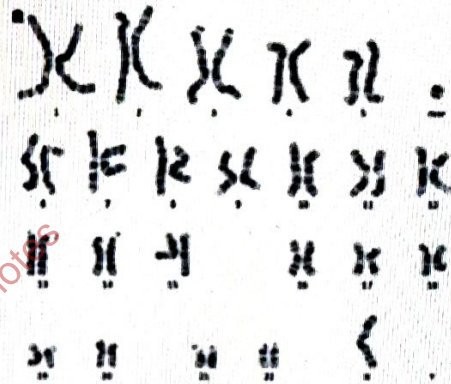
Q. A tall man presents with complaints of infertility. Examination shows gynecomastia and reduced secondary sexual characteristics. Karyotyping analysis revealed an XXY karyotype. Which of the following is not true about the condition?

Active space

- A. Levels of FSH are reduced.
- B. Testosterone levels are reduced.
- C. Plasma estradiol levels are elevated.
- D. Increased risk for breast cancer.

Q. A 21 year old girl with short stature says that her breasts have not developed properly and periods have not started as yet. She also has low posterior hair line. The karyogram from the patient is given below. Which of the following is true for the disease patient is suffering from?

- A. Endocardial cushion defect is the CVS complication.
- B. Short fourth metacarpal.
- C. Aortic valve disease.
- D. micrognathia.



@marroweditions

Active space

IMMUNITY-TYPES OF IMMUNE CELLS

Types of immunity

00:01:19

Innate and adaptive immunity.

Innate	Adaptive
Present by birth.	Acquired later on exposure to antigen.
Non specific	Specific
No memory.	memory present.
Examples include : 1. Epithelial barriers like skin, GIT. 2. Plasma proteins like C-reactive protein. 3. Neutrophils, macrophages, dendritic cells, complement.	Cells are of 2 types : 1. B lymphocytes : Provides humoral immunity and produces antibodies. 2. T lymphocytes : Involved in cell mediated immunity and help with immunity against intracellular microbes.

Pattern recognition receptors (PRR) :

This is a component of innate immunity. The receptors are present on the plasma membrane or cytoplasm or endosome. The purpose of these receptors : To recognize specific patterns on microbes.

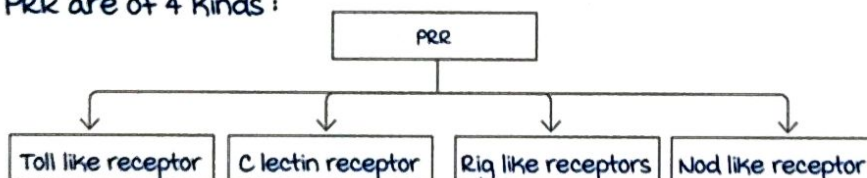
The receptors on :

The plasma membranes : Detect extracellular organism.

In the cytoplasm : Detect intracellular organism.

On the endosome : Detect ingested microbe.

PRR are of 4 kinds :



Toll like receptor (TLR) :

There are around 10 TLR's discovered, and are located on the plasma membrane.

Function :

To detect gram positive and gram negative bacteria.

C lectin receptor :

Located on the plasma membrane.

Detect the fungal glycans.

Rig like receptors :

Present on the cytoplasm and detect viruses.

Nod like receptor :

Present on the cytoplasm and detects :

N : Necrotic debris.

O : Ion transport.

D : Diabetes mellitus.

After detection of these, the NOD like receptor activates caspase 1 which then causes production of IL-1 and leads to result in inflammation and fever.

Inflammasome can be involved with NOD like receptor.

Different types of immune system cells

00:10:31

Natural Killer cell (NK) :

- Usually produced by a large granular lymphocyte.
- Constitutes 5-10% of circulating blood lymphocytes.
- It is not a B cell and a non T cell. It does not have both T cell and B cell receptors and is referred to as null cell.
- Not MCH restricted (unlike T cell).
- Function :
 - It can be part of both innate and adaptive immunity.
 - 1. Innate immunity :
 - Directly kill the virus infected cells or tumor cells.
 - 2. Adaptive immunity :
 - Antibody dependant cell mediated cytotoxicity.
- Popular markers for NK cell : CD 15 and CD 56.
 - CD 56 : Role is still not fully understood.
 - CD 16 : Detect and binds to Fc fragment of IgG on target cells and kill the cell by perforins and releasing toxins and granzymes.

- NK cells have 2 different kinds of receptors.
 1. Activating receptors : **NKG2D**.
 2. Inhibitory receptors : **CD94, CD 96**.
 These prevent self cells from being attacked by NK cells.
- Cytokines produced by NK cells :
IFN- γ : (Cytokine majorly responsible for granuloma formation). This cytokine leads to the activation of macrophages (to epithelioid cell).
- Cytokines responsible for the proliferation of NK cells :
IL-2 and **IL-15**.

B lymphocyte

00:17:57

- Constitute around **15-20%** of circulating blood lymphocyte.
- They are responsible for humoral immunity, i.e. produce antigens against **extracellular microbes**.
- These cells mature in the **bone marrow**.
- Common sites of B lymphocytes :
 1. Cortex of the lymph node.
 2. Peyer's patches in GIT.
 3. White pulp of spleen.
- In cases of B lymphocytic defect like **Bruton agammaglobulinemia**, the above mentioned sites would be **atrophic**.
 bnvssprasanth7@gmail.com
- markers for B lymphocytes :
CD 10 or **CALLA**, **CD 19**, **CD 20**, **CD 21**, **CD 22**, **CD 23**, **Ig α** (CD 79a), **Ig β** (CD 79b).
CD 79a and **CD 79b** are also signal transduction molecules present on the surface of B lymphocyte.
- Pan B cell marker : **CD 19**.
- Receptor for EBV on B cell : **CD 21**.

Infections caused by EBV :

- **Infectious mononucleosis**.
- **Hodgkins lymphoma**.
- **Burkitts lymphoma**.
- **Non hodgkins lymphoma**.

Active space

- Nasopharyngeal carcinoma.
- Post transplant lymphoproliferative disorders.

B cell receptors :

These are IgM or IgD antibody along with signal transduction molecules.

mechanism of activation of B lymphocytes :

This activation can occur by 2 pathways :

- T cell independent pathway :

This is activated when the antigen is a lipopolysaccharide. This antigen activates B cell which forms a plasma cell, which produces IgM.

- T-cell dependant pathway :

This pathway is activated when a proteinaceous antigen is present.

The B cell has CD 40 receptor & helper T-cell has CD 40L. The antigen first interacts with the CD 40L and forms IgM and IgD antibodies, which then undergo class switching mechanism to form IgG, IgA and IgE.

IgG : Crosses placenta.

IgA : Present in all body secretions.

IgE : most critical antibody in type I hypersensitivity reaction.

IgM : It has the highest molecular weight, pentameric structure and is called as millionaire's antibody.

- The helper T cell also forms IL-4 and IFN- γ .

Increased IgM secretion disorder is seen in : Waldenstrom macroglobulinemia/hyperviscosity syndrome.

Plasma cell disorder which produces abnormal immunoglobins which are monoclonal : multiple myeloma.

T lymphocytes

00:27:22

They are responsible for cell mediated immunity and constitute 60 to 70% circulating lymphocyte.

There are 2 types of T-cells :

CD 4 & CD 8 T cells (ratio 2 : 1).

This ratio is decreased in HIV patients and increased in sarcoidosis patients.

These cells mature in the thymus and are found at :

- Paracortex of the lymph node.
- Periarteriolar lymphoid sheath.
- Intraepithelial lymphocytes.

In a patient with T cell disorder, these sites undergo hyperplasia.

markers of T cell :

- CD 1, 2, 3, 4, 5, 7, 8.
- Pan T cell marker : CD 3.

T cell receptors are of 2 types :

- $\alpha\beta$: Present on 95% of cells and is a polypeptide which is MHC restricted.
- $\gamma\delta$: Present on 5% of cells and provides protection against the microbes which try to enter through the epithelial barriers.

Helper T cell	Cytotoxic T cell
CD 4+ T cell.	CD 8+ T cell.
MHC II restricted.	MHC I restricted.
1 st line of defence in the body.	2 nd line of defence in the body.
It helps the B cell in producing the antibodies.	It directly kills the infected cell by perforin granzyme mechanism.
It helps in the activation of macrophages.	

Active space

The 3 types of helper T cells :

- TH 1 : Cytokines produced are IFN- γ (signature cytokine produced by TH 1) and IL-12.
IFN- γ helps in the activation of macrophages and helps in the production of IgG antibody.
It also helps in the fight against intracellular microbes.
- TH 2 : It produces 3 other antibodies, IL-4, IL-5 and IL-13.
IL-4 : Causes increased production of IgE antibody and plays a role in activation of macrophages.
IL-5 : Helps in the production of eosinophils and helps in the activation of mast cells.
IL-13 : Activation of macrophages.
TH 2 lymphocyte helps in fighting helminthic infections.
- TH 17 : Leads to the production of 2 cytokines, IL-17 and IL 22.
They play a role in recruitment of neutrophils and macrophages.
TH 17 helps in fighting against extracellular microbes.

HYPERSENSITIVITY REACTIONS

Types of hypersensitivity reactions

00:00:52

Type I
Type II
Type III } Antibody mediated hypersensitivity reaction.

Type IV → Cell mediated hypersensitivity reaction

Type V → modification of Type 2 hypersensitivity reaction.

Type I hypersensitivity reaction

00:01:28

AKA Anaphylactic hypersensitivity (HS) reaction.

Examples :

mnemonic : **ABCD**

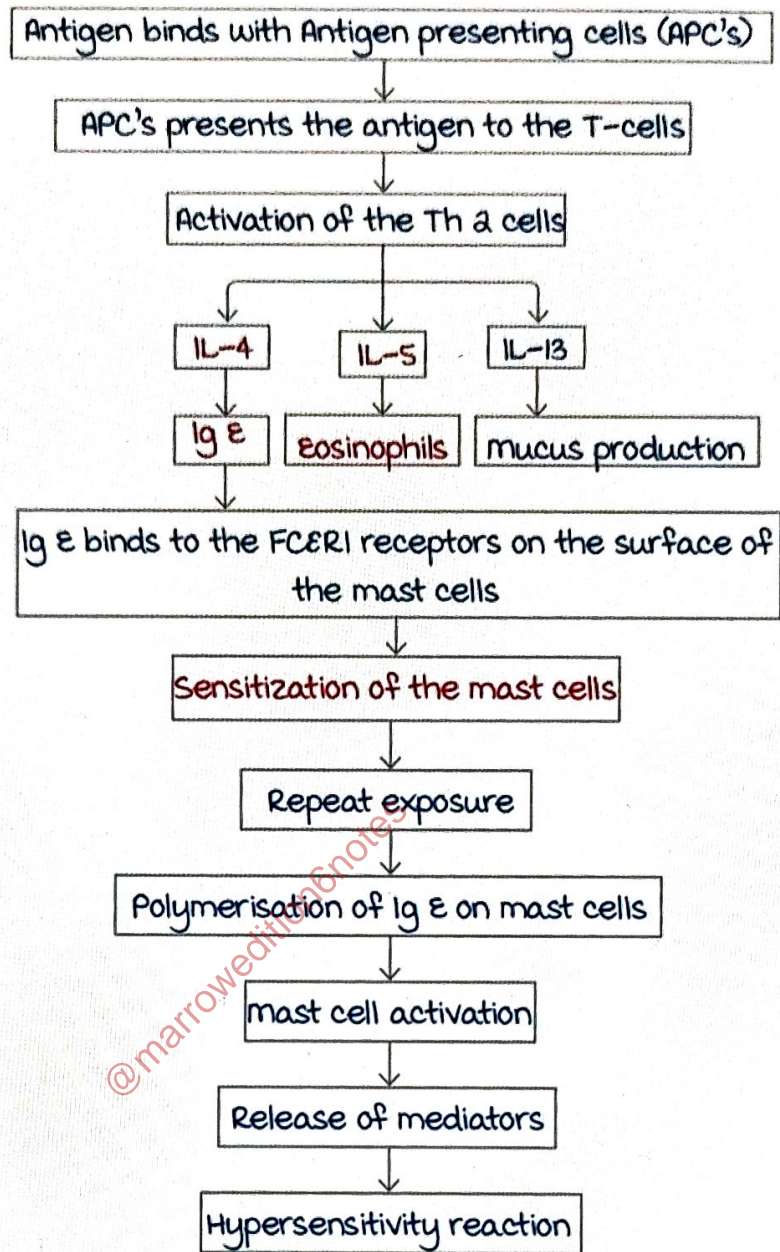
- **A**topy/**A**llergies/**A**naphylaxis (food/pollen allergies).
Atopy → Genetic determination of allergy.
Gene for atopy is located on chromosome 5.
- **B**ronchial asthma.
- **C**asoni's test (Hydatid disease).
- **D**rug reactions.
- Hay fever.
- PK reactions.
- Theobald Smith phenomena.

mechanism :

1st exposure of the antigen → mast cells get sensitised →

Repeated exposure of the antigen → Hypersensitivity reaction occurs.

Active space



Types of mediators released :

- **Preformed mediators** (from the stored granules of the mast cells) :
 1. **Histamine** (earliest mediator to be released).
 2. Proteases/enzymes.
 3. Chemotactic factors : Like C3a and C5a.
- Activation of **Phospholipase A₂** : Production of arachidonic acid metabolites like prostaglandins, leukotriens and platelet activating factor.

Phases :

- Immediate phase → Occurs within minutes.
 1. Vasodilation.
 2. Increased vascular permeability.
 3. Increased mucus production.
- Late reaction → Occurs within 2-24 hours.
 1. Fibrosis.
 2. Increased production of inflammatory cells.
 3. Epithelial damage.

most important cell in type I HS reaction → **mast cell**.

Stain for mast cell → **Toluidine blue**.

Important cell in the late phase of type I HS reaction :
Eosinophils.

most important antibody in type I HS reaction : **IgE**.

most important cytokine in type I HS reaction : **IL-4 & IL-5**.

Earliest mediator released in type I HS reaction : **Histamine**.

Type II hypersensitivity reactions

00:13:48

Antibody mediated HS reaction.

Examples :

Mnemonic : **My Blood Group Is RH Positive**.

- **Myasthenia gravis**.
- **Blood transfusion reaction**.
- **Grave's disease, Good pasteur syndrome**.
- **Idiopathic thrombocytopenic purpura (ITP), Immune hemolytic anemia**.
- **Rheumatic fever**.
- **Hyperacute graft rejection**.
- **Pernicious anemia, Post-Streptococcal Glomerulo Nephritis (PSGN)**.

Mechanism :

- **Opsonisation and phagocytosis**.
- **Inflammation and Complement activation**.
- **Antibody-dependent cell-mediated cytotoxicity (ADCC)**.

Active space

Opsonisation and phagocytosis :

It is mediated by **IgG** antibody (F_c portion of IgG is the most potent opsonin), in which IgG antibody coats the antigenic cell due to which macrophages can phagocytose these cells easily.

Examples :

- Hemolytic disease of the newborn.
- Blood transfusion reaction.
- Drug reaction.

Inflammation and complement activation :

- Occurs when the antibody is bound to the surface of the **basement membrane** or the **extracellular matrix**.
- The antibody will lead to activation of the complement factors causing neutrophil chemotaxis and then tissue injury.

Examples :

- Good pasteur syndrome.
- Glomerulonephritis.
- Graft rejection.

ADCC/Antibody dependent cell mediated cytotoxicity :

- There is no complement activation or tissue injury.
- Antibodies are produced against the cell-surface receptors.

Examples :

- Grave's disease → Anti-TSH receptor antibody.
- myasthenia gravis → Antibody against the ACh receptors.
(Recently classified under **type V HS** reaction) (Type V > II)

Type III hypersensitivity reactions

00:22:30

AKA Immune complex mediated HS reaction.

Examples → mnemonic : **SHARP**

- **S**erum sickness, **S**hick test, **S**LE (visceral lesion is type 3 HS reaction and hematological lesion is type 2 HS reaction)
- **H**enoch Schonlein purpura.
- **A**rthus reaction.

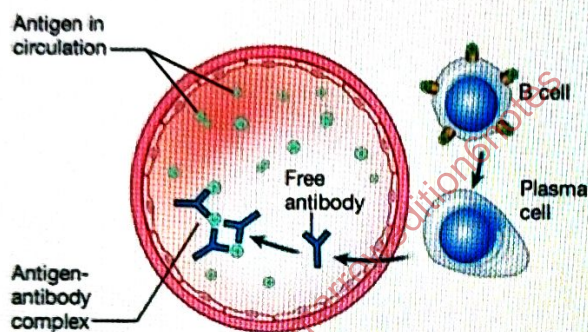
- Reactive arthritis.
- PSGN, Poly arteritis nodosa (PAN).

Formation of immune complex takes approximately 5-7 days.

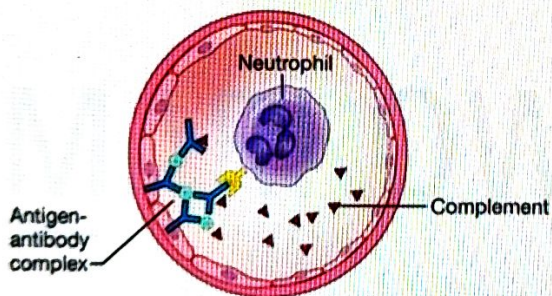
Immune complex deposition :

- most pathogenic immune complexes are small to medium-sized which usually have excess of antigens.
- Immune complexes are usually deposited in organs which have a high filtration rate like kidneys or the joints.
- Immune complex-mediated inflammation and tissue injury takes approximately 10-14 days.

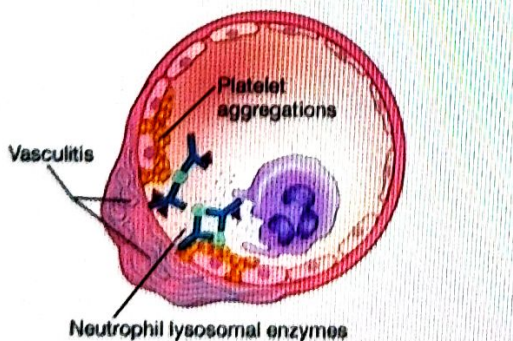
Immune complex disease- Type III Hypersensitivity Reactions



Immune complex formation



Immune complex deposition



Immune complex-mediated inflammation and
tissue injury

Active space

Type IV hypersensitivity reactions

00:28:08

AKA cell mediated HS reaction/delayed type HS reaction.

Examples :

- Granuloma formation.
- Tuberculin test.
- Lepromin test.
- Contact dermatitis.
- Sarcoidosis.
- Multiple sclerosis.
- Rheumatoid arthritis : Type 4 > type 3.
- Hypersensitivity pneumonitis : Both type 3 and type 4 HS reaction.
- Acute and chronic graft rejection.

Hyperacute graft rejection is due to type 2 HS reaction.

Clinical scenarios

00:35:08

Q. A 15-year-old healthy girl with no major medical problems notes blotchy areas of erythema that are pruritic over the skin of her arms, legs, and trunk within an hour every time she eats seafood, followed by diarrhea. These problems abate within 3 hours, and then physical examination reveals no abnormal findings. Which of the following immunologic abnormalities is she most likely to have?

- Localized anaphylaxis.
- Cell-mediated hypersensitivity.
- Complement activation.
- Hypergammaglobulinemia.
- Immune complex deposition.

Q. Twelve hours after going on a hike through dense foliage, a 40-year-old man notices a slightly raised and tender irregular reddish rash on one forearm that was not covered by clothing. This rash gradually increases in intensity for a days and then fades away after two weeks. Which of the following forms of immunologic hypersensitivity is most likely demonstrated in this patient?

- A. Type I hypersensitivity.
- B. Type II hypersensitivity.
- C. Type III hypersensitivity.
- D. Type IV hypersensitivity.

Explanation : Hypersensitivity pneumonitis occurs on exposure to foliage/ moldy hay/bird poop etc.

Q. A 30-year-old woman has experienced myalgias for the past 3 months. On physical examination she has 5/5 motor strength in all extremities. She has dullness to percussion at lung bases. A chest x-ray shows bilateral pleural effusions. Laboratory studies show a positive antinuclear antibody test at a titer of 1:1024. Her serum urea nitrogen is 30 mg/dL. A renal biopsy is performed and microscopic examination shows a granular pattern of immunofluorescence staining with antibody to complement component C1q. This pattern is most typically produced as a consequence of which of the following immunologic mechanisms?

- A. IgE coating mast cells.
- B. Antiglomerular basement membrane antibody.
- C. Antigen-antibody complexes.
- D. Macrophage release of lymphokines.
- E. Release of prostaglandins.

Explanation : SLE (visceral component). Type III reaction

Q. A 48-year-old man has had a chronic cough with fever for 2 months. On physical examination his temperature is 37.9°C. A chest radiograph reveals a diffuse bilateral reticulonodular pattern. A transbronchial biopsy is performed and microscopic examination shows focal areas of inflammation containing epithelioid macrophages, Langhans giant cells, and lymphocytes. These findings are most typical for which of the following immunologic responses?

- A. Type I hypersensitivity.
- B. Type II hypersensitivity.
- C. Graft versus host disease.
- D. Polyclonal B-cell activation.
- E. Type IV hypersensitivity, TB granuloma.

Q. A 9-year-old boy has a sore throat. A throat culture grows group A hemolytic Streptococcus. He receives antibiotic therapy. However, 17 days later he develops dark-coloured urine. Laboratory studies show 3+ blood on urinalysis. A renal biopsy is performed. On immunofluorescence staining the biopsy shows granular deposition of IgG and complement around glomerular capillary loops. Which of the following immune hypersensitivity mechanisms is most likely responsible for this pattern of findings?

- A. Type I.
- B. Type II.
- C. Type III. PSGN
- D. Type IV.

@marroweditionsnotes

Major histocompatibility complex (MHC)

00:01:26

It is A/K/A Human leucocyte antigen (HLA).

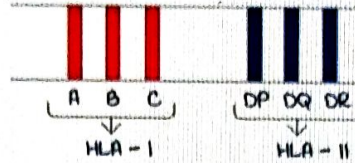
The gene is located on Chromosome 6p.

MHC is of 3 types :

MHC I.

MHC II.

MHC III : Encoded by Heat shock protein (HSP), Complement proteins, properdin. It may have a role in autoimmune diseases.



HLA I	HLA II
<ul style="list-style-type: none"> Present on all nucleated cells & platelets. Encoded by A, B, C. Presents the antigen to CD8+ T lymphocytes. Role in graft rejection. Structure : β_2 microglobulin is present. Peptide binding cleft between α_1 and α_2. 	<ul style="list-style-type: none"> Present only on antigen presenting cells (APCs - B cells, fibroblasts, dendritic cells). Encoded by DP, DQ, DR. Presents the antigen to CD4+ T lymphocytes. Role in GVHD. Structure : Peptide binding cleft is between α_1 and β_1.
<p>The diagram shows the structure of an HLA I molecule. It consists of two heavy chains, α_1 and α_2, which form a peptide binding cleft. A third chain, α_3, is attached to the α_1 chain. A β_2-microglobulin molecule is associated with the α_3 chain. The entire structure is anchored into the cell membrane.</p>	<p>The diagram shows the structure of an HLA II molecule. It consists of two heavy chains, α_1 and α_2, and two light chains, β_1 and β_2. The α_1 and β_1 chains form a peptide binding cleft. The α_2 and β_2 chains are attached to the α_1 and β_1 chains respectively. The entire structure is anchored into the cell membrane.</p>

Role of MHC

00:10:20

Paternity testing.

Prediction of incidence of autoimmune disorders.

HLA B27 : Ankylosing spondylitis.

HLA DR3, DR4 : Diabetes mellitus.

HLA DQ2, DQ8 : Celiac disease.

HLA B5, B51 : Behcet's disease.

Anthropology testing.

HLA matching in Organ transplantation and Bone marrow/ stem cell transplantation.

most important HLA which needs to be typed : HLA-DR.

All 6 loci match only in case of identical twins.

HLA A, B, DR : Should definitely match with each other (DR > B > A).

All the loci have 2 alleles each.

matching is expressed as 12/12 for all loci or 6/6 for HLA A, B, DR.

HLA matching is not done for : Cornea, Liver, Heart, Lung transplant.

Grafts and Graft reactions

00:17:28

- Isograft : Between identical twins.
- Autograft : From one part of one's own body to another.
- Allograft : Between genetically different individuals but same species.
- Xenograft : Between different species.
- Orthoptic graft : Graft is placed in the same anatomic location as the donor.
- Heterotopic graft : Different anatomic location from the donor.

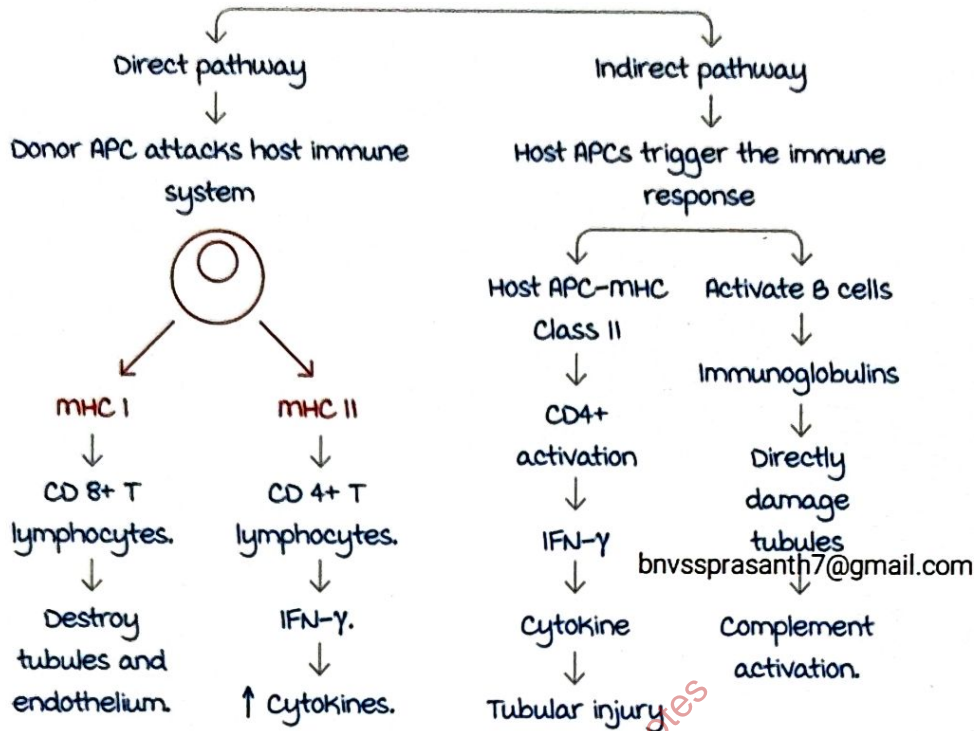
Graft reactions :

Graft rejection	Graft vs Host disease (GVHD)
Host is immunocompetent. Host cells attack graft cells.	Host is immunosuppressed. Graft cells attack host cells. Seen in bone marrow transplantation.

Mechanism of graft rejection

00:23:50

Example : Kidney transplantation.



Hyperacute graft rejection

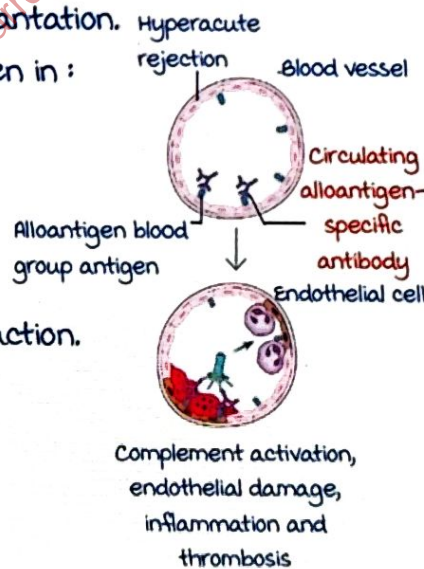
00:29:00

Occurs within minutes of transplantation.

Due to preformed antibodies seen in :

- Previous pregnancy.
- ABO & Rh incompatibility.
- Previous blood transfusion.
- Previous transplantation.

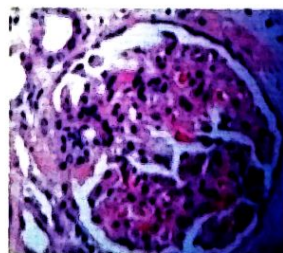
It is a type II hypersensitivity reaction.



Gross appearance of kidney :
Cyanosed, mottled, flaccid.

microscopically :

- Fibrinoid necrosis.
- microthrombi.
- Neutrophilic infiltrate.



Prevention : Donor specific antibody test should be done.

Active space

Acute graft rejection

00:36:35

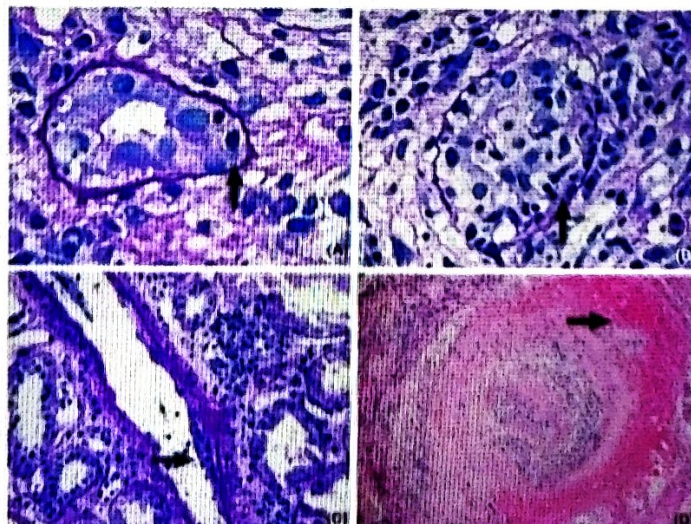
Acute cellular rejection	Acute humoral rejection
<p>mediated by CD4+ or CD8+ T cells.</p> <p>Type IV hypersensitivity reaction.</p> <p>Responsive to increasing dose of immunosuppressive drugs.</p> <p>microscopic appearance :</p> <ul style="list-style-type: none"> • Tubulointerstitial pattern : • Tubulitis : Destruction of tubules + inflammatory cells in tubules. mononuclear inflammatory infiltrate. • vascular pattern : Endothelitis 	<p>mediated by newly synthesized antibodies.</p> <p>It causes endothelial damage & complement activation.</p> <p>Type II or III hypersensitivity.</p> <p>No response to increasing dose of immunosuppressive drugs.</p> <p>Treatment : B cell depleting agents.</p> <p>microscopic appearance :</p> <ul style="list-style-type: none"> • Fibrinoid necrosis in vessels. • Deposition of C4d in peritubular capillaries. (Complement breakdown product). <p>C4d is the marker for acute humoral rejection. bnvssprasanth7@gmail.com</p>

Chronic rejection

00:44:20

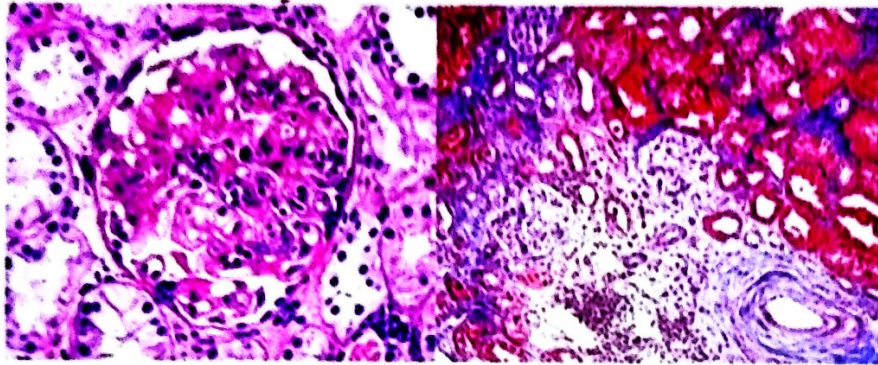
It is the m/c type of graft rejection.
 It occurs within months to years of transplantation.
 It may be cell mediated (Type IV hypersensitivity) or antibody mediated (Type II).

microscopically :



Active space

- Transplant glomerulopathy.
- Duplication of glomerular basement membrane.
- Interstitial fibrosis.
- Glomerular sclerosis.
- Tubular atrophy.



Glomerulus-inflammatory cells within the capillary loops (glomerulitis), accumulation of mesangial matrix, & duplication of capillary basement membrane.

Interstitial fibrosis and tubular atrophy. (trichrome stain), contrasted with the normal kidney. Artery-prominent arteriosclerosis

Graft vs host disease (GVHD)

00:49:33

GVHD is a complication of hematopoietic stem cell transplantation.

It is A/K/A Runt's disease in animals.

Type IV hypersensitivity reaction.

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Acute GVHD	Chronic GVHD
<p>< 100 days duration. Organs affected : Skin : Excoriation. GIT : mucosal ulceration → Diarrhoea. Liver : Jaundice.</p>	<p>> 100 days duration. Organs affected : Skin : Scleroderma. GIT : Strictures. Liver : Cirrhosis.</p>

Y linked graft rejection :

A/K/A sex linked graft rejection (Eichwald silmsen effect).

It occurs when male gives graft to a female.

(Y chromosome contains UTY gene → encodes for enzyme histone demethylase : minor histocompatibility antigen).

Active space

Complications of transplants

00:55:24

1. Infections :

CMV is the m/c following transplantation : Owl's eye inclusions.

BK polyoma virus infection : Decoy cells.

2. Graft rejection.

3. GVHD.

4. Increased risk of malignancy :

Squamous cell carcinoma (m/c) : HPV associated.

Kaposi's sarcoma : HHV-8 associated.

Non-Hodgkin's lymphoma : EBV associated.

Post-transplant lymphoproliferative disorder :

EBV associated (Poor prognosis).

MCQs :

Q. A patient has to receive liver transplant from his brother, who is not his twins. On HLA typing, HLA matched are the A, B and DRB1 locus. These siblings are considered as :

- A. matched, unrelated donors
- B. mismatched, related donors
- C. matched, related donors
- D. mismatched, unrelated donors

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Q. Which of the following statements about graft vs host disease is least correct?

- A. Occurs when host is immunocompromised.
- B. Occurs when donor cells are immunocompromised.
- C. Is also called RUNT disease.
- D. A common cause is stem cell transplantation.

Q. Acute humoral renal transplant rejection is characterized by the following except:

- A. Presence of anti donor antibodies.
- B. Necrotizing vasculitis.
- C. Interstitial and tubular mononuclear cell infiltrate.
- D. Acute cortical necrosis.

Q. most commonly involved organs in graft versus host disease are all except:

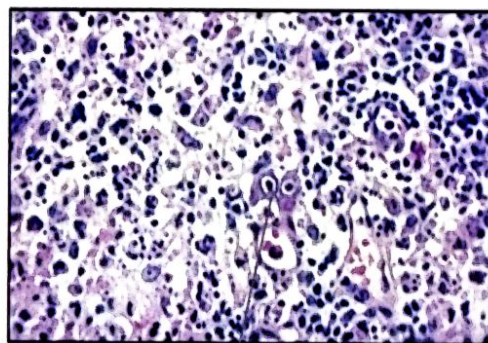
- A. Gut
- B. Liver
- C. Skin
- D. Kidney

Q. A 25 year old female with CRF receives a cadaveric renal transplant. One month later, she experienced increasing creatinine and urea levels and a renal biopsy was performed. She was treated with steroids and her renal function improved. Which of the following changes was most likely seen in the biopsy specimen before steroid therapy was initiated?

- A. Interstitial infiltration by CD3+ lymphocytes and tubular epithelial damage.
- B. Extensive fibrosis of interstitium and glomeruli with marked thickening blood vessels.
- C. Fibrinoid necrosis of renal arterioles and thrombi.
- D. Glomerular deposition of serum amyloid associated protein.

Q. A 30 year old patient who had undergone a renal transplant presented with fever and dyspnea. The histopathological examination from a lung lesion is given below. What is the most likely diagnosis?

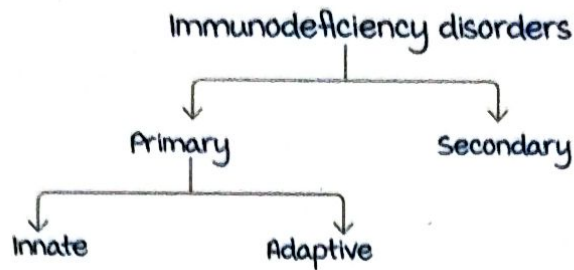
- A. Mycobacterium
- B. BK polyoma virus
- C. Herpes infection
- D. CMV



Owl's eye inclusions

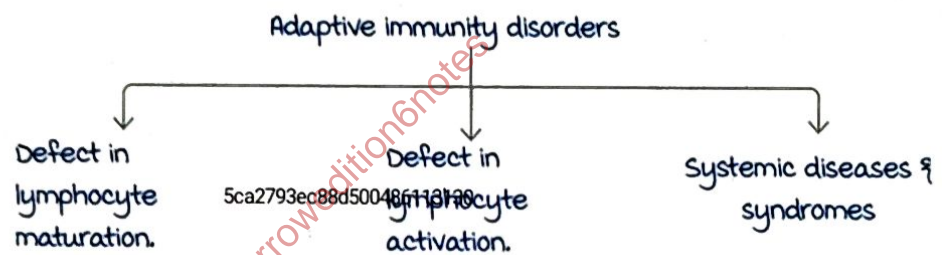
Active space

IMMUNODEFICIENCY DISORDERS



Secondary immunodeficiency causes :

- HIV.
- Cancer.
- Chemotherapy.
- **malnutrition** (most common secondary cause).



Defect in lymphocyte maturation

00:03:30

Bruton's agammaglobulinemia :

X linked recessive.

Seen in boys >> girls.

Pathogenesis :

Due to **BTK gene** (Bruton tyrosine kinase) defect → Defective maturation of B cell lymphocytes → Decreased mature B cells and plasma cells → **Defective humoral immunity.**

T cells are normal → Cell mediated immunity is intact.

Histology :

Hypoplastic / absent germinal centers.

Active space

Clinical presentation :

usually manifests after 6 months of age.

Recurrent sino-pulmonary infections / infections with enterovirus or Giardia.

Diagnosis :

Flow cytometry :

Presence/absence of surface Ig can be seen.

In bruton's agammaglobulinemia → Absence of surface Ig.

DiGeorge syndrome :

Also known as 22q 11 deletion syndrome / velocardiofacial defects.

Defect : Deletion of 22q 11 → defect in **TBX 1 gene** →

defective development of 3rd & 4th pharyngeal pouches →

Defective development of thymus & parathyroid gland →

Defective T cell development and Hypocalcemia.

mnemonic : **CATCH 22**

Cleft lip and palate.

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Abnormal facies.

T cell defect, thymic hypoplasia.

Cardiac abnormality (**MC defect** : TOF).

Hypocalcemia.

22q 11 deletion.

SCID (Severe Combined Immuno Deficiency) :

Defect in B cell, T cell and NK cell.

Pathogenesis :

2 modes of inheritance :

- X-linked recessive : (**MC**).

males >>> females.

mutation in common γ chain of cytokine receptors →

Reduced synthesis of IL 2, 4, 7, 11 and 15.

Decreased production of :

1. IL4 → Since responsible for isotype switching :
Decreased production of immunoglobulins.
2. IL7 → Decreased levels of T cell lymphocytes.

3. IL15 → Defect in NK cells.

- **Autosomal recessive** : Deficiency of **Adenosine deaminase (ADA)** → Accumulation of toxic metabolites → Destruction of B cells, T cells and NK cells.

Clinical presentation :

Can present with any kind of infection. (viral / protozoal / fungal / bacterial).

Candidial infection / diaper rashes can be seen.

Treatment :

First disease to be treated with gene therapy.

Hematopoietic stem cell transplantation.

Defect in lymphocyte activation

00:17:08

Hyper IgM syndrome :

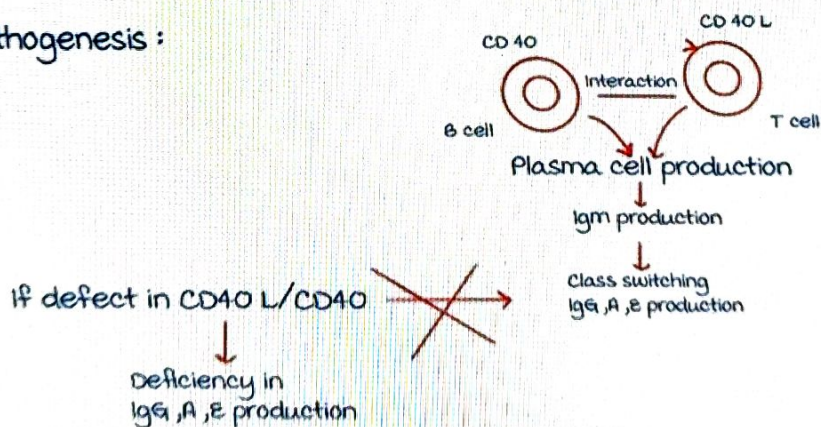
Increased production of IgM.

Decreased level of **IgG, A, E**.

X Linked Recessive disorder.

males >>> females (more common).

Pathogenesis :



most common defect in hyper IgM : CD40L defect.

2nd most common defect in hyper IgM : CD40 defect.

Therefore, **defect in class switching.**

Clinical Features :

- Increased production of IgM : **Autoimmune thrombocytopenia.**

Active space

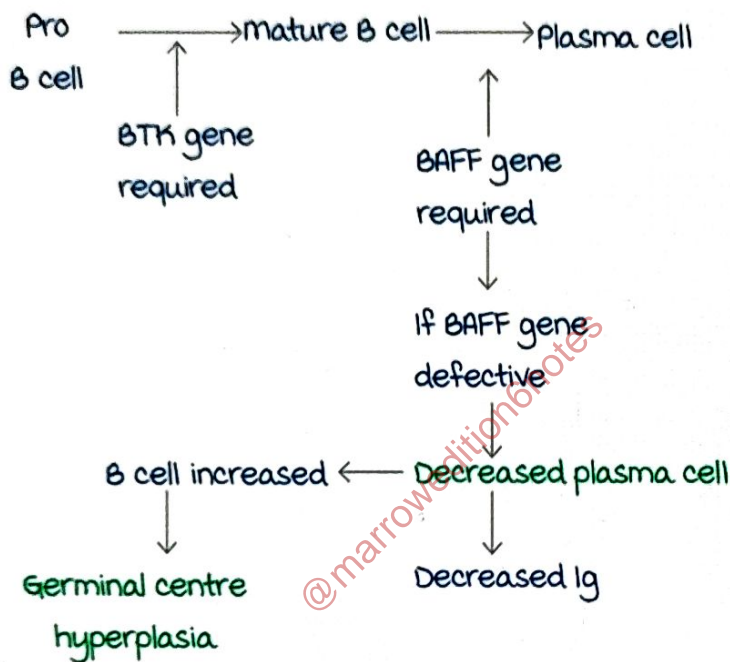
Autoimmune hemolytic anemia.
Autoimmune neutropenia.

- Decreased the level of Ig G, A, E :
Increased risk of **sinopulmonary infections**.

Common variable immunodeficiency :

It is a diagnosis of exclusion.

Pathogenesis : mutation in the **BAFF gene**.



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Isolated Ig A deficiency :

most common primary immuno deficiency disorder.

Decreased production of Ig A, G_2 , G_4 .

Clinical presentation :

Increased risk of sinopulmonary infection.

Increase the risk of **anaphylactic reaction** (can present in a person undergoing blood transfusion for the first time).

Wiscott Aldrich syndrome

00:24:58

X linked Recessive disorder.

more common in males >>> females.

Pathogenesis : **WASP gene** defect on chromosome **XP 11.2**.

Active space

Clinical Triad :

- Eczema.
- Thrombocytopenia (Decreased platelets, small sized platelets).
- Immuno deficiency due to defective T cell development.

Decreased IgM, increased IgA and normal IgG.

Mnemonic : WAITER

Wiscott, WASP gene defect.

Aldrich.

Immunodeficiency.

T cell deficiency, Thrombocytopenia

Eczema.

Recurrent infections.

Ataxia telangiectasia

00:28:19

Autosomal recessive disorder.

Incidence : male = female .

Pathogenesis :

Defect in ATM gene on chromosome 11 (WT gene for Wilms' tumor is also present on chromosome 11).

Normal ATM gene acts a DNA repair sensor and activates P53 if any DNA damage occurs.

Defective ATM gene does not activate P53 and may cause :

- Ataxia telangiectasia.
- malignancy.
- Premature aging.
- Neurodegenerative disorders.

Clinical scenarios :

Q. A male infant is born at term. No congenital anomalies are noted on examination. A year later he has failure to thrive and has been getting one bacterial pneumonia after another with both Hemophilus influenzae and Streptococcus pneumoniae cultured from his sputum. Which of the following diseases is he most likely to have?

- DiGeorge syndrome.
- Selective IgA deficiency.

- C. Epstein-Barr virus (EBV) infection.
- D. Acute leukemia.
- E. X-linked agammaglobulinemia.

Digeorge syndrome is a birth defect with abnormal facies & cleft lip and palate.

Selective IgA deficiency will not manifest after a year & doesn't produce the symptoms mentioned here.

Q. A 5 year old boy and his 4 year old brother have had recurrent pneumonia, meningoencephalitis, sinusitis, otitis, & diarrhea since infancy. Bacterial & viral agents have been implicated, as well as Pneumocystis, Cryptosporidium & Giardia. Laboratory studies show serum IgG 47 mg/dL, IgA 5 mg/dL, and IgM 671 mg/dL. Normal numbers of B and T cells are present. These children are most likely to have a mutation involving a gene encoding for which of the following?

- A. NADPH oxidase.
- B. Wiskott-Aldrich syndrome protein.
- C. Cytokine receptor common gamma chain.
- D. CD40 L.
- E. Complement C1 inhibitor.

Explanation :

IgM 671 mg/dl could suggest hyper IgM syndrome.

CD40 L is the MC defect here.

NADPH oxidase deficiency causes granulomatous disease.

Wiskott-Aldrich syndrome protein will have a history of eczema, small sized platelets etc.

Cytokine receptor common gamma chain is X linked recessive.

Complement C1 inhibitor seen in hereditary angioneurotic edemas.

Q. An 11 month old infant has had upper and lower respiratory tract infections almost continuously since the time of birth, with organisms including Pneumocystis jiroveci & Pseudomonas aeruginosa identified. The baby also has oropharyngeal candidiasis. The baby succumbs to a cytomegalovirus pneumonitis. At autopsy, the thymus is

Active space

markedly hypoplastic and lymph nodes throughout the body are small, with absent germinal centers on microscopic examination. Which of the following mechanisms is most likely to explain these findings?

- A. Adenosine deaminase deficiency.
- B. Failure of B cell maturation to plasma cells.
- C. Human immunodeficiency virus infection.
- D. Autoantibodies to both T and B lymphocytes.
- E. Failure of development of 3rd and 4th pharyngeal pouches.

Explanation :

Adenosine deaminase deficiency is seen in AR , SCID.

Failure of development of 3rd and 4th pharyngeal pouches seen in DiGeorge syndrome .

Q. A neonate born at term developed tetany soon after birth. On physical examination the infant has a heart murmur. Laboratory studies show a serum calcium of 6.3 mg/dL. Echocardiography reveals a membranous intraventricular septal defect. Within the next year, this infant has bouts of *Pneumocystis jiroveci* pneumonia, *Aspergillus fumigatus* pneumonia, and parainfluenza virus and herpes simplex virus upper respiratory infections. Which of the following abnormalities most likely explains the development of this infant's findings?

- A. Abnormal Wiskott-Aldrich syndrome protein.
- B. 22q-chromosome deletion.
- C. Reduction in CD4 lymphocytes.
- D. Defect in NADPH oxidase.
- E. Failure of B cell maturation into plasma cells.

The symptoms are suggestive of DiGeorge syndrome.

AMYLOIDOSIS

Introduction

00:01:10

Pathologic proteinaceous extracellular hyaline eosinophilic substance deposited in various tissue and organs.

misfolded protein.

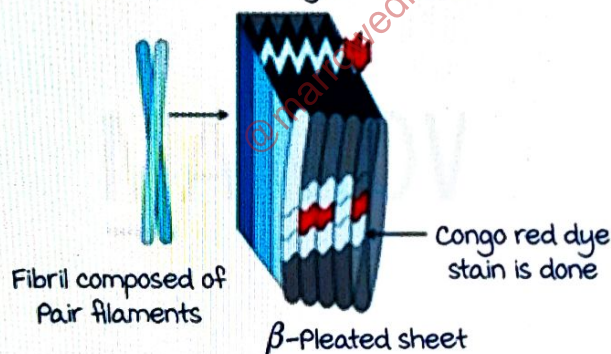
Hyaline : Pink substances.

Physical nature :

In electron microscopy, appear as non branching fibrils of indefinite length with 7.5-10 nm diameter.

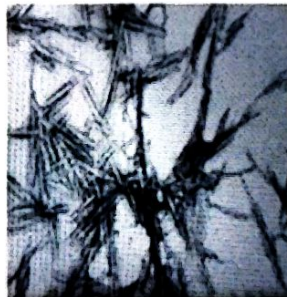
In X-ray crystallography (or infrared spectroscopy) : Cross beta pleated sheet structure (this is responsible for apple-green birefringence under polarized lens).

Structure of Amyloid material



bnvssprasanth7@gmail.com

Amyloid in electron microscopy with 7.5-10 nm diameter



Best stain : Congo red.

Chemical nature :

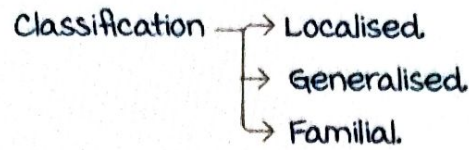
Consists of

- Amyloid protein : 95%
- P protein : 5% (depends on the disease condition)

Active space

Classification of Amyloidosis

00:07:00

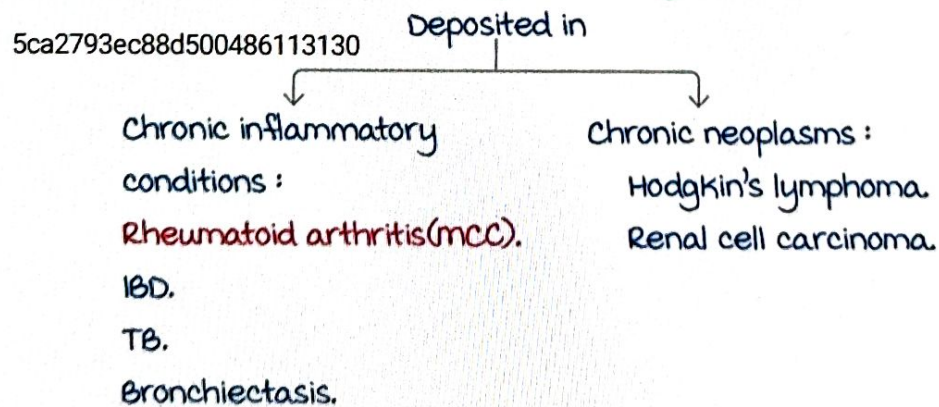


1. Localised amyloid and its protein :

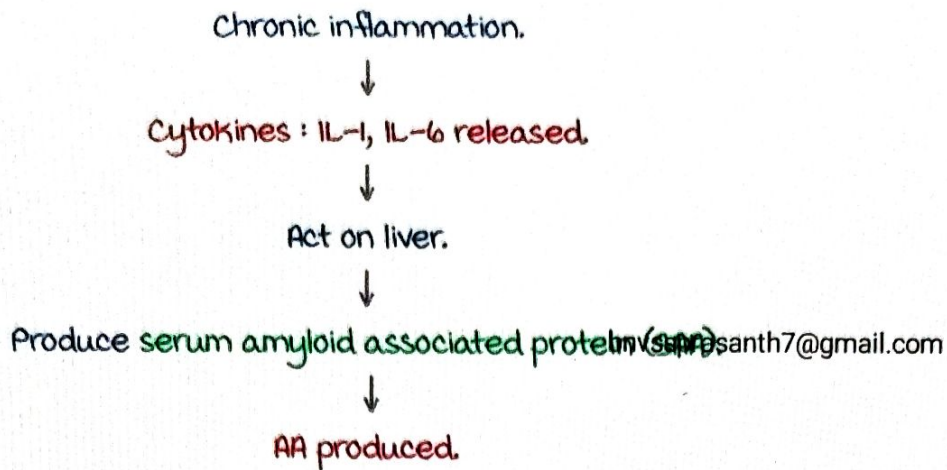
- medullary carcinoma of thyroid :
Amyloid seen is **Acal** (95% amyloid protein & 5% calcitonin protein).
- Prion disease :
Apr (95% amyloid protein & 5% prion protein).
- Type 2 Diabetes mellitus :
Amyloid islet associated pancreatic polypeptide (AIAPP).
- Alzheimer's disease :
ABeta (part of neuritic plaque).

2. Generalised : multiple organ involved.

- Primary amyloidosis :
most common type of amyloidosis.
Seen in light chain disorders like multiple myeloma.
AL (amyloid & light chain called lambda light chains commonly deposited).
In mm, **accumulation of immunoglobulins** hence light chain are precipitated.
most common cause of death : Cardiac failure.
- Secondary amyloidosis :
Also called **reactive systemic amyloidosis**.



Active space



most common cause of death : Renal failure.

3. Amyloid seen in chronic renal failure (CRF)/ long term hemodialysis :

uncommon

Dialysis membranes (DM) filters out the unwanted substances out.

Earlier the DM did not filter out β_2 microglobulin , hence it accumulated leading to A_{β_2m} amyloid formation.

Deposited in joints, tendons, median nerve leading to Carpal tunnel syndrome.

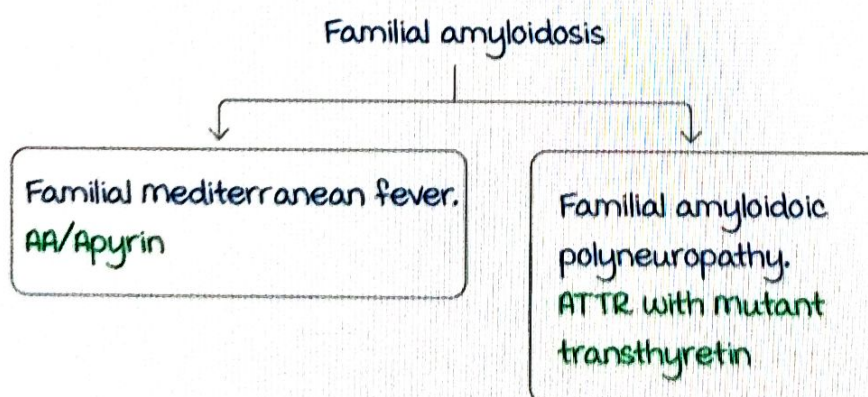
4. Senile/ Cardiac amyloidosis :

ATTR (transthyretin).

In aged/ cardiac patients normal transthyretin is deposited.

Familial amyloidosis

00:18:08



Active space

Disease / condition	Type of amyloid
Primary amyloidosis	AL
Secondary amyloidosis	AA
Familial mediterranean fever	AA/Apyrin
Familial amyloidotic polyneuropathy	ATTR (mutant transthyretin)
Senile/ cardiac amyloidosis	ATTR
CRF/ long term dialysis	Abetaam
Prion disease	Apr
Diabetes mellitus	AIAPP
medullary ca thyroid	Acal
Alzheimer's disease	Abeta

Diagnosis :

Biopsy(Bx) site :

If localised : Bx taken form that localised tissue.

If generalised : Bx

- Abdomen fat pad aspirate (best).
 - Rectal (not best as painful & invasive).
 - Gingival (usually macroglossia seen)
- Abdomen fat pad > rectal > gingival.

Bx is H & E stained :

In kidney Bx

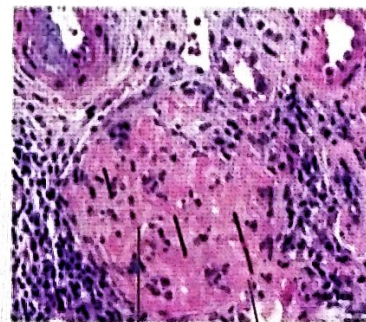
Extracellular : pink.

Glomeruli with pinkish material that is amyloid.

Eosinophilic hyaline pink :

Can be sclerosis/fibrin/

hyaline : So to differentiate



Amyloid Glomeruli

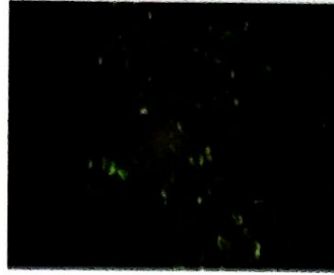
5ca2793ec88d500486113130

Active space

Amyloid Congo red is used.

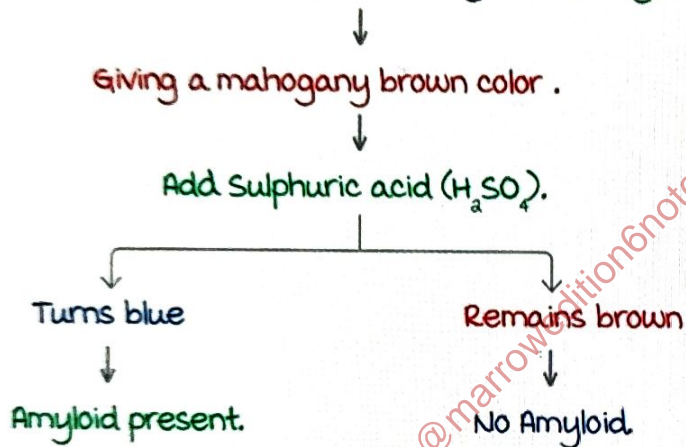
Stains used :

- Congo red (Best).
Congo red gives Salmon pink colour but under polarised lens apple-green birefringence seen.
- PAS :
Appear magenta colour.
- methyl violet/ crystal violet : metachromatic stain.
- Thioflavin S } shows immunofluorescence.
- Thioflavin T }



Gross stain :

Paint the cut surface of the organ with Lugol's iodine.



Organ involvement of amyloid

00:28:46

Gross :

Any organ with amyloidosis. → Organomegaly with waxy appearance.

Organs :

- Kidney (most common organ affected).

Affects : Glomeruli,

Tubules,

mesangium (earliest to involve).

Clinically manifested with nephrotic syndrome.

- Liver :

Earliest part affected is space of Disse.

Produces pressure atrophy of hepatocytes cirrhosis.

Active space

- Heart :
Earliest part affected is subendocardium.

Can lead to :

- Arrhythmia
- Heart Failure.

Restrictive Cardiomyopathy.

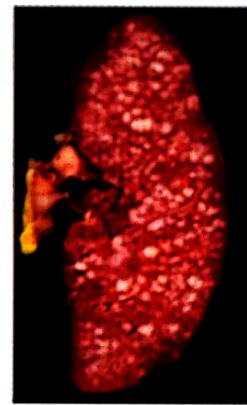
Normal ATTR deposited.

- Skin : bnvssprasanth7@gmail.com
Pinch Purpura.
- Spleen :
Spleen contains follicles and in between follicles sinuses are there.



Sago spleen :
If follicles/white pulp are affected → produce whitish nodules looking like sago grains.

Lardaceous spleen :
If sinuses /red pulp affected → Large geographical map like areas



Sago spleen

Important MCQ's

00:35:12

- most common biopsy site : Abdomen Fat Pad .
- Best stain for amyloid : Congo Red
- Gross stain for amyloid : Lugol's Iodine & Sulphuric Acid
- m.c. organ affected : Kidney.
- m.c. cause of death in Primary amyloidosis : Cardiac failure.
- m.c. cause of death in secondary amyloidosis : Renal failure.

Active space

- Liver : part affected earliest → Space of disse.
- Spleen : appearance → Sago spleen & Lardaceous spleen.

Q. Elderly diabetic on hemodialysis. Which amyloid is likely to be deposited?

- A. AA.
- B. Abeta.
- C. Transthyretin.
- D. **Beta 2 microglobulin.**

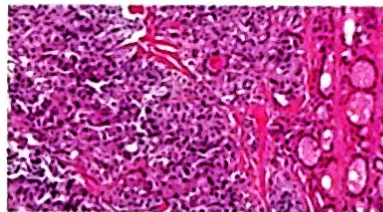
bnvssprasanth7@gmail.com

Q. A 55 year old man has developed progressive renal failure for the past 5 years. Microscopic examination of a renal biopsy shows extensive glomerular and vascular deposition of pink amorphous material on H&E staining. This material demonstrates apple-green birefringence under polarized light after Congo red staining. Immunohistochemical staining of these deposits is positive for lambda light chains. Which of the following conditions is most likely to be present in this man?

- A. Rheumatoid arthritis .
- B. Tuberculosis.
- C. Systemic lupus erythematosus.
- D. **Multiple myeloma.**
- E. Alzheimer disease.

Q. A 45 year old woman developed a thyroid swelling. The microscopic image from the swelling is shown. What type of amyloid deposits are seen in this condition?

- A. AL.
- B. ATTR.
- C. **AAL.**
- D. Apyrin.



Active space

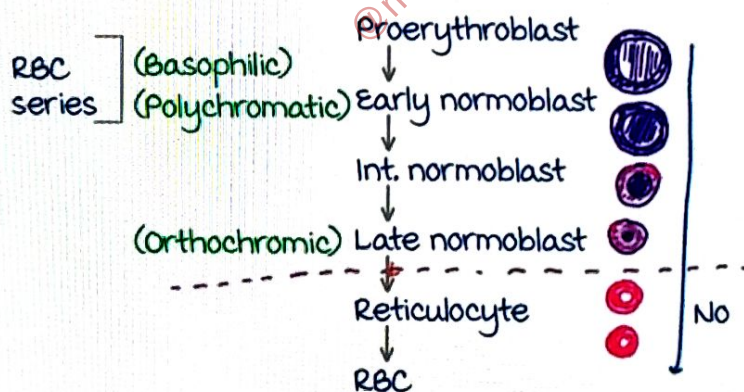
Q. A 60 year old woman has developed crippling arthritis over the past 20 years. On physical examination the arthritis primarily involves her hands and feet, with marked joint deformities characterized by ulnar deviation and swan-neck deformities of her fingers. She has an irregular heart rate. Laboratory studies show that her rheumatoid factor titer is markedly elevated, but her antinuclear antibody test is negative. A rectal biopsy shows submucosal deposition of pink amorphous material that stains positively with Congo red. Which of the following precursor proteins most likely gave rise to these deposits?

- A. Serum amyloid-associated protein.
- B. Lambda immunoglobulin light chains (AL).
- C. Transthyretin (ATTR).
- D. Amyloid precursor protein (A β).
- E. Beta-2-microglobulin (A β am in CRF).

RBC : INTRODUCTION AND HYPOPROLIFERATIVE ANEMIA

RBC introduction :

- Normal size of an RBC : 7-8 μm .
- To identify if the RBCs are macrocytic, microcytic or normal, the size of an RBC is compared with the nucleus of lymphocyte.
- Life span of RBC : 120 days.
- RBCs do not have nucleus and has a central 1/3rd pallor.
- Biconcave shape of RBS is maintained by the protein spectrin.
- RBCs originate from Haemopoietic Stem Cells (HSC) which develop into the Common myeloid Progenitor (CMP) cells.
- HSC \rightarrow CMP \rightarrow Proerythroblast \rightarrow Early normoblast (basophilic normoblast) \rightarrow Intermediate normoblast (polychromatic normoblast) \rightarrow Late normoblast (orthochromic normoblast) \rightarrow Reticulocyte \rightarrow RBC formed.



- Basophilic appears blue, polychromic normoblast will be both pinkish and bluish, and late normoblast is pinkish color (orthochromic).
- As we move along from proerythroblast (largest cell), the cell size **decreases** and nuclear size also **decreases**. At the stage of reticulocyte, the RBC has no nucleus.
- Reticulocyte is the first cell which is non-nucleated and takes 1-2 days to mature into an RBC.
- Hemoglobin production starts at the stage of

proerythroblast and can only be seen on electron microscopy.

- In intermediate normoblast, the hemoglobin is visible through a light microscope.

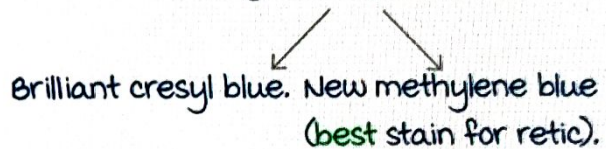
Reticulocyte

00:11:09

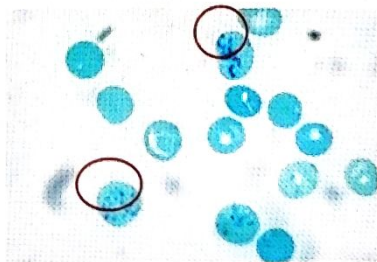
It is the immediate precursor of RBC and the 1st precursor with no nucleus.

Normal retic count is : 0.5-1.5%.

Special stain used to view a reticulocyte : Supravital stain.



- Supravital stain stains the living state or living structure of a cell (in this case RNA).
- Basic steps : Sample on glass slide → Add stain → Incubate for 30 mins → Count the reticulocytes on smear.
- Because it stains living structures, the slide has to be analyzed on time and never delayed.
- The blue filaments in the reticulocytes are the RNA, i.e., there is a reticulum (meshwork) of RNA and hence named as reticulocyte.



Another blue stain to be remembered is Prussian Blue or Pearl Stain done for hemosiderin or iron.

↑ Retic count : Reticulocytosis.

↓ Retic count : Reticulocytopenia.

Causes of Reticulocytosis	Causes of Reticulocytopenia
Acute & chronic blood loss	Bone marrow suppression
Hemolytic anemia	Aplastic anemia
Response to treatment in Iron or Vit. _B ₁₂ anemia	Megaloblastic anemia

Active space

Corrected reticulocyte count :

Retic count with degree of anemia taken into consideration.

$$\text{Corrected Retic Count} = \frac{\text{Retic \%} \times \text{Patient Hb}}{\text{Normal Hb for that age}}$$

Also note that, Hct \rightarrow Hb % \times 3.

Reticulocyte Production Index (RPI) :

$$\text{RPI} = \frac{\text{Corrected Reticulocyte Count}}{\text{maturation time.}}$$

Normal maturation time is 1-2 days.

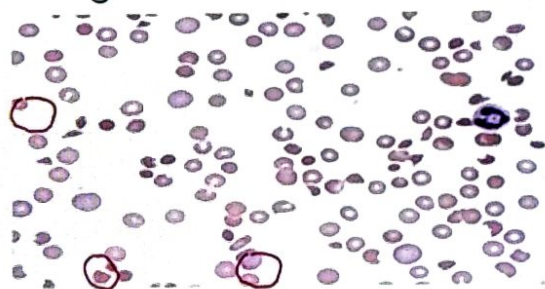
Packed cell volume and maturation time are related as such :

PCV	maturation Time
45	1 day
35	1.5 days
25	2 days
15	2.5 days

These numbers are important to find the RPI for a given PCV.

Reticulocyte on a peripheral smear :

Polychromasia (pinkish bluish hue) on a peripheral smear is usually a reticulocyte.



RBC Indices

00:27:45

- **mCV :**
mean Corpuscular volume, signifies the size/volume of the RBC. Normal value : 82-96 fl (80-100 fl)
 $\text{mCV} = \text{PCV} / \text{RBC count.}$
- **mCH :**
mean Corpuscular Hemoglobin. It is the average volume of Hb in a single RBC.
 $\text{mCH} = \text{Hb} / \text{RBC count.}$
Normal mCH = 27-32 pg.

Active space

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- MCHC : mean Corpuscular Hemoglobin Concentration.
i.e. Average volume of Hb in a given volume of packed red cells.

$$MCHC = MCH/MCV.$$

Normal MCHC : 33-37gm/dL.

Raised MCHC is seen in Hereditary spherocytosis, because of water loss, and concentration of more Hb in smaller spherical RBC.

Normal MCHC is seen in megaloblastic anemia due to B₁₂ deficiency.

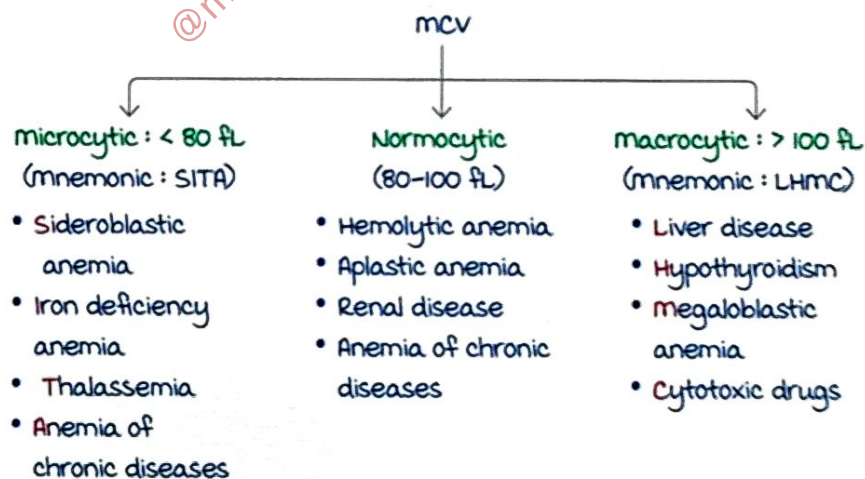
- RDW :
Red Cell Distribution width. Normal RDW : 11.5 - 14.5%.
Indicates the co-efficient of variation of red cell size or degree of anisocytosis.

RDW helps to differentiate iron deficiency anemia from thalassemia. microcytic anemia is seen in both of them.

Iron deficiency anemia : RDW raised.

Thalassemia : RDW normal.

Anemia Based on MCV :



Anemia based on MCH :

- Hypochromic anemia : < 27 pg.
- Normochromic anemia : 27-32 pg.

Variation in red cell size : Anisocytosis.

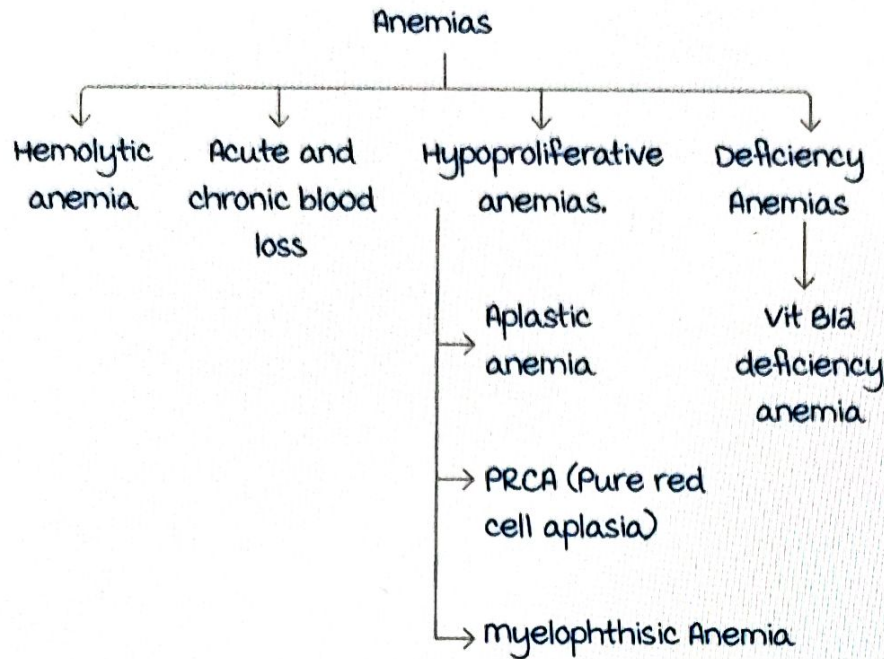
Variation in red cell shape : Poikilocytosis.

Active space

Anemias

00:51:07

Decreased Hb/red cell mass/hematocrit.



Hypoproliferative anemias :

Aplastic anemia :

- Generalized bone marrow suppression seen. Decreased Hb, TLC, platelet counts & reticulocytes.
- Causes : Inherited or acquired.
 - a. Acquired causes are more common like drugs, chemicals, viruses (Parvo virus B19, HIV, Hep B,C).
 - b. Inherited causes : Fanconi's anemia - DNA repair defect, Diamond Schachman Syndrome, dyskeratosis congenita (short telomeres).
- Clinical presentation : Pallor, fatigue, increased infections, bleeding tendencies.
- Spleen is **never** involved in aplastic anemia, therefore **splenomegaly** is never seen.
- Lab tests : Pancytopenia and reticulocytopenia seen. On peripheral smear : Normocytic normochromic anemia with pancytopenia.
- Next step :
 - BMA (Bone marrow aspiration) : **Dry tap** is seen because there are few cells in the bone.
- Therefore, IOC : Bone marrow biopsy.

Active space

Increased fat and decreased cellularity is seen (space that cells used to occupy is now replaced by fat).

Bone marrow biopsy showing increased fat and reduced cells.



Also note that fat naturally increases with increase in age.
Normal cellularity in bone marrow = $100 - \text{age}$ of patient.

Causes of dry tap on bone marrow aspiration (IOC is biopsy) :

1. Aplastic anemia.
2. myelofibrosis.
3. Hairy cell leukemia.
4. AML-M7.
5. myelophthisic Anemia.
bhvsprasanth7@gmail.com

Treatment :

Stem cell transplantation.

GM-CSF.

Severe aplastic anemia :

Criteria for severe aplastic anemia :

- Bm cellularity $< 25\%$.
- Any 2 of the following :
 - a. Platelet count is $< 20,000/\mu\text{L}$.
 - b. Corrected reticulocyte count $< 1\%$.
 - c. Absolute Neutrophil Count (ANC) $< 500/\mu\text{L}$.

Very Severe aplastic anemia :

All above criteria but ANC $< 200/\mu\text{L}$.

Pure Red Cell Aplasia

01:06:39

Reduced erythroid precursors, Hb & retic count.

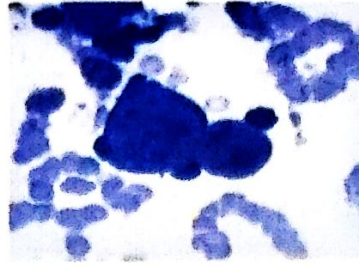
w

Causes :

Inherited : Diamond Blackfan syndrome.

Acquired : Parvovirus B19, thymoma, large granular lymphocytic leukemia, certain B cell disorders.

In Parvovirus B19, the erythroid precursors show dog ear erythroid precursors seen.



myelophthitic Anemia :

Anemia caused by a space occupying lesion of the bone marrow like metastatic cancer, any granulomatous lesion of the bone.

RBC profile in these patients are tear drop shaped also called as dacryocytes.

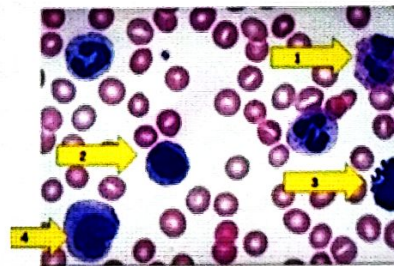
The smear will show a leucoerythroblastic blood picture (immature cells appear in blood because of no space in the bone).

Q. A 65-year-old female is diagnosed with pure red cell aplasia and a mediastinal mass. Which of the following can be the likely cause? bnvssprasanth7@gmail.com

- A. Thymoma
- B. Non Hodgkin lymphoma
- C. Bronchogenic Ca.
- D. Germ cell tumour.

Q. Identify the cells marked in the given image

- A. Lymphocytes.
- B. monocytes.
- C. Eosinophils.
- D. Basophils.
- A-3,B-4,C-1,D-2
- A-2,B-4,C-1, D-3
- A-2,B-4,C-3,D-1
- A-2,B-1,C-4,D-3



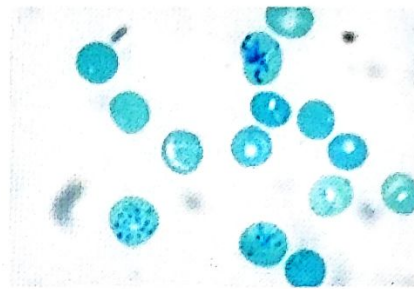
Q. Which of the following conditions will have the least chances of a dry tap on bone marrow aspiration?

- A. Hairy cell leukemia.
- B. Follicular lymphoma.
- C. AML-M7.
- D. myelodysplastic syndrome.

Active space

Q. A 59-year-old woman with a history of chronic kidney disease comes to the physician for a 3-month history of easy fatiguability. Physical examination shows subconjunctival pallor. Her haemoglobin concentration is 8.9 g/dL, mean corpuscular volume is 86 μm^3 , and serum ferritin is 110 ng/mL. Treatment with erythropoietin is begun. A peripheral blood smear is obtained one week after treatment. A photomicrograph of the smear after specialized staining is shown. The prominent colour of the intracellular structure in some of the cells is most likely the result of staining which of the following?

- A. Ribosomal RNA.
- B. Golgi apparatus
- C. mitochondria.
- D. Nuclear remnant.
- E. Lysosomes.



Q. An 18-year-old male presented to the OPD with gum bleeding, fever for the past 2 months. General examination showed pallor and the systemic examination was unremarkable. Laboratory examination revealed Hb level : 3 gm/dl, TLC : 1500, Platelets 15000. Further examination shows a low reticulocyte count, and bone marrow examination revealed fatty streaks and absent megakaryocytes but no immature cells. What is the likely diagnosis?

- A. Acquired aplastic anemia.
- B. PNH
- C. myelodysplastic syndrome.
- D. Tuberculosis.

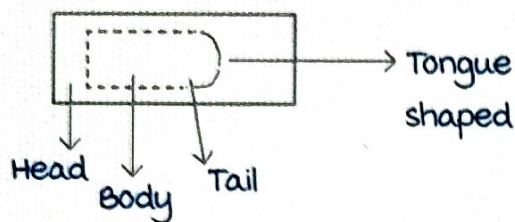
PERIPHERAL SMEAR EXAMINATION

Peripheral smear

00:02:48

A drop of blood is put on a glass slide. Another glass slide (spreader) is put at 30-45 degrees on the first slide and blood is spread to produce a tongue shaped smear. Smear has 3 parts: Head, body and tail.

Parts of a peripheral smear :



Staining peripheral smear :

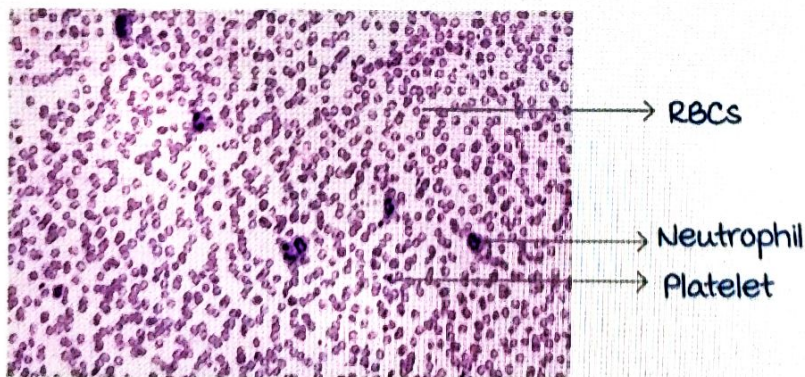
Romanowsky stain is most commonly used.

Common Romanowsky stains : Leishman, Giemsa, Wright and Jenner.

Components : methylene blue (basic dye), Eosin Y (acidic component).

Supravital stains : Brilliant cresyl blue and new methylene blue (best) are used to stain reticulocytes.

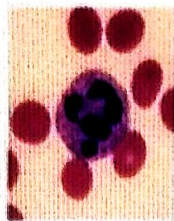
Normal peripheral smear :



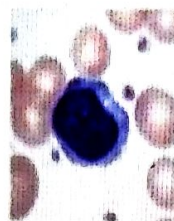
RBCs : Small cells with pallor in central 1/3rd, size of RBCs

are compared with nucleus of a lymphocyte (7-8 microns). If smaller → microcytic and larger → macrocytic.

Types of WBCs :



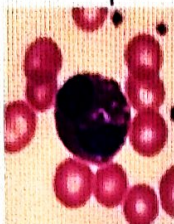
Neutrophil



Lymphocyte & Platelets



monocyte



Eosinophil



Basophil

Neutrophils : 3-5 lobes. Few small granules in cytoplasm.
2 lobes : Pseudo pelger huet cells seen in myelodysplastic syndrome.

>5 lobes : Hyper segmented neutrophil seen in megaloblastic anemia (Vit B12, folic acid deficiency).

Lymphocytes : No granules in cytoplasm.

monocytes : Big cell with kidney shaped nucleus. No granules in cytoplasm.

Eosinophils : Brick red granules in cytoplasm, Bilobed nucleus. Increases in allergic reaction

Basophils : Purplish granules obscure the entire nucleus.

RBC abnormalities on peripheral smear

00:09:52

microcytic hypochromic (MCV < 80 fL) :

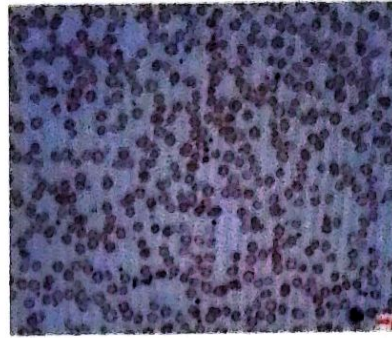
Small RBCs with >1/3rd central pallor.

S : Sideroblastic anemia; lead poisoning.

I : Iron deficiency anemia.

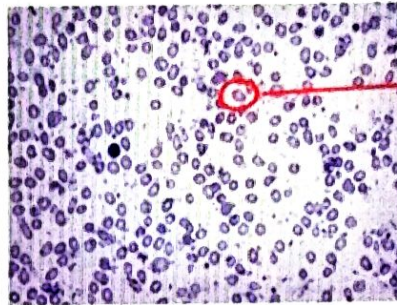
T: Thalassemia
 A: Anemia of chronic disease.

The image shows anisocytosis (variation in size) and pencil cell. High RDW, low MCV and low MCH.



macrocytic (mcv >100 fl):
 Bigger, oval cell with no central pallor.

- L: Liver disease.
- H: Hypothyroidism.
- m: megaloblastic anemia (vit B₁₂/folate deficiency).
- C: Cytotoxic drugs.



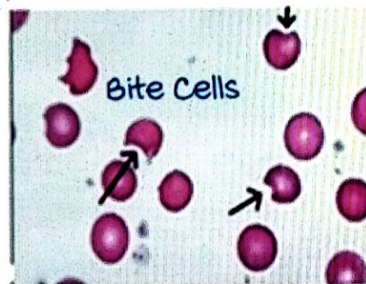
bnvssprasanth7@gmail.com

Pencil cells:

- Iron deficiency anemia.

Bite cells:

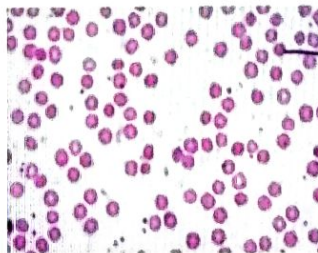
- G6PD deficiency.



Spherocytes:

Small cells (spherical) with no central pallor.

- Hereditary spherocytosis.
- Autoimmune hemolytic anemia (most common cause).
- Blood transfusion reactions.
- Burns.

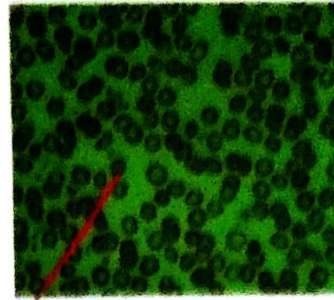


Active space

Burr cells/Echinocytes :

RBCs with blunt projections.

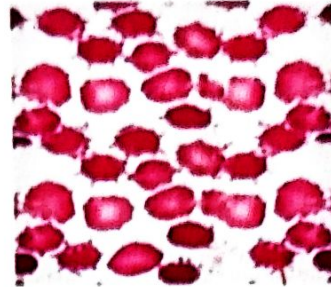
- Chronic renal failure.
- Uremia.
- Liver disease



Spur cell/acanthocytes :

RBCs with sharp projections.

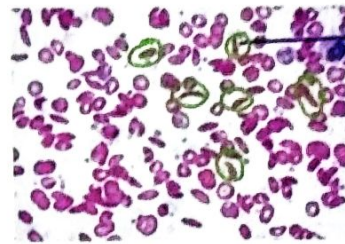
- Abetalipoproteinemia.



Sickle cells :

- Sickle cell anemia.

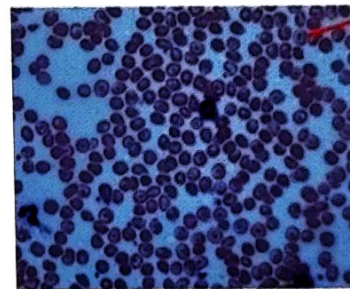
bnvssprasanth7@gmail.com



Target cells/codocytes :

Looks like a target.

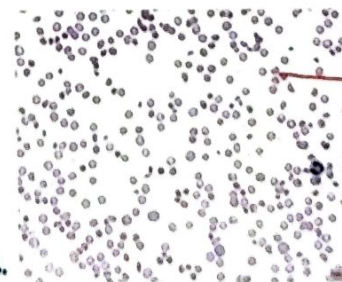
- Thalassemia (most common).
- Liver disease.
- Iron deficiency anemia.



Schistocytes (helmet) cells :

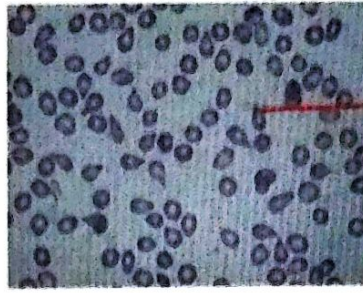
Fragmented red cells.

- microangiopathic hemolytic
- Anemia : HUS, TTP and DIC.
- Prosthetic cardiac valves.
- mechanical disruption of RBCs.



Tear drop cells/dacrocytes :

- myelofibrosis.
- myelodysplastic syndrome.
- myelophthisic anemia.
- Leucoerythroblastic blood picture.

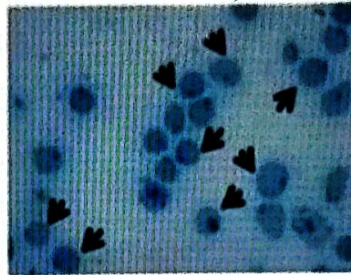


Heinz bodies :

Seen on new methylene blue stain.

Denatured hemoglobin.

- G6PD deficiency.

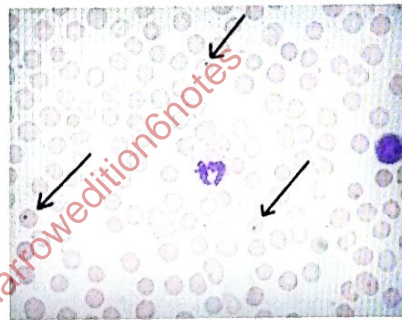


Heinz Bodies
New methylene Blue stain

Howell Jolly bodies :

Remnant of nucleus.

- Asplenia.
- megaloblastic anemia.
- Thalassemia.



Pappenheimer bodies :

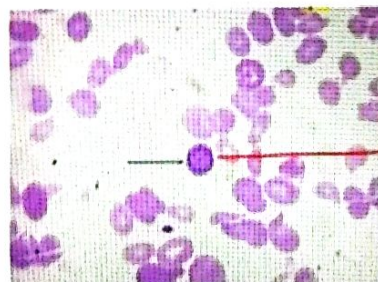
multiple, composed of iron.

- Sideroblastic anemia.

Cabot ring :

Figure of 8/ring configuration. Formed by microtubules.

- megaloblastic anemia (vit B12/folate deficiency).
- Thalassemia.



Active space

Rouleaux formation :

Stack of coin appearance.

- multiple myeloma (conditions with high ESR/ blood viscosity).

Polychromasia :

Neither pink nor purple.

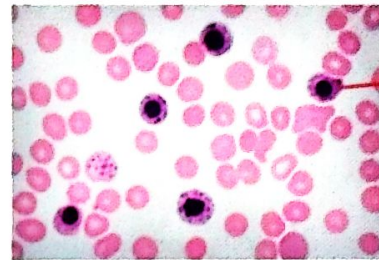
- Hemolytic anemia (Reticulocytes).

Basophilic stippling :

Bluish colored dots.

2 types :

- Fine : Seen in megaloblastic anemia, thalassemia.
- Coarse : Seen in sideroblastic anemia.



Stomatocytes :

Slit like space in RBCs.

- Hereditary stomatocytosis.

WBC changes

00:19:44

Hypersegmented neutrophils : >5 lobes.

- megaloblastic anemia due B12/folate deficiency.

Bilobed neutrophils.

myelodysplastic syndrome.

Toxic granules : Neutrophil with coarse granules, separate nuclei.

- Sepsis.

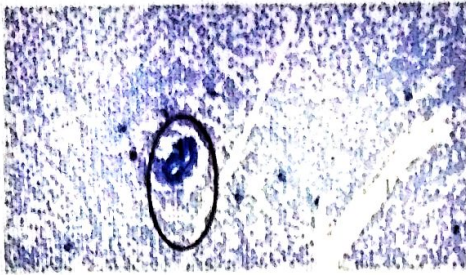
Dohle bodies : Patches of dilated endoplasmic reticulum.

- Sepsis.

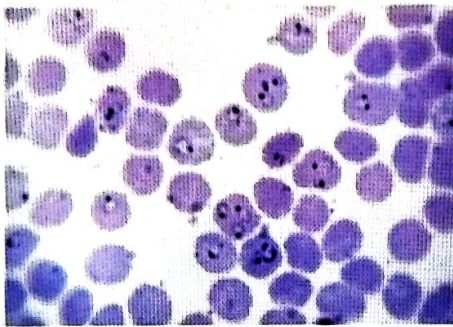
Active space

Infections :

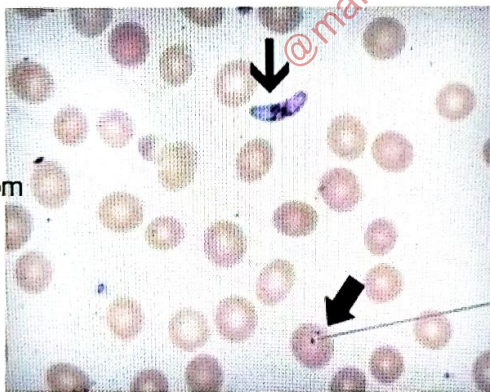
microfilaria :



malaria : Ring form of Plasmodium vivax



malaria : Gametocyte of Plasmodium falciparum.



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Ring form

Clinical case discussions

00:30:38

Q. A 17 year old male presented with fatigue

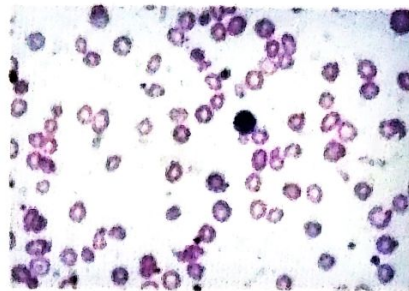
Lab tests : Hb - 9gm%

- mcv - 67 fl
- mch - 20pg
- mchc - 16 gm/dl
- RBC count - $3.9 \times 10^6/u$
- WBC - 6000/L

Active space

- PLC - 3.5 lakhs

Impression ???



Hb, mcv, mch, mchc, RBC counts are low.

WBC and platelet counts are normal.

Peripheral smear shows microcytosis, hypochromia and anisocytosis.

Impression : Iron deficiency anemia.

Confirm by iron profile : Serum iron, serum ferritin, total iron binding capacity.

Start on iron therapy. monitor treatment by reticulocyte count.

Q. A 15 year old male, with history of weakness since few months

- Hb - 6gm/dl
- mcv - 105 fl
- Reticulocyte count < 1%
- WBC - 1000/w
- Plc - 90,000/w

Impression ??

Hb, reticulocyte count, WBC, platelet counts are low.

mcv is high.

Impression : Pancytopenia

Evaluate the cause.

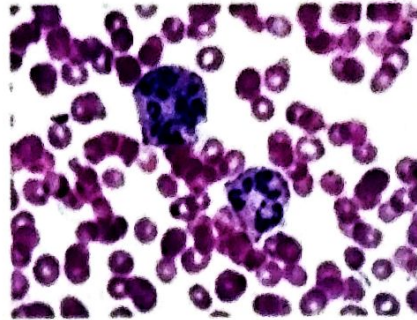
- Aplastic anemia.
- Leukemia.
- Paroxysmal nocturnal hemoglobinuria (reticulocyte count is high).
- megaloblastic anemia due to B12 deficiency.

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Active space

Q. A 32 year old female, vegetarian present with fatigue

- Hb - 7 gm%
- TLC - 1300/UL
- PLC - 1 lakh
- MCV - 132fl
- MCH - 28 pg
- MCHC - 39 gm/dl



Impression ??

Hb, TLC, platelet count are low.

MCV is high.

MCH and MCHC are normal.

Peripheral smear shows macro ovalocytes with hypersegmented neutrophils.

Impression : megaloblastic anemia due to B12 deficiency.

@marroweditionsnotes

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Active space

MICROCYTIC HYPOCHROMIC ANAEMIA

Mnemonic : SITA.

Sideroblastic anaemia.

Iron deficiency anaemia.

Thalassemia.

Anaemia of chronic disorders.

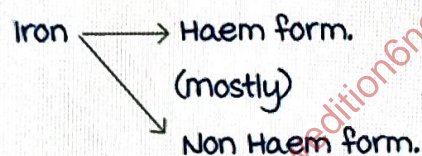
Iron deficiency anaemia (IDA)

00:01:42

most common of the nutritional deficiencies.

Daily requirement : 10-20 milligrams/day.

Rich source :



In the Haem form : 80% is in haemoglobin.

20% is in enzymes or others.

Iron is seen in Fe^{2+} (ferrous) & Fe^{3+} (ferric) form.

Iron absorbed from duodenum as Fe^{2+} form only.

most common site of iron absorption : Duodenum.

Storage form of Iron :

1. Ferritin.
2. Haemosiderin.

Transportation form of iron : **Transferrin**.

Causes of IDA

00:06:41

- Decreased intake :
 - Poverty.
 - Low socioeconomic status.

MICROCYTIC HYPOCHROMIC ANAEMIA

Mnemonic : **SITA**.

Sideroblastic anaemia.

Iron deficiency anaemia.

Thalassemia.

Anaemia of chronic disorders.

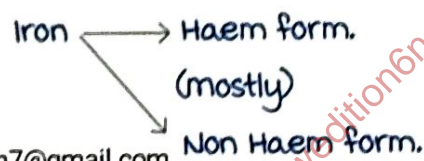
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1. Ferritin.
2. Haemosiderin.

Transportation form of iron : **Transferrin**.

Causes of IDA

00:06:41

- Decreased intake :
 - Poverty.
 - Low socioeconomic status.

- Increased demand :
 - Puberty.
 - Pregnancy.
 - Lactation.
- Impaired absorption :
 - In certain malabsorption syndromes.
 - Vit B12 deficiency causing worm : Fish tape worm (Diphyllobothriasis).
- Chronic blood loss :
 - GI/colon malignancy.
- Worm infestation : *Ancylostoma duodenale* (hookworm).

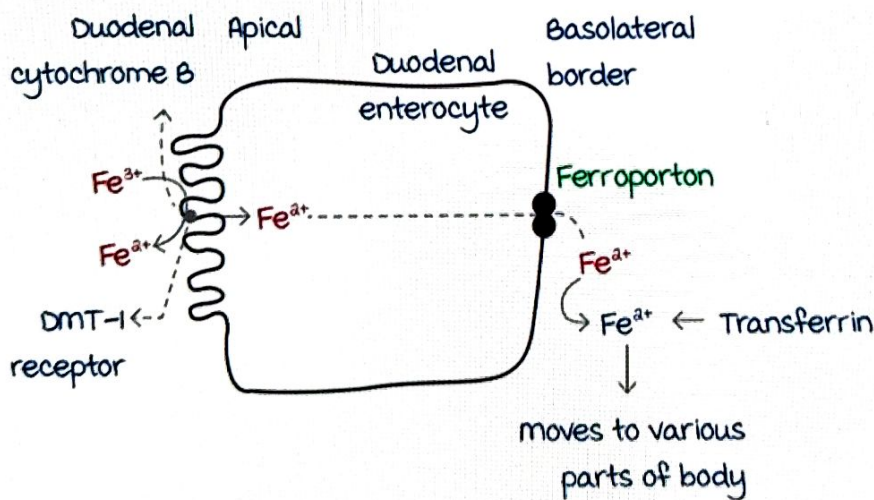
Factors :

Increasing iron absorption : Acidic pH , Vit. C /Ascorbic acid, amino acids, citric acid.
 Decreasing iron absorption : Alkaline pH, tannates , phytates, tea (only have iron after 30 mins).

Mechanism of iron absorption

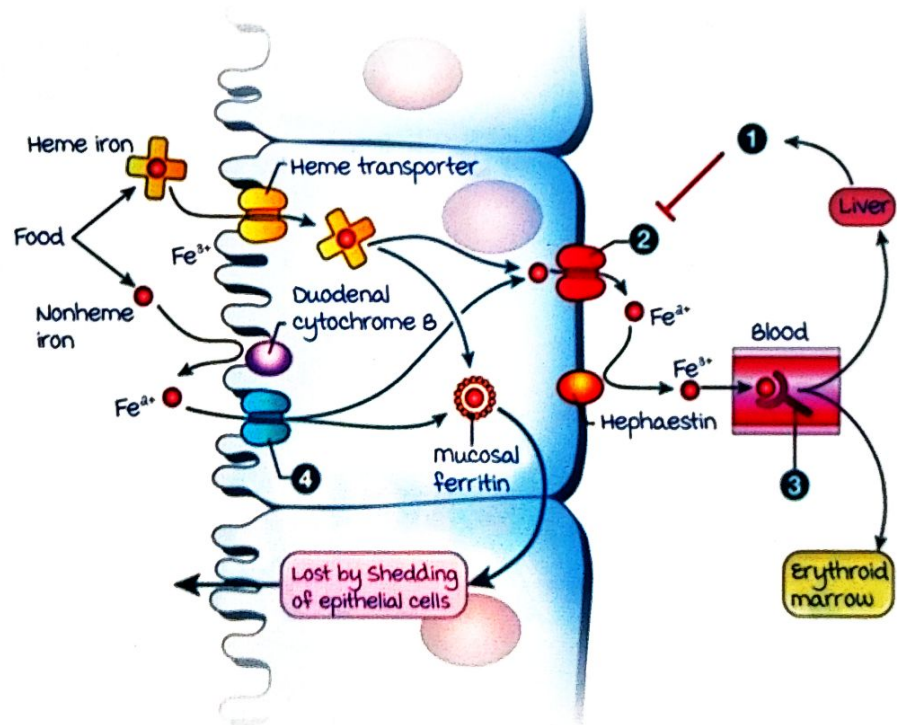
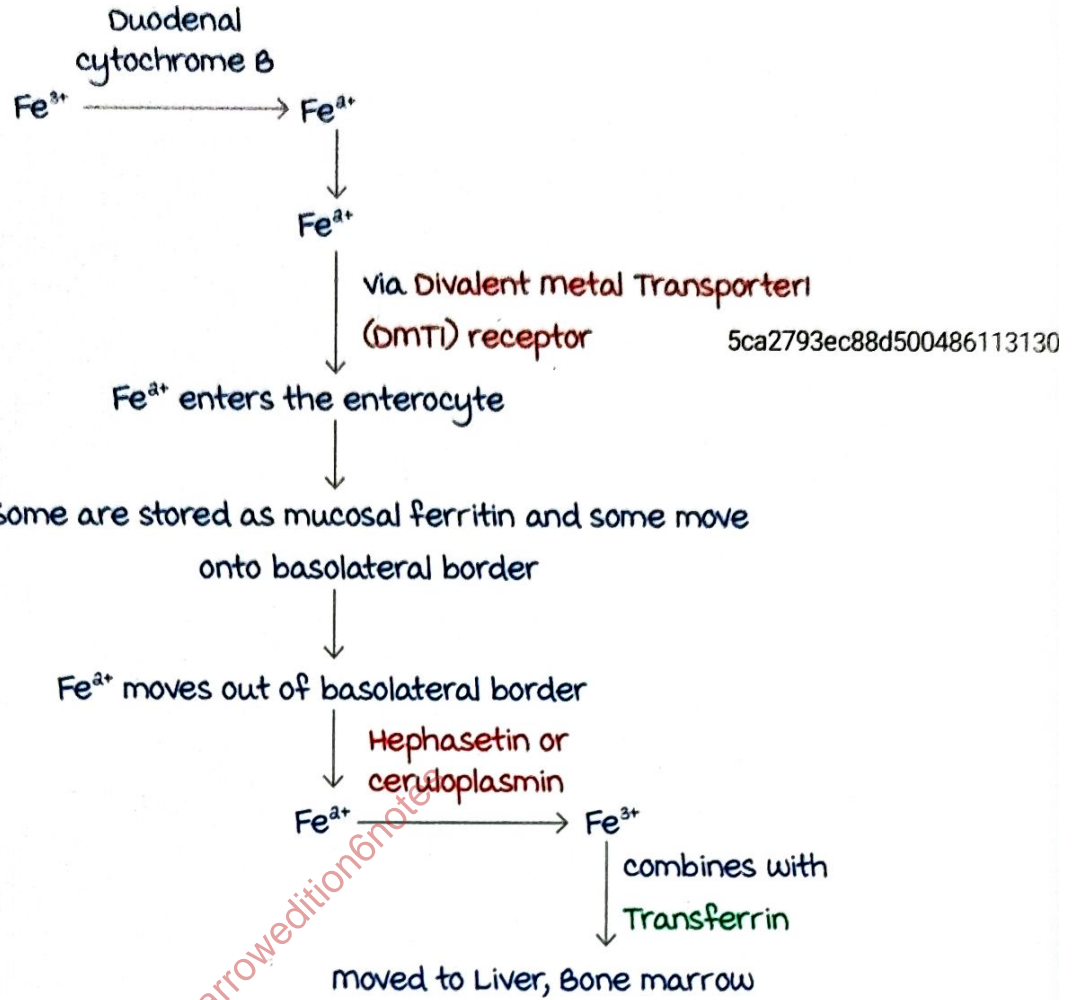
00:12:34

Haem iron directly enters as Fe^{2+} .
 Non-Haem mostly in form of Fe^{3+} and is converted to Fe^{2+} .

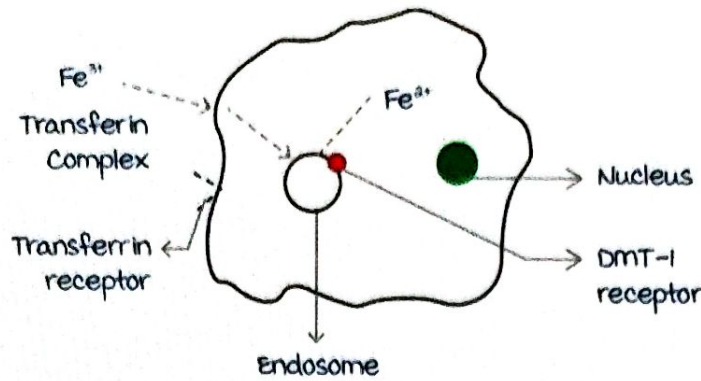


Active space

At the villi of duodenum,



Active space



At the bone marrow,

In erythroid precursors, iron & porphyrin combines forming hemoglobin.

Erythroid precursors have transferrin receptors to which Fe³⁺-Transferrin complex binds and enters the enterocyte & into the endosomes.

In the enterocyte, iron is stored in endosome having DMT-1 receptor which will again convert to Fe²⁺ and comes out.

DMT 1 is present in :

Placenta.

macrophages.

Erythroid precursors.

Transferrin (TF)

00:21:37

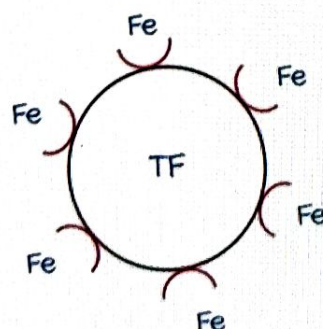
Transporting molecule for iron.

1 TF can combine with 6 molecules of iron ideally.

But clinically 1 TF combines with only 2 Fe molecules implying Transferrin saturation (TS) = $\frac{2}{6} \times 100 = 33\%$.

Early erythroid precursors (EP) have more TF receptors (TFR) but in late EP, TFR sheds off.

Soluble TFR ratio (STFRc) : measure of erythroid activity of bone marrow.



Hepcidin (H)

00:25:39

Hep : Hepatocytes. Cidin : Inhibitor.

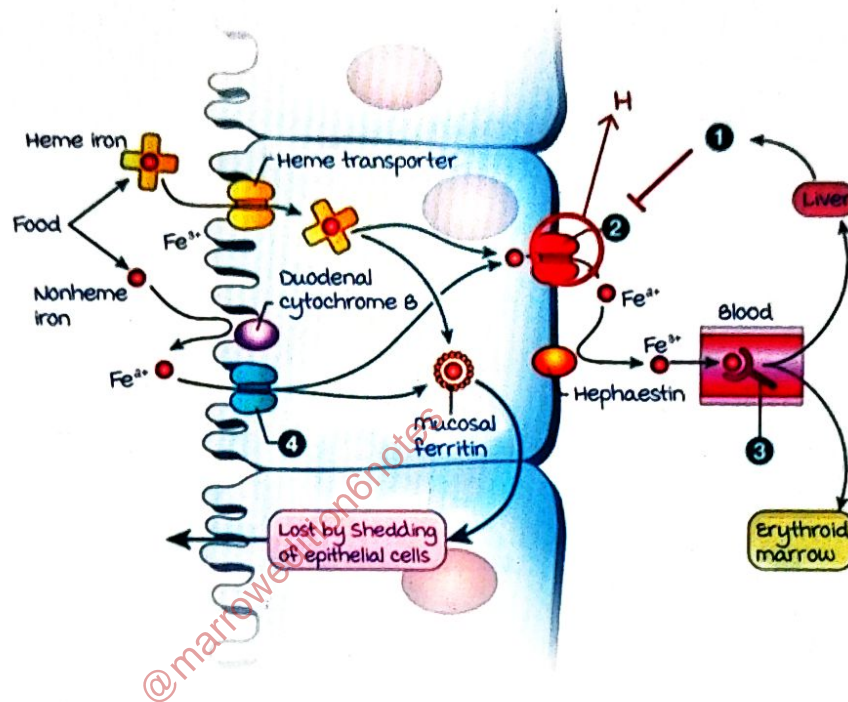
Produced by liver.

master regulator of iron.

Inhibits Fe absorption.

Increase in H, implies decrease in iron and vice versa.

Mechanism :



H binds to Ferroportin and degrades it.

↓
Fe not released

↓
Decrease in serum iron

↓
microcytic hypochromic anaemia.

H is an acute phase reactant, hence increased during inflammation.

Genes Regulating H :

HFE } mutation causes hemochromatosis by iron overload.
HJV }

Tmprss 6 : Seen in Iron Refractory Iron Deficiency Anaemia (IRIDA).

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Active space

Clinical features

00:31:52

usually seen in middle aged women.

- Pallor.
- Fatigue.
- Dyspnoea.
- Palpitation.
- Angular stomatitis, cheilitis.
- Koilonychia (spoon shaped nails).
- PICA: H/o child eating mud, clay.
- Plummer Vinson Syndrome or Peterson Brown Kelly Syndrome.
- Triad:
 - Fe deficiency anemia.
 - Esophageal webs.
 - Atrophic glossitis.



Investigations:

1. CBC:

Hb low

RBC mass low

TLC normal


Platelet count ideally normal but clinically seen as thrombocytosis/increased called as reactive thrombocytosis.

MCV, MCH, MCHC all are decreased.

RDW: Indicator of anisocytosis.

Variation in RBC size seen so increased.

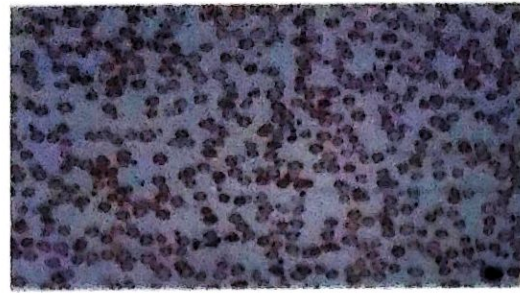
2. Peripheral smear:

- microcytic hypochromic RBC: Smaller cell and have more than one third central pallor.
- Pencil cells. 
- Anisopoikilocytosis.

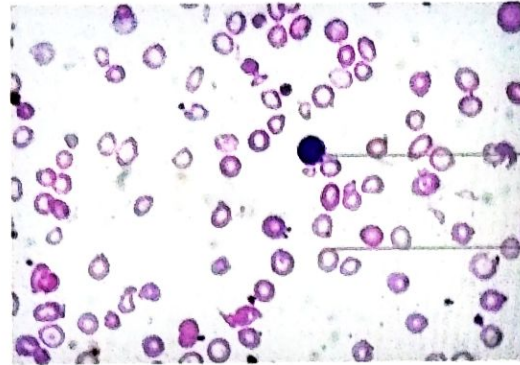
RBC size is 7-8 microns and is comparable to the size of nucleus of small lymphocyte.

Active space

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microcytic hypochromic
RBCs
→
Reactive thrombocytosis



→ Small lymphocyte nucleus
→ microcytic hypochromic
RBCs

Iron studies

00:41:22

Expensive so can be skipped for patients in lower social economic strata and can start on Fe therapy.

- Serum Fe decreased.
- Serum ferritin decreased :
Ferritin is the storage form. It is the sensitive test and one of the earliest parameters to decrease.
- S. TIBC : Binding capacity of iron to receptor.
Here, No Fe implying increased S.TIBC.
- Transferrin Saturation (TS) : Decreases
- Free erythrocyte Protoporphyrin level : Increases.
- Bone marrow Fe : **Golden Standard Test.**
Stain for iron hemosiderin in bone marrow is Prussian blue or Pearl stain. Decrease in stainable iron.
It is not done as is painful and invasive.
- **STFRc assay to Log Ferritin ratio :**
> 1.5 : indicator of FDA.
STFRc increases and ferritin decreases in FDA.
- STFRc Assay : Sensitive test.

Order of sensitivity test : STFRc assay to Log Ferritin ratio >
STFRc assay > S. Ferritin.

Stages of anaemia :

Stage 1 : **Decrease in storage.**

Decrease in Ferritin.

Stage 2 : **Iron Deficient erythropoiesis.**

Stage 3 : **Iron Deficiency Anaemia.**

Peripherals Smear finding is seen stage 3.

Treatment : Iron therapy

monitoring is done with **Retic count** :

First indicator to increase.

Starts increasing in 5 to 7 days of iron therapy.

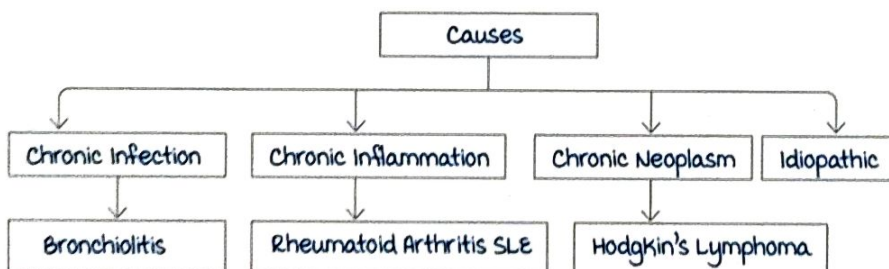
CBC peripherals smear done and started on iron therapy → Then after a week → Investigation for iron studies done and also retic count → If count increases continue Fe therapy.

mentzer Index : $\frac{MCV}{RBC \text{ Count.}}$

> 13 : FDA
< 13 : **Thalessemia** } Aids in differentiating both.

Anaemia Of Chronic Disease (AOCD)

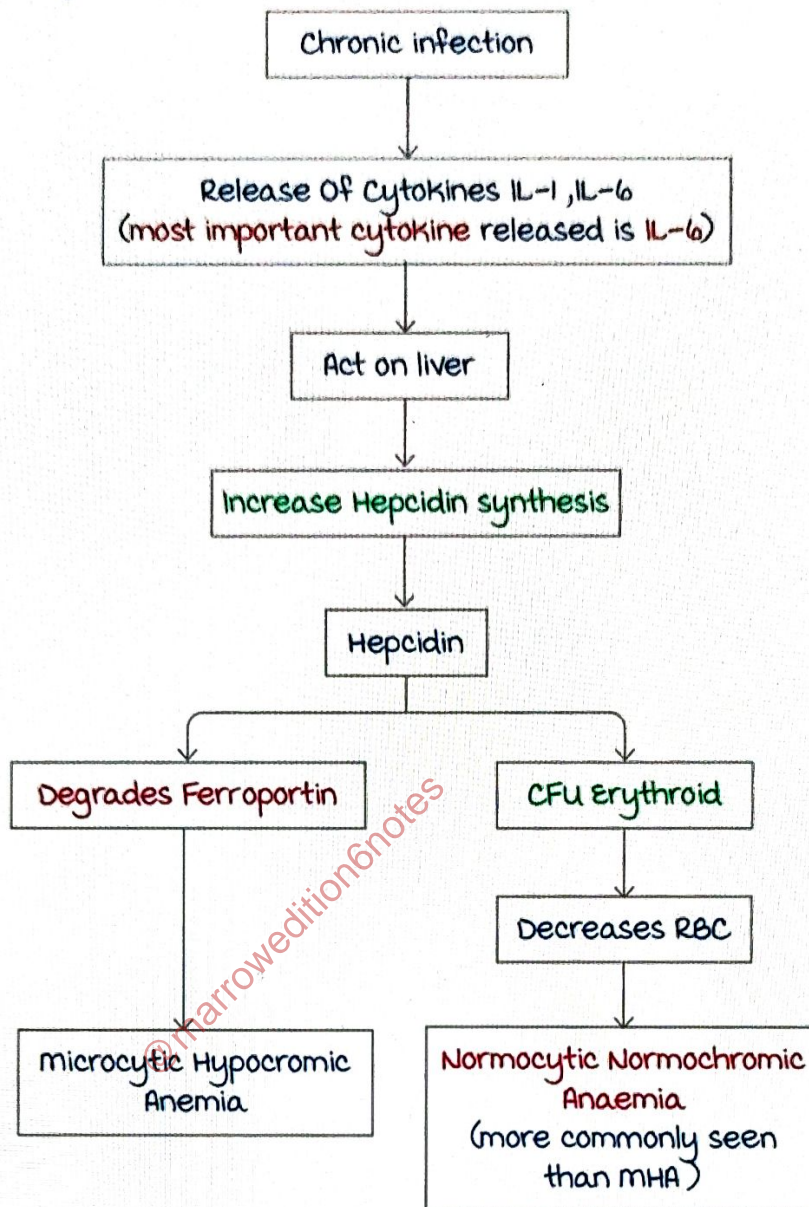
00:51:22



Active space

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Pathogenesis :



Investigation :

- Hb, TLC, Platelet count decreases.
- MCV MCH MCHC normal or decreased.
- Peripherals smear shows normocytic normochromic anemia sometimes as normocytic hypochromic anemia.
- Iron profile :
 Serum iron : Decreases.
 Serum Ferritin : Increases as stores of Fe increased and would not get released.
 S.TIBC : Decreased as it is a measured with Serum Ferritin and is overloaded with Fe.

Active space

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- STFRc assay to Log Ferritin ratio :

<1.5 implies AOCD.

S. Ferritin is high.

Treatment : Treat the cause.

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Sideroblastic anaemia(SA)

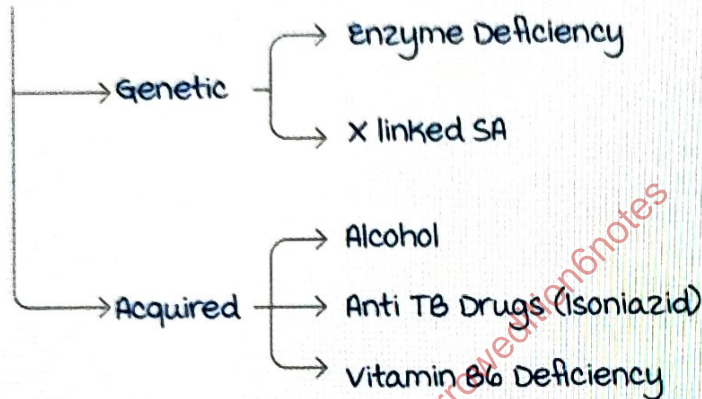
00:58:18

Opposite to iron profile study in FDA.

Sider is iron & Blastic is immature precursors.

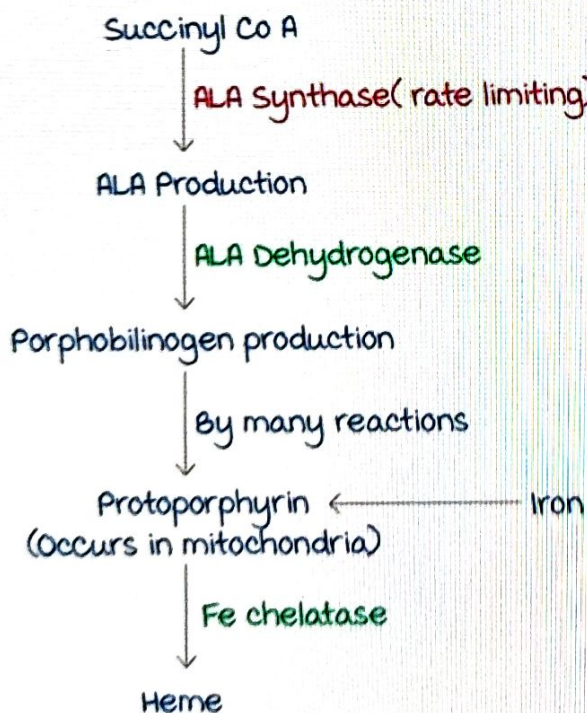
Excessive iron in immature precursors but cannot be utilised by EP for Hb synthesis.

Causes:



Pathogenesis :

Hb Synthesis Pathway :



Active space

If defect in ALA Synthase/ ALA Dehydrogenase/

Ferrochelatase :

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No protoporphyrin formed but Fe accumulation increases.

No Hb is formed.

Vit B6 is needed in Hb synthesis.

Alcohol is mitochondrial poison.

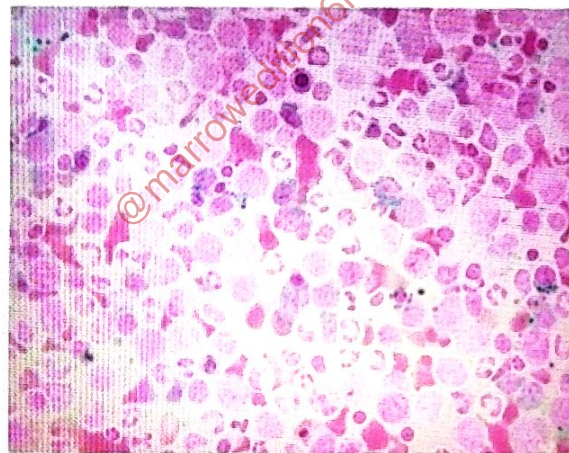
- Bone marrow aspiration :

Ringed Sideroblast :

mitochondria is located in perinuclear area and Fe keeps accumulating in the mitochondria.

more than 5 iron granules in perinuclear location and covering one third of nucleus is called as Ringed Sideroblast.

Seen using Prussian blue stain.



- CBC :

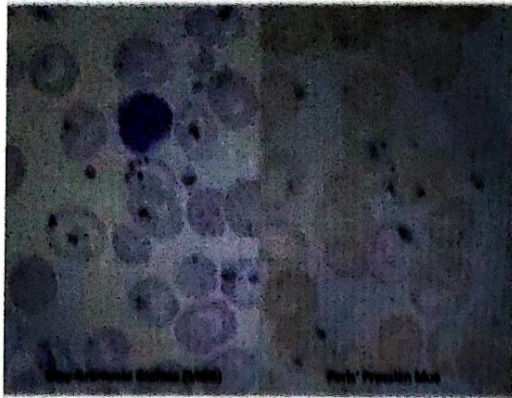
Hb decreased.

TLC , platelet count normal.

MCV MCH MCHC decreased in MHA.

- Peripheral smear :

1. Pappenheimer bodies : Iron in mature RBCs.
2. microcytic hypochromic red Cells.
3. Coarse basophilic stippling.



→ Pappenheimer bodies

- Iron profile :
 Serum Iron increases.
 Serum Ferritin increases.
 S. TIBC decreases.
 TS increases.

Treatment :

Phlebotomy.
 Iron chelaters.

	S. Iron	S. Ferritin	S.TIBC	P/S	Extra
Iron deficiency anaemia	Dec	Dec	Inc	Pencil cells	RDW inc mentzer Index >13
Sideroblastic anaemia	Inc	Inc	Dec	Ringed sideroblasts Coarse Basophilic Stippling	
Anaemia of chronic disease	Dec	Inc or normal	Dec		ESR inc
Thalassemia	N	N	N	N	HbA2 inc mentzer index <13

HbA2 Inc → >3.5 → Thalassemia trait.

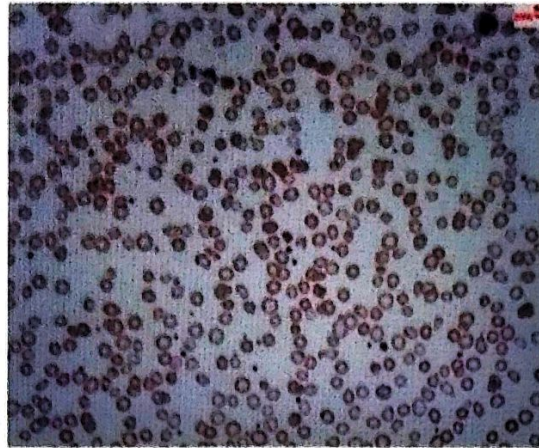
Scenario 1 :

Q. A 17 year old male presented with fatigue. Lab tests w:

- Hb : 9 gm%.
- MCV : 67 fl.
- MCH : 20pg.
- MCHC : 16 gm/dl.
- RBC count : $3.9 \times 10^6/\mu\text{L}$.
- WBC : 6000/L.

Active space

Answer : MH cells present s/o FDA. Advise Fe studies.



- Q. Which of the following is used in the treatment of sideroblastic anemia?
- A. Vitamin B12.
 - B. Vitamin B6.
 - C. Vitamin B1.
 - D. Iron.
- Q. Which of the following is not involved in iron metabolism:
- A. Hpcidin.
 - B. Transferrin.
 - C. Transferrin.
 - D. Ceruloplasmin.

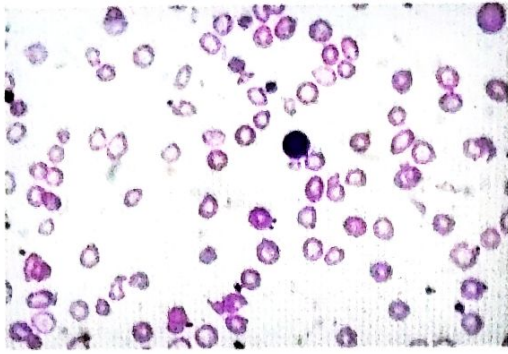
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Scenario 2 :

- Q. A 33 year old woman presents with dysphagia and atrophic glossitis. The peripheral smear from this patient is given below. which of the following lab findings is consistent with her likely diagnosis?

Options	S.Ferritin	TIBC	Transferrin saturation
1	High	Normal	High
2	Normal	Normal	High
3	Low	High	Low
4	High	Low	Low

Active space



Answer :

Pencil cells present.

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MH are red cells present thrombocytosis present
in female

Dysphagia and atrophic glossitis present. Suggestive FDA
classical presentation of Plummer Vinson disease.

- Q. In iron deficiency anaemia, all of the following are increased except
- A. Transferrin saturation.
 - B. RBC protoporphyrin.
 - C. TIBC.
 - D. Ferritin soluble receptors.

@marroweditionsnotes

Active space

MEGALOBLASTIC ANAEMIA

- megaloblastic anemia (MA) is a type of macrocytic anemia.
- megaloblast = Large, blast = Immature.

INDEX

- MA due to vitamin B12 deficiency.
- MA due to Folate deficiency.
- Pernicious anemia.

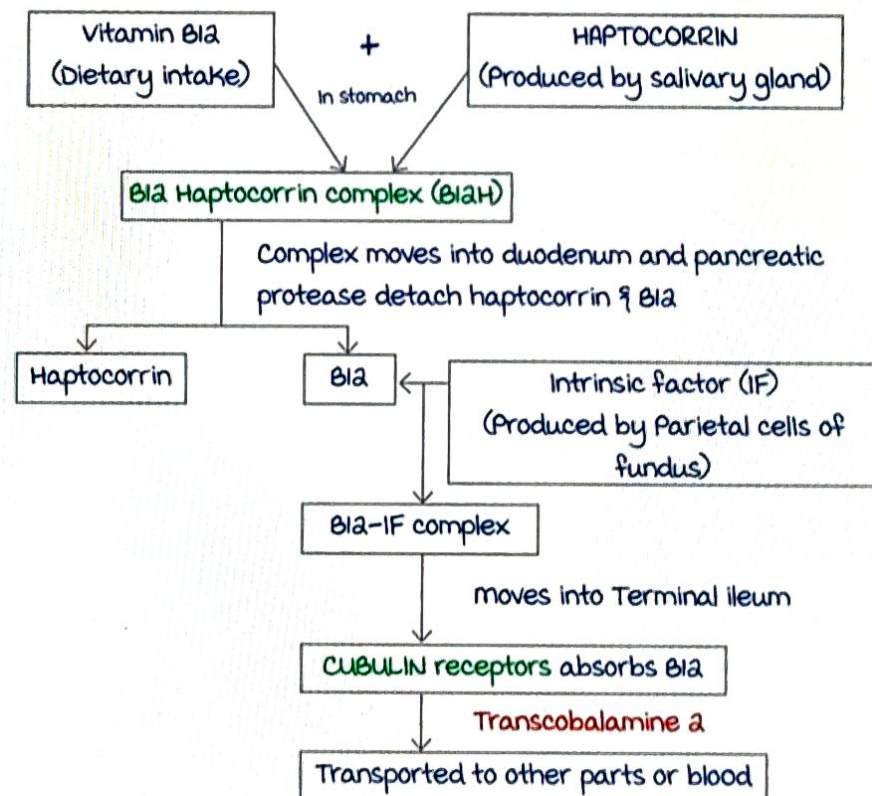
MA Due To Vitamin B12 Deficiency

00:00:48

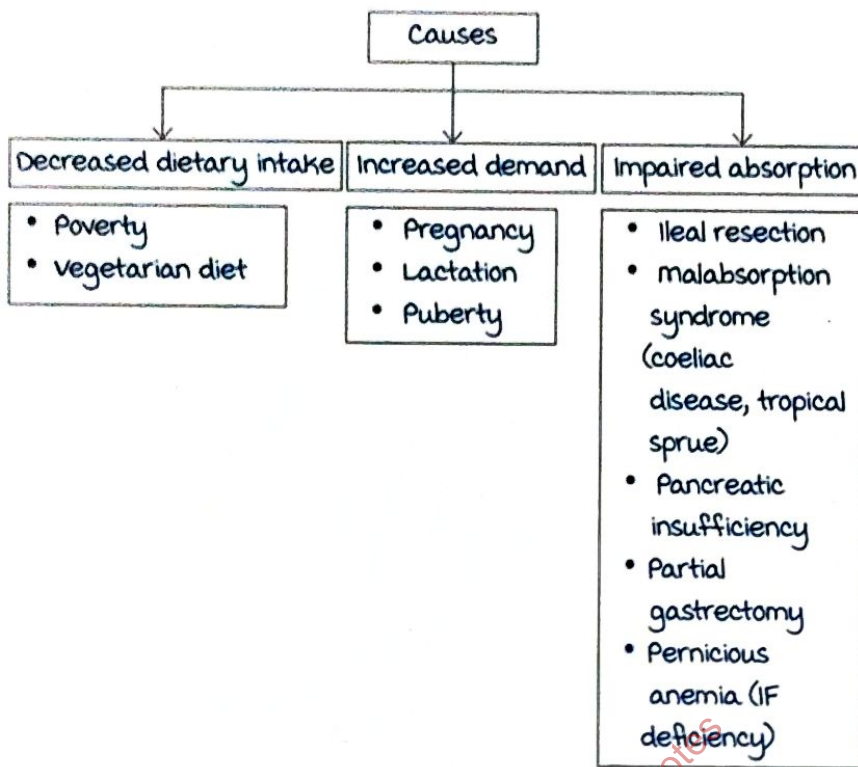
Vitamin B12 (B12) :

- Also called as Cyanocobalamine.
- Water soluble (non toxic).
- Daily requirement is 2-3mcg.
- Source : Dairy products (milk), egg, fish, meat (usually deficient in vegetarian people).

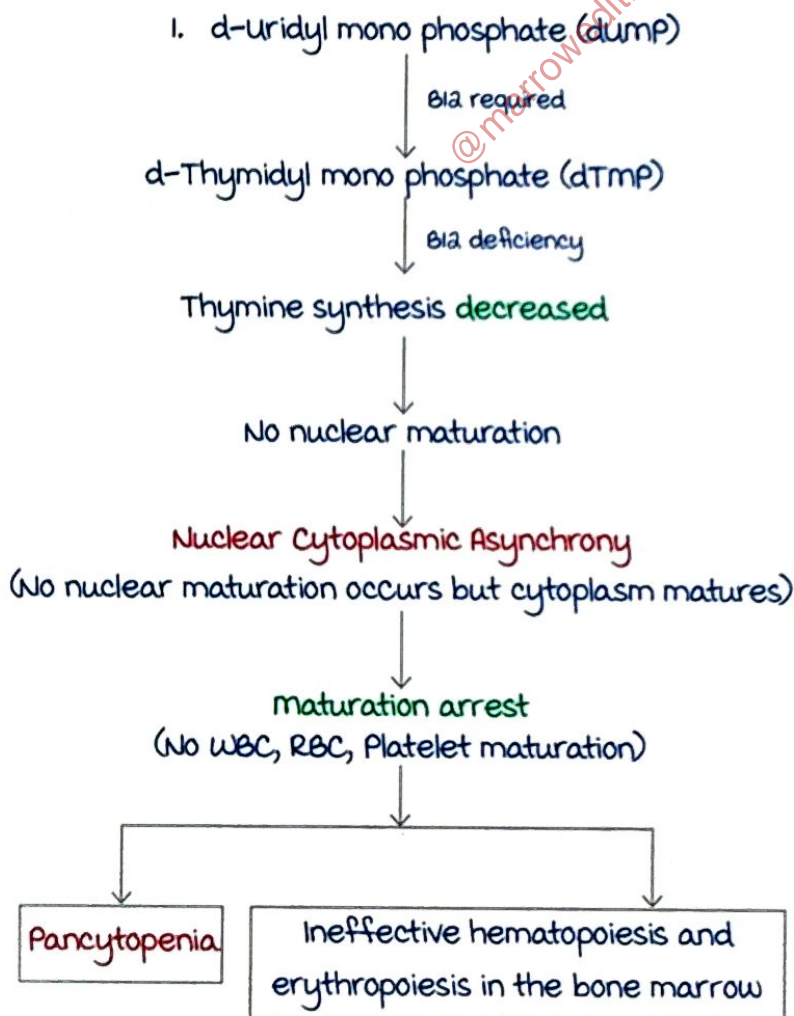
Mechanism of action :



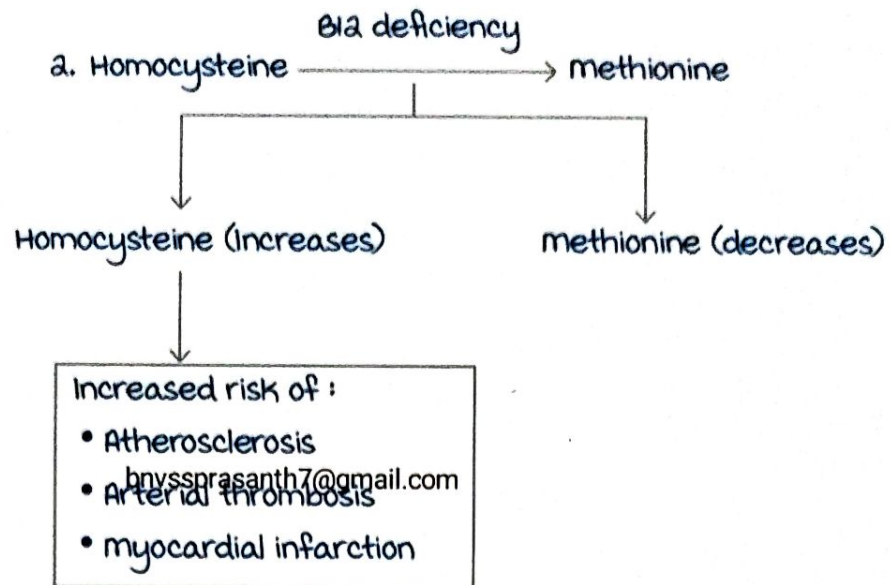
- The site of maximum B12 absorption is Terminal ileum.
- Transportation molecule is Transcobalamin 2.



Biochemical reaction catalyzed by B12 :



Active space



3. methyl malonyl CoA $\xrightarrow{\text{B12}}$ Succinyl CoA.
- Succinyl CoA is a component of neuronal lipids and myelin sheath, so its decrease causes **neurological complications**.

Clinical features :

- Pallor.
- Fatigue.
- Jaundice.
- Splenomegaly.
- Neurological complications like **Sub Acute Combined Degeneration of Spinal Cord**.

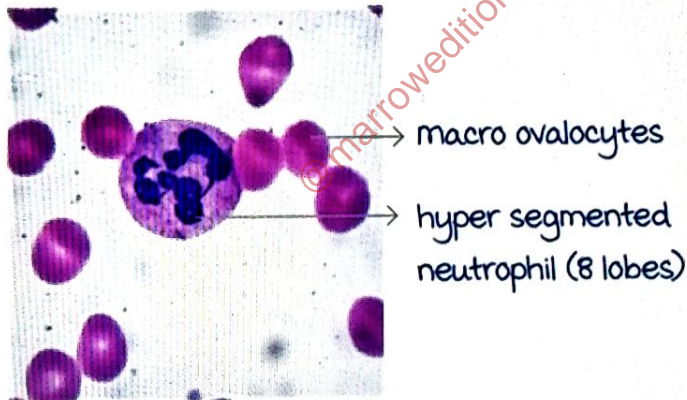
Investigation :

- CBC :
 1. Decrease in hemoglobin, total count, platelet count. (Pancytopenia)
 2. Increase in MCV (as large RBC) : >100 femtolitres.
 3. Increase in MCH as they appear hyperchromic.
 4. Normal MCHC (as size of RBC and hemoglobin also increase).
- Peripheral smear (PS) :
In RBCs -

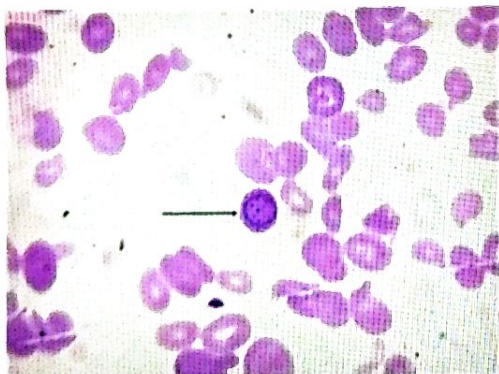
1. macro ovalocytes :
 - Large oval RBC and macrocytic, (sometimes no central pallor).
 - **Earliest finding.**
2. CABOT rings :
 - Appear like either figure of 8 or round ring shaped formed by microtubules .
3. HOWELL JOLLY BODY : Remnant of nucleus formed due to ineffective hematopoiesis / erythropoiesis.
bnvssprasanth7@gmail.com
4. Fine Basophilic stippling : RBC appear to have fine bluish dots .

In WBCs : Hyper segmented neutrophils (>5 lobes) is seen

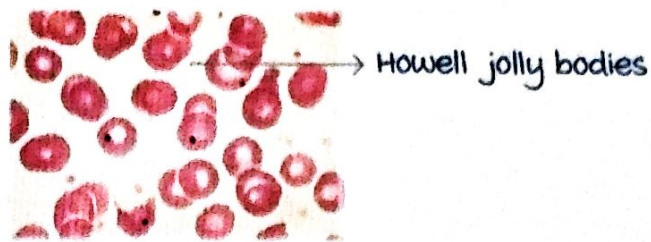
- **Criteria for diagnosis** : >5% neutrophils with 5 or more lobes.
- Single neutrophil with 6 or more lobes.



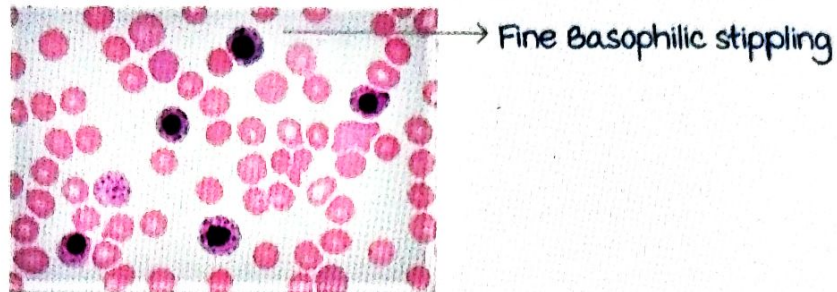
Hyper segmented neutrophil (8 lobes)



Cabot ring



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- Bone marrow aspirate :
 1. Erythroid hyperplasia
 2. Reversal of m : E ratio.
 3. Shows large immature precursors of RBCs, WBCs, megakaryocyte.
 4. Large Erythroid precursor with sieve like chromatin (thin chromatin that have not matured) called megaloblast.
 5. Giant metamyelocytes and band forms.
 6. Giant megakaryocytes.

- Biochemical investigations :
 1. Vitamin B12 assay.
 2. Serum Homocysteine levels.
 3. Serum methyl malonyl CoA.
 4. Serum LDH (as ineffective erythropoiesis).
 5. Reticulocyte count (because there is general bone marrow suppression of all lineages).

- Treatment :
 - a. Dietary intake of food rich in Vitamin B12.
 - b. Supplementation with Vitamin B12 tablets (Neurobion forte).
 - c. Intramuscular Vitamin B12 injections.

MA Due To Folate Deficiency Anemia

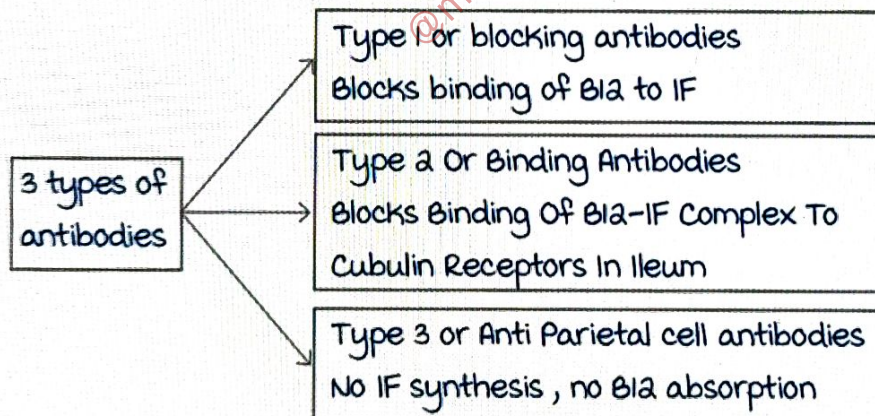
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- Derived from word Folia meaning green leafy vegetables (spinach, broccoli).
- Overcooking destroys folate.
- In pregnancy folate deficiency causes **Neural Tube Defect**.
- Site of folate absorption : **Jejunum**.
- Deficiency is seen in alcoholics.
- **Neurological complications** seen in B12 deficiency are **absent here as folate doesn't help in myelination**. If B12 deficiency is treated with folate anemia improves as Folate helps in Thymine synthesis but neurological symptoms worsen.

Pernicious Anemia

00:37:09

- **Type 2 Hypersensitivity Reaction** (Antibody mediated).
- **Autoimmune reaction**.
- Pathogenesis :



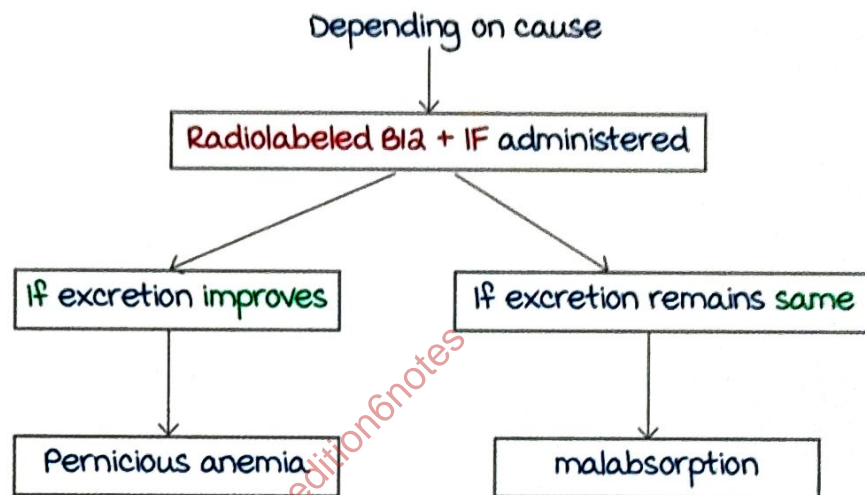
- Clinical features :
 1. Pallor.
 2. Fatigue.
 3. **Beefy tongue**.
 4. **Atrophic Glossitis**.
 5. Increased risk of **autoimmune disorders**.
 6. Increased risk of **Gastric Adenocarcinoma**.
 7. **Fundic gland atrophy**.

Active space

Schilling's Test

00:41:16

- Not used nowadays.
- Not for diagnosis but to identify cause of B12 deficiency anemia.
- Radio-labelled vitamin B12 administered and depending on its excretion in 24 hours.
 - > 8% → Normal
 - < 8% → Vitamin B12 deficiency



EXTRA POINTS :

- Site of maximum absorption of iron : Duodenum.
- Coarse basophilic stippling seen in Sideroblastic anemia.
- In myelodysplastic syndrome- **Hypossegmented neutrophil (Pseudo Pelger-Huet)** is seen .
- Pancytopenia is seen in
 1. Aplastic anemia .
 2. MA due to B12 deficiency.
 3. Leukemia.

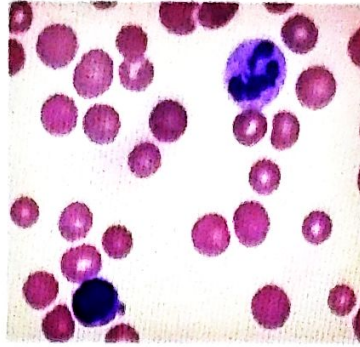
Clinical scenario :

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Q. A 55-year-old man comes to the physician because of the Balance problems. He follows a vegan diet And does not eat meat, eggs or dairy. Physical examination shows pale oral mucosa and conjunctivae. Neurological examination shows normal strength in all extremities and decreased sense to vibration in his lower extremities bilaterally. A Peripheral blood smear is shown. which of the following enzymes is most likely impaired?

A. Succinate dehydrogenase.

- B. Pyruvate carboxylase.
- C. methylmalonyl coA.
- D. Dopamine beta hydroxylase.



Answer : methylmalonyl coA.

Q. A 51-year-old man has become increasingly fatigued for the past 10 months. On physical examination there are no abnormal findings. Laboratory studies show his Hgb - 9.2, Hct - 27.9% , MCV -132 fl, Platelet count - 242,000/microliter, \uparrow WBC count 7590/ microliter. Which of the following morphological findings is most likely to be present on examination of his Peripheral blood smear?

- A. Hypersegmented neutrophils.
- B. Nucleated red blood cells.
- C. Blasts with Auer bodies.
- D. Hypochromic, microcytic RBCs.
- E. Schistocytes.

Answer : Hypersegmented neutrophils > Nucleated red blood cells.

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HEMOLYTIC ANEMIA : PART 1

Hemolytic anemia

00:01:16

Caused by excessive destruction of RBCs.

Hemolytic anemia can produce two types of defects :

- Intracorpuseular defects : Defect inside RBCs.
- Extracorpuseular defects : Defect outside RBCs.

Intracorpuseular defects : Either hereditary or acquired.

Hereditary :

- Red cell membrane defects : Hereditary spherocytosis, hereditary elliptocytosis.
- Enzyme deficiencies : G6PD deficiency, pyruvate kinase deficiency, hexokinase deficiency.
- Hemoglobinopathies : Sick cell anemia, thalassemia.

Acquired :

Paroxysmal nocturnal hemoglobinuria (only acquired intracorpuseular defect).

Extracorpuseular defects : Either immune mediated or non - immune mediated.

Immune mediated : Autoimmune hemolytic anemia.

Non immune mediated : Infections like malaria.

Hemolysis can be intravascular or extravascular.

Intravascular hemolysis	Extravascular hemolysis
Hemolysis occurs inside a vessel	Hemolysis occurs outside a vessel Example : Liver, spleen
Hepatomegaly or splenomegaly are usually absent	Hepatomegaly or splenomegaly are usually present
Serum haptoglobin is reduced	Serum haptoglobin is usually not decreased
Hemoglobinuria and hemosiderinuria are seen	Hemoglobinuria and hemosiderinuria are absent

Active space

General clinical features

00:12:10

Triad :

Pallor.

Jaundice.

Splenomegaly.

usually, unconjugated bilirubin is increased

Chronic hemolysis : Increased risk of gallstones (pigment gallstones).

Splenomegaly and gallstones are seen in extravascular hemolysis.

General lab investigations :

- Hemoglobin : Decreased.
- MCV, MCH and MCHC : usually normal.
- Peripheral smear : Specific for anemia.
- Liver function tests : Deranged, increased bilirubin.
- Reticulocyte count : Increased.
- Serum haptoglobin : Decreased (intravascular hemolysis).
- Hemoglobinuria (intravascular hemolysis).
- Hemosiderinuria (intravascular hemolysis).
- Serum LDH : Increased.

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Hereditary spherocytosis

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75% of cases : Autosomal dominant.

m = F.

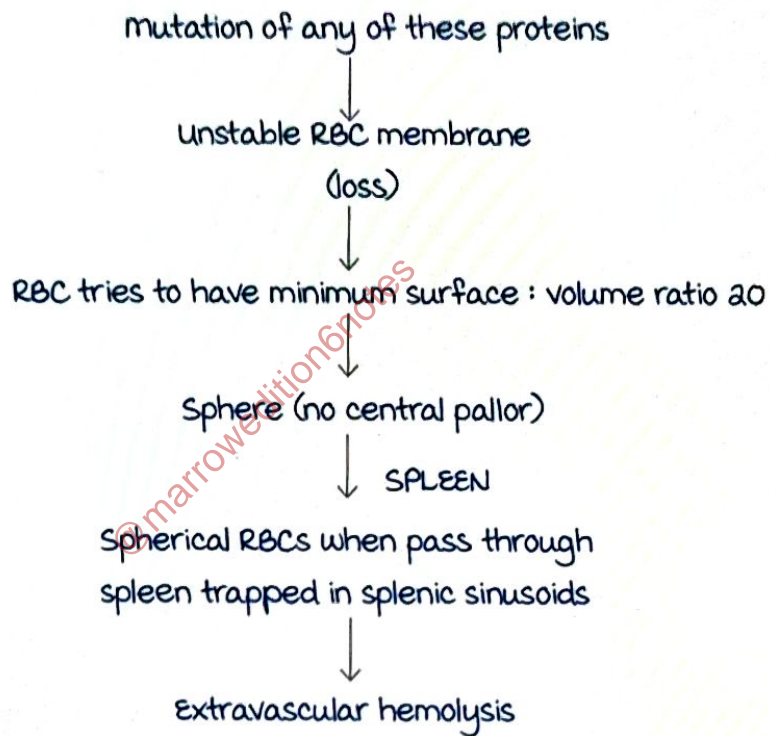
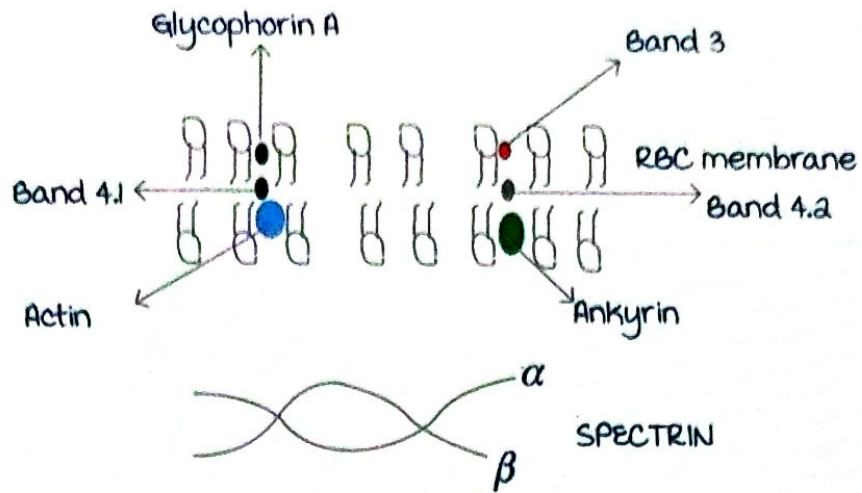
Pathogenesis :

Spectrin (contains alpha and beta chains) is responsible for biconcave shape/stability of RBCs.

membrane bound proteins :

- Ankyrin, band 4.2, band 3.
- Actin, band 4.1 and glycophorin A.

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Life span of normal RBCs : 120 days.

Life span of RBCs in hereditary spherocytosis : 10 to 20 days.

- Increase in MCHC is seen in hereditary spherocytosis. It is due to loss of K⁺ and water due to dehydration.
- most important/common protein defective in hereditary spherocytosis is ankyrin.
- Protein defect not seen in hereditary spherocytosis : Glycophorin A.
- most abundant protein in RBC membrane : Glycophorin A.

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- Spectrin mutations :
Common in hereditary elliptocytosis.
Produces most severe defects.

Clinical features :

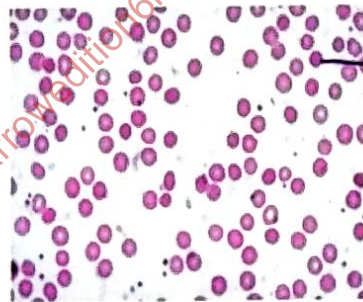
Pallor, jaundice, splenomegaly, increased risk of gallstones.

Aplastic crisis : Seen with parvovirus B19 infection.

Hemolytic crisis : Caused by EBV virus.

Lab tests :

- Hemoglobin : Decreased.
- TLC, platelet count : Normal.
- MCV, MCH : usually normal (MCV can be low).
- MCHC : Increased.
- RDW : Increased.
- Reticulocyte count : usually increased (It decreases in aplastic crisis).
- Peripheral smear :
Presence of spherocytes.
(small RBCs with no central pallor).



Other causes of spherocytes : Autoimmune hemolytic anemia (most common cause), burns, blood transfusion reactions.

Screening test : Osmotic fragility test.

RBCs of patient are suspended in increasing concentrations of normal saline.

Principle : Normal RBCs (biconcave) are isotonic with 0.9% NaCl : RBCs swell and rupture if suspended in increasing concentrations of normal saline (Normally, starts at 0.5% NaCl and completes by 0.3% NaCl)

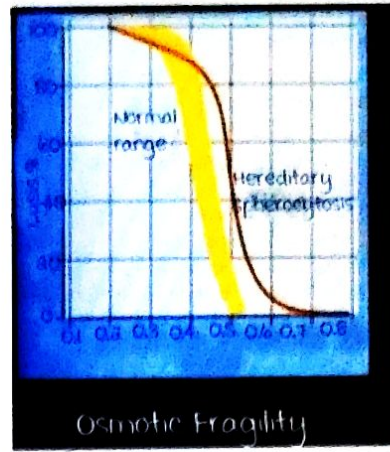
Spherocytic RBCs are fragile. They burst quickly at must lower concentration. The osmotic fragility curve shifts to the right.

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Osmotic fragility curve shifts to the left in thalassemia. Nestroft test is based on this principle.

Confirmatory test for hereditary spherocytosis: EMA binding test done by flow cytometry.



Treatment :

Splenectomy (spherocytic RBCs stay but anemia is corrected)
Peripheral smear of splenectomy patients show Howell jolly bodies.

G6PD deficiency

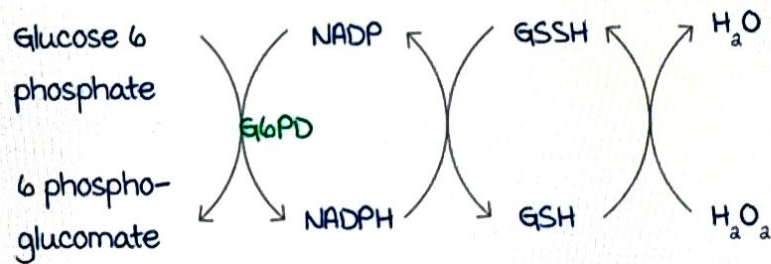
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X-linked recessive inheritance.
m >>> F.

Pathogenesis :

Hexose monophosphate shunt (HMP shunt) :



Deficiency of G6PD causes increase in hydrogen peroxide (H₂O₂) in a cell.

H₂O₂ is a free radical → Oxidative stress in a cell → RBC lysis.

Conditions causing hemolysis in G6PD deficiency :

- Chronic infections : Pneumonia
- Drugs : Antimalarials (primaquine).
- Fava beans (Favism).

G6PD deficiency is more common in people of African and

Active space

Mediterranean descent.

G6PD deficiency provides protection against *Plasmodium falciparum*.

Oxidative stress can lead to intravascular hemolysis, or it can lead to cross-linking of sulfhydryl groups in hemoglobin → Denaturation of hemoglobin → **Heinz bodies**.

When RBCs with Heinz bodies pass through spleen → Splenic macrophages try to pluck these Heinz bodies → membrane loss → **Bite cells** → **Extravascular hemolysis**.

Clinical features :

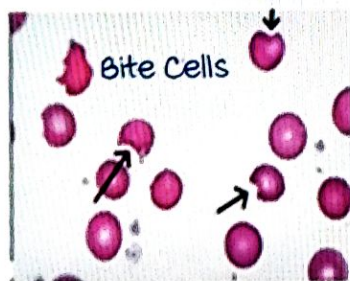
Episodic pallor, jaundice, hemoglobinuria (only when there is oxidative stress).

Splenomegaly and gallstones are features of chronic anemia. They are usually absent as hemolysis is episodic.

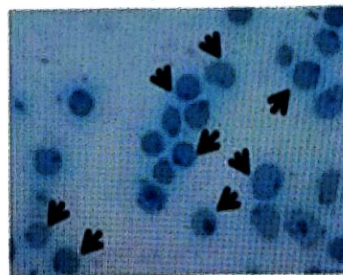
Lab investigations :

- Hemoglobin : Decreased.
- TLC, platelet count : Normal.
- MCV, MCH, MCHC : Usually normal.
- Reticulocyte count : Increased.
- Increase in urinary bilirubin.
- Peripheral smear : Bite cells or degmacytes and Heinz bodies.

Bite Cells



Heinz Bodies
New methylene blue stain



Heinz bodies are not seen on Romanowsky stain. They are seen on supravital stains like crystal violet or new methylene blue.

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Other tests for G6PD deficiency :

1. methemoglobin reduction test.
2. G6PD enzyme assay.

Hemolysis occurs more in older RBCs.

Treatment : Avoid oxidative stress.

Paroxysmal nocturnal hemoglobinuria (PNH) 00:57:26

Only **acquired** intracorpuscular defect.

Defect is at the level of stem cells.

Pathogenesis :

Normally, **PIG A** (phosphatidyl inositol glycan A) gene that synthesizes GPI anchored proteins :

- CD55 (DAF : Decay accelerating factor).
- CD 59 (MIRL : membrane inhibitor of reactive lysis).
- C 8 binding protein.

These GPI anchored proteins decrease the activity of complement (complement regulatory proteins).

In PNH, there is a mutation in PIG A gene → **Decreased** synthesis of GPI anchored proteins (CD 55, CD 59, C8 binding protein) → **Increased** complement activity → **Complement mediated hemolysis** → **Intravascular hemolysis**.

Complement also damages endothelium and leads to **thrombosis**.

most common/important protein defective in PNH is **CD59 (MIRL)**.

Clinical features :

- Pancytopenia.
- **Nocturnal hemoglobinuria** (seen in **25%** of cases) :
Blood pH decreases in sleep and increases complement activity.
- **Thrombosis** : most common cause of disease related death in PNH.
most commonly it presents as **hepatic vein thrombosis**.

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Complications :

- Acute myeloid leukemia.
- myelodysplastic syndrome.
- Aplastic anemia.

Diagnosis :

- Hemoglobin, TLC, platelet count decreased.
- Peripheral smear :
Normocytic normochromic anemia + pancytopenia.
- Reticulocyte count : Increased.
- Increased unconjugated bilirubin.

Other tests :

1. Ham's test / acidified serum lysis test.
2. Sucrose lysis test.
3. Flow cytometric evaluation of CD55 and CD59 : Best test.

Treatment :

Stem cell transplantation : **Best treatment.**

Eculizumab : Complement inhibitor.

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@marroweditionsnotes

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