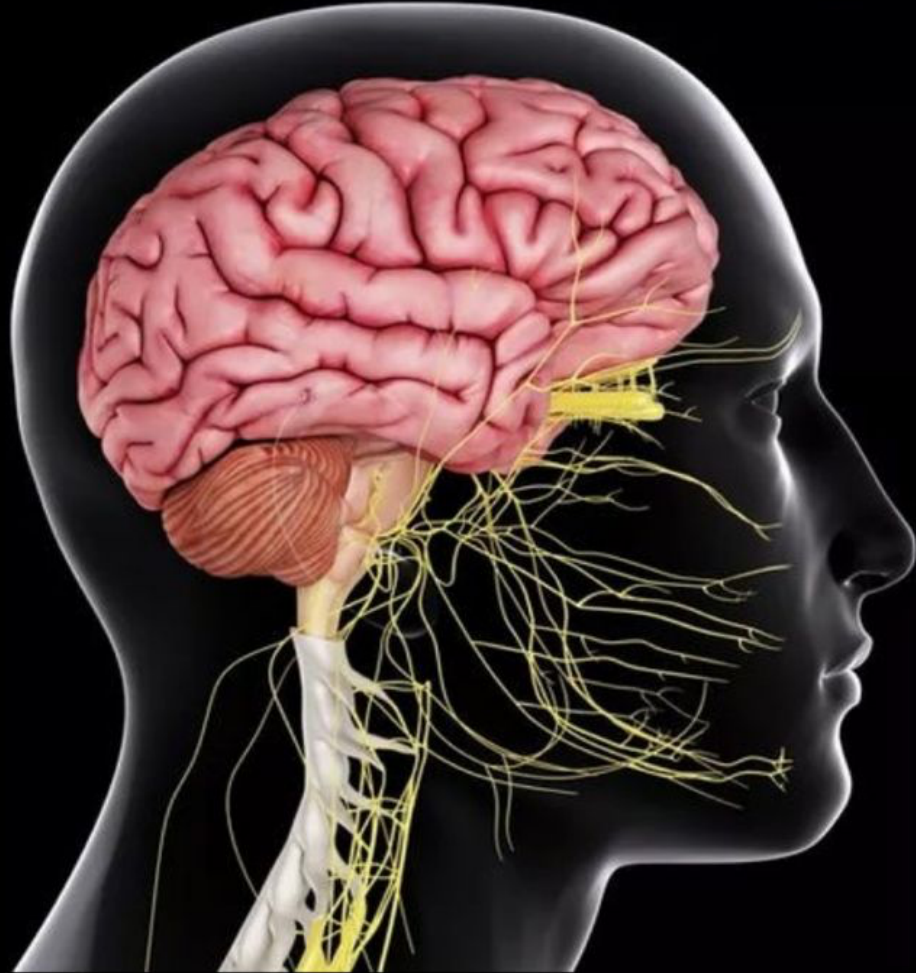


وَقُلْ رَبِّ زِدْنِي عِلْمًا



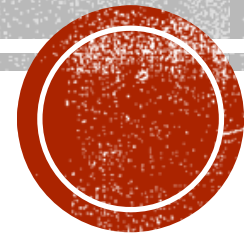
CENTRAL NERVOUS SYSTEM

SUBJECT : patho

LEC NO. : lec 1

DONE BY : sara nafi'

CNS LECTURE 1



Markers of Neuronal Injury, Edema, Herniation, and Hydrocephalus

Dr. Dua Abuquteish

MARKERS OF NEURONAL INJURY



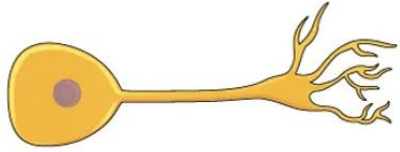
SIGNIFICANT FEATURES IN CNS PATHOLOGY

- ✓ Extremely susceptible to increased intracranial pressure (I.C.P)
- ✓ Highly susceptible to ischemia & hypoxia
- ✓ Site of lesion may be more important than its nature
- ✓ Selective vulnerability of defined structures to disease processes
- ✓ There is no regeneration (**gliosis** not fibrosis)

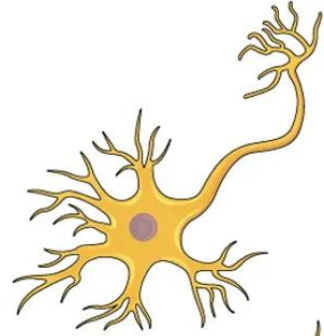
↳ no regeneration



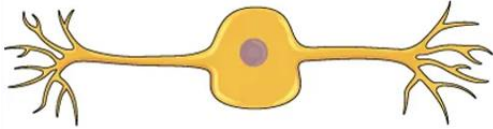
شماره بین تانجیسی



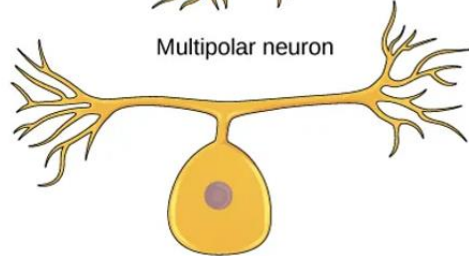
Unipolar neuron



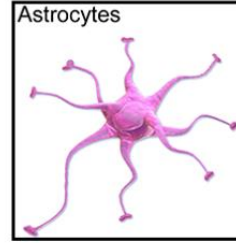
Multipolar neuron



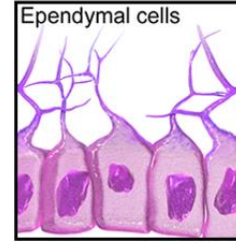
Bipolar neuron



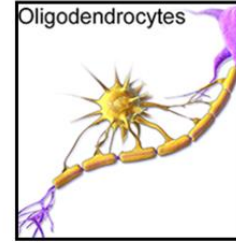
Pseudounipolar neuron



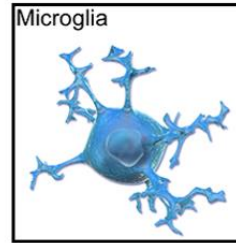
Astrocytes



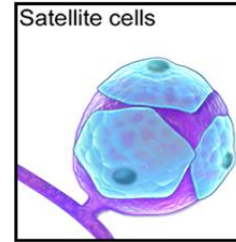
Ependymal cells



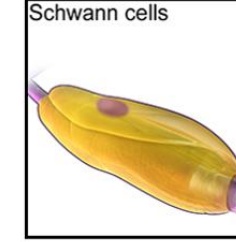
Oligodendrocytes



Microglia



Satellite cells



Schwann cells

Neurons

neurons are broadly divided into four basic types: unipolar, bipolar, multipolar and pseudounipolar.

www.majordifferences.com

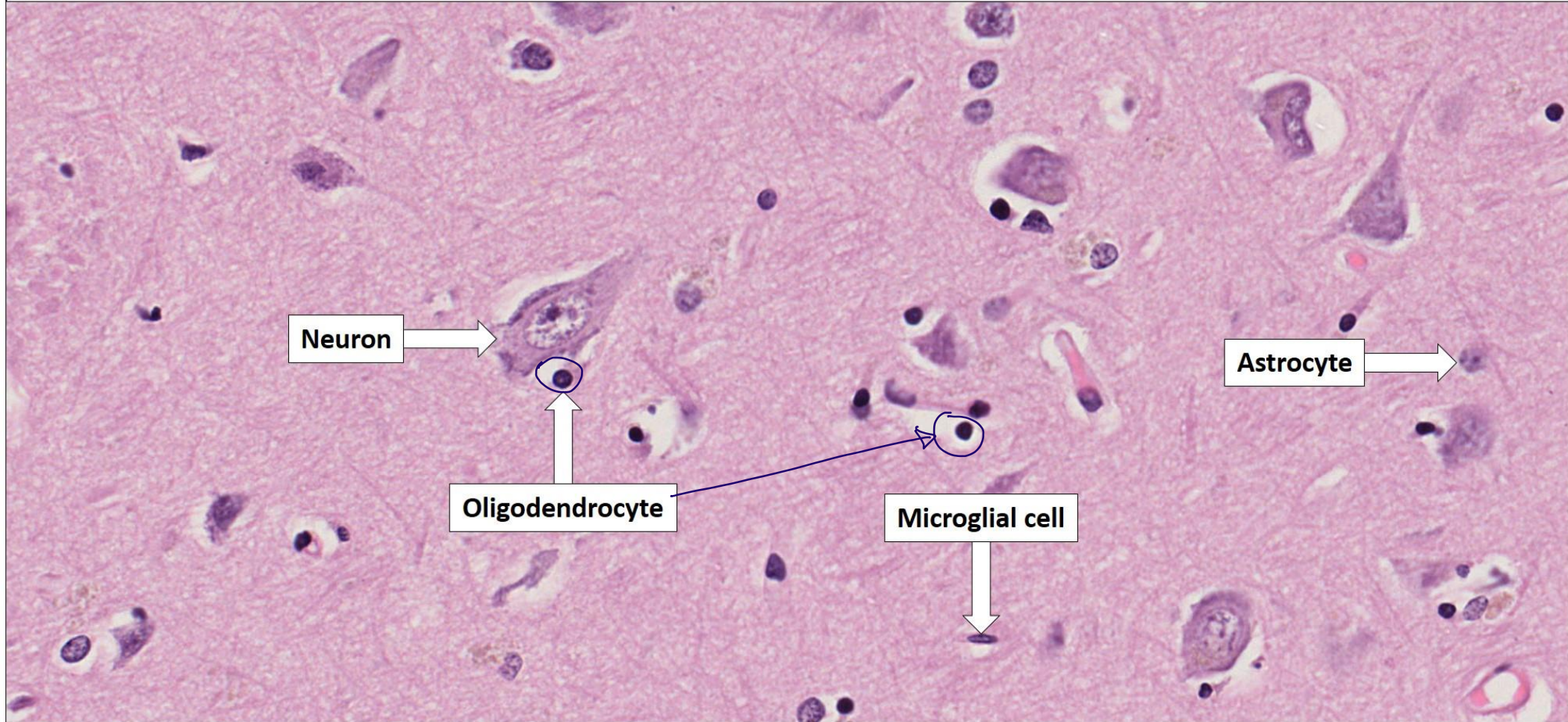
Glia

CNS Astrocytes, Ependymal cells
Oligodendrocytes, Microglia, Satellite cells, Schwann cells *PNS*

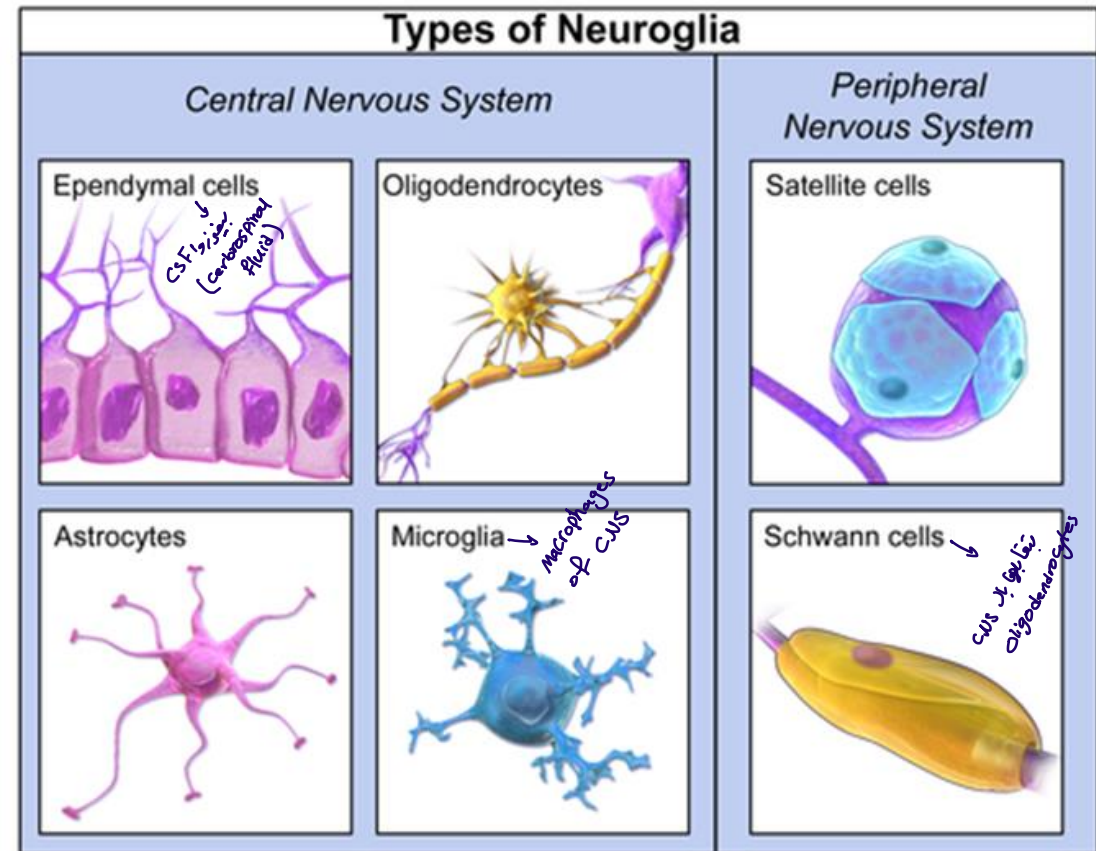
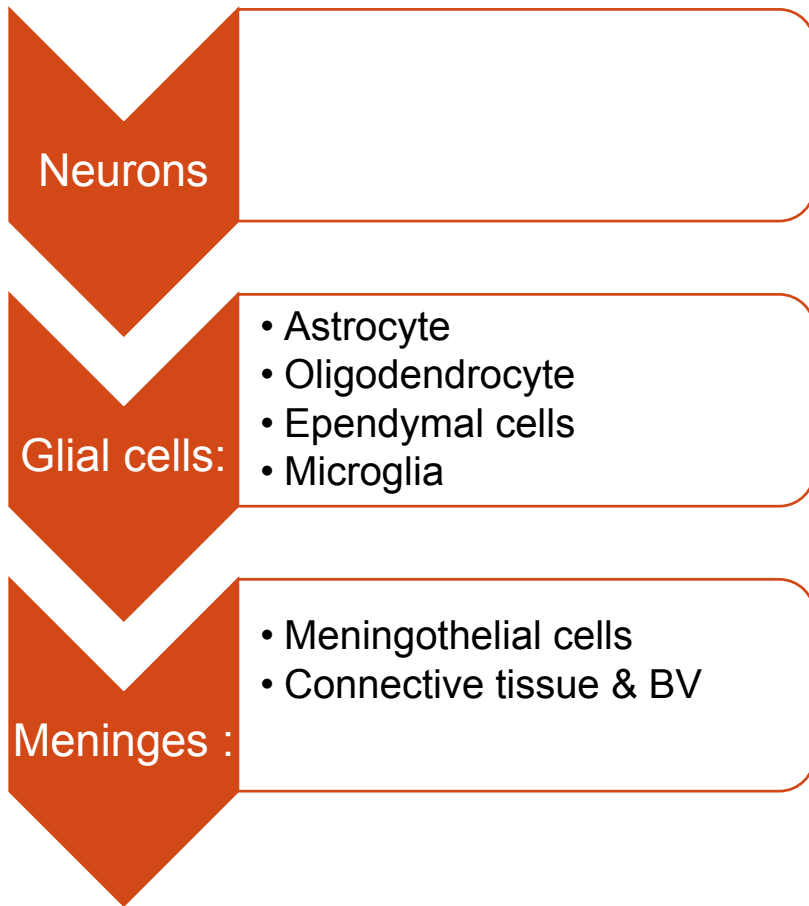


Cerebral cortex (High power):

Neurons typically have large, pale nuclei with prominent nucleoli. The non-neuronal (glial) cells in the grey matter include oligodendrocytes (hyperchromatic, round nuclei and abundant clear-appearing cytoplasm), astrocytes (paler, more elongated nuclei and usually scant cytoplasm), and microglial cells.



CELLS OF CNS

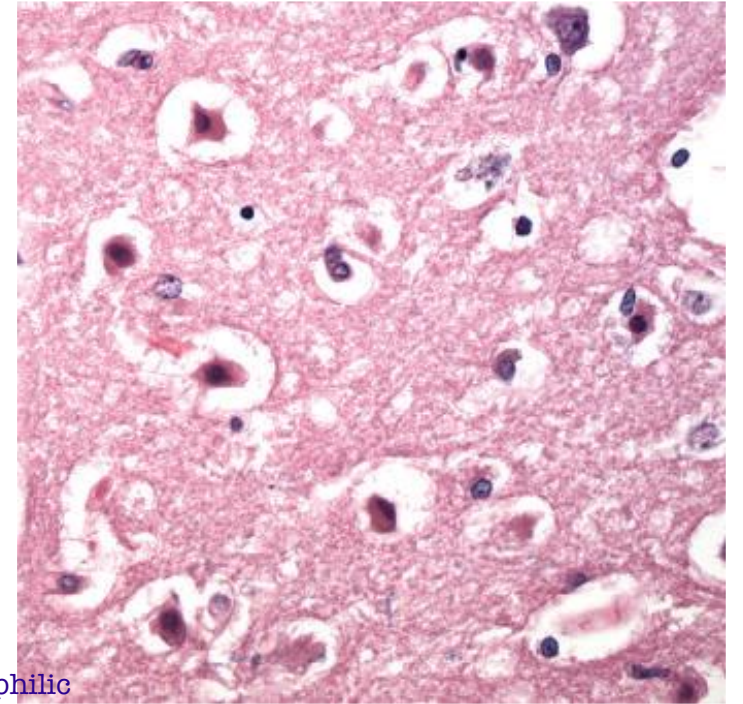


PATTERNS OF NEURONAL INJURY

Acute hypoxic/ischemic injury in cerebral cortex:

Within 12 hours of an irreversible hypoxic/ischemic insult, **acute neuronal injury** becomes evident even on routine H & E staining.

The necrotic neuronal cell bodies & their nuclei are shrunken & pyknotic, loss of Nissl substance, and prominently eosinophilic, so-called "**red neurons**"



dying cells due to acute insult

it's basophilic

مشان هيك لما نخسرها
بتصير الخلية

↳ red neurons

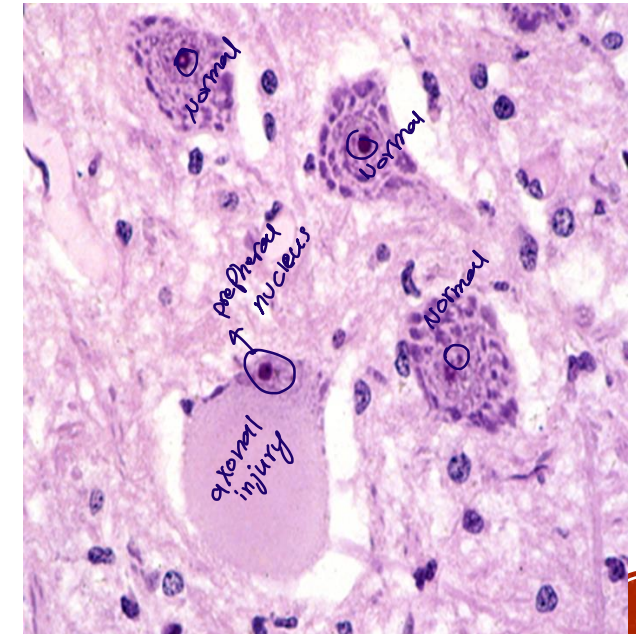
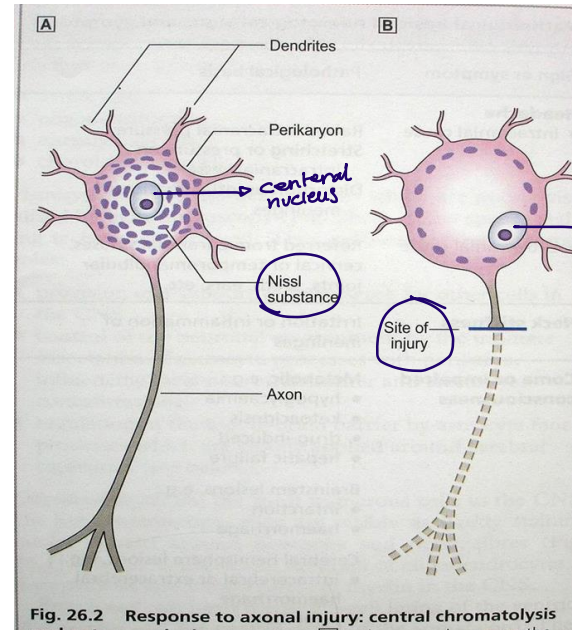


PATTERNS OF NEURONAL INJURY

Axonal reaction:

Features of axonal injury:

Axonal injury also leads to cell body enlargement & swelling and dispersion of Nissl substance from the cell center to the periphery (**central chromatolysis**)



CHRONIC OR SUBACUTE INJURY

- Degeneration and neuronal loss & replacement by **gliosis** in progressive diseases.

بدل ال fibrosis

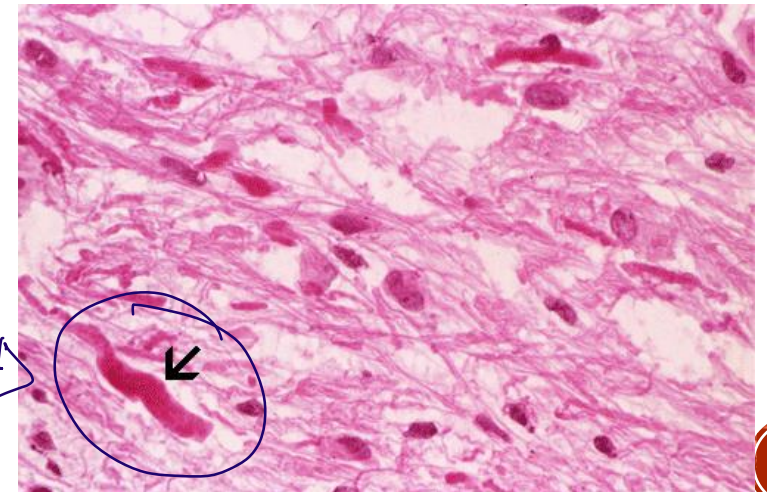
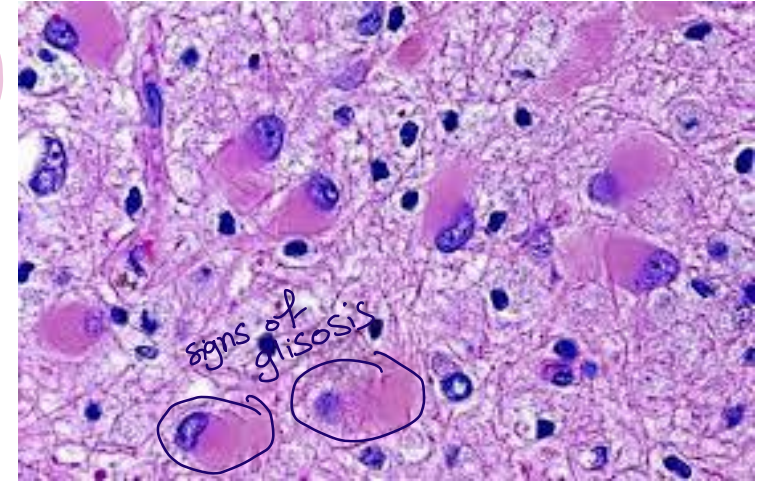
بجانب مكان ال injury
hypertrophy
hyperplasia

▪ **Astrocytes** are the principal cells responsible for repair and scar formation in the brain, a process termed gliosis. In response to injury, astrocytes undergo both hypertrophy and hyperplasia.



ASTROCYTE INJURY AND REPAIR

- In response to injury, astrocytes undergoes (**Gliosis**)
- The nucleus enlarges, the nucleolus becomes prominent, and the cytoplasm becomes bright pink hue (gemistocytic astrocyte)
- Unlike elsewhere in the body, fibroblasts participate in healing after brain injury to a limited extent
- Rosenthal fibers are thick, elongated, brightly eosinophilic protein aggregates found in astrocytic processes in chronic gliosis and tumors

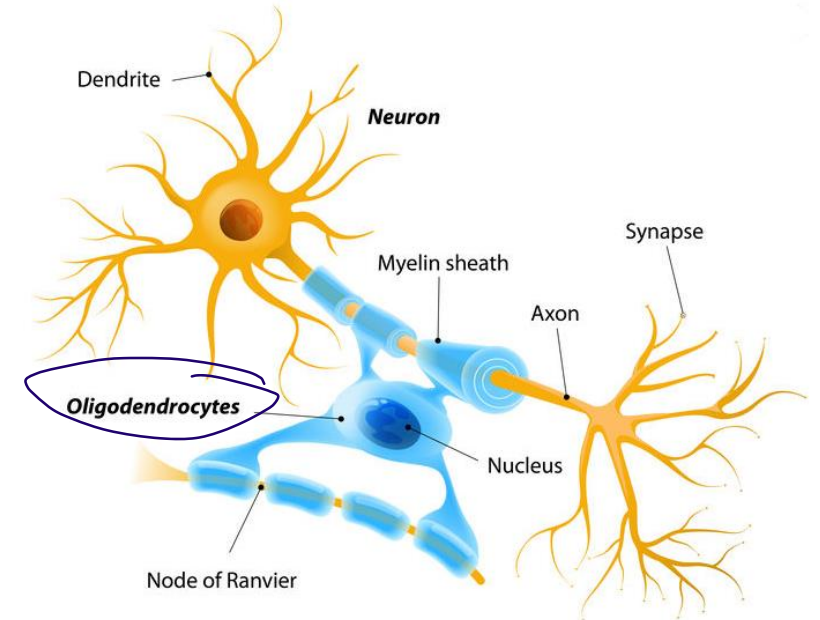


في Tumors
بمشيمة بوردو

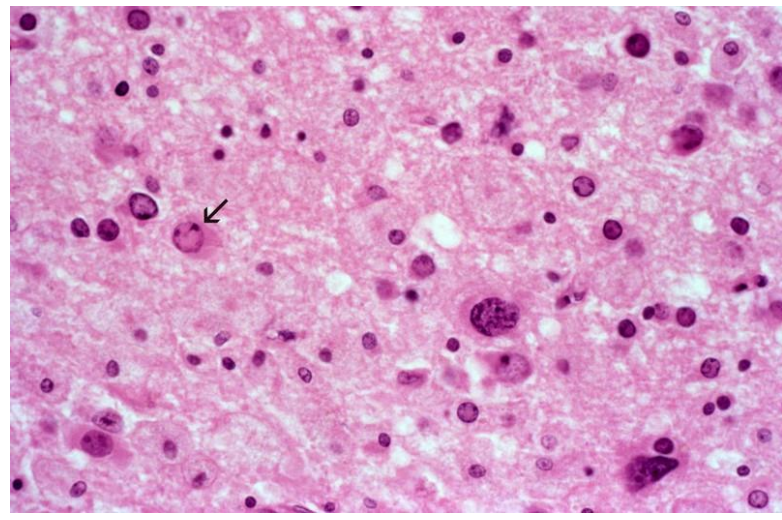


OLIGODENDROCYTES

- Produces myelin, exhibit a limited spectrum of changes in response to injuries.
- For example: in progressive multifocal leukoencephalopathy, viral inclusions can be seen in oligodendrocytes.

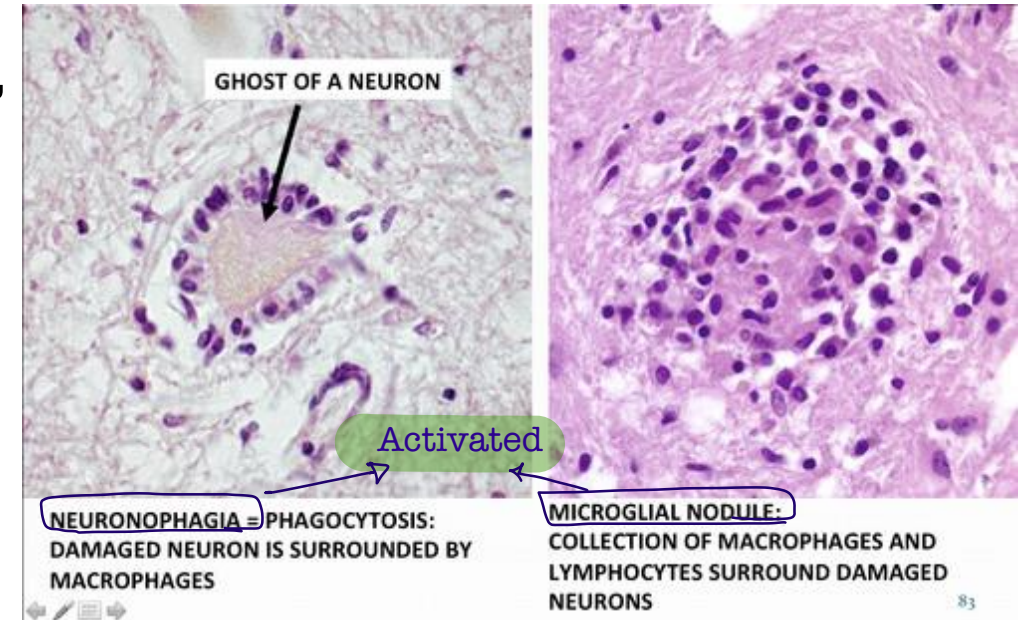


موتان سنجي ← فقط هيا هيا
باللبن ←
Leuko ← Disease
myelin sheath



MICROGLIAL CELLS

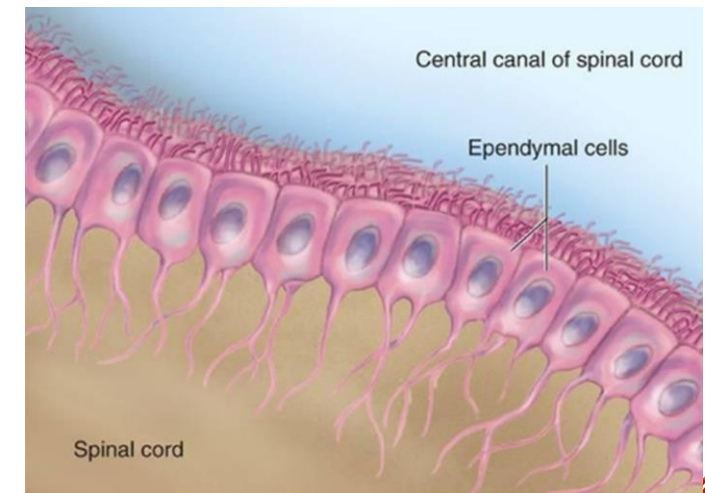
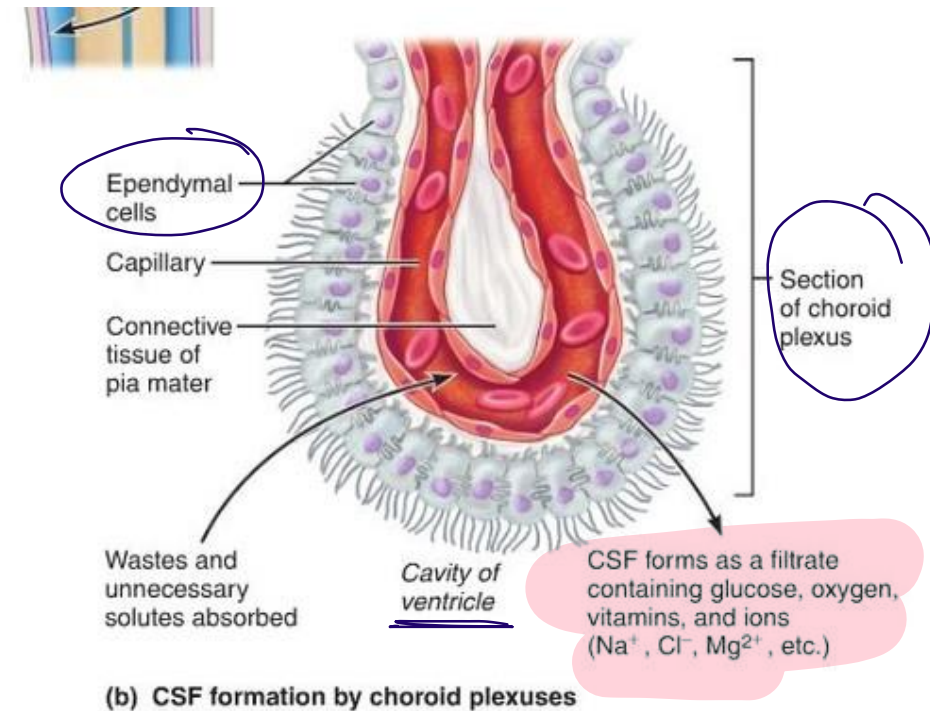
- **Microglial cells** are long-lived cells that function as the **phagocytes of the CNS**.
- When activated by tissue injury, infection, or trauma, they proliferate and become more prominent histologically.
- Microglial cells have the appearance of **activated macrophages** in areas of demyelination, organizing infarct, or hemorrhage
- Aggregates of elongated microglial cells at sites of tissue injury are termed **microglial nodules**
- Collections around and phagocytosing injured neurons (**neuronophagia**).



EPENDYMAL CELLS

ciliated cells lining
choroid plexus ←

- Ependymal cells line the ventricular system and the central canal of the spinal cord.
- Certain pathogens, particularly cytomegalovirus (CMV), can produce extensive ependymal injury.
- Choroid plexus is in continuity with the ependyma, and its specialized epithelial covering is responsible for the secretion of cerebrospinal fluid (CSF).



- The brain and spinal cord are encased within the skull and spinal canal, with nerves and blood vessels passing through specific foramina. These rigid structures provide little room for brain expansion in diseases.
- **An increase in the volume of the skull contents (Brain, CSF, & blood) → increase in intracranial pressure (ICP)**
- Increased ICP compromises the blood supply to the brain, resulting in decreased brain perfusion and serious/fatal consequences.

Causes of increase ICP:

- Generalized cerebral edema ↑Brain volume
- Hydrocephalus (↑CSF)
- Hemorrhages (↑Blood)
- Ischemia → leads to cerebral edema
- Masses (tumors)

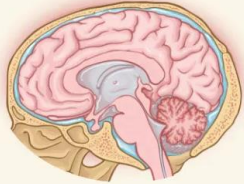
↓
 Mass effect: obstruction:
 Edema ↑CSF



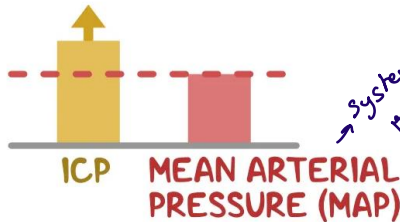
حزرم نؤخو
بسنكل عام

↑ INTRACRANIAL PRESSURE (ICP)

⚠ MEDICAL EMERGENCY ⚠



CAN RESULT from:
HEAD INJURY, BLEEDING in BRAIN,
TUMORS, INFECTIONS, EXTRA FLUID
in BRAIN, STROKE



WHEN ICP > MAP,
BRAIN CAN NO LONGER
RECEIVE ENOUGH O₂

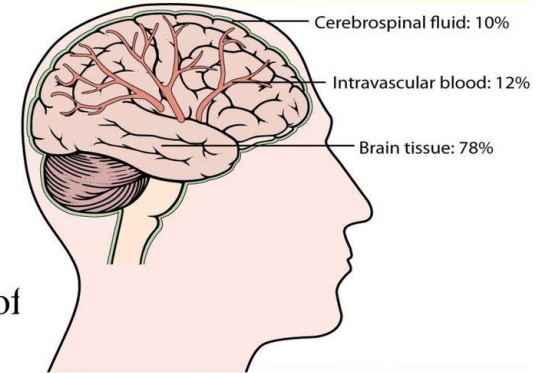
↳ SYMPATHETIC NERVOUS SYSTEM ACTIVATES

↳ THEN PARASYMPATHETIC NERVOUS SYSTEM ACTIVATES

□ ICP is usually measured in the lateral ventricles, with a normal pressure of 0 to 15 mm Hg.



- brain tissue (1400 g)
- Blood 75 mL
- CSF 75 mL
- the three components are in a state of equilibrium



↑ ICP → >15

Symptoms of ICP

Headache
Nausea and Vomiting
Altered Level of Consciousness
Vision Problems

* Papilledema → check the eyes for optic disc

Seizures

Cushing's Triad (a late sign) → activation of parasympathetic system

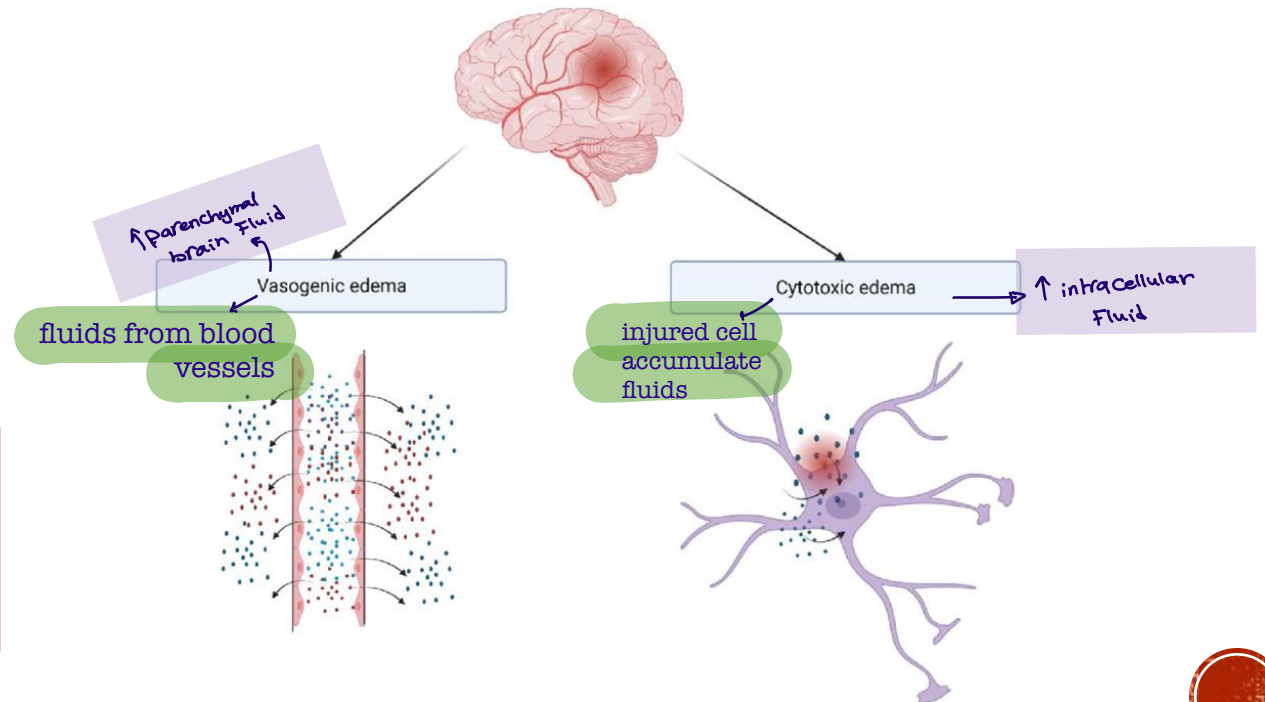
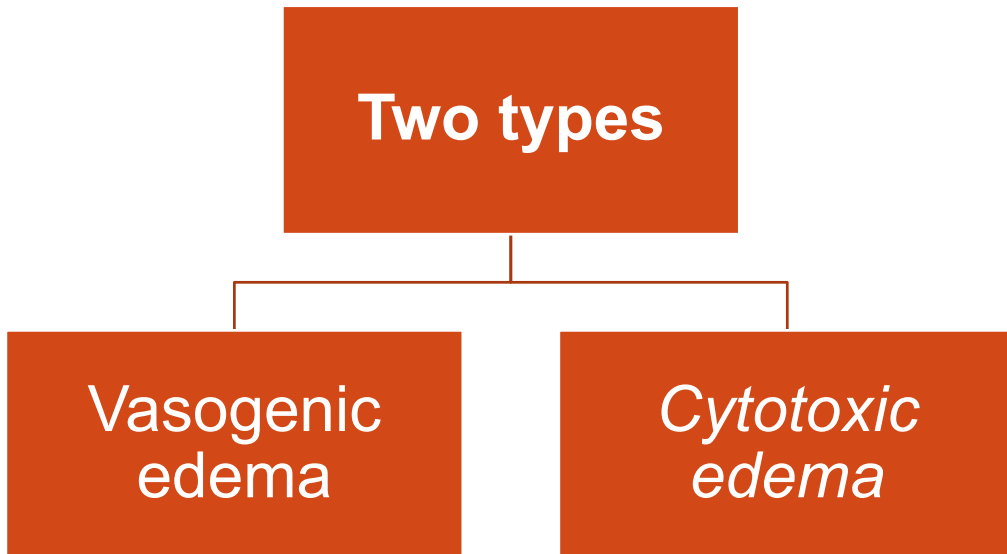


CEREBRAL EDEMA



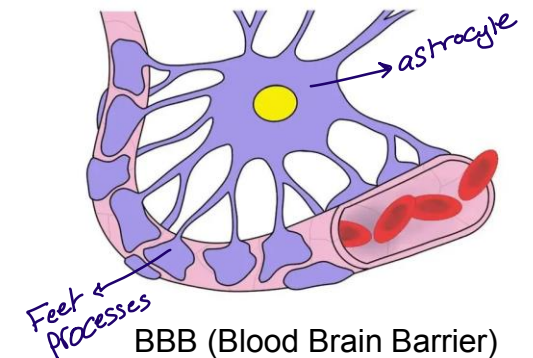
CEREBRAL EDEMA → ↑ ICP

- Cerebral edema is the accumulation of **excess fluid within the brain parenchyma**
- Cerebral edema commonly present with neurological symptoms, and if severe it can cause ↑ ICP, herniation, and death

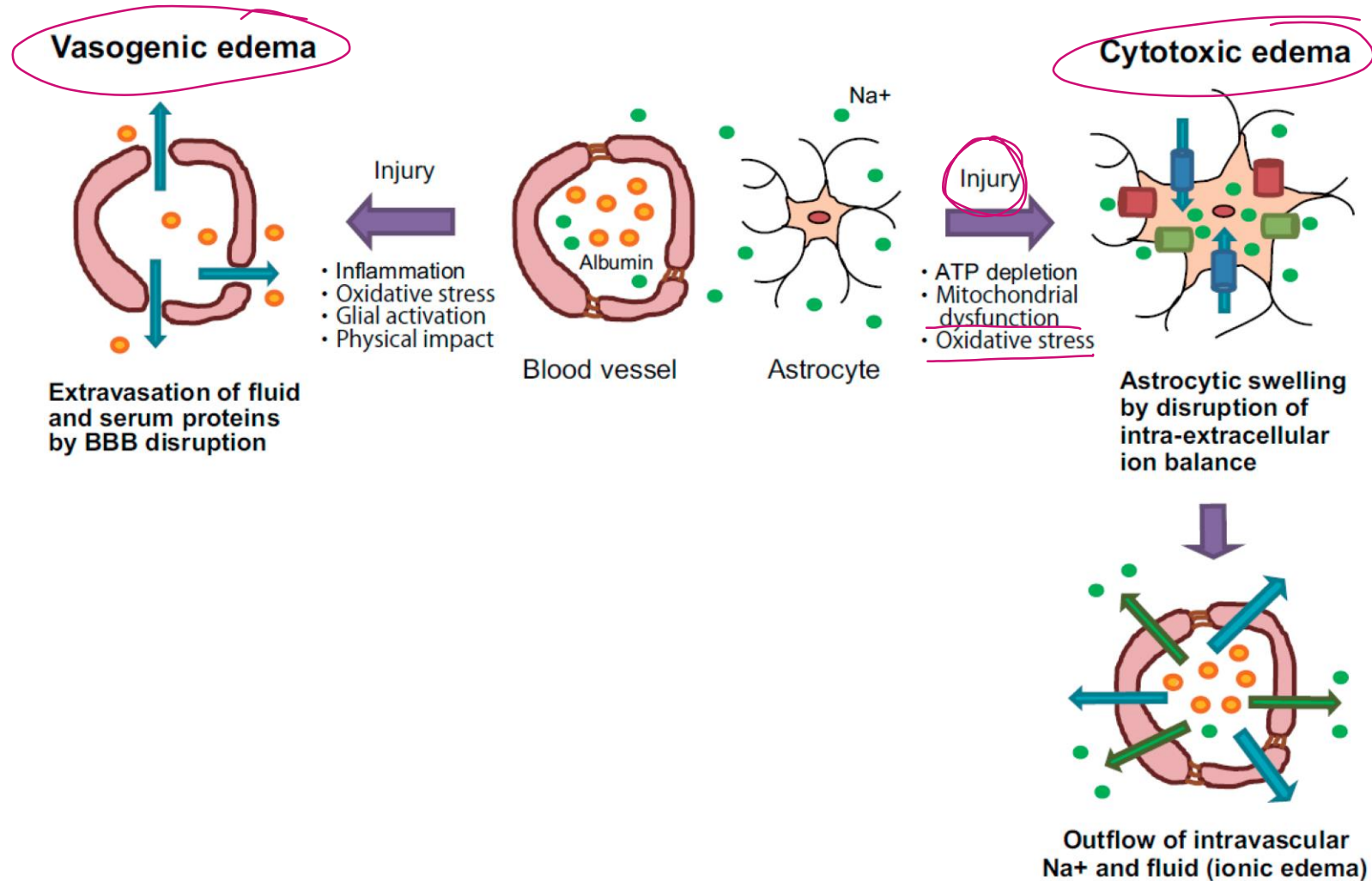


CEREBRAL EDEMA

- **Vasogenic edema** occurs when the integrity of the normal blood-brain barrier is disrupted, allowing fluid to shift from the vascular compartment into the extracellular spaces of the brain.
- Vasogenic edema can be localized (e.g., inflammation or in tumors) or generalized.
- **Cytotoxic edema** is an increase in intracellular fluid secondary to neuronal and glial cell injury, as might follow generalized hypoxic or ischemic insult or exposure to toxins.

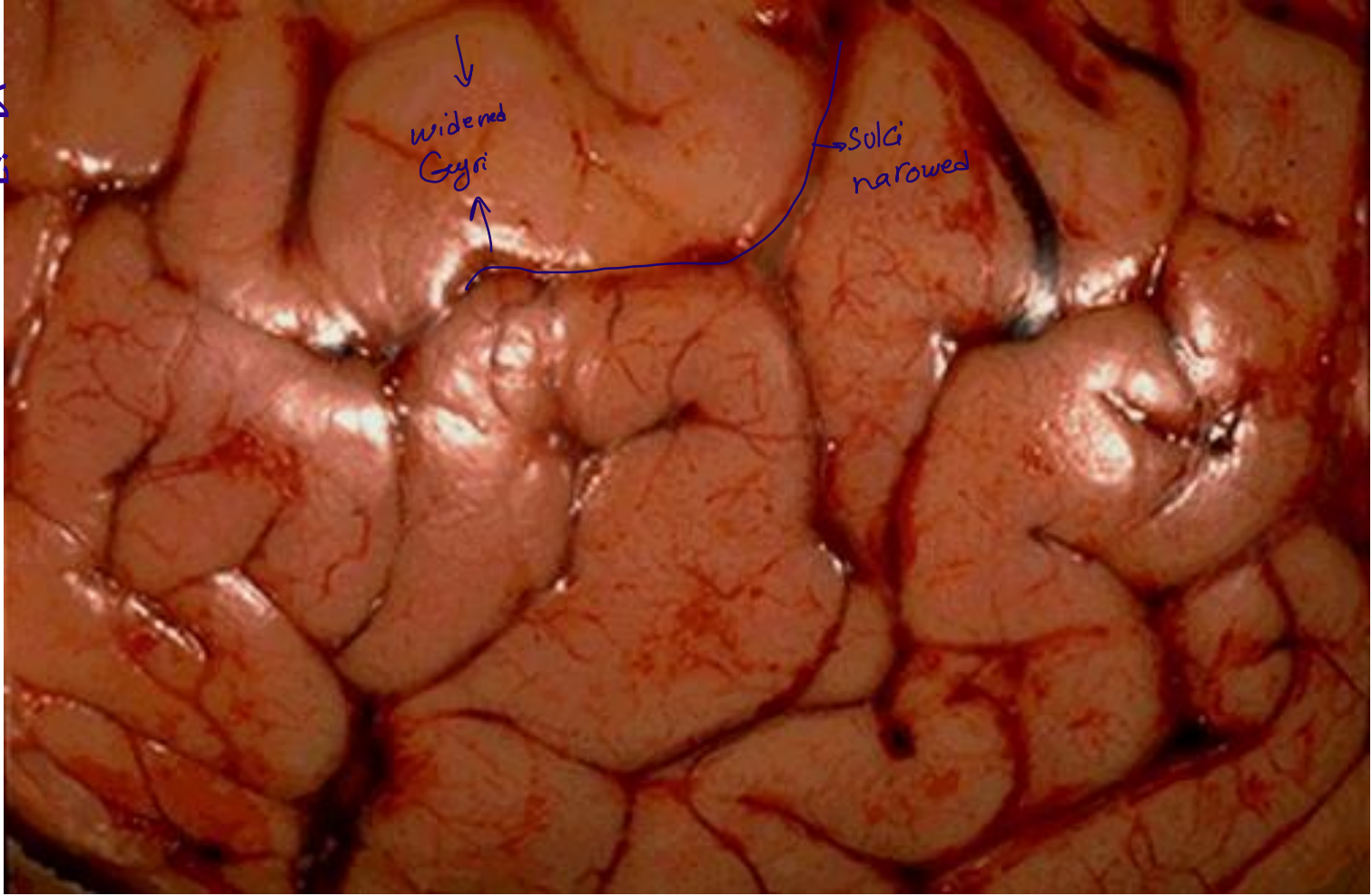


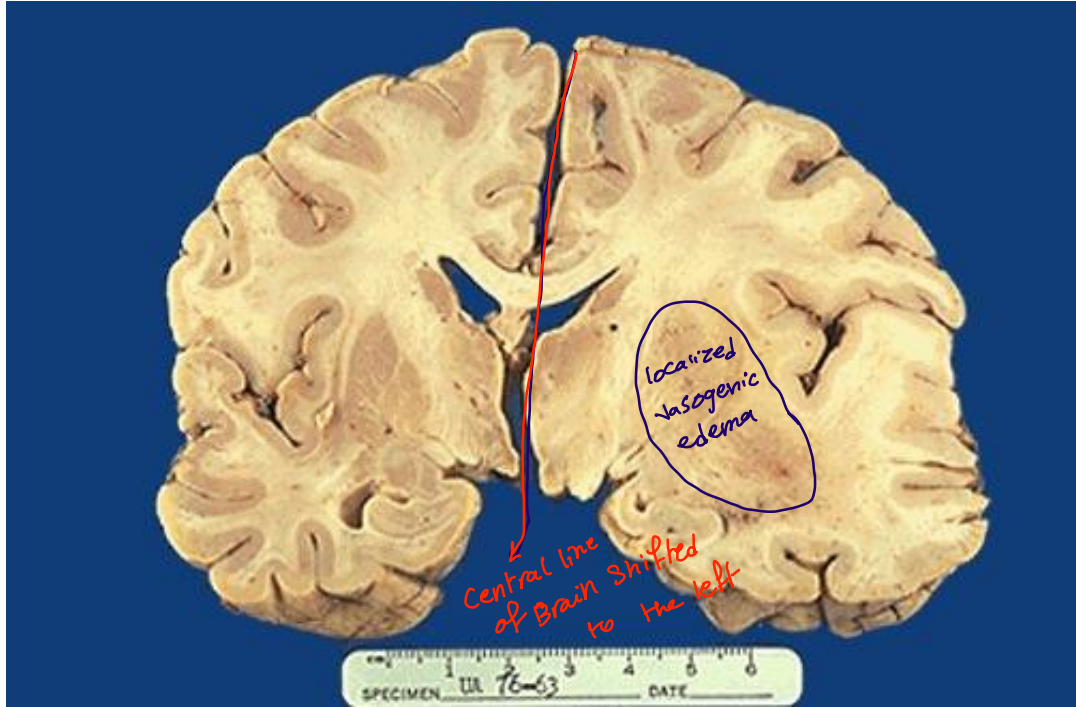
CEREBRAL EDEMA



The surface of the brain with cerebral edema demonstrates widened gyri with a flattened surface. the sulci are narrowed.

كل هي الصور راح
تدخل باللاب





There is cerebral edema seen at the right which obscures the structures. There is a shift of the midline to the left. Multiple small metastases were the cause for the edema in this case



Acute brain swelling is serious. Swelling of the left cerebral hemisphere has produced a shift with herniation of the uncus of the hippocampus through the tentorium, leading to the groove seen at the white arrowhead.



HYDROCEPHALUS

→ ↑ CSF volume within the ventricular system

- Choroid plexus produces CSF within the ventricles
- CSF then circulates through the ventricular system and flows through the foramina of Luschka and Magendie into the subarachnoid space, where it is absorbed by arachnoid granulations (arachnoid villi). → then drain CSF to the venous system
- The balance between rates of CSF generation and resorption regulates CSF volume.
- **Hydrocephalus is an increase in the volume of the CSF within the ventricular system.**

accunulation of csf ← اي مشكلة بتصير فيهم بصير في ; لهذا مهمين للcsf

في balance بين

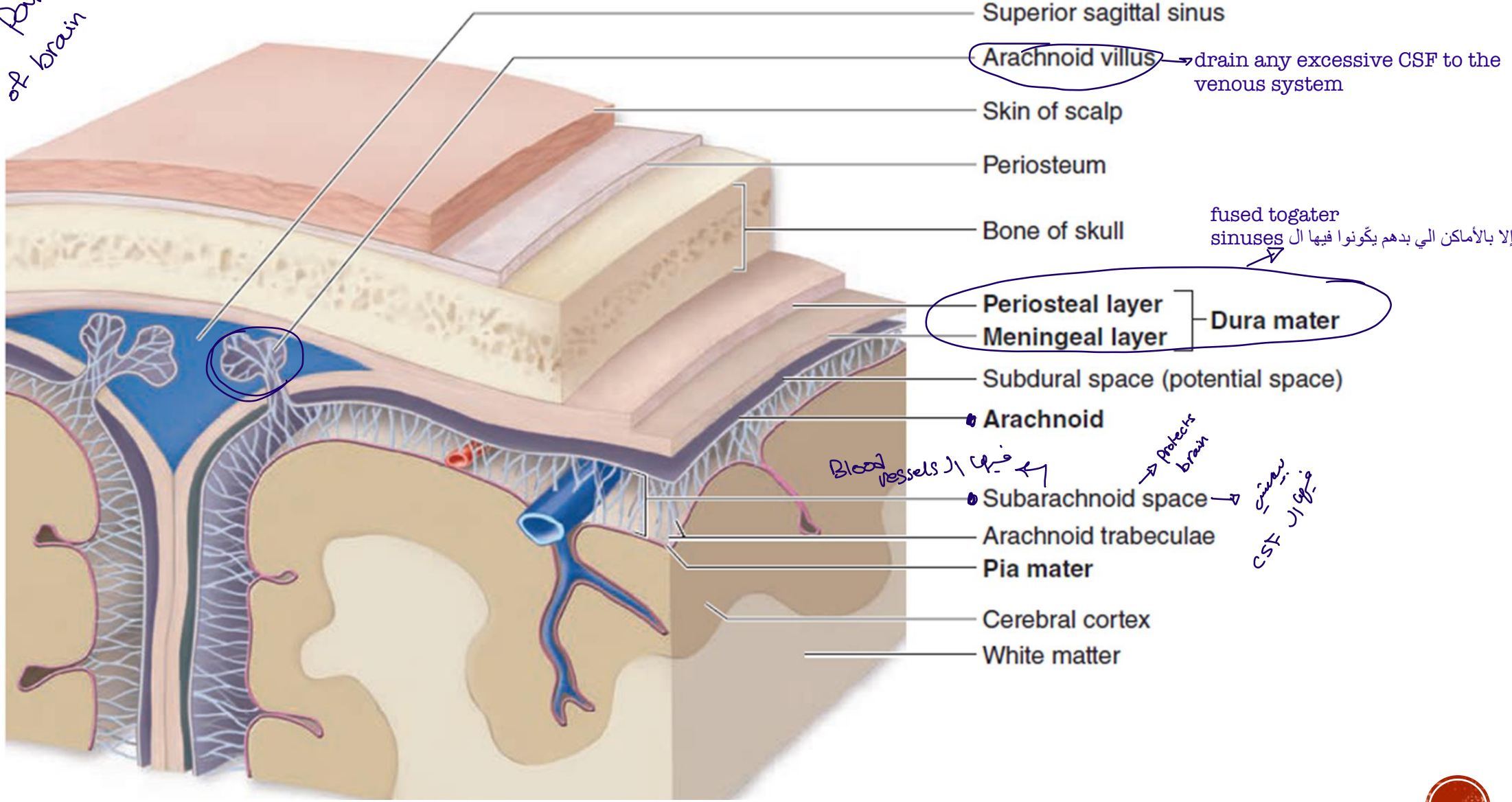
$\frac{\text{CSF generation}}{\text{CSF resorption}}$

Causes:

- Impaired flow or resorption of CSF
- Overproduction of CSF (e.g. some tumors of choroid plexus)

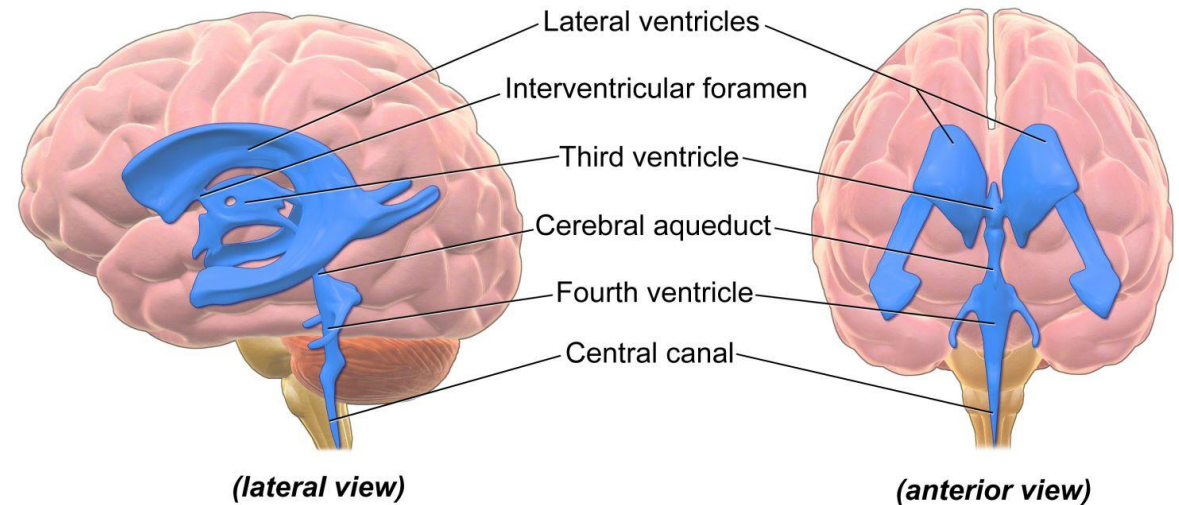
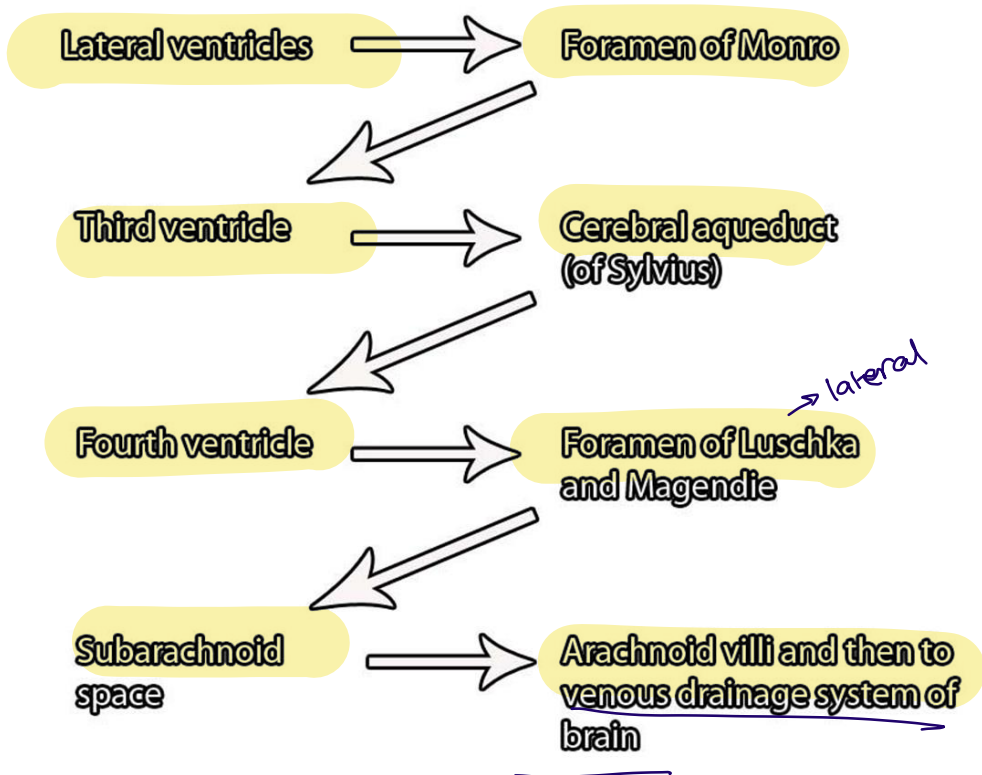


Upper Part
of brain



عربي prep
Pathway of CSF

Cerebrospinal Fluid Pathway



المختصر
↓

Anything obstruct the pathway of CSF will lead to **accumulation** of CSF in the Arachnoid villi an subarachnoid **which** will lead to hydrocephalus



HYDROCEPHALUS -TYPES

في شي عمل obstruction قطع الطريق

- **Non-communicating hydrocephalus:** there is obstruction to CSF flow within the ventricular system, then a portion of the ventricles enlarges while the remainder does not. → in tumors, Congenital problems: Atresia

No obstruction

inflammation, meningitis
Fibrosis → خاصة اذا كنته
arachnoid villi بل

- **Communicating hydrocephalus:** it is usually caused by reduced CSF resorption, and the entire ventricular system is enlarged

↳ ↑CSF volume

not clinical significant
the problem is the underlying neurological disease

- **Hydrocephalus ex vacuo:** a compensatory increase in CSF volume may occur secondary to a loss of brain volume (e.g., infarction, neurodegenerative disease). In such settings, the hydrocephalus is of no clinical significance.

* communicating → flow within the ventricles

: obstruction فلما يصير

• are they still

• أو هل في شي مسكر الطريق بينهم!

مشان أقرر communicating or non communicating

① disease like Alzheimer, dementia

② loss of brain volume (tissue)

③ Atrophy

④ ventricles become dilated

⑤ ↑CSF



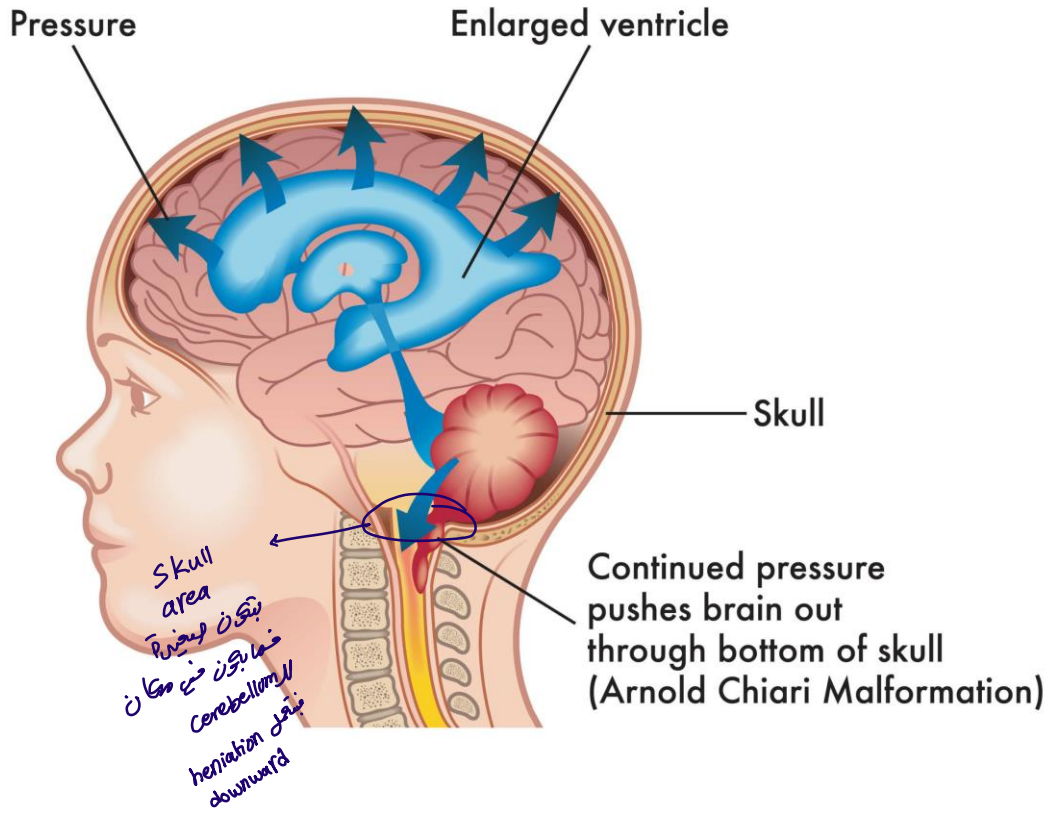
NON-COMMUNICATING HYDROCEPHALUS

- **Foramen of Monro obstruction:** dilation of one or both lateral ventricles.
- **The aqueduct of Sylvius** obstruction (e.g., atresia, hemorrhage, or tumor) and lead to dilation of both lateral ventricles, as well as the third ventricle.
- **Fourth ventricle obstruction** leads to dilatation of the aqueduct, as well as the lateral and third ventricles (e.g., Chiari malformation).
- **The foramina of Luschka and foramen of Magendie** may be obstructed due to congenital malformation (e.g., Dandy-Walker malformation).



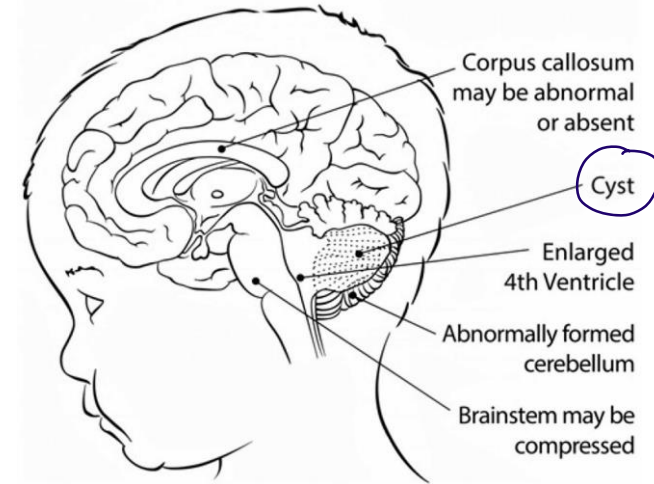
Chiari malformation

↳ malformation in skull development



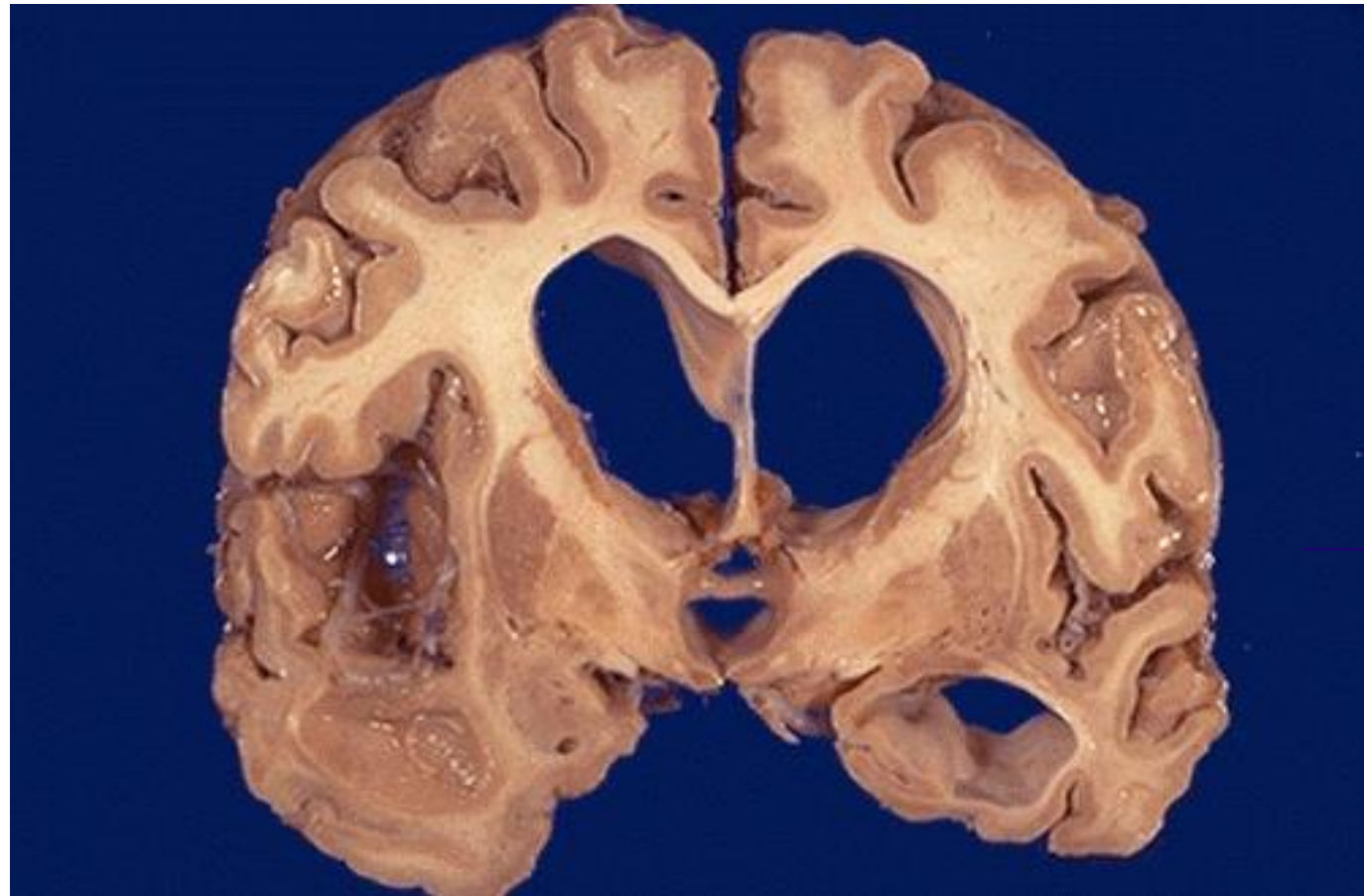
↳ become a cyst
Congenital failure of cerebellar vermis to develop so that 4th ventricle is massively dilated and cerebellum is absent; often accompanied by hydrocephalus

Dandy Walker Syndrome



Note the marked dilation of the cerebral ventricles in hydrocephalus.

Hydrocephalus can be due to lack of absorption of CSF or due to an obstruction to flow of CSF



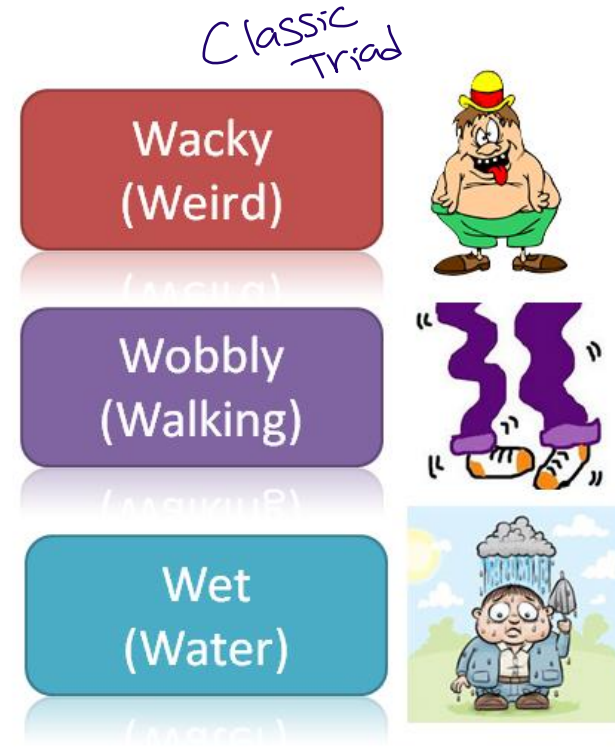
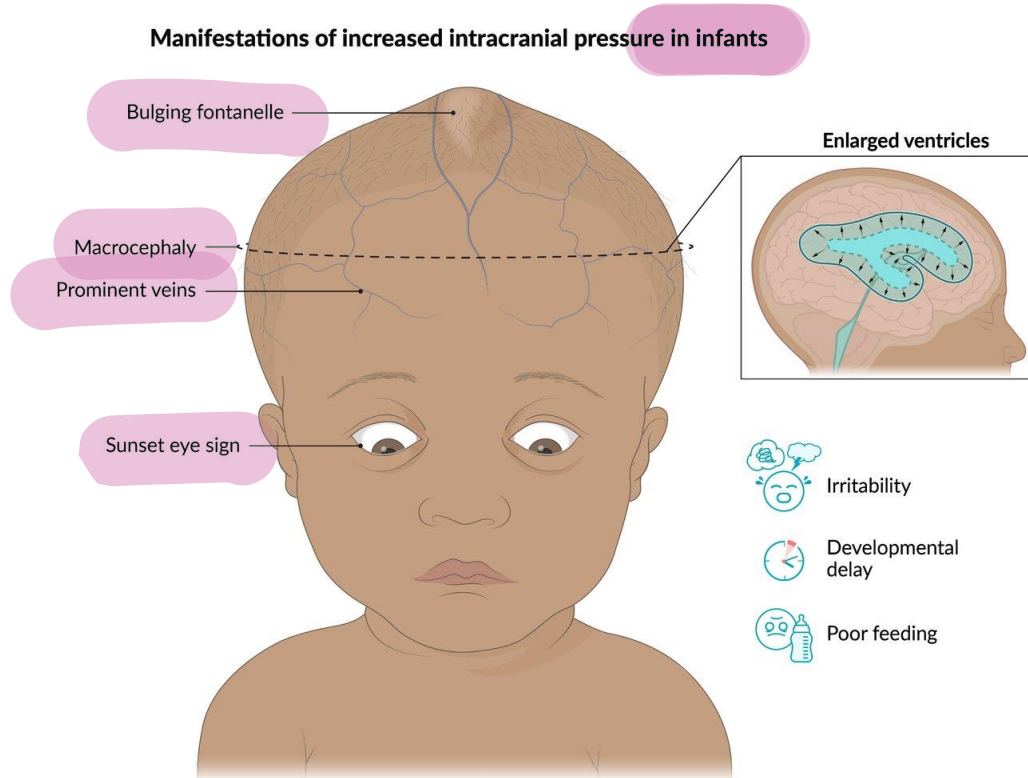
→ dilation of the lateral ventricle



If hydrocephalus develops in infancy before closure of the cranial sutures, the head enlarges.



Symptoms of hydrocephalus in babies and adults



Urinary incontinence (“wet”), gait instability (“wobbly”), and cognitive changes (“wacky”).



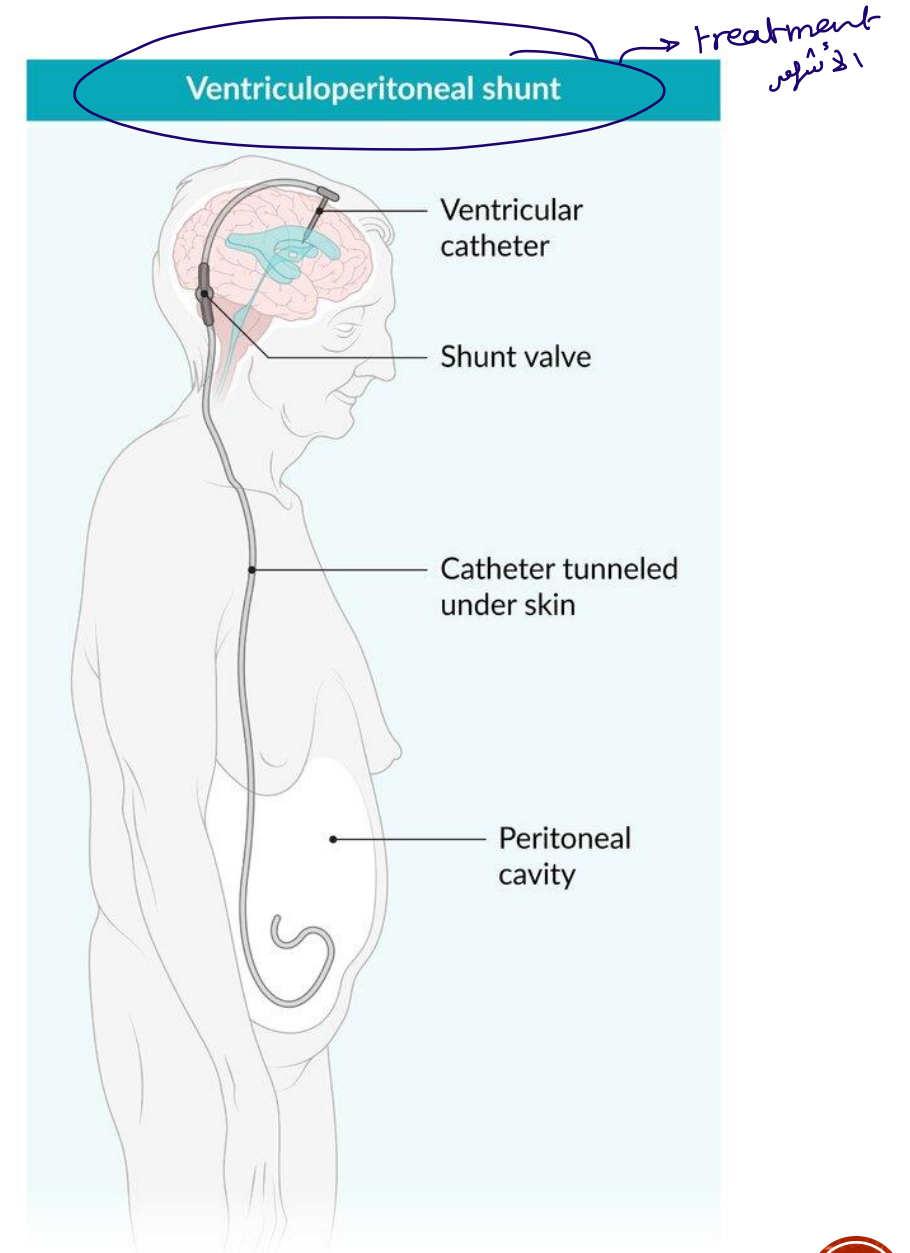
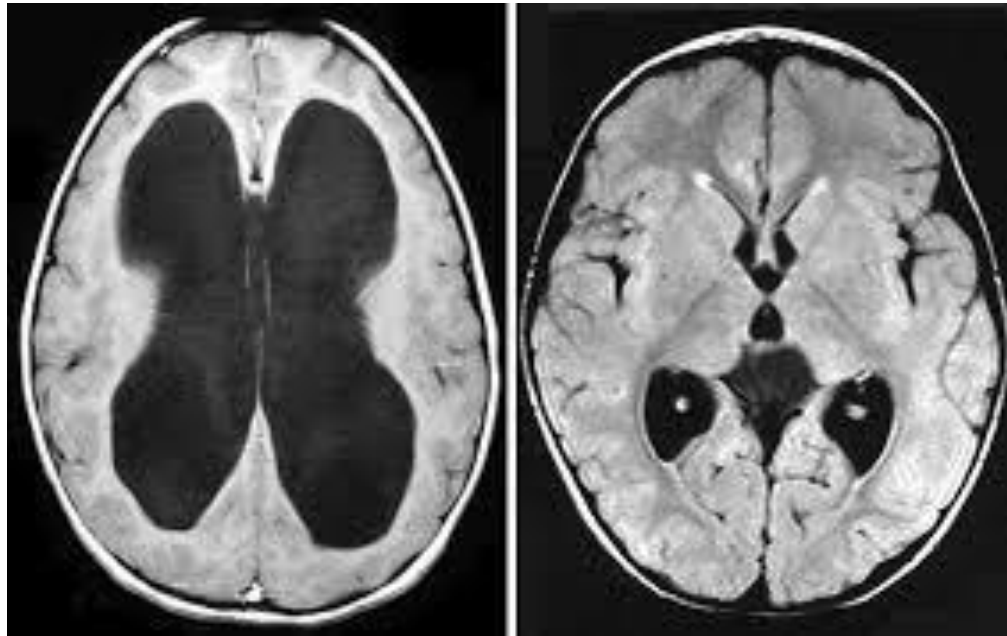
Overview of hydrocephalus

	Pathophysiology	Clinical features	Diagnosis
Communicating hydrocephalus	<ul style="list-style-type: none"> ↑ CSF production ↓ CSF absorption 	<ul style="list-style-type: none"> Typical findings of raised ICP <ul style="list-style-type: none"> Headache, nausea, and vomiting Papilledema Abducens nerve palsy Abnormal gait Impaired consciousness Cushing triad (irregular breathing, widening pulse pressure, bradycardia) <p><i>Handwritten note: Symptoms ↑ ICP</i></p>	<ul style="list-style-type: none"> Ultrasonography (children ≤ 18 months of age) MRI or CT (older children and adults)
Noncommunicating hydrocephalus	<ul style="list-style-type: none"> Obstructed passage of CSF from the ventricles to the subarachnoidal space 		
Normal pressure hydrocephalus (NPH) → very chronic	<p>The brain will try to reserve the ICP ← <i>Chronic</i></p> <ul style="list-style-type: none"> ↓ CSF absorption 	<ul style="list-style-type: none"> Classic triad <ul style="list-style-type: none"> Wet: urinary incontinence Wacky: dementia Wobbly: frequent falls, broad-based gait with short, shuffling steps (gait apraxia) <p><i>Handwritten note: No symptoms of ↑ ICP</i></p>	<ul style="list-style-type: none"> MRI (initial test) CSF tap test
Hydrocephalus ex vacuo → very chronic	<ul style="list-style-type: none"> Loss of brain tissue 	<ul style="list-style-type: none"> Symptoms of the underlying condition (e.g., Alzheimer disease, Pick disease) 	<ul style="list-style-type: none"> Cortical atrophy may be prominent on imaging.



Diagnosis and treatment

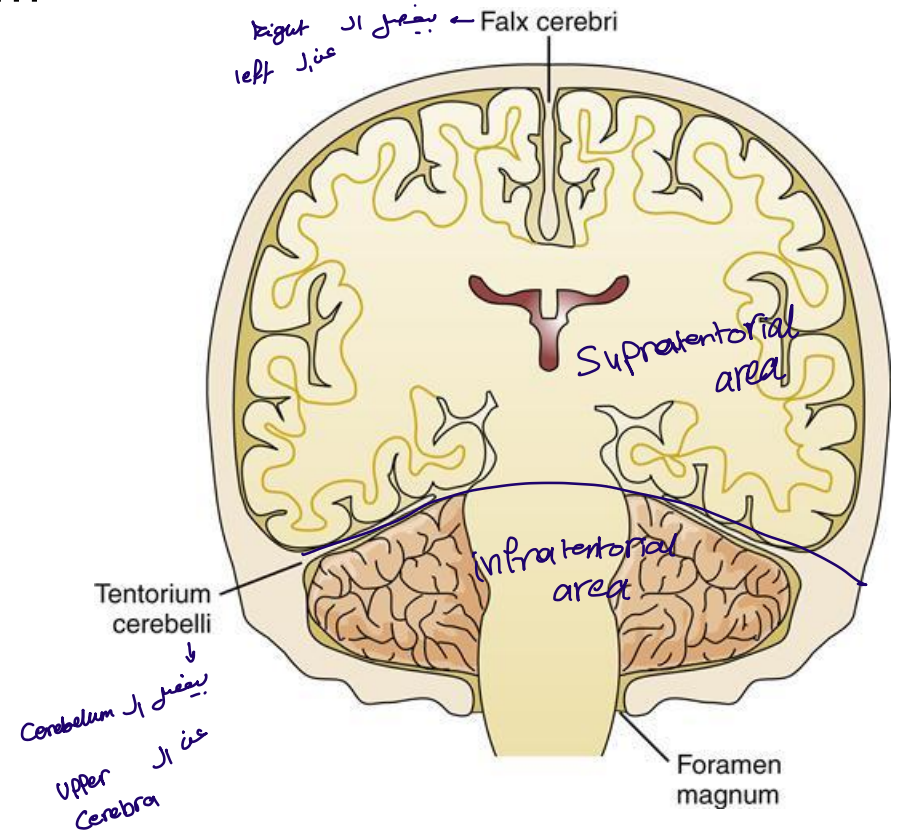
CT and MRI scans



Brain is very narrowed area so any increase in the pressure may lead to displacement of one area to another area

HERNIATION

- ❑ Herniation is the displacement of brain tissue from one compartment to another in response to increased ICP
- ❑ The intra-cranial compartment is divided by rigid dural folds (falx and tentorium).
- ❑ If the pressure is sufficiently high, portions of the brain are displaced across these rigid structures. This herniation often leads to compromise of the blood supply to compressed tissue, producing infarction, swelling, and further herniation.



THERE ARE THREE MAIN TYPES OF HERNIATION

من 5-6 انواع
Herniation
سه اصنایع تاخذ
3 =

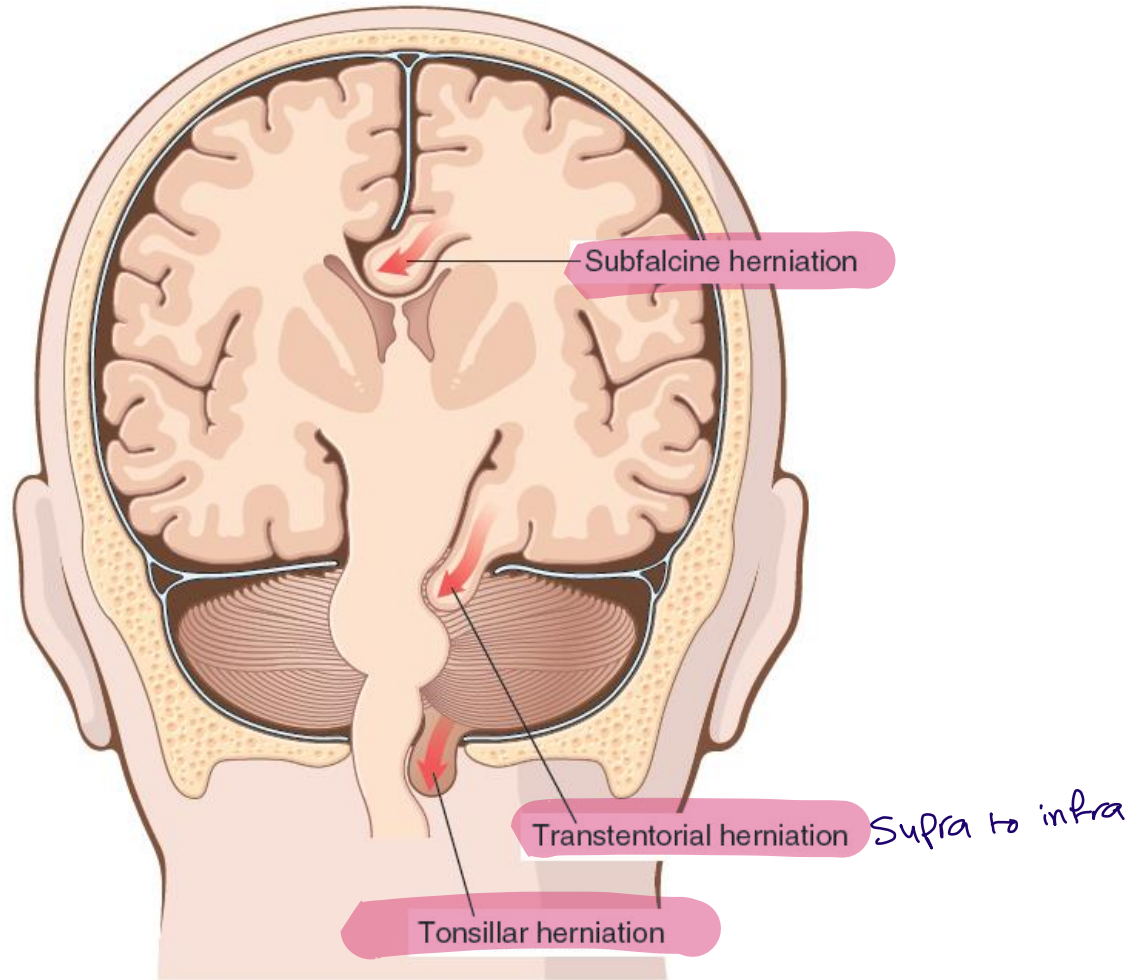


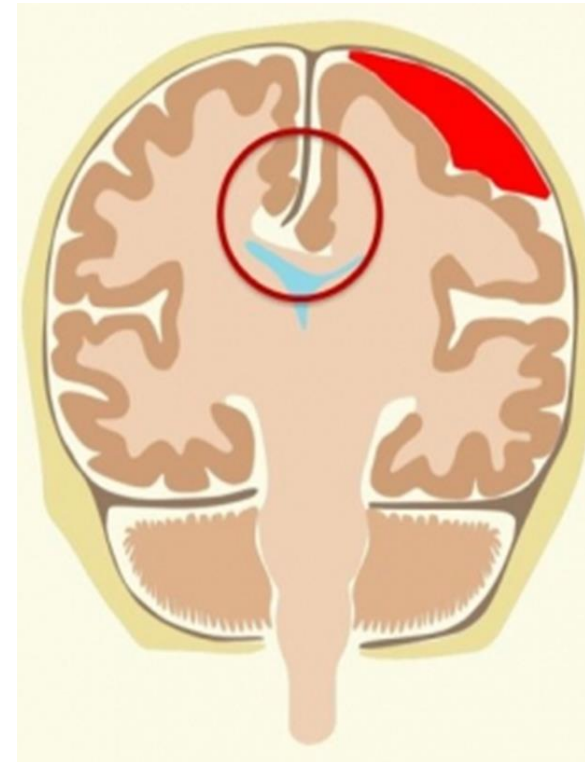
Fig. 23.4 Herniation syndromes. Displacement of brain parenchyma across fixed barriers can be subfalcine, transtentorial, or tonsillar (into the foramen magnum).



1. SUBFALCINE (CINGULATE) HERNIATION

← الاسم الثاني لها

- Herniation of Cingulate gyrus under falx cerebri into the subfalcine space
- Compression of branches of Anterior Cerebral Artery causes cerebral infarction



2- *trans*

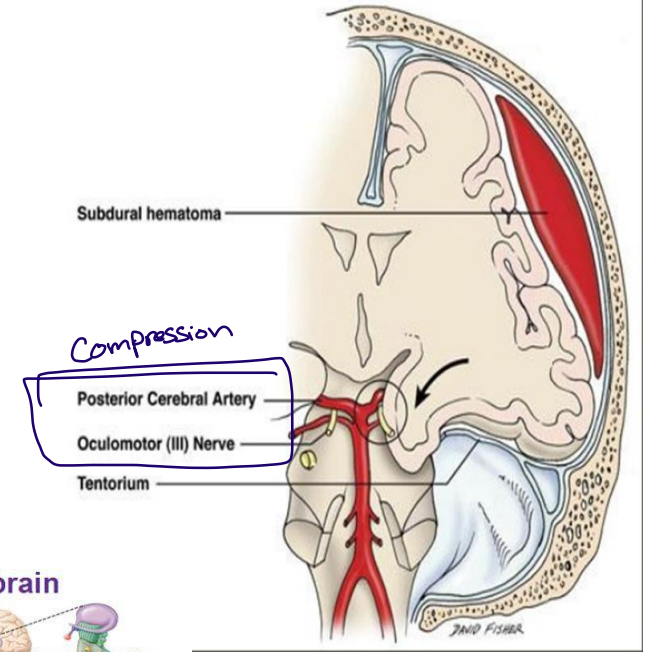
2- TRANSTENTORIAL (UNCINATE) HERNIATION

الأسوأ من

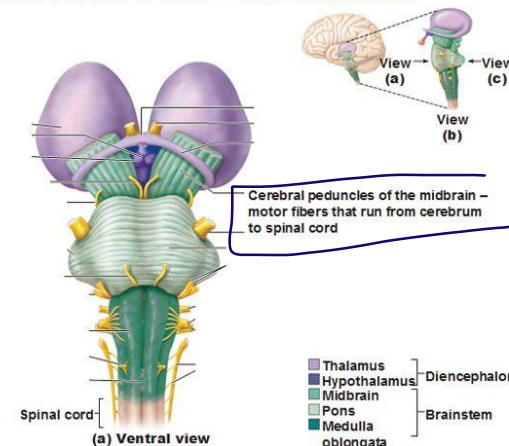
- Herniation of medial temporal lobe through tentorium.
- 3rd Cranial Nerve compression lead to ipsilateral dilated pupil & impaired eye movement
- Pressure on Posterior Carotid Artery lead to Occipital infarction, including visual cortex
- With further displacement of the temporal lobe, pressure on the midbrain may compress the contralateral cerebral peduncle against the tentorium, resulting in hemiparesis ipsilateral to the side of the herniation.

منها بفوق المكان

منها لفوق

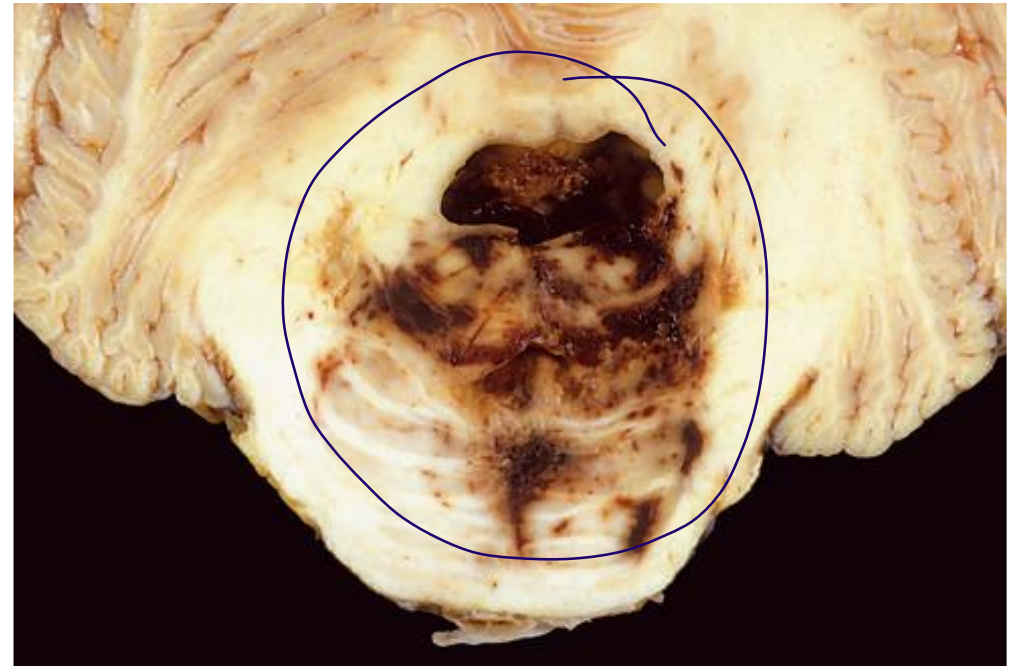


The Brain Stem- The Midbrain



2- TRANSTENTORIAL (UNCINATE) HERNIATION

- Progression of transtentorial herniation is often accompanied by linear or flame-shaped hemorrhages in the midbrain and pons, termed **Duret hemorrhages**

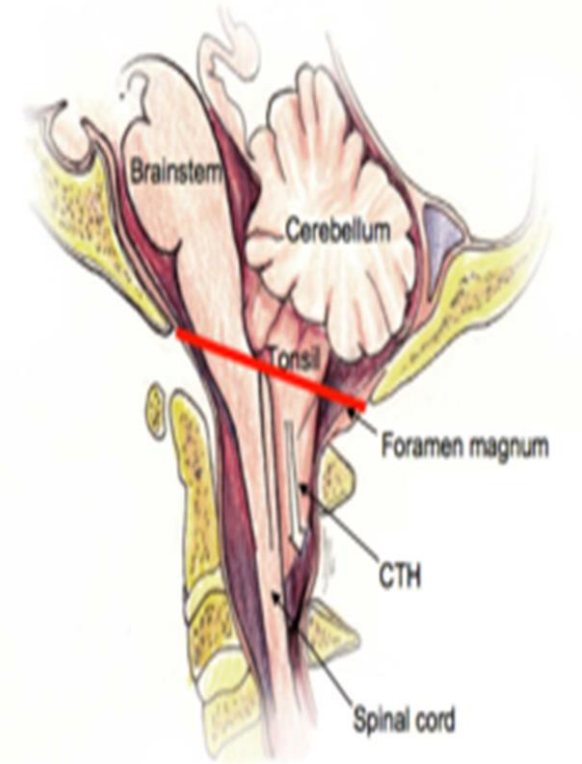


Duret hemorrhage. As mass effect displaces the brain downward, there is disruption of the vessels that enter the pons along the midline, leading to hemorrhage.



3- TONSILLAR HERNIATION → very Fatal

- *Tonsillar herniation* refers to displacement of the cerebellar tonsils through the foramen magnum.
- This type of herniation causes brain stem compression and compromises vital respiratory and cardiac centers in the medulla, and is often fatal.



DEVELOPMENTAL ANOMALIES

Two types → $\frac{1}{2}$ ← hydrocephalus

① Chiari

② Dandy-walker

Neural tube defects :

▪ Anencephaly: absence of skull and brain

← العرجيد اليه
legal to do abortion

- There is no cure or standard treatment for anencephaly and the prognosis for patients is death.
- Most anencephalic fetuses do not survive birth, accounting for 55% of non-aborted cases. Infants that are not stillborn will usually die within a few hours or days after birth



DEVELOPMENTAL ANOMALIES

Neural tube defects :

▪ Spina bifida :

- Failure of posterior vertebral arch to close

(A) ▪ Spina bifida occulta (asymptomatic) →

(B) ▪ Spina bifida - cystic protrusion of underlying tissue :

1. Meningocele - meninges protrude
2. Meningomyelocele - meninges and spinal cord protrude

