

Tumors of the Central Nervous System-1



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what is the difference between grading & staging?

Staging: TNM { Tumor size
Nodal involvement
Mets.
grading: differentiation

Central nervous system tumors



Central nervous system tumors According to the 2016



MoH Cancer Incidence Report CNS tumors are the 10th most common cancer in Jordan But the 2nd most common cancer among Jordanian children (20% of all pediatric tumors)

leukemia is the first common type among children



CNS tumors include intracranial and intraspinal tumors

Unique features of the Nervous system tumors



Metastasis: More common than primary brain tumor, which rarely metastasize



Precursors: No morphologically evident premalignant or in situ stages like carcinomas.



In Children: ~70% of CNS tumors arise in posterior fossa



Anatomical location: Influence outcome independent of histologic classification due to local effects.



Pattern: Tumors with low grade histologic features can infiltrate widely → serious clinical deficits In adults



~70% of CNS tumors arise above the tentorium.

WHO Classification of Tumors of the CNS



WHO Classification of Tumors series are authoritative reference books for the histological & molecular classification of tumors.



Classification of tumors of the CNS has been based on the concept of “histogenesis” & Grading on the basis of histologic criteria to predict tumors behavior.



2000 & 2007 classifications considered histological features and genetic changes that underlie the tumorigenesis. (Genetics were supplementary information)



The 2016 CNS WHO presents major restructuring of the diffuse gliomas, medulloblastomas & other embryonal tumors, defining tumors by both histology & molecular features

2016 WHO Classification of Tumors of the CNS



*The most common tumor in CNS → mets

*The most common primary tumor in CNS → glioma

هذا هو مبدأ التصنيف .. * ألقم أنتي من اللي فوق

Main histologic category (cell of origin)



01

Gliomas: long been classified as astrocytomas, oligodendrogliomas, and ependymomas

02

Neuronal tumors: composed of cells with neuronal characteristics

03

Embryonal (Primitive) neoplasms: have “small round cell” appearance reminiscent of normal progenitor cells in the developing CNS.

04

Others: Lymphoma, meningioma, germ cell tumors, metastasis

Gliomas

* تومور مغز
tumor

- The most common primary tumor of the brain
- They arise from a progenitor cell(not mature) that differentiates down one of the cellular lineages.
- Many subtypes typically occur in certain anatomic regions, with characteristic age distribution & clinical course.

Gliomas

Grades

(1) → localized

(≥2) → infiltrative

(4) → necrosis, microvascular proliferation

Highlights of WHO 2021 classification is the incorporation of molecular features, specifically IDH gene mutations and deletion on chromosomes segments 1p/19q

• Simplified classification of adult type diffuse gliomas into 3 groups:

- Astrocytoma, ^{→ good prognosis} IDH mutant, WHO grade 2 – 4
- Glioblastoma, ^{→ bad prognosis} IDH wildtype, WHO grade 4
- Oligodendroglioma, IDH mutant and 1p / 19q co-deleted, WHO grade 2 - 3 .

Localized astrocytomas; of which the most common are the pilocytic astrocytomas

Isocitrate Dehydrogenase Mutations / IDH1 and IDH2



Where?

Astrocytoma, glioblastoma and oligodendrogliomas



Function

lead to increased production of 2-hydroxyglutarate → interferes with the activity of several enzymes that regulate gene expression



Importance

In diagnosis & prognosis (significantly better prognosis than tumor)



Testing

Immunohistochemistry for IDH1
DNA sequencing for IDH1 and IDH2

cellularity
Atypia
mitotic figure

Astrocytoma, IDH mutant, WHO grade

2 - 4 (infiltrative)

← كيف عرفنا؟
من خلال الهيئة المناعية

- Arise from astrocytes
- Most frequent in 30s-50s.
- Usually found in the cerebral hemispheres.
- Signs & symptoms: seizures, headaches, & focal neurologic deficits related to the anatomic site of involvement.

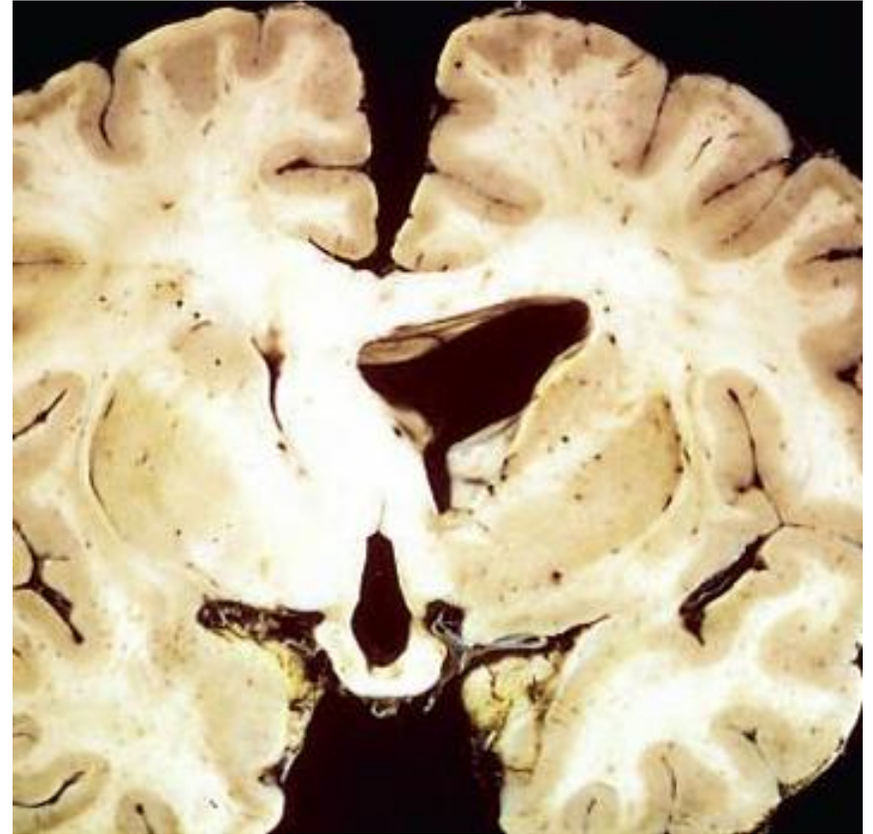
→ Not characteristic
في الدماغ

Astrocytoma, IDH mutant, WHO grade 2 - 4

- On the basis of histologic and molecular features, they are stratified into three groups (WHO grade).
 - No WHO grade 1 for infiltrating astrocytomas.
 - These grade correlates well with the clinical course & outcome (prognosis).
 - Pathogenesis: - As the name indicate, driver mutations in isocitrate dehydrogenase (IDH) gene 1 or, less frequently IDH2. - Inactivating mutation in p53 and ATRX genes
-

WHO grade II and III

- Poorly defined, gray, infiltrative (beyond grossly evident margins) tumors that expand & distort the invaded brain without forming a discrete mass.

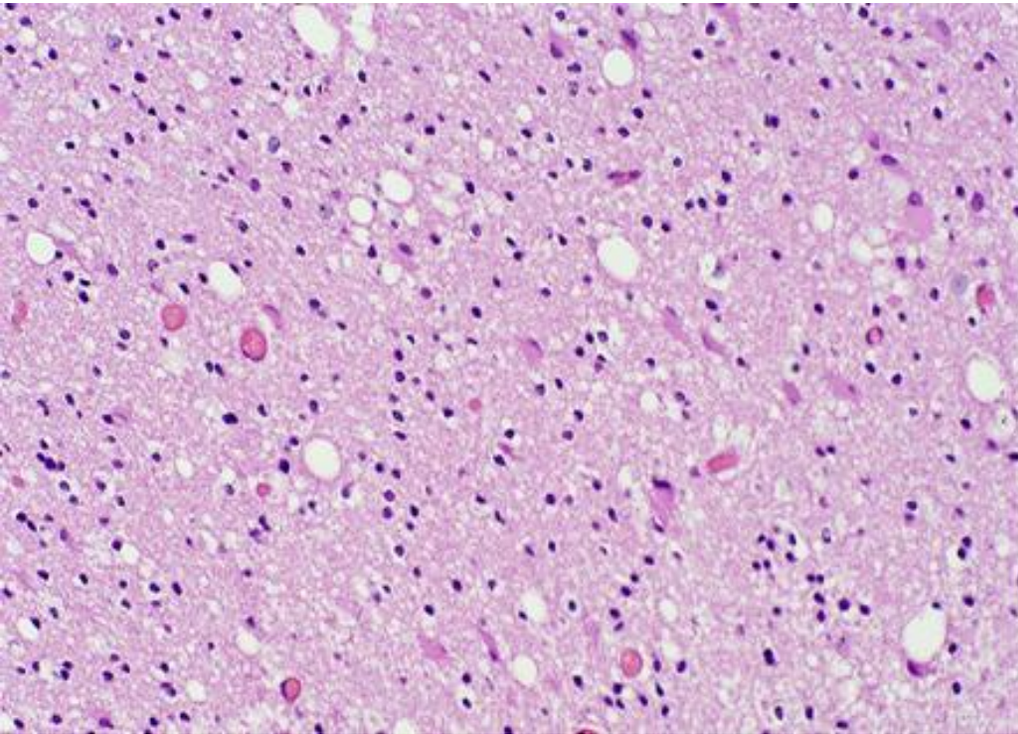


↑↑ cellularity

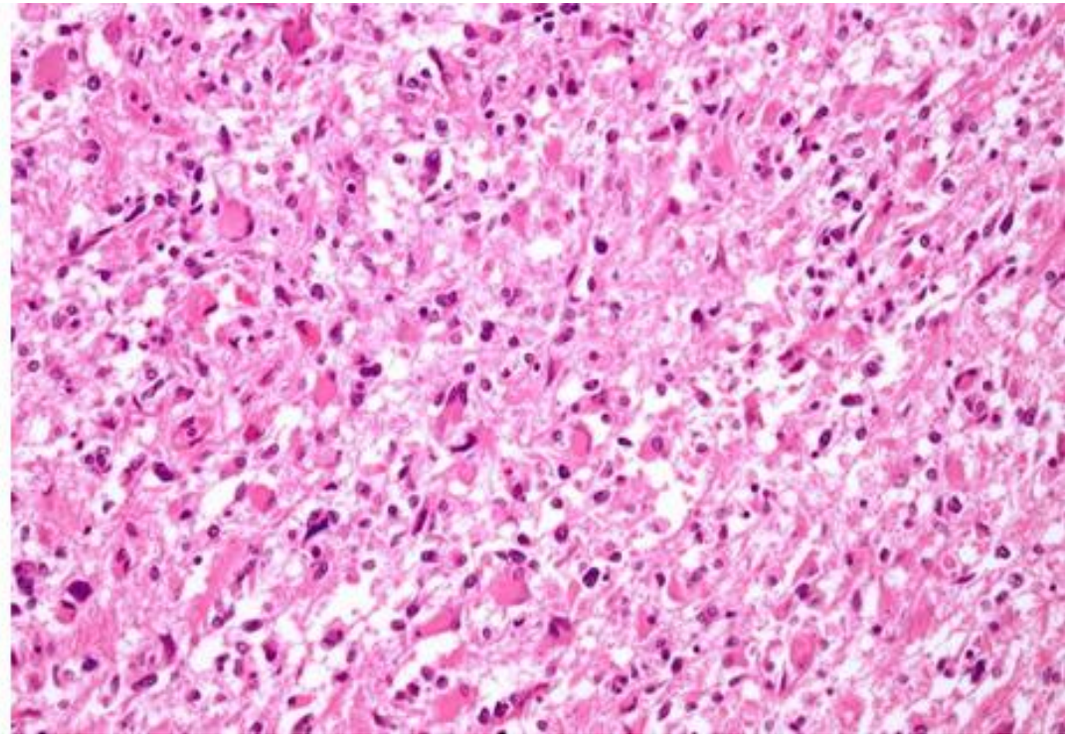
Astrocytoma (WHO grade 2) - microscopic

- low-grade (WHO grade II) astrocytomas are characterized by a mild to moderate increase in the number of glial cell nuclei, some what variable nuclear pleomorphism.
- Notice how the margin is not distinct between the tumor & the normal adjacent brain at the right





Grade 2



Grade 3, cellularity

Astrocytoma WHO 2 vs WHO 3 –Microscopic: grade 3 shows more densely cellular & have greater nuclear pleomorphism; more mitotic figures.

2 + 3 → No necrosis, No Microvascular proliferation
↳ only seen in grade 4

IDH-mutant astrocytoma - Clinical

- can be static for several years.
 - The mean survival is more than 5 years.
 - Clinical deterioration invariably occurs and is usually due to the emergence of a more rapidly growing tumor of higher histologic grade.
 - Median overall survival
 - > 10 years, grade 2
 - -5-10 years, grade 3
 - 3 years, grade 4
-

Glioblastoma IDH-wild-type (WHO grade 4)

**aggressive*

**Grade 4*

- The most common malignant glioma, 50% of adult gliomas.
 - They arise originally grade 4 (previously: primary glioblastoma).
 - Previously called glioblastoma multiforme (GBM)
 - Very poor prognosis.
-

Glioblastoma IDH-wild-type - Pathogenesis

Most imp: - common location
- age
- prognosis

Harbor multiple genetic alterations → acquisition of cancer hallmarks

- a) Evasion of senescence (telomerase mutations or mutations that lengthen of telomeres)
 - b) Escape normal growth controls (biallelic deletion of CDKN2A, which encodes the cyclin-dependent kinase inhibitor p16)
 - c) Activation of growth factor signaling pathways (EGFR or PDGFR gene amplification).
 - d) Resistance to apoptosis (TP53 mutation).
-

Glioblastoma IDH-wild-type - Clinical

- Affects older patients in their 6th to 8th decades of life.
- Sites: cerebral hemispheres (temporal, parietal, and frontal lobes; basal ganglia and thalamus).
- Develop rapidly, most patients presenting with seizures, neurocognitive impairments, nausea, vomiting, & occasionally severe pulsating headache.
- Butterfly glioma: Rapid infiltration of the corpus callosum with subsequent growth in the contralateral hemisphere → a bilateral symmetrical lesion
- Prognosis is very poor; even with treatment (resection, radiotherapy, and chemotherapy), the median survival is only about 15 to 18 months)

non specific

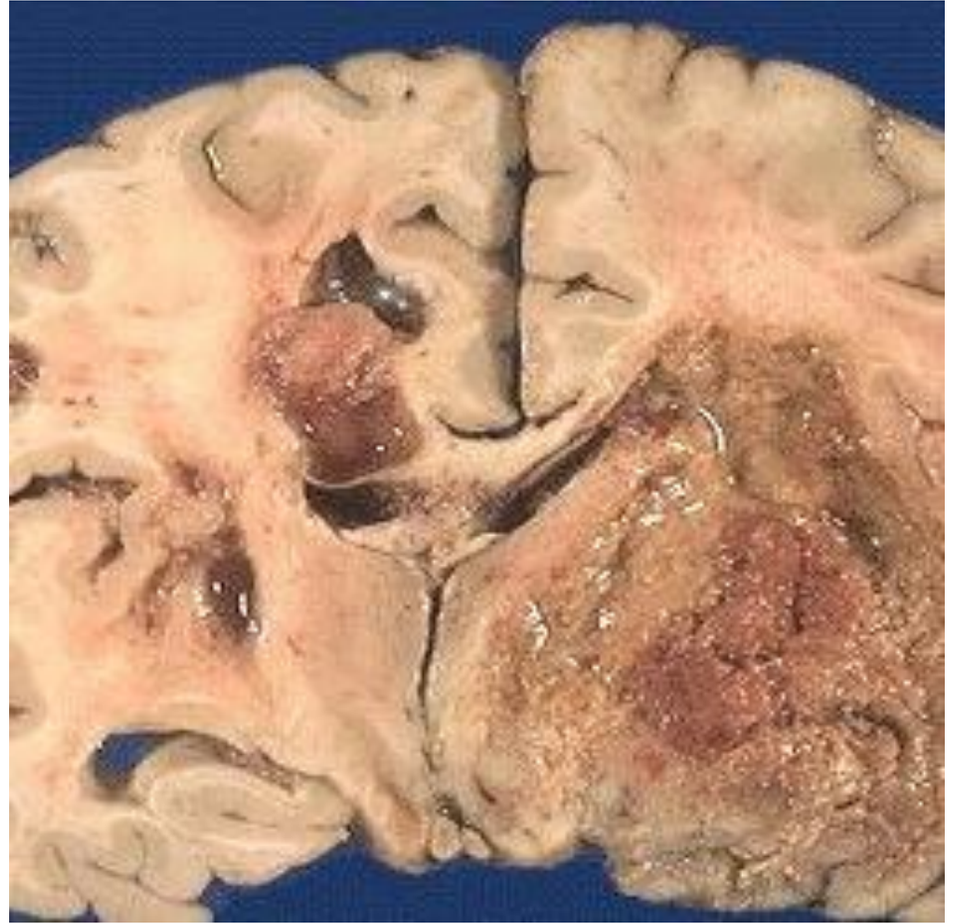
Glioblastoma (WHO grade 4) - MRI

Imaging studies most often reveal a ring enhancing lesion, abnormal vessels that are “leaky,” + abnormally permeable blood-brain barrier (BBB) → contrast enhancement on imaging studies.



Glioblastoma (WHO grade 4) - Gross

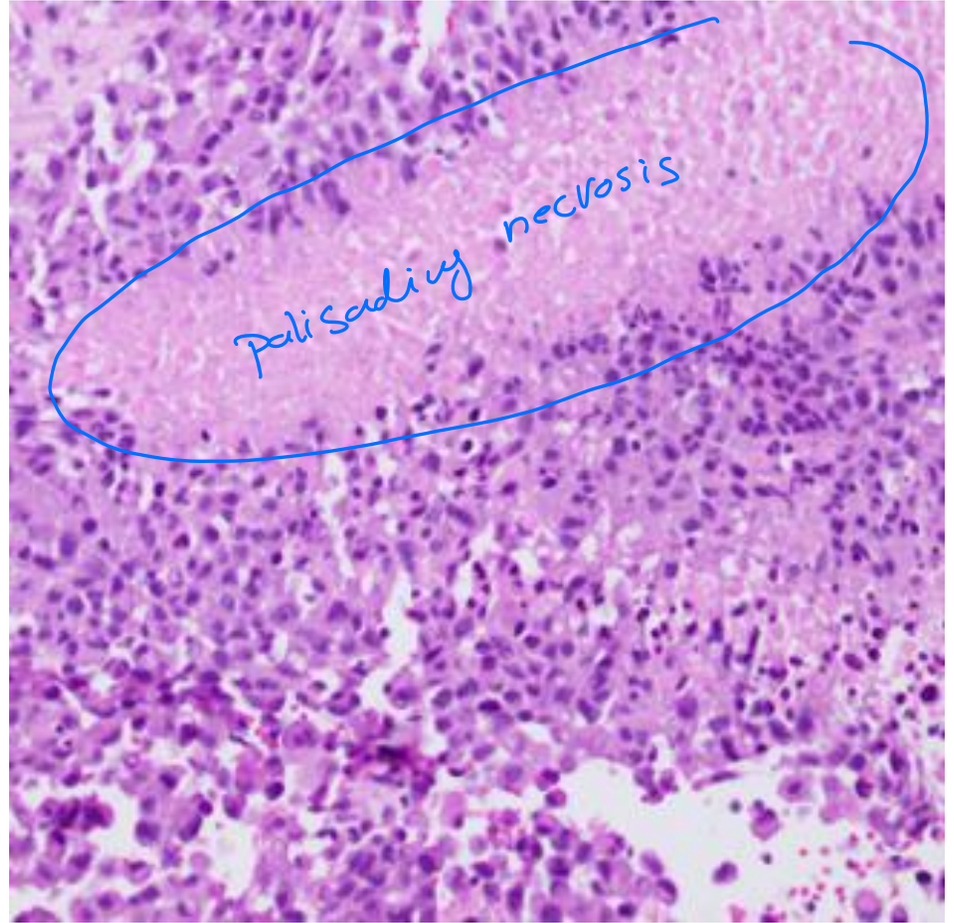
- Characteristic variation from adjacent normal area.
- Soft & yellow (tissue necrosis), regions of cystic degeneration & hemorrhage.



infiltrative

Glioblastoma (WHO grade 4) - Microscopic

- Histologic appearance varies widely (hence: multiforme).
- Cellular features similar to that of astrocytoma – grade 4 , either:
 1. as well as Necrosis (commonly present as wavy bands of necrosis with palisaded tumor cells along the border)
 2. or Microvascular proliferation



Glioblastoma (WHO grade IV) - Microscopic

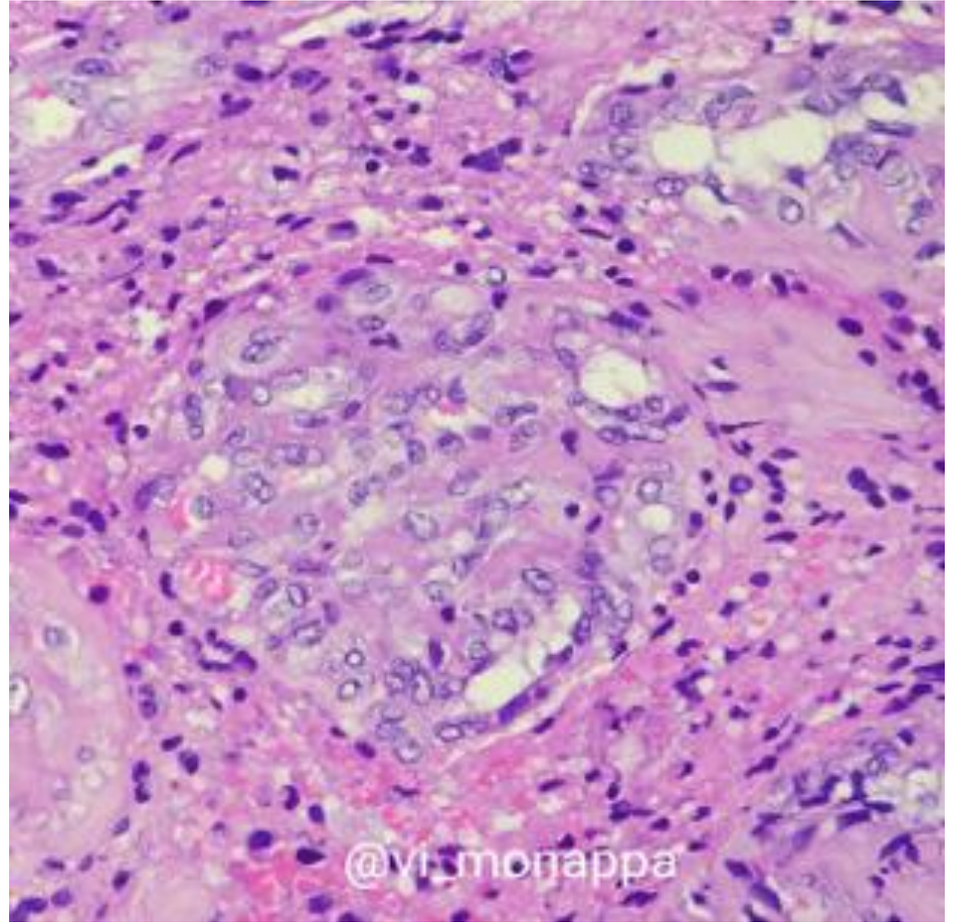
صارت تشبه ال glomeruli تبعث ال kidney من ال proliferation

2. or Microvascular proliferation

(forming tufts that bulge into the lumen
→ ball-like structures “ glomeuloid”
bodies)

* A 60-year-old man has a ring-enhancing brain lesion on MRI.
The tumor is IDH -ve ,What is the expected prognosis?
Glioblastoma

If IDH +ve wild-type the prognosis will be
Astrocytome grade 4



Oligodendroglioma

Age

30s-40s

5% to 15% of gliomas

01

Location

In the cerebral hemispheres, mainly in **white matter** in frontal or temporal lobes.

02

Genetics

- IDH1/IDH2 mutations
- 1p and 19q co-deletions

03

Gross

Infiltrative tumors, form gelatinous, gray masses & may show **cysts, focal hemorrhage, & calcification.**

04

WHO grading (2 & 3)

grade 3 is a more aggressive, higher cellularity, nuclear anaplasia, more mitoses, & microvascular proliferation.

05

لأنه ما عندي Grade 4 بالذات (Microvascular proliferation)
Grade (2,3) -

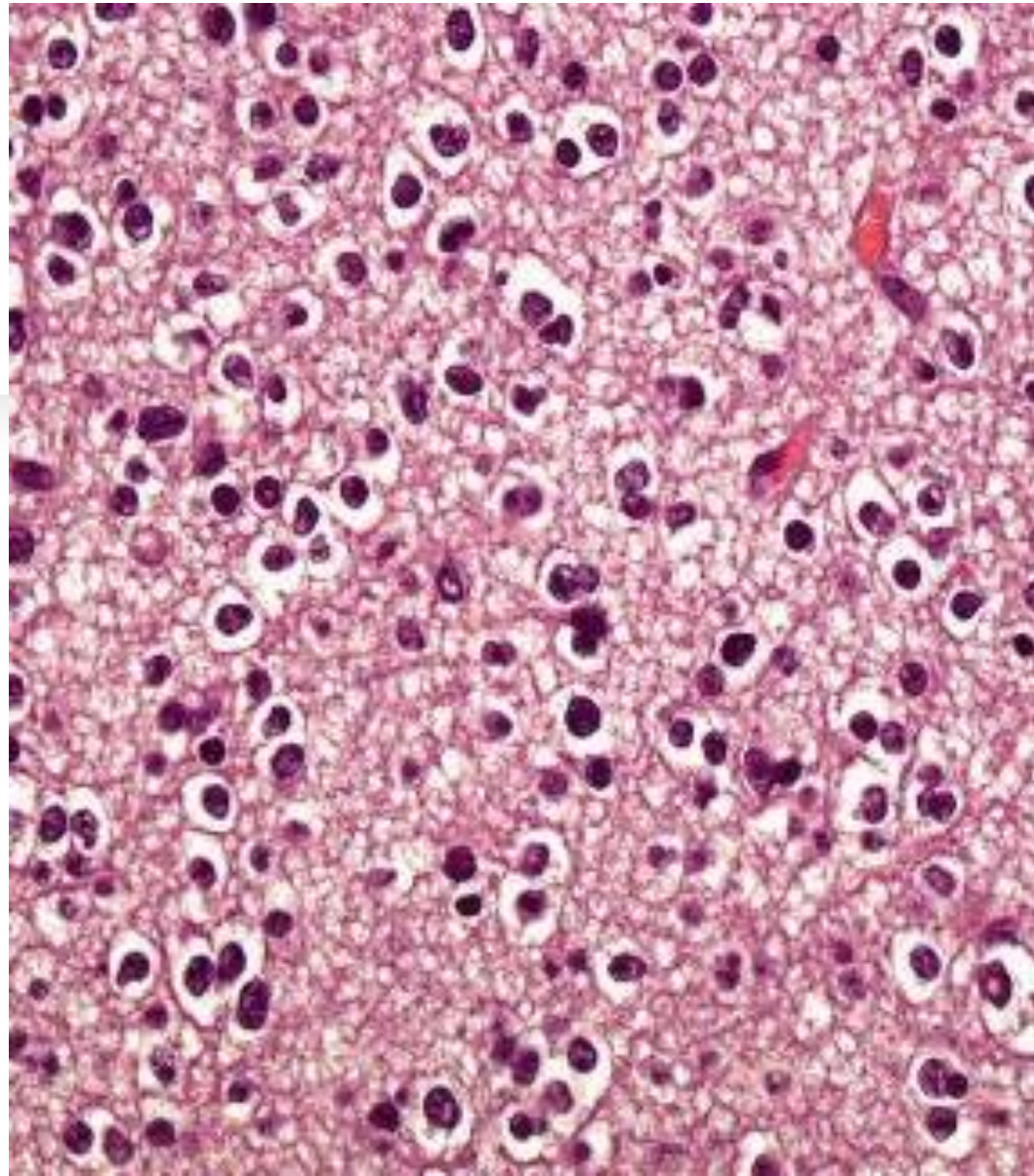
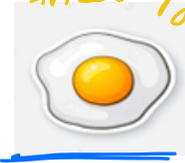
Prognosis

+ Best prognosis among diffuse gliomas.
+ surgery, chemo, & radio
average survival of 5 to 10 years

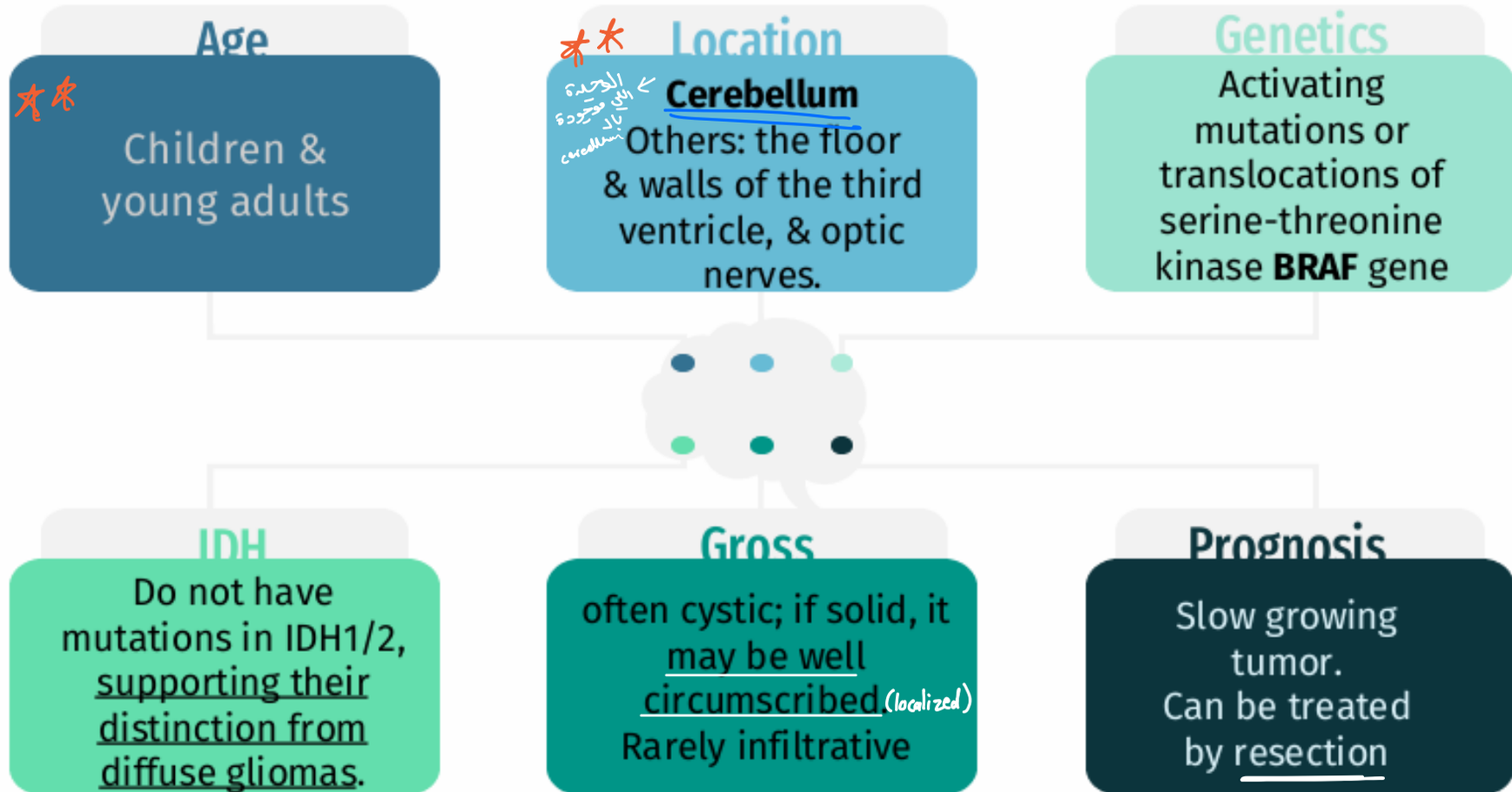
06

Oligodendroglioma- microscopic

- Sheets of regular cells with spherical nuclei containing finely granular chromatin (similar to normal oligodendrocytes) surrounded by a clear halo of vacuolated cytoplasm *fried egg*
“fried egg”
- Contains a delicate network of anastomosing capillaries.
- Calcification, in 90% of these tumors



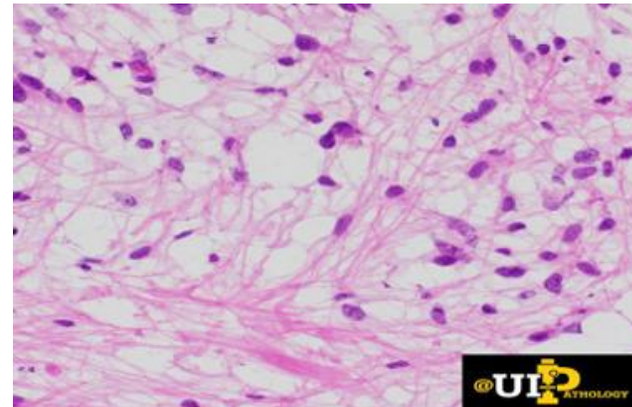
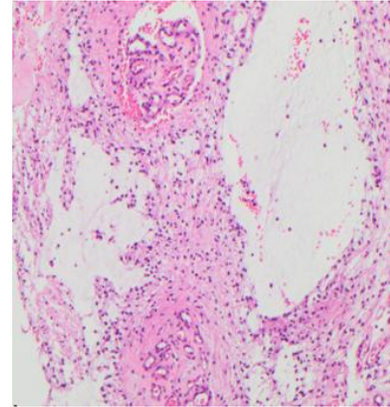
Localized astrocytoma- Pilocytic Astrocytoma (WHO grade I)



Pilocytic Astrocytoma (WHO grade 1) -microscopic

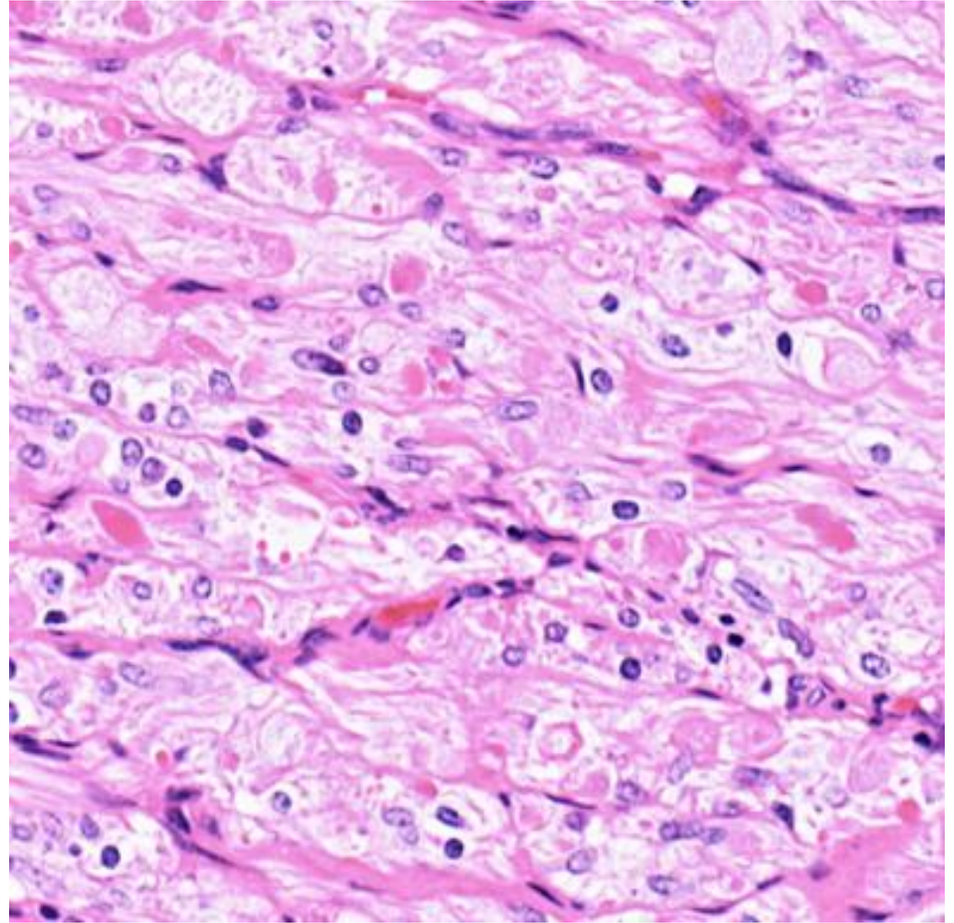
The tumor is composed of:

- Bipolar cells with long, thin “hair-like” (pilo) processes.
- Rosenthal fibers.
- Eosinophilic granular bodies.
- Microcysts often present.
- Necrosis & mitoses are rare.



Pilocytic Astrocytoma (WHO grade I) -microscopic

- Rosenthal fibers are thick, elongated, brightly eosinophilic, irregular structures that occur within astrocytic processes
- Rosenthal fibers are typically found in regions of longstanding gliosis and some brain tumors.



Ependymoma (WHO grade 2,3)

- The clinical outcome for completely resected supratentorial and spinal ependymomas is better than for those in the posterior fossa.
 - most often arise next to the ependymal lined → ventricular system.
 - first 2 decades of life, typically occur near the fourth ventricle.
 - In adults, the spinal cord is their most common location.
 - 5% to 10% of the primary brain tumors in 1st two decades.
 - spinal cord site is particularly frequent in the setting of neurofibromatosis type 2.
-

Ependymoma-Gross

+ localized
* non-Communicating ←

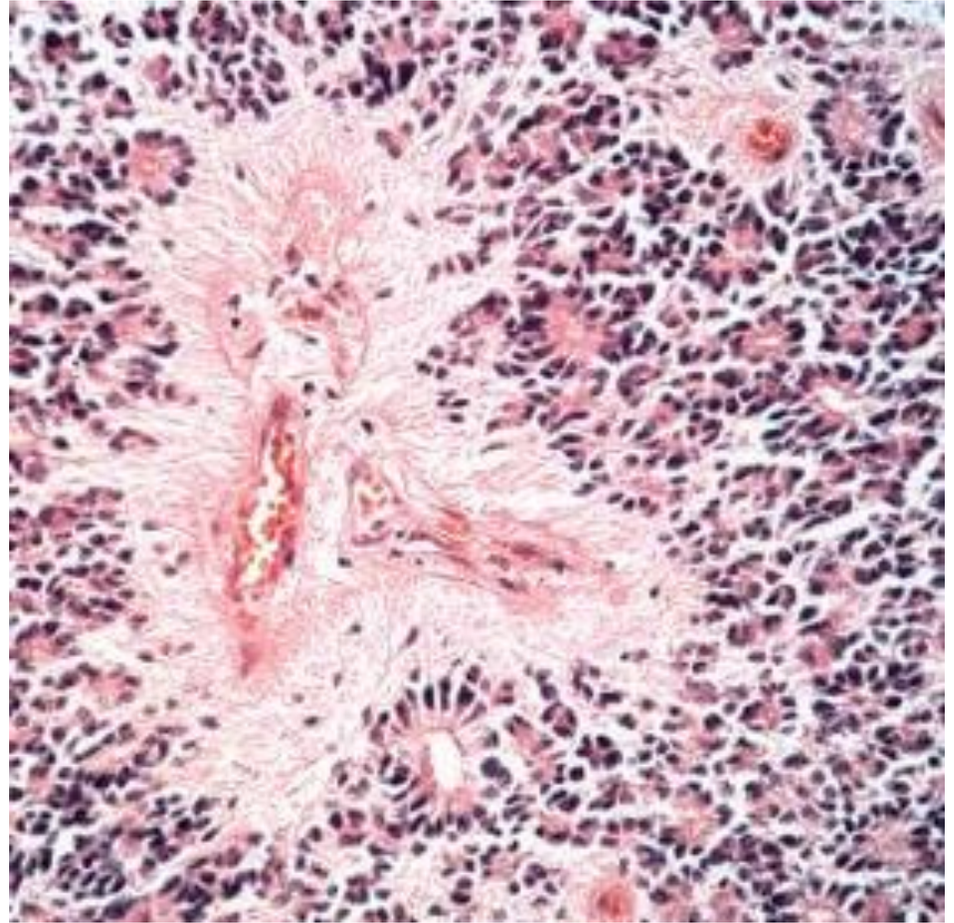
- In the fourth ventricle, ependymomas typically are solid or papillary masses extending from the ventricular floor.



Ependymoma- microscopic

→ No Grade 4

- Cells with round to oval nuclei & abundant granular chromatin.
- Dense fibrillary background.
- Cells may form round or elongated structures (rosettes, canals).
- Or more frequently present are perivascular pseudo-rosettes in which tumor cells are arranged around vessels.
- Anaplastic ependymomas (WHO grade 3) : cellularity, mitosis, & necrosis.



THANK YOU

