CENTRAL NERVOUS SYSTEM TUMORS(1)

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CNS TUMORS:

may arise from the cells of the coverings (meningiomas), the brain cells (gliomas, neuronal tumors), or other CNS cell populations (primary CNS lymphoma, germ cell tumors), or they may originate elsewhere in the body (metastases).

• Can involve the brain or spinal cord



EPIDEMIOLOGY:

• INCIDENCE:

- The annual incidence of CNS tumors in the U.S \rightarrow
 - 24 /100,000 for intracranial tumors , 1/3 malignant
 - 1-2/100,000 for intraspinal tumors

• Metastases are more common than primary brain tumors.



Characteristic features of CNS tumors:

Premalignant stage: NO premalignant or in situ lesions.

> Metastasis is rare!

- Even the most highly malignant gliomas <u>rarely spread</u> outside of the CNS.
- but the brain is **not comparably protected** against the spread of distant tumors.

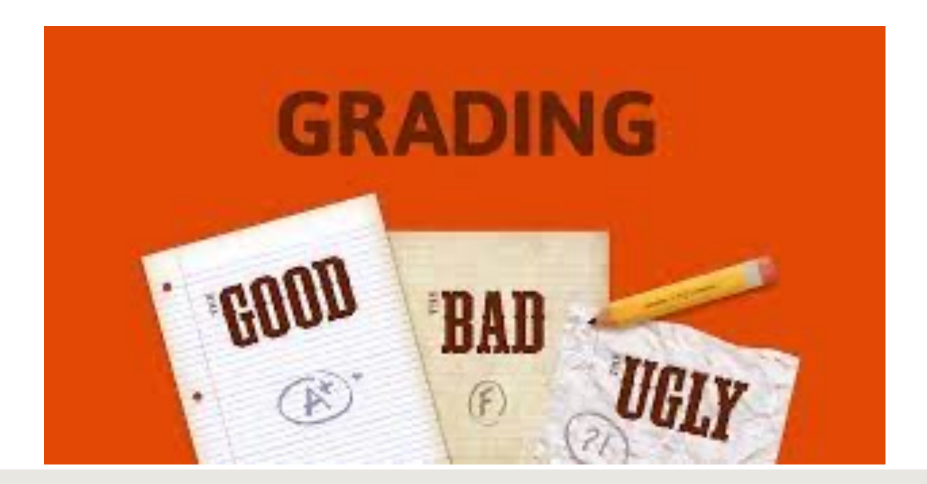
Characteristic features of CNS tumors:

- Growth pattern (infiltrative or not) and tumor location strongly influence the prognosis:
 - Even low-grade lesions may infiltrate large regions of the brain, leading to serious clinical deficits, inability to be resected, and poor prognosis.
 - The anatomic site of the neoplasm can influence outcome independent of tumor type or grade due to local effects

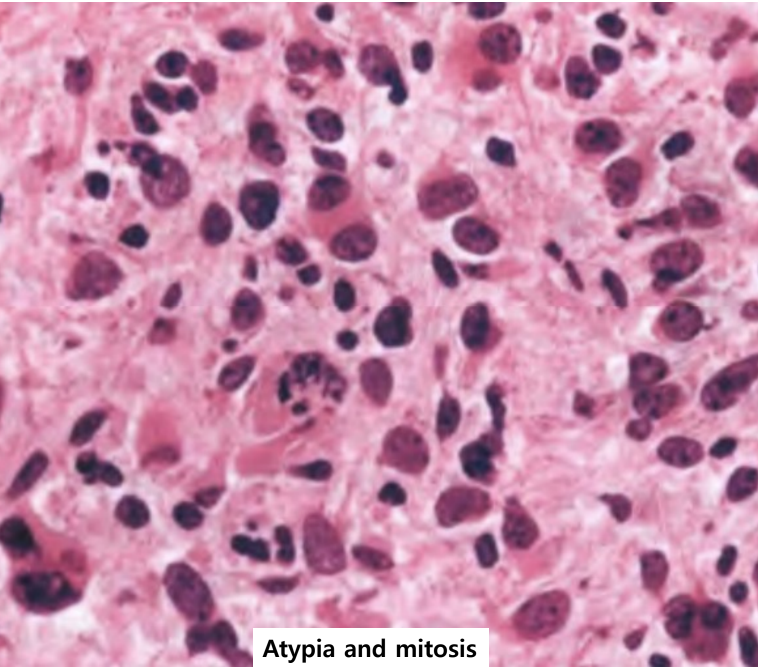
LOCATION LOCATION LOCATION LOCATION



Histologic grading of CNS tumors



The histologic grading of CNS tumors depends on:



Microvascular proliferation

necrosis 🚪

Grade 1 lesions:

- low proliferative activity
- Can be cured after surgical resection alone.

Example: pilocytic astrocytoma (PA), subependymal giant cell astrocytoma (SEGA), choroid plexus papilloma, myxopapillary ependymoma

Grade 2 lesions:

- low proliferative activity
- usually infiltrative and often recur
- Some grade II entities tend to progress to higher grades of malignancy.

Examples: astrocytoma, IDH- mutant, grade 2, oligodendroglioma, IDH- mutant and 1p/19q-codeleted, grade 2

• grade 3 lesions:

- clear histological evidence of malignancy(nuclear atypia and Higher proliferative activity (mitosis)).
- In most settings, patients receive radiation and/or chemotherapy.

Examples: astrocytoma, IDH- mutant, grade 3, oligodendroglioma, IDH- mutant and 1p/19q-codeleted, grade 3.

• grade 4 lesions:

- cytologically malignant, mitotically active, rapid proliferation, necrosisprone neoplasms
- associated with rapid pre- and postoperative disease evolution and fatal outcome.
- Widespread infiltration of surrounding tissue and a risk of craniospinal dissemination.

examples: Glioblastoma, IDH-wildtype, medulloblastoma, pineoblastoma, and most embryonal neoplasms

WHO grades of select CNS tumours		Desmoplastic infantile astrocytoma and ganglioglioma	
Diffuse astrocytic and oligodendroglial tumours		Papillary glioneuronal tumour Rosette-forming glioneuronal tumour	
Diffuse astrocytoma, IDH-mutant	Ш	Central neurocytoma	i
Anaplastic astrocytoma, IDH-mutant	ü	Extraventricular neurocytoma	Ш
Glioblastoma, IDH-wildtype	IV	Cerebellar liponeurocytoma	11
Glioblastoma, IDH-mutant	IV	Tumours of the pineal region	1
Diffuse midline glioma, H3 K27M-mutant	IV	Pineocytoma	II or III
Oligoden droglioma, IDH-mutant and 1p/19q-codeleted		Pineal parenchymal tumour of intermediate differentiation	
Anaplastic oligodendroglioma, IDH-mutant and		Pineoblastoma	IV
1p/19q-codeleted	III	Papillary tumour of the pineal region	II or III
Other astrocytic tumours		Embryonal tumours	
Pilocytic astrocytoma	1	Medulloblastoma (all subtypes)	IV
Subependymal giant cell astrocytoma	1	Embryonal tumour with multilayered rosettes, C19MC-	IV
Pleomorphic xanthoastrocytoma		altered	
Anaplastic pleomorphic xanthoastrocytoma	III	Medulloepithelioma	IV
Ependymal tumours		CNS embryonal tumour, NOS	IV
Subependymoma	L	Atypical teratoid/rhabdoid tumour	IV
Myxopapillary ependymoma	1	CNS embryonal tumour with rhabdoid features	IV
Ependymoma	Ш	Tumours of the cranial and paraspinal nerves	
Ependymoma, RELA fusion-positive	ll or III	Schwannoma Neurofibroma	
Anaplastic ependymoma	III	Perineurioma	
Other gliomas		Malignant peripheral nerve sheath tumour (MPNST) I	I, III or IV
Angiocentric glioma	1		
Chordoid glioma of third ventricle		Meningiomas Meningioma	
Choroid plexus tumours		Atypical meningioma	
Choroid plexus papilloma	1		
Atypical choroid plexus papilloma		Anaplastic (malignant) meningioma	
Choroid plexus carcinoma	III	Mesenchymal, non-meningothelial tumours	
Neuronal and mixed neuronal-glial tumours		Solitary fibrous tumour / haemangiopericytoma	I, II or III
Dysembryoplastic neuroepithelial tumour	1	Haemangioblastoma	
Gangliocytoma	1	Tumours of the sellar region	
Ganglioglioma	1	Craniopharyngioma Granular cell tumour	
Anaplastic ganglioglioma	III	Pituicytoma	
Dysplastic gangliocytoma of cerebellum (Lhermitte-Duclos)	1	Spindle cell oncocytoma	

Pediatric CNS tumors:

- 20% of all pediatric tumors.
- Childhood CNS tumors differ from those in adults in:

>Location:

> 2/3 infratentorial in kids (posterior fossa)

> 2/3 supratentorial in adults (cerebral hemispheres above tentorium)

> Mutation profile & histologic subtype:

- Kids: medulloblastoma, pilocytic astrocytoma, ependymoma
- Adults: glioblastoma, metastases, meningiomas, diffuse gliomas constitute most gliomas in adults(including astrocytomas and oligodendrogliomas).



CLASSIFICATION OF CENTRAL NERVOUS SYSTEM TUMORS



• For nearly a century, the classification of brain tumors has been done according to their **microscopic similarities** with what's thought to be their cell of origin (based on the light microscopic appearance, the immunohistochemical expression of proteins, and the electron microscopic assessment of ultrastructural features).

 The 2016 classification breaks with this nearly century-old tradition and <u>incorporates well-established</u> <u>molecular parameters</u> into the <u>classification.</u> • the classification includes diagnostic categories that depend on genotype.

 The 2016 WHO classification implemented the <u>combined phenotypic-genotypic diagnostics based on histologic features & tumor genetic</u> profile (integrated diagnoses)

The 2016 classification helped improving treatment protocols and predicting prognosis.

genetic alterations in gliomas:

- 1- Mutations in isocitrate dehydrogenase (IDH) genes:
- observed as an <u>early</u> event in gliomagenesis
- Seen in astrocytomas and oligodendrogliomas
- Gain of function Mutation affection IDH1 codon 132 or IDH2 codon 172.
- The most frequent is IDH1 R132H mutation (83-91%) of IDH mutant gliomas
- IDH2 mutation: R172K is the most frequent IDH2 mutation

- ✓ Can be detected by <u>immunohistochemical stains and molecular studies</u>:
 - IDH1-R132H immune stain
 - IDH sequencing for IDH1 codon 132 and IDH2 codon 172

 ✓ Gain of function mutation → lead to increased production of 2hydroxyglutarate (oncometabolite) → interferes with the activity of several enzymes that regulate gene expression → DNA hypermethylation & maintaining the cells in stem cell-like physiological states → self- renewal and tumorigenesis

2- whole arm Co-deletion of 1p and 19q chromosomal segments:

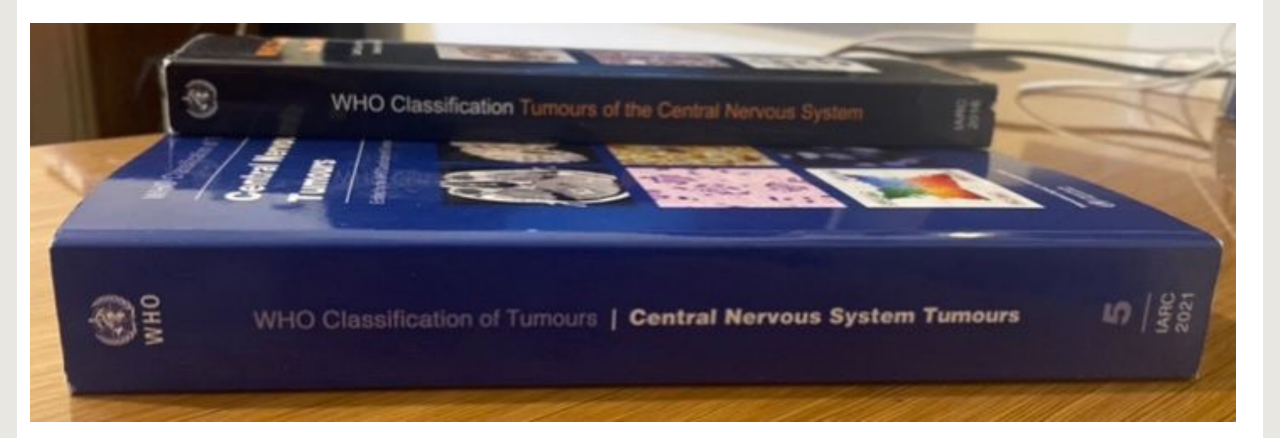
• Diagnostic of oligodendrogliomas in the presence of IDH mutation.

The vast majority of IDH mutant and 1p/19q co-deleted oligodendroglioma
 →carry TERT promotor hotspot mutations

• **TERT promotor hotspot mutations**: telomerase stabilization, cellular immortalization and proliferation

- 3- ATRX and P53 loss of function mutation:
- Both occur in IDH mutant astrocytomas
- **ATRX mutation** induces abnormal telomeres maintenance mechanism known as **"alternative lengthening of telomeres"**
- ATRX mutation is Mutual exclusive with the activating promoter mutation of the TERT gene (1p/19q codeletion)
- **P53 mutation:** enable tumor cell survival
 - ATRX → associated with genomic instability → induces P53 dependent cell death → mutation in P53 helps these cells to survive.





Diffuse astrocytoma, MYB or MYBL1-altered

Astrocytoma, IDH-mutant Diffuse midline glioma, H3 K27-altered Chordoid glioma

Astroblastoma, MN1-altered ZFTA

Supratentorial ependymoma, C11orf95 fusion-positive

Embryonal tumor with multilayered rosettes

Malignant melanotic nerve sheath tumor

Solitary fibrous tumor

Mesenchymal chondrosarcoma (formerly a subtype)

Adamantinomatous craniopharyngioma (formerly a subtype)

Papillary craniopharyngioma (formerly a subtype)

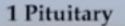
Pituicytoma, granular cell tumor of the sellar region, and spindle cell oncocytoma (grouped Pituitary adenoma / PitNET

Primary intracranial sarcoma, DICERT-mutant

Pituitary blastoma

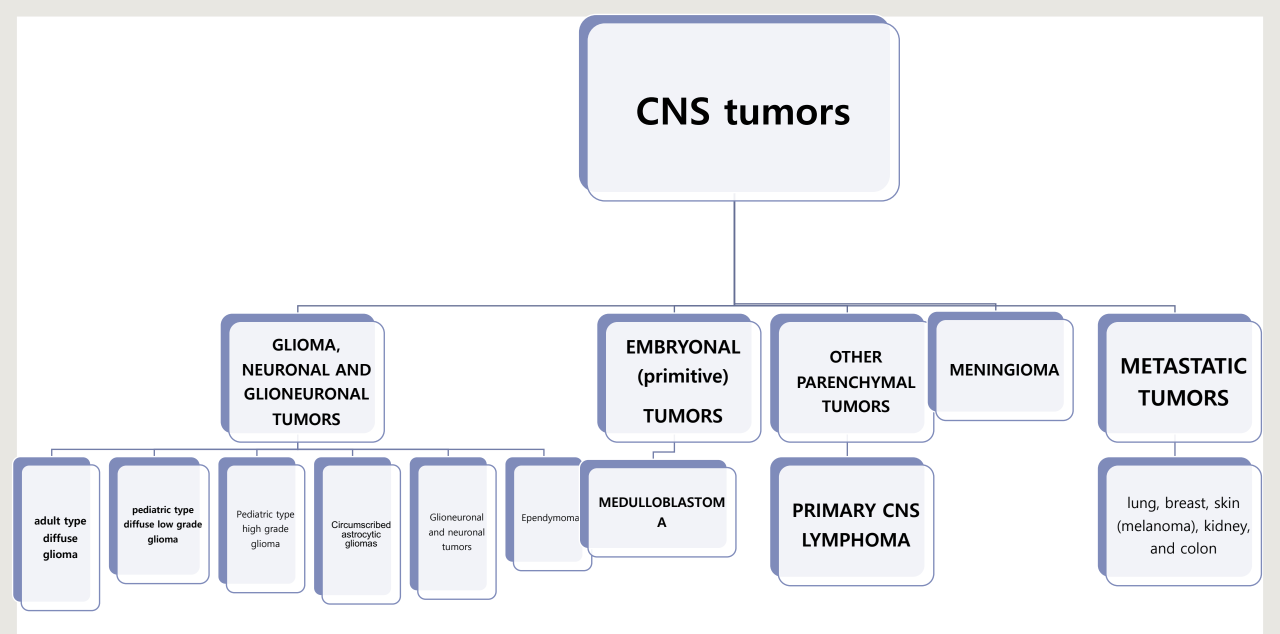
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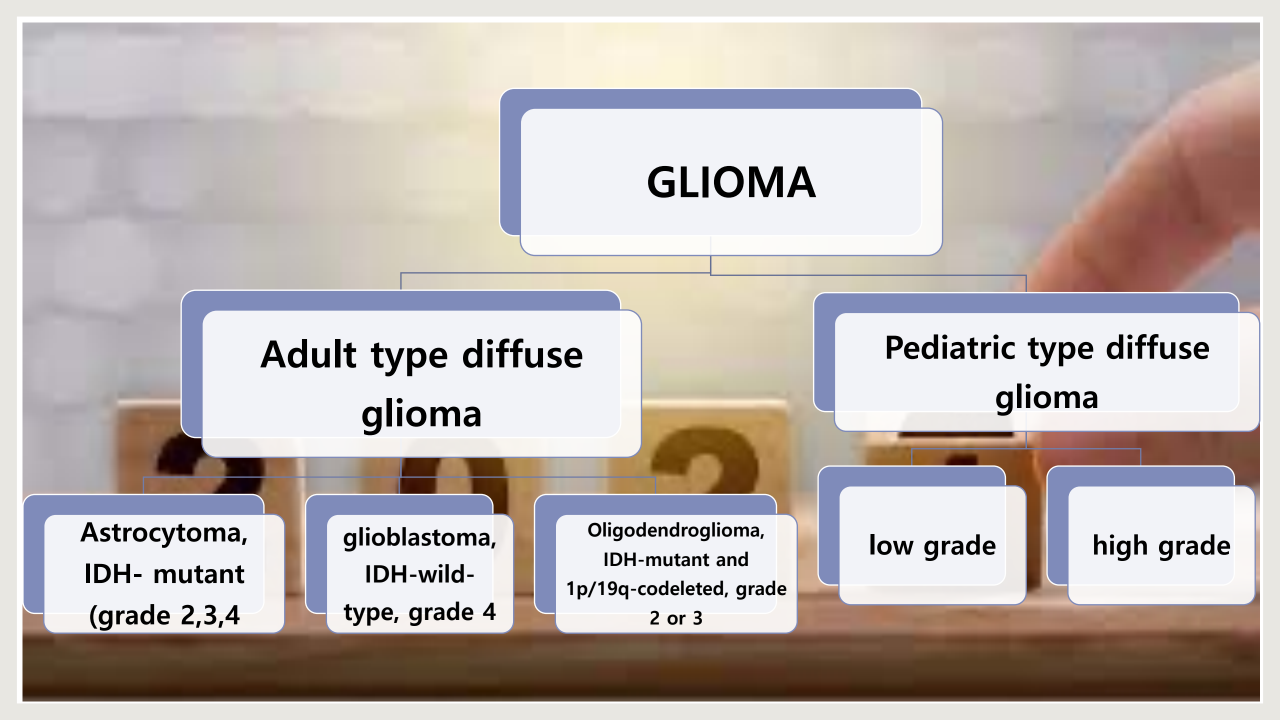
13 with Revised Terminology





Glior	nas	WHO 2016	Gliomas.	Glioneuronal and Neuronal Tumours WHO 2021
2.1:	Diffus	e astrocytic and oligodendroglial tumours	,	2.0.0.1: Introduction to gliomas, glioneuronal tumours, and neuronal tumours
		Introduction	2.1: Glion	nas, Glioneuronal and Neuronal Tumours
	2.1.2: 2.1.3: 2.1.4: 2.1.5: 2.1.6: 2.1.7: 2.1.8:	Diffuse astrocytoma, IDH-mutant 2.1.2.1: Gemistocytic astrocytoma, IDH-mutant Diffuse astrocytoma, IDH-wildtype Diffuse astrocytoma, NOS Anaplastic astrocytoma, IDH-mutant Anaplastic astrocytoma, IDH-wildtype Anaplastic astrocytoma, NOS Glioblastoma, IDH-wildtype 2.1.8.1: Giant cell glioblastoma 2.1.8.2: Gliosarcoma 2.1.8.3: Epithelioid glioblastoma		Adult-type diffuse gliomas 2.1.1.1: Astrocytoma, IDH-mutant 2.1.1.2: Oligodendroglioma, IDH-mutant and 1p/19q-codeleted 2.1.1.3: Glioblastoma, IDH-wildtype Paediatric-type diffuse low-grade_gliomas 2.1.4.1: Diffuse astrocytoma, MYB or MYBL1-altered 2.1.4.2: Angiocentric glioma 2.1.3.5: Polymorphous low-grade neuroepithelial tumour of the young 2.1.5.1: Diffuse low-grade glioma, MAPK pathway-altered Paediatric-type diffuse_high_grade_gliomas 2.1.2.1: Diffuse midline glioma, H3 K27-altered 2.1.2.2: Diffuse hemispheric glioma, H3 G34-mutant
	2.1.11:	Glioblastoma, IDH-mutant Glioblastoma, NOS Diffuse midline glioma, H3 K27M mutant Oligodagdroglioma, IDH mutant and 1p/19g codelated		2.1.2.3: Diffuse paediatric-type high grade glioma, H3 wildtype and IDH wild type 2.1.3.14: Diffuse midline glieme, EGER mutant (fermerly Bithelemie glieme, EGER mutant) 2.1.2.4: Infant-type hemispheric glioma
	2.2.1: 2.2.2: 2.2.3: 2.2.4: 2.2.5: 2.2.6:	Oligodendroglioma, IDH-mutant and 1p/19q-codeleted Oligodendroglioma, NOS Anaplastic oligodendroglioma, IDH-mutant and 1p/19q-codel Anaplastic oligodendroglioma, NOS Oligoastrocytoma, NOS Anaplastic oligoastrocytoma, NOS		Circumscribed astrocytic gliomas 2.1.3.1: Pilocytic astrocytoma 2.1.3.2: High-grade astrocytoma with piloid features 2.1.3.3: Pleomorphic xanthoastrocytoma 2.2.0.4: Subependymal giant cell astrocytoma 2.2.0.1: Chordoid glioma 2.2.0.2: Astroblastoma, MN1-altered
2.3:	2.3.1: 2.3.2:	astrocytic tumours Pilocytic astrocytoma 2.3.1.1: Pilomyxoid astrocytoma Subependymal giant cell astrocytoma Pleomorphic xanthoastrocytoma	2.1.4:	Glioneuronal and neuronal tumours 2.1.3.7: Ganglioglioma 2.1.3.9: Desmoplastic infantile ganglioglioma / Desmoplastic infantile astrocytoma 2.1.3.10: Dysembryoplastic neuroepithelial tumour 2.2.0.3: Diffuse glioneuronal tumour with oligodendroglioma-like features and nuclear clusters 2.2.0.5: Papillary glioneuronal tumour





Astrocytoma, IDH- mutant

Phenotype: It Is a diffusely infiltrating glioma

Genotype:

- IDH1 or less frequently IDH2 mutation.
- Inactivating mutation in TP53 and/or ATRX
- absence of 1p/19q codeletion

• Age at diagnosis: 40–60 year old.

• Location: cerebral hemispheres +/- cerebellum, brainstem, or spinal cord.

- Presentation:
 - seizures, headaches, and focal neurologic deficits related to the anatomic site of involvement.
 - Clinically: static for years or Progressive.

The prognosis gets poorer as the grade increases

- On the basis of histologic features astrocytomas, IDH- mutant are stratified into three groups:
 - astrocytomas, IDH- mutant, grade 2, median survival is >10 years.
 - astrocytomas, IDH- mutant grade 3, median survival is 5-10 years
 - astrocytomas, IDH- mutant grade 4, median survival is 3 years.

 NO grade 1 astrocytoma, IDH- mutant, because by convention grade 1 implies benign behavior and all diffuse gliomas are considered malignant

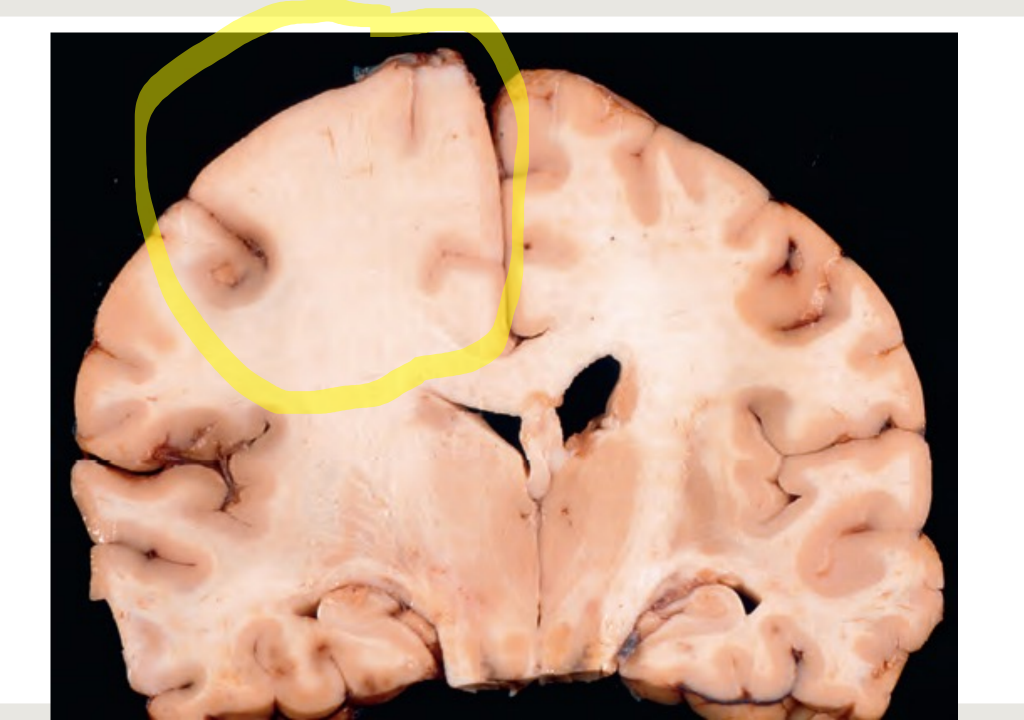
Morphology, macroscopic:

Grade 2 &3:

- poorly defined, infiltrative tumors
- expand and distort the invaded brain
- NO discrete mass, Infiltration beyond the grossly evident margins.

Grade 4:

- poorly defined, infiltrative tumors
- lacks large areas of central necrosis and hemorrhage seen in IDH-wild-type GBM



Diffuse astrocytoma, IDH- mutant, WHO grade 2, Microscopic:

- The transition between neoplastic and normal tissue is indistinct
- tumor cells infiltrate normal tissue many centimeters from the main lesion.

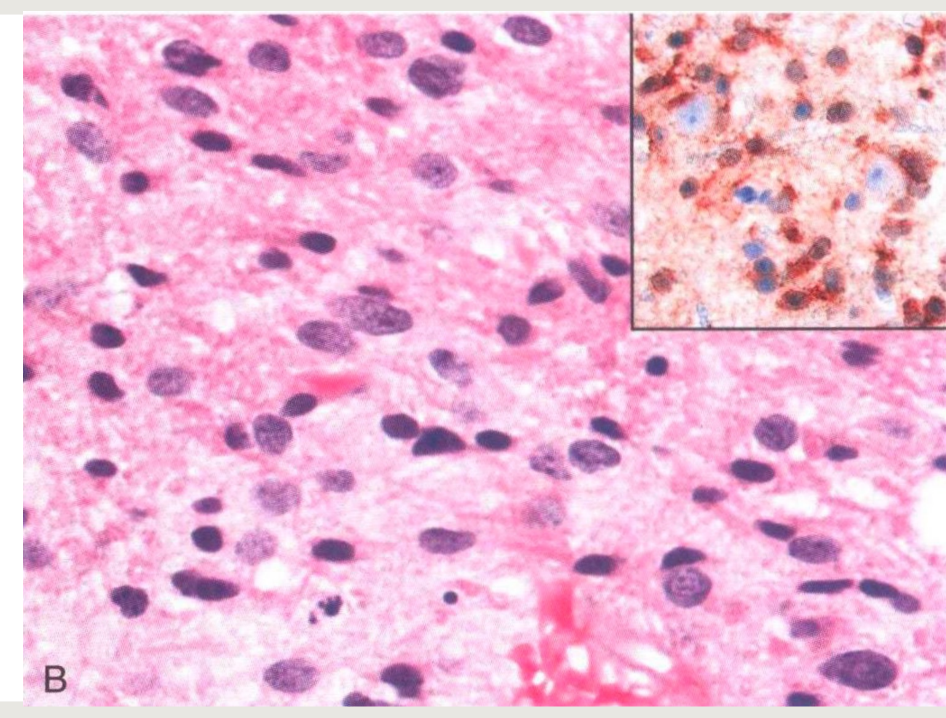
- Hypercellular (compared to normal white matter): <u>mild to moderate</u> increase in the number of glial cell nuclei.
- Cytologic atypia:
 - ≻ mild
 - enlarged, elongated or irregular hyperchromatic nuclei
 - > No prominent atypia

+ **fibrillary background** made of a network of fine astrocytic cell processes

- **<u>NO or rare</u>** Mitotic activity
- <u>NO</u> necrosis
- <u>NO</u>microvascular proliferation

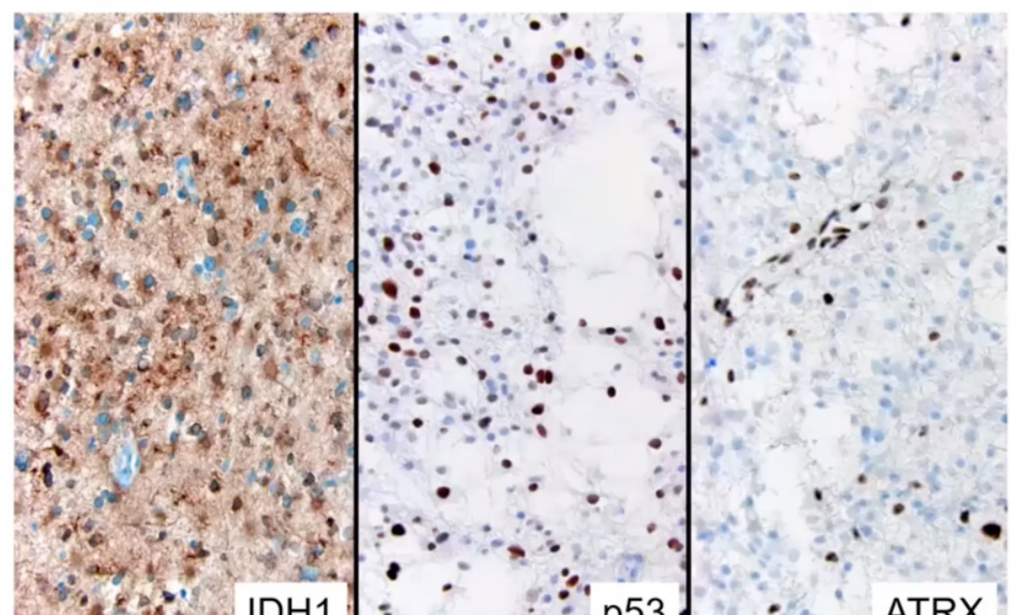
Enlarged irregular nuclei embedded within fbrillar matrix of the brain

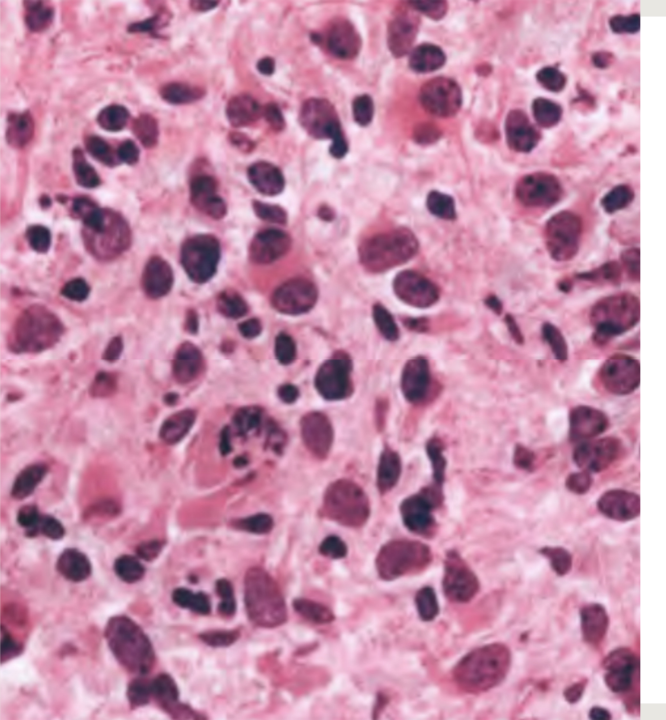
Inset: IDH1 immune stain is positive in tumor cells



GFAP :glial fibrillary acidic protein

Astrocytoma, IDH-mutant, CNS WHO grades 2-4





Astrocytoma, IDH-mutant, grade 3:

- More densely cellular
- More nuclear pleomorphism
- mitotic figures are present
- ✤ <u>NO</u> necrosis
- ✤ <u>NO</u>microvascular proliferation

Astrocytoma, IDH-mutant, grade 4:

• Same as grade 3 with <u>Microvascular proliferation and/or necrosis</u>

The presence of homozygous deletion of CDKN2A &/or CDKN2B
 → astrocytomas, IDH- mutant, grade 4 (EVEN IF THE
 HISTOLOGY SUGGESTS A LOWER GRADE).

