DIAGNOSIS AND MANAGEMENT OF MIDLINE POSTERIOR FOSSA TUMORS IN CHILDREN



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INTRODUCTION

- Medulloblastoma
- Ependymoma
- Astrocytoma
- Brainstem glioma
- Choroid plexus papilloma
- Dermoid

Medulloblastoma

- Bailey and Cushing in 1925 first used the term medulloblastoma.
- One of the most common tumors of posterior fossa (20 – 25 % all pediatric brain tumors)
- 5 –7 yrs median age of diagnosis.
- 2 4 and 6 –8 yrs: two peaks in children

Medulloblastoma

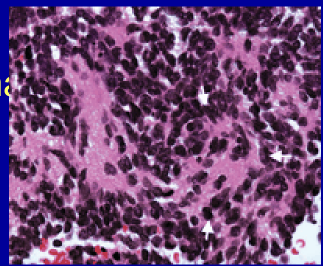
Histologic subtypes:

Classical medulloblatoma

Desmoplastic medulloblastoma

Medullomyoblastoma

Melanotic medulloblastoma



Large-cell medulloblastoma: Very poor outcome

Medulloblastoma....origin

Debatable:

- Origin from remnant of cells of the external granular layer of the cerebellum.
- Transformation of normal undifferentiated progenitor cells of superior medullary velum which migrate to the external granular layer.

- Hydrocephalus : Raised ICP
 - Behavioral change, listlessness, irritability, vomiting, and decreased social interactions.
 - Headache, especially in the morning.
 - Double vision.
 - Head tilt: tonsillar herniation below the foramen magnum.
 (Can result from trochlear nerve palsy caused by direct tumor compression)

Cerebellar symptoms

Brain stem involvement

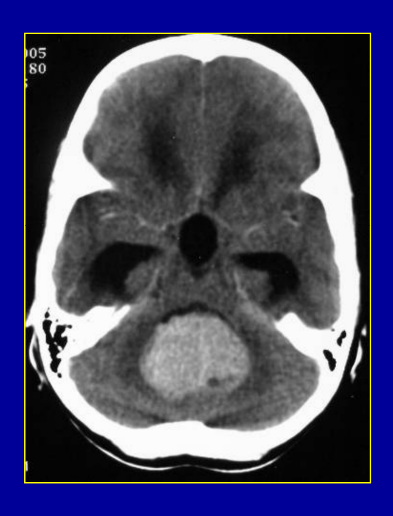
Leptomeningeal dissemination

- Physical:
 - Increasing head circumference, full anterior fontanalles with widely split cranial sutures.
- Fundus examination
 - Papilledema can be present in as many as 90% of patients.

- Extraocular examination
 - Diplopia and lateral gaze paresis
 - Fourth cranial nerve palsy (should be considered in any patient with a head tilt)
 - Nystagmus
- Cerebellar signs (ataxia > unilateral dysmetria)

Radiology.....CT





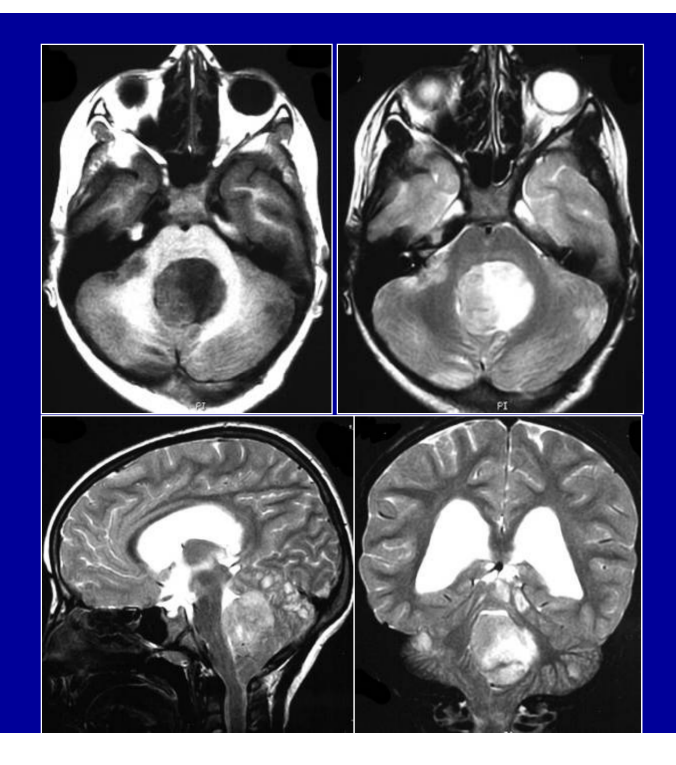
NCCT

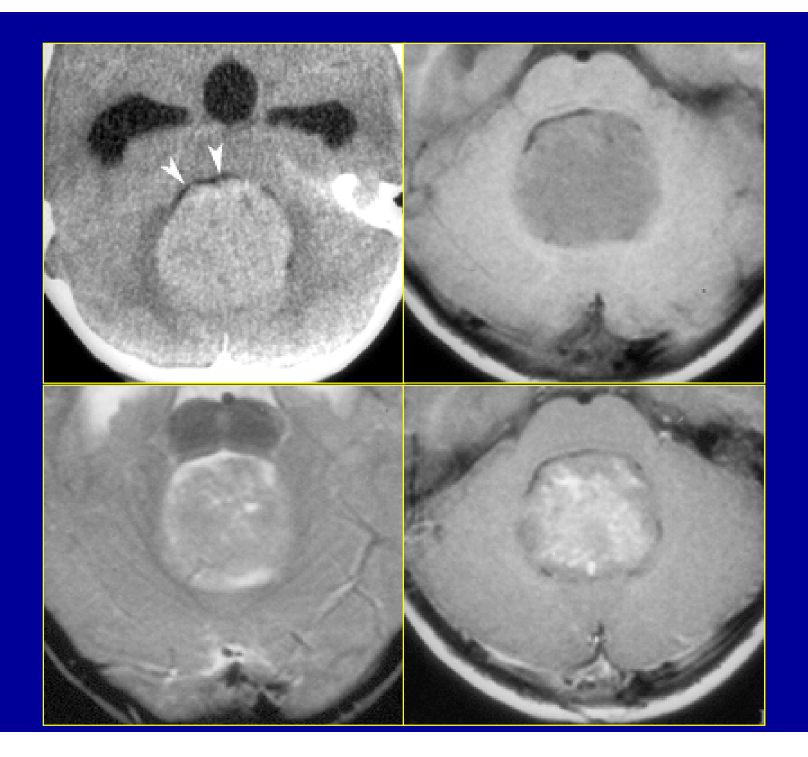
CECT

Radiology.....MRI

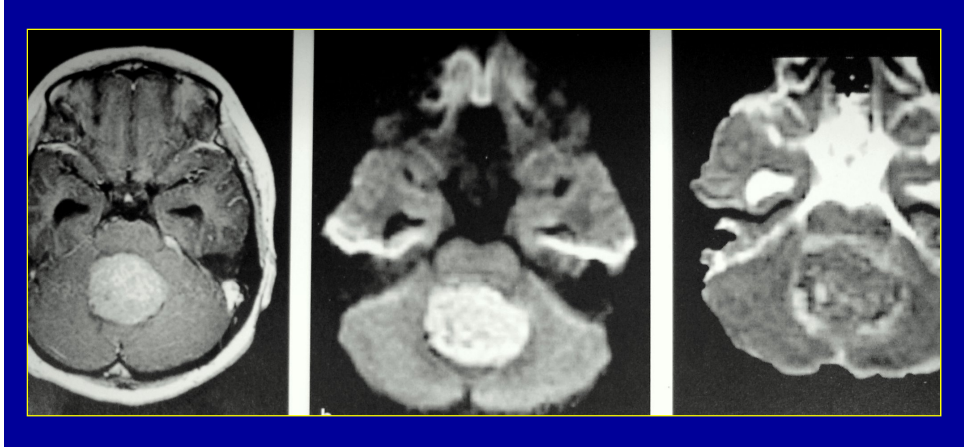
- Homogeneous enhancement (may be absent in about 15 20 %)
- DWI shows restricted diffusion with increased ADC.
- MRI spine : Should be done at time of diagnosis.
- BEST: prior to surgery. If not possible Should be delayed for at least 2 weeks after surgery.

M E L L O B A S M

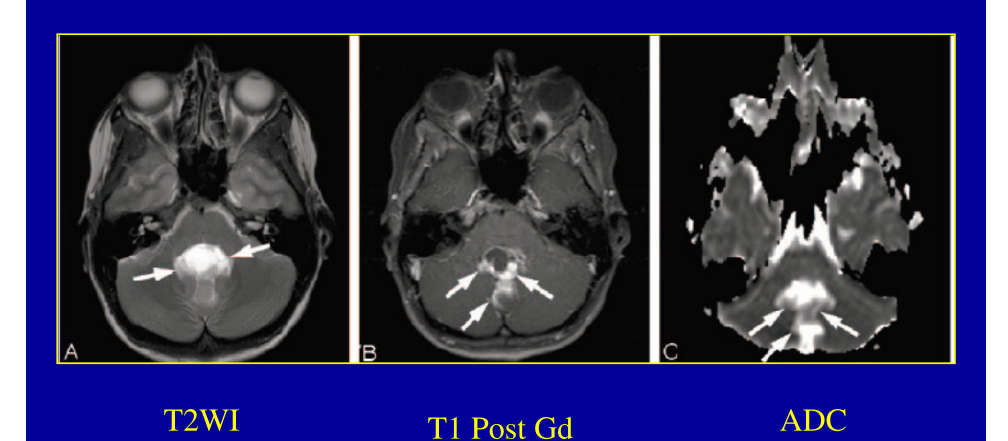


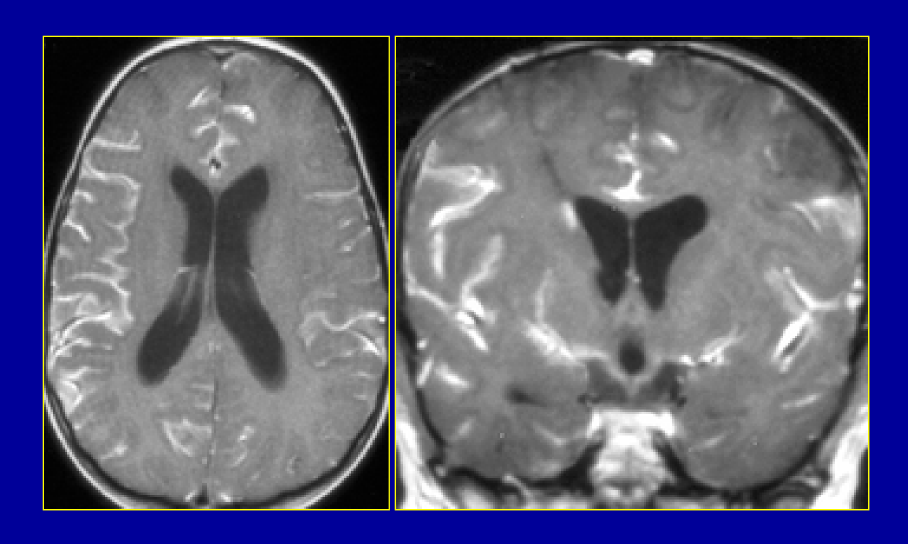


DWIMedulloblastoma



DWIEpendymoma





Leptomeningeal Dissemination

MRI SPINE



Radiology.....

Skeletal imaging

 Metastasis to the bone must be considered in any child with medulloblastoma and bone pain.

 A skeletal survey helps elucidate lytic or sclerotic lesions.

DiagnosisCSF cytology

- No standardized method: HOW and WHEN ??
 - Lumbar puncture
 - Ventricular drain
 - Cisterna magna at the time of surgery from the for cytologic analysis.

Stage	Feature		
Tumor stage			
Tl	Less than 3 cm diameter; limited to vermis, roof of fourth ventricle, or hemisphere		
T2	More than 3 cm diameter; invades one adjacent structure or partially fills fourth ventricle		
T3a	Invades two adjacent structures or completely fills fourth ventricle with extension into cere-		
	bral aqueduct, foramen of Luschka, or foramen of Magendie		
T3b	Arises from floor of fourth ventricle or brain stem; fourth ventricle completely filled		
T4	Spreads to involve cerebral aqueduct, third ventricle, midbrain, or upper cervical spinal cord		
Metastasis stage			
M0	No evidence of metastasis		
M1	Tumor cells in CSF		
M2	Gross nodular seeding of brain CSF spaces		
M3	Gross nodular seeding of spinal CSF space		
M4	Extraneural spread		

Modified Chang's Staging for medulloblastoma

Staging.....

- Within 48 hours of surgery, a Gd MRI.
 - Staging.
 - Assess residual tumor size prior to the onset of enhancing reactive gliosis.
- Staging is dependent upon :
 - · extent of resection,
 - radiographic evidence of tumor spread,
 - and CSF cytology.

Current staging of medulloblastoma

- Standard Risk
- Posterior fossa
- No metastasis
- < 1.5 cm² residual
- Undifferentiated

- High Risk
- Posterior fossa with intracranial or spinal dissemination.
- Extra neural metastasis
- > 1.5 cm² residual
- Differentiated

Diagnosis....genetics

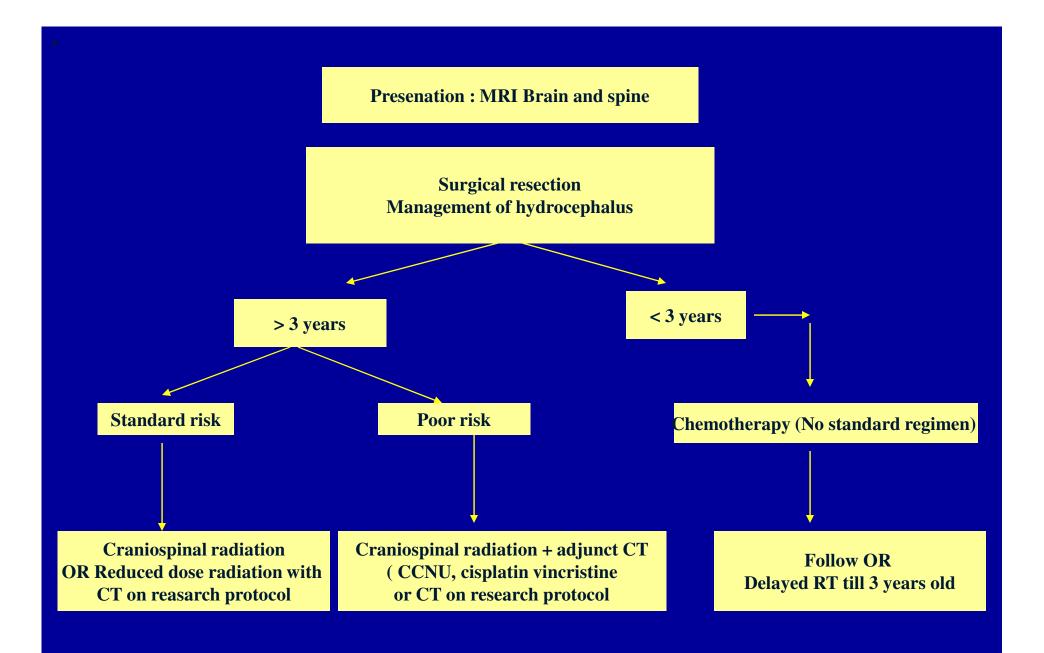
- Routine use: Controversial.
- Correlation between aneuploid DNA content and a better prognosis.
- 17qi an isochromosome : Most common
- C-ERB2 poor outcome
- Neurotropin growth factor receptor (TrkC) expression: associated with better outcome.

Risk factors associated with outcome for medulloblastoma

- Good Prognosis
- Females Sex
- Gross total resection
- No metastasis
- Desmoplastic histology
- Increased apoptosis index
- Hyperdiploidy
- High TRKC expression

Poor Prognosis

- Younger age
- Subtotal resection
- Metastasis
- Large-cell anaplastic histology
- Elevated Ki-67/MIB index
- Aneuploidy
- Elevated ERB2 expression
- Isolated 17p LOH
- Elevated expression and amplification of MYCC
- Up regulation of PDGFR
- Over expression of calbindin-D28k



Management algorithm for medulloblastoma

Hydrocephalus

 The majority of children with posterior fossa tumors have hydrocephalus at the time of presentation.

 There is no consensus regarding the management of HC in these children

Hydrocephalus

- Treatment options:
 - Ventriculoperitoneal shunt
 - Perioperative EVD
 - Endoscopic third ventriculostomy
 - Direct surgical resection

Hydrocephalus.....

 Recent studies have shown that ultimately 17 to 40% of children have uncontrolled hydrocephalus and require shunt placement during the postoperative period; and that this predominantly occurred within the 1st postoperative month.

 An expectant policy in these subgroup who ultimately require a shunt place them at risk of developing intracranial hypertension ,an increased rate of CSF leakage, and pseudomeningocele formation, prolonged hospitalization. HydrocephalusFactors predicting patients at risk of requiring placement of a shunt postoperatively

- Younger age at diagnosis
- The severity of hydrocephalus prior to resection of the tumor
- Midline localization
- Incomplete tumor removal
- Use of substitute dural grafts during closure
- CSF infection
- Persistent pseudomeningocele
- An analysis of factors determining the need for ventriculoperitoneal shunts after posterior fossa tumor surgery in children.
 - Neurosurgery 34:402-408, 1994
 - Pediatr Neurosurg 20:240-247, 1994

Management..... Surgery

- Gross Total Resection, if possible.
- Brainstem damage should be avoided.
- Resolution of natural CSF pathways.
- Tumor adheres to the floor of the fourth ventricle, precluding gross total resection.(1/3 rd of cases)
- Sugar coating subarachnoid spread.

Management...... Radiotherapy

- SURGERY alone: NOT CURATIVE
- RADIOTHERAPY: cornerstone of adjuvant therapy.
- 54 to 58 Gy to the primary site with 35Gy to the entire craniospinal axis

Management...... Radiotherapy

Complications of radiotherapy:

- lowered intelligence quotient (IQ),
- small stature, endocrine dysfunction,
- behavioral abnormalities,
- secondary neoplasms
- white matter necrosis.
- Reduction in IQ and neurobehavioral function.

Radiotherapy and chemotherapy trials

SIOP and the (German) Society of Paediatric Oncology (SIOPII) Bailey et al. Med Pediatr Oncol 25:166178, 1995	Patients with low-risk medulloblastoma were randomized to receive or not receive CT as well as randomized to reduced-or standard-dose neuraxis RT treatment groups.	Patients receiving a reduced craniospinal axis dose of 2500 cGy had a worse mean survival rate when compared with those treated with a dose of 3500 cGy (5-year event-free survival [EFS] 55.3% and 67.6 respectively; p = 0.07). In a subgroup analysis, the addition of a chemotherapy regimen produced a negative effect on survival in patients who received reduced doses of craniospinal axis radiation (p = 0.0049).
French Society of Pediatric Oncology Journal of Clinical Oncology 23,4726- 34;2005	Standard-Risk Medulloblastoma Treated by Adjuvant Chemotherapy Followed by Reduced-Dose (25 Gy) Craniospinal Radiation Therapy	The overall survival rate and 5-year recurrence-free survival rate were 73.8% ± 7.6% and 64.8% ± 8.1%, respectively
CCG multicenter randomized trial (CCG-921) Zeltzar et al. J Clin Oncol 17:832845, 1999	Compared the 8-in-1 chemotherapy regimen both before and after radiotherapy with a combination of vincristine, CCNU, and prednisone (VCP) after radiotherapy	Chemotherapy with VCP was superior to the 8-in-1 regimen in patients with medulloblastoma, with a 5-year PFS rate of 63% compared with 45%, respectively (p = 0.006).
CCG and Pediatric Oncology Group Deutsch et al. Pediatr Neurosurg 24:167177, 1996	Standard-risk patients were randomized to receive standard dose of craniospinal axis radiation (3,600 cGy in 20 fractions) or reduced dose (2,340 cGy in 13 fractions).	The study was closed before patient accrual was complete because of an increased number of recurrences in the low-dose treatment group (31% compared with 15% recurrence, respectively, at 16 months)
Randomized phase III study (CCSG-942) Evans AE et al. J Neurosurg 72:572-582, 1990	Compared standard radiotherapy with or without the addition of vincristine, CCNU, and prednisone in patients with newly diagnosed medulloblastoma	Overall, there was no significant difference in EFS rates between the radiotherapy group (52%) compared with the combined radiotherapy/chemotherapy group (57%). Despite this, in the subset of children with extensive disease (stage M_{1-3} or T_{3-4}), the 5-year EFS rate was improved in the group undergoing chemotherapy (46% compared with 0% respectively; p = 0.006).

Management..... Hyperfractionated radiotherapy

- Delivery of higher doses of radiation without increased toxicity.
- The typical hyperfractionated radiotherapy schedule consists of twice-daily fraction sizes of 100 to 120 cGy to a total dose of 7200 to 7800 cGy.

Management..... Chemotherapy

- Delay the onset of radiation therapy in young children (< 3 years)
- Increase in survival rates in high-risk children with medulloblastoma
- Patients with recurrent or advanced disease
- Reduction in the RT dose to the neuraxis in patients with nondisseminated disease

Management..... New studies

- Sensitizing the tumor to irradiation with the concomitant use of chemotherapy.
- Presurgical chemotherapy to treat patients prior to surgery.
- Intraventricular administration of cytotoxic agents,
- Newer drug combinations, and
- Immunotherapy based on genetics analysis

Management...... Recurrent Medulloblastoma

• Recurrences: 30 to 40% of patients

 Chemotherapy: limited due to chemo resistance in those patients who have previously undergone CT

Redosing with RT avoided due to radiation necrosis.
 (Local RT using stereotactic techniques can be used can palliative)

Management...... Recurrent Medulloblastoma

 High-dose chemotherapy with autologous SCR or autologous BMR: subject of intense investigation.

Stem cell rescue involves harvesting autologous bone marrow or preferably, peripheral stem cells by using pheresis techniques and subsequently reinfusing them after provision of high-dose myeloablative chemotherapy.

Int J Legal Med. 2001;114(6):331-7

Substantial toxicity:

Death, serious infection, and venoocclusive disease.

Management...... Recurrent Medulloblastoma

 Though data suggests longer EFS. (In the absence of RCT, the interpretation of the results remains limited)

Management..... Prognosis

- 5 year recurrence-free survival rates: 55% 67%.
- Even after a good response to surgery and radiation, recurrence is common.
 - Most common site: PRIMARY TUMOR SITE
- Bone: most common site of systemic metastasis; followed by regional lymph node.



Surgical resection Management of hydrocephalus

CSF - VE

CSF +VE

Cranial RT – 56Gy / 30# / 6 wks.

(36 Gy/20# followed by a boost of 20Gy /10 #)

Spinal RT – 30 Gy / 20# / 4 wks.

Concurrently with cranial RT)

Dose of spinal RT 36Gy/30#/6 weeks

Cerebellar Mutism

 Cerebellar mutism was first reported in 1979 by Hirsh after a posterior fossa tumor resection.

- Also known as posterior fossa syndrome
- Approximately 10 -1 5 % of children undergoing posterior fossa surgery for tumor.

Cerebellar Mutism

- Decreased or absent speech, irritability, hypotonia, ataxia.
- Onset: Immediate or delayed.
- Virtually all cases of mutism will occur within the first week of surgery (50% within the first two days)
- Most cases resolves in a week or two.(longest 52 months) with return of functional speech.

Factors associated with the development of mutism

- Posterior fossa surgery for tumor.
- Children
- Midline tumor location
- Cerebellar vermal incision
- Large tumor size (> 5cm)
- Medulloblastoma

Cerebellar Mutism.... Pathophysiology.

UNKNOWN. However not emotional.

 Focal decreased cerebral and cerebellar blood blow leading to decreased cell functioning in particular areas, <u>dentate-thalami-cortical</u> <u>pathway</u> causing dysfunction. SPECT studies have lead support to this theory

Cerebellar Mutism.... Outcome

Speech almost always returns.

 The speech is virtually always becomes functional for communication,

Cerebellar Mutism.... intervention

- Speech therapy
- Assisting in some form of nonverbal communication
- Reassurance: usual course of cerebellar mutism and what to expect in the recovery.
- Practicing tongue and lip movements before speech returns

Brain Stem Gliomas

 Brainstem tumors comprise 10–20% of all pediatric central nervous system tumors.

 Once considered uniformly fatal; the perspective has changed now.

Clinical hallmark

- Bilateral long tract signs
- Bilateral multiple contiguous cranial nerve palsies.
- Horner's syndrome
- Inter Nuclear Ophthalmoplegia

BSG.....Classification

 The most recent classification system by Choux et al based on both CT and MRI imaging

- Type I Diffuse
- Type II Intrinsic, focal
- Type III Exophytic, focal
- Type IV Cervicomedullary

Pediatric Neurosurgery. New York, Churchill Livingstone, 2000, pp 471–491.

BSG.....

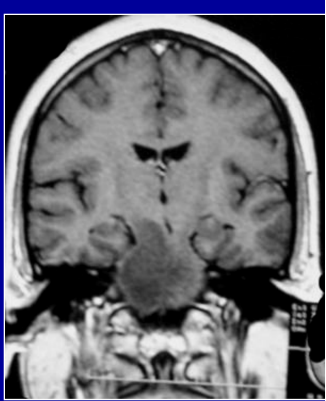
: Diffuse brainstem gliomas

- Appro. 75% of all tumors
- Hypointense on CT
- No significant enhancement on MRI.
- Characterized by diffuse infiltration and swelling of the brainstem.
- Typically, are malignant fibrillary astrocytomas (WHO grade III or IV).

Diffuse Brainstem Glioma







BSG.....

: Focal intrinsic tumors (cystic/solid)

- Sharply demarcated from surrounding tissue on MRI and are associated with less brainstem edema.
- Majority of these lesions are low grade gliomas (WHO I or II).
- Contrast enhancement : variable

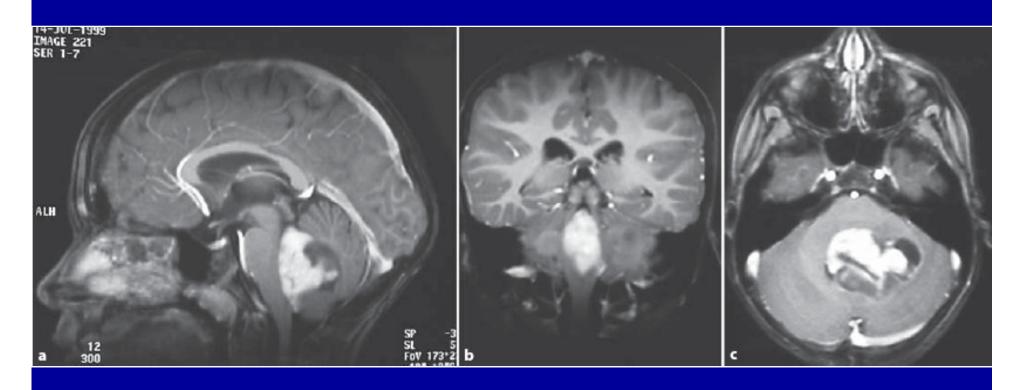
Focal Medullary BSG

BSG.....

Exophytic tumors that arise from the subependymal glial tissue of the fourth ventricle and mostly grow dorsally or laterally.

 MRI characteristics similar to type II lesions, and histologically, these lesions are usually low-grade lesions (WHO I or II) like type II lesions.

Dorsal Exophytic BSG



Post Gd T1W Post Gd

BSG.....

lesions are cervicomedullary brainstem gliomas.

- Imaging, histology and behavior: similar to intramedullary spinal cord gliomas.
- Majority are low-grade, non-infiltrative tumors.

BSG....Clinical

- Repeated vomiting with failure to thrive.
- School-aged children : a decline in school performance.
- Cranial neuropathies can develop and produce subtle changes.
- A history of dysphonia or changes in voice pitch and tone.
- Frequent upper-respiratory infections

BSG.....Management

- Biopsy: only for indeterminate lesions as no therapeutic benefit is gained by sampling lesions that behave and appear like diffuse gliomas.
- Stereotactic biopsy: can provide diagnostic tissue.
- Not without risk:
- Damage to the cranial nerves and long tracts.
- The HPE may not necessarily correlate with clinical prognosis. (Tissue heterogeneity)

Management

 A patient with a clinical presentation and imaging consistent with a diffuse glioma: NO BENEFIT from surgery.

Corticosteroids/ RT may provide temporarily benefit.

 A large phase III trial demonstrated no benefit for the use of hyperfractionated radiation in children newly diagnosed with diffuse brainstem glioma.

Mandell LR, Int J Radiat Oncol Biol Phys 1999; 43: 959–964.

Management...Focal BSG

Location	Approach
Dorsal midbrain (tectum mesencephali)	Supracere be llar infratentorial
Ventral midbrain	Pterional trans-Sylvian
Lateral midbrain	Subtemporal transtentorial
Ventrolateral pons (cerebellopontine angle)	Retromastoid retrosigmoid
Dorsal pons and medulla oblongata	Midline suboccipital transventricular (through the fourth ventricle)
Lower medulla oblongata and cervicomedullary junction	Midline suboccipital and C1 laminectomy

Management.....Postoperative Course

- Postoperative treatment and monitoring : on the location
- Patients who have had a CSF diversion procedure: monitor for reemergence of signs and symptoms of hydrocephalus.
- Tumors of the pons carry the worst prognosis because the majority are diffuse gliomas. (survival rates are low with a 1-year survival of 35–46% and 3-year survival of 11–17%.

Pediatr Neurosurg 1996; 24: 9-23.

Management.....Postoperative Course

- The postoperative course of focal medullary neoplasms depends on the tumor type.
- Dorsal exophytic tumors treated with surgery have an excellent prognosis with a 92% long-term survival some series.
 - Pediatr neurosurg 1994; 20: 2–10
- Pollack et al. reported a long-term survival of 94% in their series of 18 patients.
 - J Neurosurg 1993; 78: 859–863

Management.....Postoperative Course

 However, significant lower cranial nerve dysfunction can occur and may need prolonged postoperative ventilation or a feeding gastrostomy postoperatively.

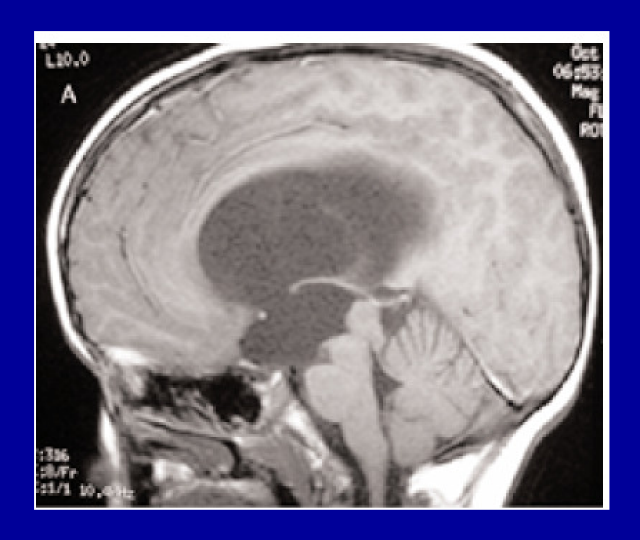
BSG....Role of stereotactic radiosurgery

Authors & Year	No. of Patients	Peripheral Dose (mean/range)	Tumor Location (no. of patients)	FU Period	Tumor Outcome (no. of patients)	Clinical Outcome (no. of patients)
Kihlstrum et al., 1994	7	14–30 Gy	tectum	6.9 yrs	disappearance (4), regression (2), progression (1)	improved (4), no change (2), sequelae (1)
Fuchs et al., 2002	12	12 Gy/9–20 Gy	midbrain (5), pons (6), medulla (1)	41 mos	absent or smaller (6), no change (5), larger (1)	improved (4), stable (3), worse (2), dead (3)
Yen et al., 2007	20	12.8 Gy/4–18 Gy	midbrain (16), pons (3), medulla (1)	78.0 mos	disappearance (4), regression (12), progression (4)	improved (8), stable (9), worse (3)

Tectal plate gliomas

- Unique subset of brainstem gliomas.
- Presents with late onset obstructive hydrocephalus that can be confused with benign aqueductal stenosis.
 - Tectal gliomas are believed to be low-grade astrocytic tumors that usually follow a benign clinical course.
 - VP shunts or ETV for CSF diversion.

MRI



AIIMS Protocol

Radical Radiotherapy with concurrent chemotherapy.

60 Gy/30#/ 6 wks.

Ependymoma

- Ependymomas are glial tumors that arise from ependymal cells within the CNS.
- WHO grade I: Myxopapillary ependymoma and subependymoma;
- WHO grade II: Ependymoma (with cellular,papillary and clear cell variants)
- WHO grade III : Anaplastic ependymoma.
- Who grade IV : Ependymoblastomas

Ependymoma

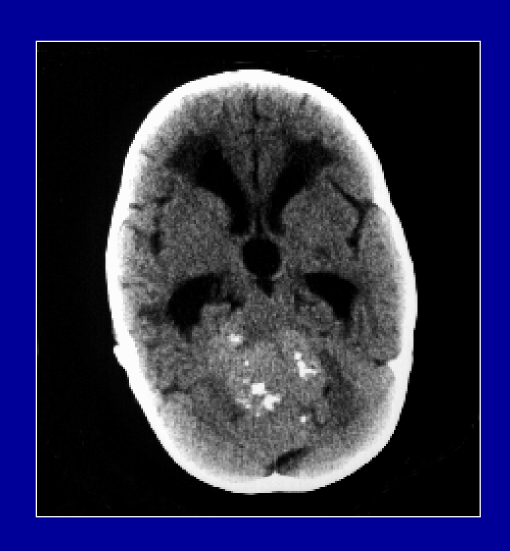
 In children: 90% of ependymomas are intracranial, majority of these occurring in the posterior fossa usually arising from the roof of the fourth ventricle

• In adults: 75% of ependymomas arise within the spinal canal, with a significant minority occurring intracranially in the supratentorial compartment.

Ependymoma Imaging

CT: Typically isodense with heterogenous enhancement

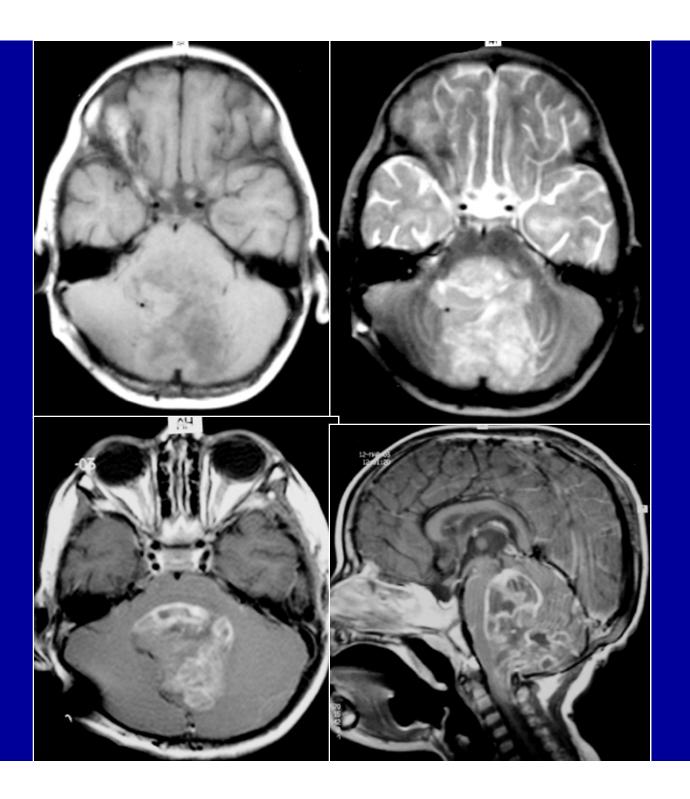
Calcification:
common (can be seen in one half of cases)



Ependymoma.....MRI

- On MRI, heterogeneous secondary to necrosis, hemorrhage and calcification.
- Heterogenous contrast enhancement
- Plastic ependymomas.
- Extension to the cerebellopontine angle is characteristic of ependymomas
- Commonly found intraventricularly
- Calcification common (appro.45% of cases)

E P E N D Y M O M A



Ependymoma.....

- Staging: No conventional staging criteria.
- Postoperative MRI is recommended within 48 hours of tumor resection to assess presence of residual tumor and to facilitate adjuvant treatment planning.

Ependymoma.....Surgery

: most significant

factor associated with increased survival in almost every large series of pediatric ependymoma.

- Aggressive primary resection,
- Immediate second look surgery if a post-operative residual tumor is identified and
- Re-surgery at time of recurrence.

Ependymoma...Role of Radiotherapy

 Post-operative radiation recommended for patients older than 3 years.

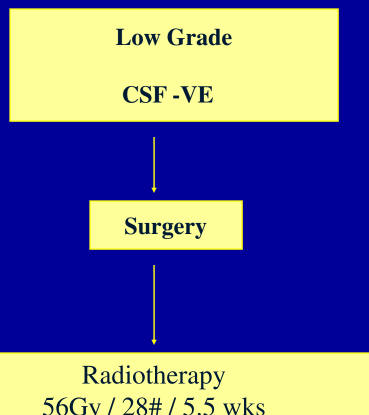
 Stereotactic radiosurgery: therapeutic option in patients with residual, unresectable or recurrent tumor.

Ependymoma...Role of Chemotherapy

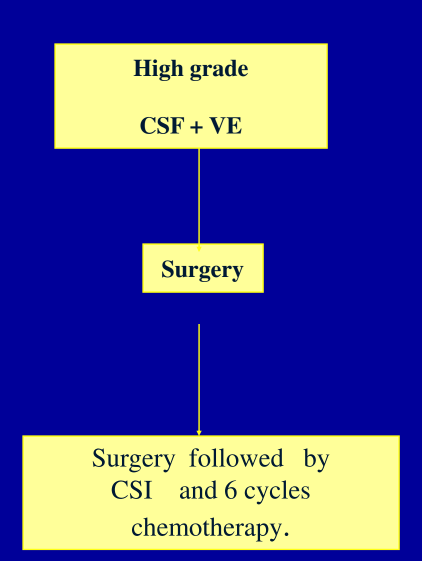
- May be useful < 3 years : Delay cranial radiation.
- Childhood intracranial ependymomas : in general chemo-resistant

over-expression of the multi-drug resistance-1 gene and the 06-methylguanine-DNA methyl transferase.

AIIMS Protocol



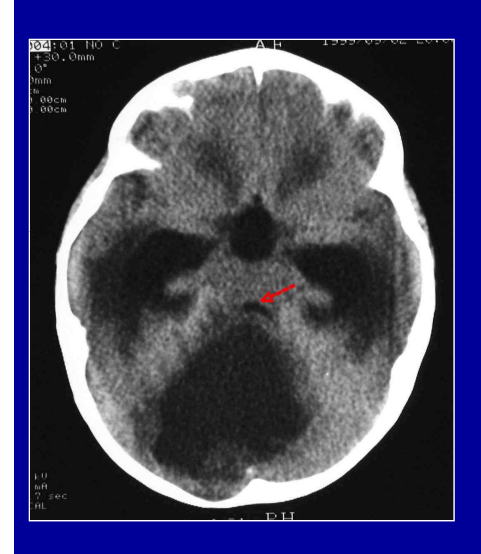
56Gy / 28# / 5.5 wks (50 Gy followed by a boost of 6 Gy)



Pilocytic astrocytoma

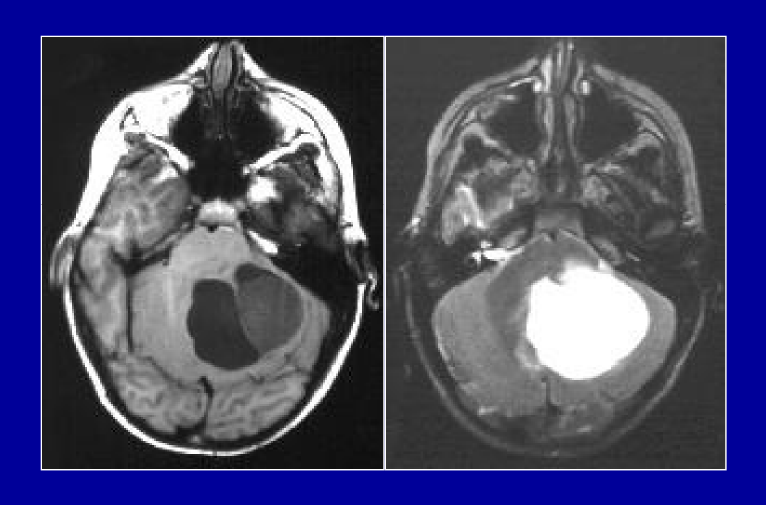
- Pilocytic astrocytoma is the most common pediatric central nervous system glial neoplasm
- Exceptional benign biologic behavior : extremely high survival rate 94% at 10 years
- Most patients present in the first 2 decades
- Surgical resection is the treatment of choice.

Pilocytic astrocytoma....NCCT + CECT





Pilocytic astrocytoma....MRI



Pilocytic astrocytoma....MRI







Pilocytic astrocytoma....MRI

Four predominant imaging patterns:

Mass with a nonenhancing cyst and an intensely enhancing mural nodule (21%)

Mass with an enhancing cyst wall and an intensely enhancing mural nodule (46%)

Necrotic mass with a central nonenhancing zone (16%), and

Predominantly solid mass with minimal to no cystlike component (17%)

Pilocytic astrocytoma....

 Surgical resection of cerebellar pilocytic astrocytomas is considered the treatment of choice.

 Radiation therapy is strictly avoided, given its risk of causing significant morbidity in children younger than 5 years of age.

Pilocytic astrocytoma....

since the surrounding cyst occurs as a simple reactive change in most cases.

Resection of the cyst wall: Controversial??

NO STATISTICAL DIFFERENCE IN SURVIVAL

has been noted in patients who have undergone resection of the cyst wall compared with those in which the cyst is left alone.

Pilocytic astrocytoma.... Prognosis

- EXCELLENT: 10-year survival rate: up to 94%

has a much better prognosis, with stable neurologic status and long term survival.

Pilocytic astrocytoma....Recurrence

- Can occur many years after surgery
- Repeat surgery : Desired treatment
- Radiotherapy can be avoided if complete resection possible.
- Residual / Unresectable recurrence : RT preferably SRS.

Choroid Plexus Papilloma

- CPP are benign neoplasms of the choroid plexus.
- Lateral ventricles: most common location in children.
- 4-6% of the intracranial neoplasms in children younger than 2 years.
- 12-13% of intracranial neoplasms in children younger than 1 year.

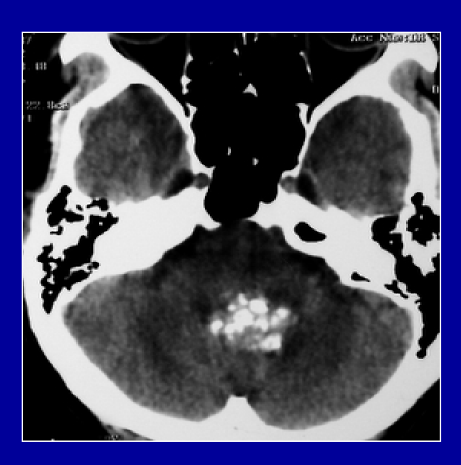
Choroid Plexus Papilloma.....Clinical

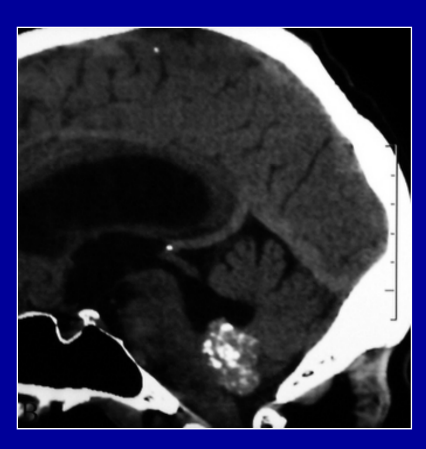
Hydrocephalus and raised ICT

The tumor itself can cause mass effect.

, possibly because of derangement of reabsorption mechanisms or blockage at other sites in the ventricular system.

Choroid Plexus Papilloma.....Radiology



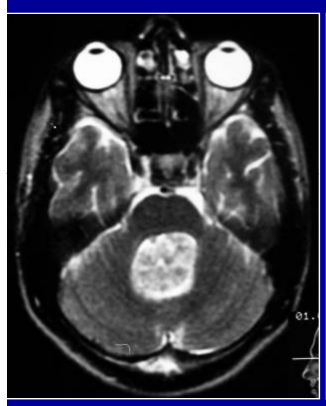


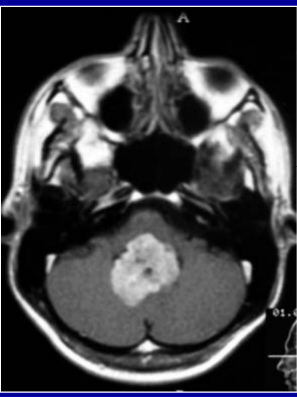
Choroid Plexus Papilloma.....Radiology

On MRI: intermediate-tostrong intensity on both T1- and T2 - weighted images with dense enhancement.

Choroid plexus carcinoma appears more heterogeneous than the papilloma and often shows adjacent parenchymal invasion or surrounding edema.

Choroid Plexus Papilloma.....Radiology







Choroid Plexus Papilloma...Management

 Treatment of hydrocephalus must be considered both before and after any surgical procedures.

An acute increase in ICP: V P Shunt.

Hydrocephalus often resolves following removal of the mass.

Choroid Plexus Papilloma...Management

- Total surgical resection is the goal.
- Complete removal: generally curative in CPP.
- Even in choroid plexus carcinoma, total resection leads to the best possible outcome.
- Adjuvant CT and RT have been demonstrated to increase survival in the treatment of choroid plexus carcinoma, although gross total resection remains the primary treatment.

Dermoid cyst

- Congenital ectodermal inclusion cysts.
- Extremely rare, constituting fewer than 0.5% of primary intracranial tumors.
- Midline sellar, parasellar, or frontonasal regions : most common sites.
- Posterior fossa (vermis or within the 4th ventricle)

Dermoid cyst

- Origin: .(inclusion of ectodermally committed cells at the time of neural tube closure (3rd–5th week of embryogenesis.)
- Glandular secretion and epithelial desquamation.
- Growth can lead to rupture of the cyst contents, causing a chemical meningitis that may lead to vasospasm, infarction, and even death.

Dermoid cyst

- Well defined, lobulated, "pearly" mass of variable size.
- Characteristically, the cyst contains thick, disagreeable, foul-smelling, yellow material due to the secretion of sebaceous glands and desquamated epithelium.
- The cysts may also contain hair and/or teeth

Dermoid cyst....MRI

Same imaging characteristics as fat

Hyperintense on T1WI and do not enhance

Heterogeneous signal intensity on T2WI



CONCLUSIONS

- Pilocytic astrocytoma bears the best outcome.
- Management of hydrocephalus still remains controversial.
- Though surgery and RT remains the treatment of choice for medulloblastoma; optimal cranispinal radiation dose remains debatable.
- Outcome for brainstem gliomas remains dismal.

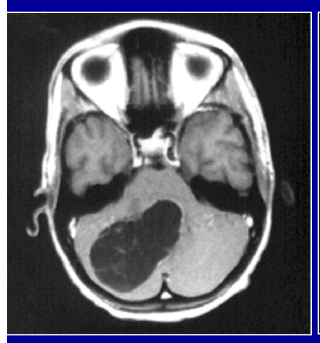
Any questions?

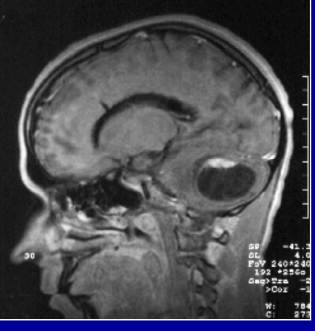
What is the diagnosis?



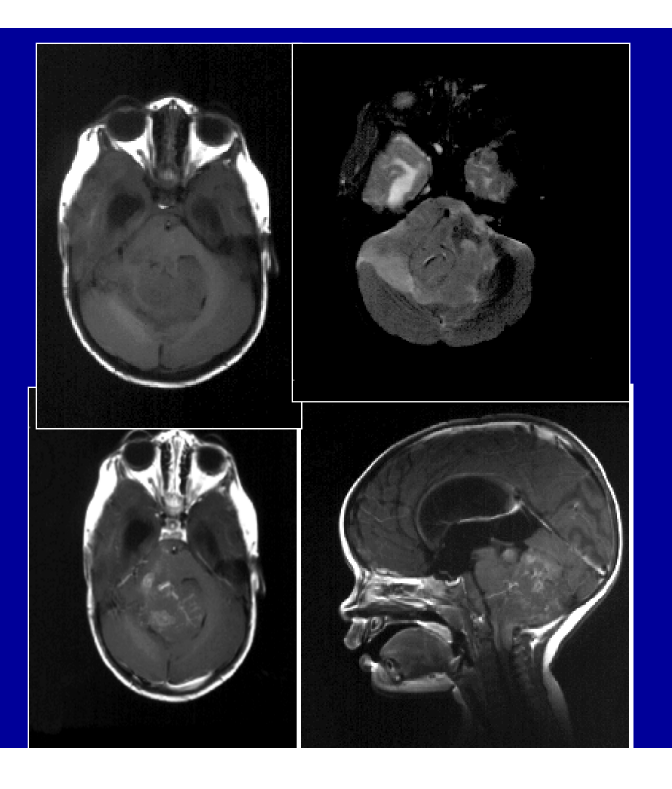


What is the diagnosis?

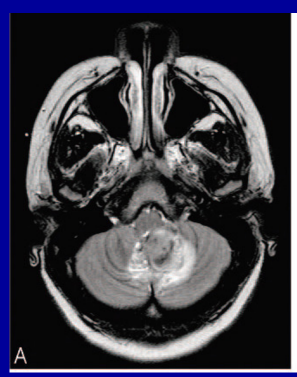


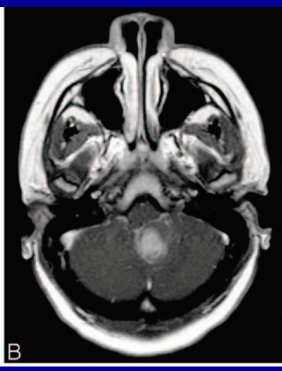


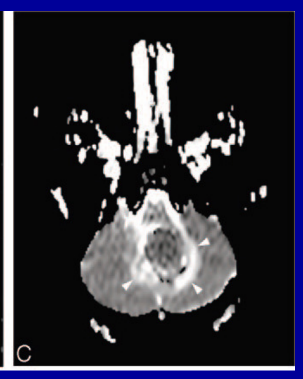




What is it??





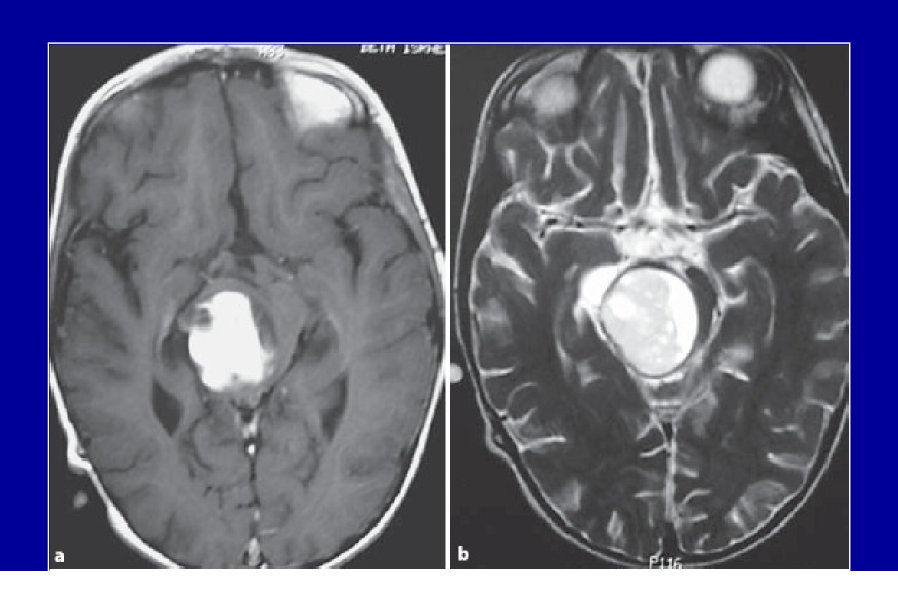


T 2 Axial

Post contrast

ADC Map

What is the tumor ??



What is it ???

