Pineal region tumors: A retrospective analysis

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Abstract
Pineal region tumors are rare comprising 0.4% to 0.1% of all primary tumors of the central nervous system1,2 and constitute 3% to 11% of childhood brain tumors.3-5 These tumors are classified into tumors of germ cell origin, which clearly account for the majority of tumors in this region and those originating from pineal parenchymal cells.1 The latter include pineoblastomas, pineocytomas, and tumors of glial origin. These retain the potential for neuronal or glial differentiation.1,6 Approximately three-fourths of tumors in this area are malignant with the propensity for seeding(3,7,8,9,10). Retrospectively we have analyzed 47 cases of pineal region tumors and have discussed the clinical features, histopathology, management and outcome.

Keywords: Pineal tumors, Central nervous system, Tumors of glial origin.

Introduction
The clinical syndromes associated with posterior third ventricular tumors relate directly to normal anatomy as well as tumor histology. Headache, nausea, vomiting and alteration of mental status are the usual presenting features. These symptoms are due to obstructive hydrocephalus caused by compression of the aqueduct of sylvius. Compression of superior colliculi either directly or due to tumor invasion results in the syndrome of vertical gaze palsy or Parinaud’st syndrome. In pineal germ cell tumors-precocious puberty, diabetes insipidus or hypogonadism may be the presenting features. In some of these tumors surgery offers cure, while others need adjuvant therapy. Adequate tissue sample for histological diagnosis is important for treatment planning. Before the development of contemporary microsurgical techniques, surgical therapy was associated with an operative mortality of 30% to 70% and morbidity of up to 65%.1-11-16 Most neurosurgeons performed only ventricular shunting for obstructive hydrocephalus followed by fractionated radiation therapy, which allowed an overall mortality of less then 5% and a five year survival rate of 60 to 75%.17-19,20 Recent advances in Neuroradiology [computed tomography (CT), magnetic resonance imaging (MRI)], micro neurosurgical techniques, modern neuroanaesthesia and postoperative intensive care have led to more encouraging surgical results. The mortality of direct surgery has been reduced to less than 5% and morbidity to a minimal level.16,21,26 Presently surgical extirpation of pineal tumors is the main approach for pineal region tumors. Chemotherapy coupled with irradiation is needed to treat malignant pineal tumors.

Remissions have been reported in many cases in which mostly cisplatin based chemotherapy in conjunction with radiation therapy was delivered.27-32 Retrospectively we have analyzed 47 cases of pineal region tumors and have discussed the clinical features, histopathology, management and outcome.

Materials and Methods
This is a retrospective study carried out in the department of surgery, Career Institute of medical sciences and hospital, I.I.M road Lucknow, (U.P).

Data was obtained from departmental records, patient files, operation notes and follow up files. From these records patient’s demographic profile, pre and post-operative neurological status, management, complications, hospital course and follow up was analyzed. Appropriate statistical methods were used to analyse the data.

Result
Observation
Forty seven cases of posterior third ventricular tumors were reviewed. There were 35 males and 22 females between the age group of 1-87 years. Majority of cases were in their second and third decade. Male to female ratio was 2.9:1.
Table 1: Study of the symptomatology

| S. No. | Symptom                  | Numbers | %    |
|-------|--------------------------|---------|------|
| 1     | Headache                 | 42      | 89.3 |
| 2     | Vomiting                 | 22      | 46.8 |
| 3     | Visual deterioration     | 16      | 34   |
| 4     | Diplopia                 | 8       | 17   |
| 5     | Generalized seizures     | 6       | 12.7 |
| 6     | Memory and behavioral changes | 7   | 14.8 |
| 7     | Altered sensorium        | 8       | 12.7 |

Table 2: Neurological deficits

| S. No. | Symptom              | Number | %   |
|-------|----------------------|--------|-----|
| 1     | Hemiparesis          | 5      | 10.6|
| 2     | 6th paresis          | 6      | 12.6|
| 3     | Perinaud’s syndrome  | 27     | 57.4|
| 4     | 7th nerve paresis    | 3      | 6.3 |
| 5     | 3rd nerve paresis    | 2      | 4.2 |
| 6     | Altered sensorium    | 6      | 12.6|
| 7     | Gait ataxia          | 11     | 23.4|

Table 3: Radiological findings

| S. No. | Diagnosis            | Total | MRI | CT Scan |
|-------|----------------------|-------|-----|---------|
|       |                      |       | T1W | T2W     | Others                               |                        |
| 1     | Epidermoid           | 12    | Hypointense | Hyper intense | Non contrast enhancing ADC* similar to brain | Hypodense, non enhancing |
| 2     | Cavernoma            | 2     | Mixed signal surrounded by hemosiderin ring | Low signal more prominent | Homogenous enhancement | Isodense, contrast enhancing |
| 3     | Arachnoid cyst       | 2     | Hypointense | Hyper intense | Non contrast enhancing, ADC* similar to water | Resemble to CSF, nonenhancing |
| 4     | Pinealocytoma        | 4     | Hypo to iso intense | Hyper intense Occasional area of hypo intense on T2W | Contrast enhancing | Isodense enhancing |
| 5     | Pinealoblastoma      | 5     | Hypo to iso intense | Hyper intense | Contrast enhancing | Isodense enhancing |
| 6     | Astrocytoma          | 1     | iso to hypointense | Heterogeneous | Irregular enhancement | Mixed density |
| 7     | Glioblastoma         | 1     | Heterogenous ment | Heterogeneous | Inhomogenous contrast enhance | Mixed density |
| 8     | Immature teratoma    | 3     | Mixed signal | Mixed signal | Mixed contrast enhancing | Mixed density |
| 9     | Germinoma            | 2     | Isointense | Isointense | Contrast enhancing | Mild hyper dense and homogenously contrast |
Surgery was performed in 35 cases (74.4%). Out of these 35 cases, 24 cases required ventriculoperitoneal shunting prior to surgical excision. In 30 cases infratentorial supracerebellar approach was used and in 4 cases transcortical transventricular approach was used (Table 4). One patient was operated endoscopically and ETV and cystoventriculostomy was performed.

Gross total excision was achieved in 17 cases. Near total excision in 6 cases and subtotal excision was performed in 12 cases.

Complications
Postoperative complications were observed in 9 cases (18.3%). CSF leak and pseudomeningocele were present in 2 and 1 case respectively. They were managed by lumbar CSF drainage. Transient up gaze palsy was seen in 3 cases and bilateral 6th and up gaze palsy was seen in one case. These cases improved subsequently on steroid therapy. Operative site extradural hematoma and seizure were noted in one case each and were treated successfully.

Discussion
Pineal region tumors are uncommon deep seated tumors of brain and comprise 1% or less of all intracranial neoplasms.1,2 These tumors are approximately ten times more common in children as compared to adults and constitute 3% to 11% of childhood brain tumors3,5 and commonly seen in 2nd decade of life.33,34 In our series 61.7% cases were present in 2nd and 3rd decade. Approximately three-fourths (75%) of Pineal tumors are malignant in nature with the tendency for seeding.3,7-10 A higher incidence of pineal region tumors in Asian countries compared to Western countries has been reported. In the Brain Tumor Registry of Japan (BTRJ), there were 38,273 primary brain tumors except those of unknown histology (1123 cases) registered in the period between 1984 and 1993. There were 807 pineal region tumors (with 104 unknown histology) who were registered in BTRJ. Of these pineal region tumors, germ cell tumors had highest frequency (70.3%), followed by pineal parenchymal tumors (12.2%), pineocytoma 7.8% and pineoblastoma 4.2%. Limited to germ cell tumors germinoma had highest incidence (68.0%), followed by pineal parenchymal tumors (12.2%), pineocytoma 7.8% and pineoblastoma 4.2%. Limited to germ cell tumors germinoma had highest incidence (68.0%), followed by teratoma (including malignant teratoma) with frequency of 14.7% in pineal region.35 In Ojemann et al, Regis et al and Alexander N. Konvalav et al series, germ cell tumor were commonest pineal region tumor.33,34,36 Gliomas were second commonest tumors in the series of Regis et al36 and Alexander N. Konovalav et al34 but pineal: parenchymal tumors were second most common in Ojemann
In a study from India, Pragati Kumar et al found highest incidence of pineal parenchymal tumors 33.8% followed by gliomas in 37%. They found germ cell tumors in only 7.4% and miscellaneous tumors in 16.6%.37 In the present series, benign lesions were present in 16 cases (34%) and malignant lesions were present in 31 (65.9%) cases. In benign lesions, epidermoid tumors were commonest (12 cases). In malignant lesions, Germ cell tumors were most common (16 cases), followed by pineal parenchymal tumors (8 cases) and gliomas (7 cases). Majority of pineal region tumors present with symptoms of intracranial hypertension because of obstructive hydrocephalus due to compression or direct infiltration of the aqueduct of sylvius. Approximately 90% of patients with pineal region tumors have symptoms of intracranial hypertension at the time of presentation.3 In our series 89.3% cases had symptoms of raised intracranial features. Headache was present in 42 cases (89.3%) and it was associated with vomiting in 22 cases (46.5%). Perinaud’s syndrome was seen in 27 (57.4%) cases. In a study carried out by Alexander N. Konovalov et al found eye movement disorders in 76% cases 30. Magnetic resonance imaging and Computed tomography are important diagnostic tools in the detection of these tumors. CT scan easily detects tumor calcification. MRI is the investigation of choice for pineal region tumors. It gives accurate information about size, extent of tumor, anatomical relations to surrounding neurovascular structures. Establishing histological nature of the tumor and extent of tumor resection are important prognostic factors. These two factors appear dominant in predicting outcome.25,38-40 Before the development of microsurgical techniques, treatment strategy usually involved ventricular drainage followed by fractionated radiation therapy because of high operative mortality of 30% to 70% and morbidity of up to 65% after surgery.11-16 Ventricular drainage followed by fractionated radiation therapy had overall mortality of less than 50 and a S-year survival rate of 60 to 75 year.17,18 Nowadays with the introduction of modern microsurgical techniques 24 Y and technology, development of neuroanesthesiology and neuro-radiological techniques, surgical treatment of pineal region tumors can be carried out. The mortality of direct surgery has been reduced to under 5% and morbidity to a minimal level.16,21-26

In the present series, ventriculoperitoneal shunting was performed 33 cases (70.2%). In 9 cases, only VP shunt was done and in 24 cases definite surgery was done after VP shunt. Nowadays the endoscopic third ventriculostomy is good alternative to normalize CSF circulation for hydrocephalus in pineal region tumors. In patients with malignant germ cell tumors or pineoblastomas, there is risk of peritoneal metastasis following shunting.42,43,49 Infection, malfunction or peritoneal metastasis associated with ventriculoperitoneal shunts are eliminated by third ventriculostomy. In the present series ETV and biopsy was done in 3 cases. ETV itself can potentially facilitate dissemination of the neoplasm.44,45 Haw and Steinbok47 reported metastatic deposit of the pineal germinoma at the ventriculoscope tract. There was no dissemination in ETV group in the present series.

Presently, surgical excision remains the treatment of choice for pineal region tumors and a wide variety of lesions can be safely excised.7,23-26,48,49 Many approaches for removal of pineal region tumors can be adopted. Supracerebellar-infratentorial and occipital-transtentorial approaches are commonly used surgical approaches for this location.3a In lg7l, Stein popularized the infratentorial supracerebellar approach, originally described by Krause.

Germinomas are highly radiosensitive and radiation is the primary treatment modality. There is no significant difference between the outcome of patients irradiated after direct surgery and those receiving radiotherapy alone.8,17,50 The patients with definite radiologic features do not require direct surgery or stereotactic biopsy for pathologic confirmation of diagnosis and can be irradiated directly. In the present series direct radiotherapy was given to 11 cases on the basis of radiological diagnosis of germinoma and these patients responded very well. Three cases were lost to follow up. The tumor disappeared in other with no recurrence in available follow up.

Aggressive tumors like malignant germ-cell tumors, pineal parenchymal cell tumors tend to invade surrounding structures and have a risk of CSF dissemination.27,29-32,51 There is controversy about the extent of radiation. At present craniospinal irradiation is indicated only in patients with clinical or radiologic evidences of dissemination of tumor because myelopathy is a serious late complication of craniospinal irradiation.38,52-54 In the present series, one case of pinealocytoma had leptomeningeal seeding. Platinum-based multiagent chemotherapy has improved the outcome in patients with nongerminomatous germ cell tumors and anaplastic pinealocytomas.30,31,43,55,56 The benefits of this approach are still debated,29,51 although patients with pineoblastomas are often treated with adjuvant systemic chemotherapy after craniospinal irradiation. The incidence of pineal region epidermoid tumors is approximately 3% to 4% of all intracranial epidermoids.57 The incidence of epidermoid tumors in the 26 pineal region varies from 3.4% to 10% in different series.57,58 In present series, epidermoid was present in 12 cases (255%) out of 47 cases. Exact pathogenesis of epidermoids in general is still under discussion. Defect in the cleavage of the neural tissue from
the cutaneous ectoderm, embryonic inclusions, differentiation from multipotential cell rests, and epithelial remnants are the various other mechanisms implicated in the origin of epidermoid tumor. The seeding of the subarachnoid space by the irritant component of the epidermoid cyst results in aseptic chemical meningitis. The incidence of recurrence after radical excision is uncommon.57-60

Pineal region cavernomas and arachnoid cysts are rare and only case reports are available in the literature.61-63 In present series 2 cases of arachnoid cysts and 2 cases of cavernomas were noted. There was no recurrence in available follow up.

Conclusion
Pineal tumors are a heterogeneous group of mass lesions originating in and around the pineal gland and represent a spectrum of neoplasms" Direct surgical excision remains the treatment of choice for pineal region tumors with minimum mortality and morbidity. Complete excision should be the goal for majority of these lesions. For germinomas, radiation therapy good long term recurrences free survival and outcome. Endoscopic biopsy and ETV offers histological diagnosis and CSF diversion in many of such lesions and further treatment can be planned accordingly.

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Conflict of Interest
None.

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