Diagnosis and management challenge of a granular cell astrocytoma of the pineal region: case report

Abad Cherif El Asri, MD,1 Hassan Baallal, MD,1 Youssef Zoubeir, MD,2 Mohamed Sinaa, MD,2 Abderrahman Albourzidi, MD,2 Miloudi Gazzaz, MD,1 Ali Akhaddar, MD,1 Mohamed Boucetta, MD,1 and Brahim El Mostarchid, MD1

Departments of 1Neurosurgery and 2Pathology, Military Hospital, Rabat, Morocco

Granular cell astrocytoma (GCA) is a rare type of infiltrative brain tumor with most reported cases occurring in the suprasellar region. A pineal localization is extremely rare, with only 4 previously reported cases in the literature. The authors describe the case of a 16-year-old boy who developed signs of increased intracranial pressure and Parinaud syndrome. Cranial CT and MRI revealed a well-demarcated and enhanced mass in the pineal region accompanied by obstructive hydrocephalus. Subtotal resection was performed via a subtemporal approach. A histological diagnosis of GCA was made. Three years after surgery, the patient was alive and well without adjuvant therapy, and serial MRI showed no signs of progression of a small residual tumor. After a thorough review of the different epidemiological, clinical, and imaging features; treatments; and prognoses of GCAs in other intracranial localizations, the authors analyzed features of this tumor in the pineal region.

http://thejns.org/doi/abs/10.3171/2014.10.PEDS13388

KEY WORDS granular cell tumor; astrocytoma; pineal region; surgery; oncology

Case Report

History and Examination

A 16-year-old previously healthy boy presented with a 1-month history of excruciating headaches associated with vertigo and occasional episodes of nausea and vomiting. He had abnormal extraocular movements with incomplete Parinaud syndrome and moderate bilateral papilledema. There was no other neurological deficit. Cranial CT demonstrated a mass of the pineal gland, including ganglioneuromas, gangliogliomas, and gliomas.5,6 Granular cell astrocytoma (GCA) is thought to represent an unusual astrocytoma originating from the pineal gland.3,10,15 There have been only 4 previously reported cases of GCA in the pineal region.5,10,15 In this report, we describe a surgical case of GCA in the pineal region. After a thorough review of the different epidemiological, clinical, and imaging features; treatments; and prognoses of GCAs in other intracranial localizations, we analyzed features of this tumor in the pineal region (Table 1).

http://thejns.org/doi/abs/10.3171/2014.10.PEDS13388

KEY WORDS granular cell tumor; astrocytoma; pineal region; surgery; oncology

http://thejns.org/doi/abs/10.3171/2014.10.PEDS13388

KeY words granular cell tumor; astrocytoma; pineal region; surgery; oncology

ABBREVIATIONS EMA = epithelial membrane antigen; GCA = granular cell astrocytoma; GCT = granular cell tumor; GFAP = glial fibrillary acidic protein.


INCLUDE WHEN CITING Published online February 20, 2015; DOI: 10.3171/2014.10.PEDS13388.

DISCLOSURE The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.
Approximately 70% of the tumor was resected; the rest was closely apposed to the basal and internal veins.

Posttreatment Course

Histological examination showed the specimen to be moderately cellular with marked nuclear pleomorphism and foamy cytoplasm. Numerous bizarre multinucleated giant cells but no mitosis, necrosis, or endothelial proliferation were observed (Fig. 2). Glial fibrillary acid protein (GFAP), epithelial membrane antigen (EMA), macrophage marker CD68, PAS, and S100 were immunohistochemically positive. The tumor cells were negative for PAS-diatase, anti-neurofilaments, human chorionic gonadotropin, $\alpha$-fetoprotein, and placental alkaline phosphatase antibodies. Consequently, the tumor proved to be a GCA.

Postoperatively, mild right paresis was newly observed, but there was no other neurological deficit, and the patient did not require care in his daily life. Because the tumor was considered benign, he did not receive any adjuvant therapies. Cranial MRI studies at 6 months, 1 year, and 3 years after treatment showed no signs of growth of the residual tumor (Fig. 3).

Discussion

Granular cell astrocytoma, which has also been called "intracerebral granular cell tumor" or "astrocytoma with granular cell differentiation," is a rare type of infiltrative brain tumor. It was first described by von Luthy and Klinger, in 1951, as a granular cell tumor (GCT) in the

---

**TABLE 1. Literature summary of GCA cases**

<table>
<thead>
<tr>
<th>Authors &amp; Year</th>
<th>Age (yrs)</th>
<th>Sex</th>
<th>Onset of Symptoms</th>
<th>MRI Findings</th>
<th>Surgery</th>
<th>CSF Shunt</th>
<th>RT</th>
<th>FU &amp; Outcomes</th>
<th>CSF Shunt</th>
<th>MRI Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Snipes et al., 1992</td>
<td>25, F</td>
<td>6 yrs</td>
<td>Headache</td>
<td>Pineal tumor with 3rd ventricular obstruction</td>
<td>Occipital transtentorial approach/occipital transtentorial approach/occipital transtentorial approach</td>
<td>Yes</td>
<td>56 Gy</td>
<td>8 yrs; no signs of recurrence</td>
<td>None</td>
<td>Total resection</td>
</tr>
<tr>
<td>Nitta et al., 2001</td>
<td>30, F</td>
<td>Suddenly</td>
<td>Rt paresis followed by loss of consciousness</td>
<td>2.0 × 1.5 × 1.3–cm partially Gd-enhanced mass in pineal region</td>
<td>Occipital transtentorial approach/reddish, soft, &amp; vascular</td>
<td>Yes</td>
<td>None</td>
<td>7 yrs; no signs of recurrence</td>
<td>None</td>
<td>Total resection</td>
</tr>
<tr>
<td>Ohta et al., 2010</td>
<td>67, F</td>
<td>Progressive</td>
<td>Headache &amp; gait disturbance</td>
<td>3.0 × 3.4 × 4.0–cm low-intensity lesion &amp; heterogeneous contrast enhancement</td>
<td>Occipital transtentorial approach/reddish, soft, &amp; vascular</td>
<td>No</td>
<td>45 Gy</td>
<td>1.5 yrs; possible residual tumor in right temporoparietal region; patient alive well</td>
<td>None</td>
<td>Total resection</td>
</tr>
<tr>
<td>Present case</td>
<td>16, M</td>
<td>1 mo</td>
<td>Parinaud syndrome, raised intracranial tension</td>
<td>3.0 × 3.5 × 4.0–cm partially heterogeneous contrast enhancement</td>
<td>Subtemporal approach/reddish, gritty fibrotic, &amp; vascular</td>
<td>Yes</td>
<td>None</td>
<td>3 yrs; no signs of progression</td>
<td>None</td>
<td>Subtotal resection</td>
</tr>
</tbody>
</table>

Approximately 70% of the tumor was resected; the rest was closely apposed to the basal and internal veins.

FIG. 1. Preoperative sagittal (A) and axial (B) T1-weighted MR images showing a 3 × 3 × 4–cm homogeneously enhanced lesion in the pineal body and third ventricle wall. Sagittal T2-weighted MR image (C) and axial FLAIR image (D) showing hyperintensity of the adjacent structures.
central nervous system. Most reported cases occur in the suprasellar region. The pineal localization is extremely rare, with only 4 previously reported cases in the literature.9,10,15

The histogenesis of a GCT is debated. For tumors that arise from the peripheral nervous system, Schwann cells appear to be the most likely cells of origin.15 The source of a pineal region GCT may differ from its extracranial counterpart. For GCTs in the sellar area, the pituicyte, a modified astrocyte and the principal posterior pituitary cell, is considered the cell of origin.3,12,14 Some authors now consider the GCT as a hamartomatous lesion; and other authors, as tumors of uncertain histogenesis.13 To date, no recognized common cell accounts for all types of GCT.

The age of tumor occurrence in 4 cases, including our patient, was under 40 years, and only 1 patient was 67 years of age. The cases to date occurred in 3 males and 2 females. Symptoms were generally related to the compression of adjacent structures. The most frequent symptoms, including in our case, were dysfunctions of eye movements (Parinaud syndrome) and signs of rising intracranial tension caused by hydrocephalus (Table 1). Sudden-onset diplopia or visual loss, short-term history of confusion, headache, and vomiting have been described by Snipes et al.15 and Nitta et al.9 The duration of symptoms has ranged from sudden onset (5 days) to 4 years. Spontaneous intratumoral hemorrhage and intraventricular hemorrhage have also been reported in suprasellar region GCAs.3,14

The imaging findings of GCTs are relatively nonspecific given the broad range of attenuation and signal intensity of most common pineal region masses: germinoma, meningioma, pineocytoma, and glioma.3,8,10,14,15 In our case and in the case reported by Ohta et al., the tumor presented as a well-demarcated, low-intensity to isointense lesion in the pineal body and third ventricle wall with homogeneous contrast enhancement.16 In these cases the CT and MRI characteristics were similar to the appearance of a large meningioma. Snipes et al. described the CT scanning features of a case with a cystic component.15 On the other hand, a GCA in the pineal region has often been associated with increased signal intensity in the adjacent thalami on MRI, while GCAs elsewhere promote virtually no edema in the adjacent brain structures. The presence of calcification, which was reported by Snipes et al., was exceptionally reported in other localizations.15

Granular cell tumor is generally a surprising finding in that it is seldom considered in the preoperative differential diagnosis of a pineal region lesion.3,8,10,15 They display a distinctive histological appearance characterized by the presence of neoplastic cells with large eosinophilic and granular cytoplasm.4,10,12,13 At the periphery of the tumor, gradual transition into classic astrocytoma is frequently found, although it can be subtle.6–9,12 Tumors can be entirely composed of granular cells, but more often granular cells coexist with conventional infiltrating astrocytoma. Immunohistochemical studies show that most GCAs are positive for GFAP and S100.5,7 Negative GFAP immunostaining was reported in a few articles.1,3,8,10,12,13 As in our case, GCAs are positive for CD68, PAS, and EMA.

The treatment of pineal region tumors remains one of the major intellectual and technical challenges facing neurosurgeons. The choice of the surgical approach depends...
on the size and histological nature of the tumor and the neurosurgeon’s habits. In our case, we used the subtemporal approach to achieve subtotal resection of the tumor. The parietooccipital interhemispheric transtentorial route was performed in the case described by Nitta et al., while the occipital transtentorial approach was used by Snipes et al. for subtotal resection in 2 cases and by Ohta et al. for total resection in 1 case.9,10,15 The firm and vascular nature of GCTs often encountered intraoperatively, along with the lack of an obvious dissection plane between the tumor and normal brain and adjacent vessels, often prohibits their gross-total resection.3,9,10,15 An attempt at subtotal resection of this slow-growing tumor may be justified in certain patients in whom gross-total removal was determined to be too risky at the time of surgery. In only 1 reported case, the tumor was considered soft and could be aspirated after radiation therapy.15 Nitta et al. found no signs of recurrence for 7 years postoperatively without any adjuvant therapy.15 For this reason, we did not administer adjuvant therapy to our patient.

The natural history of a GCT is poorly understood.3,9,12 Although no systematic studies of this tumor have been undertaken to date, GCTs of the neurohypophysis and pineal region, unlike GCAs of the cerebral hemisphere, seem to be typically benign and indolent despite occasional invasion and recurrence.2,3,10 Follow-up evaluation in our case, in the 4 previously reported pineal cases, and in cases in all other intracranial localizations has shown that subtotal resection is effective for symptomatic tumors.5,8,10,12,15 The role of radiation therapy for this benign, slow-growing tumor remains unclear. Outcomes analysis of the published cases treated using subtotal resection and with available follow-up data showed no difference in recurrence between patients who received radiation treatment and those who did not. Glazer et al. described 1 patient who underwent 3 weeks of postoperative radiation therapy consisting of 32 Grays after subtotal resection.4 The patient died 3 months postoperatively. No radiation effect on the tumor was noted at autopsy. Therefore, in light of the tumor’s slow growth, radiation therapy for these lesions remains controversial.3 However, adjuvant radiation treatment may be reasonable for tumors with atypical features.11

Conclusions

A GCT should be considered in the differential diagnosis of an enhancing lesion in the pineal region. This tumor is commonly firm and vascular. However, the surgeon must be prepared to face a hemorrhagic unresectable tumor. The goal of decompression of the surrounding structures via subtotal resection may be reasonable. Although this tumor has been known as an aggressive astrocytic entity, our review of the pineal localization revealed a relative benign tumor prognosis. Thus, in our opinion, GCA in the pineal region must be managed with less aggressive therapeutic management.

Acknowledgment

We thank Sophie Domingues-Montanari for her help with English editing.

References


Author Contributions

Conception and design: El Asri, Baallal, Boucetta, El Mostarchid. Acquisition of data: El Asri, Sinha, Gazzaz. Analysis and interpretation of data: El Asri. Drafting the article: Sinha. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: El Asri.

Correspondence

Abad Cherif El Asri, Department of Neurosurgery, Military Hospital, Rabat 10100, Morocco. email: abad20031@hotmail.com.