Case Report
Intra-suprasellar paraganglioma radiographically resembling a craniopharyngioma: a case report and literature review

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Abstract: Paragangliomas arising in the sellar region are extremely rare. Most of the cases presented as adenoma or meningioma. Here, we described a case of a 40-year-old woman with sellar paraganglioma presented as craniopharyngioma. She was admitted to our department with the complaint of gradually decreasing visual acuity and slightly disturbance of menstruation for 6 months. Computed tomography (CT) and magnetic resonance imaging (MRI) scan showed an intra- and supra-sellar expanding lesion which was regarded as craniopharyngioma before operation. Histological examination was diagnostic of a paraganglioma. We review the literature and discuss the characteristic feature of sellar region paragangliomas.

Keywords: Paraganglioma, sellar, craniopharyngioma

Introduction
Neoplasm of the sellar region are entities with a large differential diagnosis. Pituitary adenoma, craniophargioma and meningioma are the most common lesion in this region. Paragangliomas are tumors derive from the neural crest cells; they include neoplasm of the adrenal medulla and paraganglia. They may occur at multiple locations within the body. Paragangliomas which arise from sellar and parsellar area are very rare and there have been reported only a few cases in the literature up to date. Here we report a further case of sellar region paraganlioma presenting as craniopharyngioma.

Case report
A previously healthy 40 year-old women presented with gradually decreased visual acuity accompanied with slightly disturbance of menstruation for 6 months. Neurological examination revealed bitemporal hemianopsia. Laboratory examination findings were within normal limits except slightly elevated PRL level. Computed tomography (CT) revealed a high-dense intra-suprasellar cystic lesion (Figure 1). Magnetic resonance imaging (MRI) showed the cyst cavity as a hyper-intense area in T1-weighted images and slightly enhanced with gadolinium (Figure 2). The operation was performed through subfrontal approach. An yellow colloid mass inside the cyst and part of the cyst wall were removed. The optic nerve compression was released. After the operation the patient recovered her visual acuity and menstruation. External beam radiation at a dose of 50 Gy was applied to the residual tumor. During 3-year postsurgical follow-up, the patient remained in good health.

Pathological findings
Under operation microscope, the tumor was red cyst with highly vascularized cystic wall, yellow colloid mass and a piece of confirm mass inside the cyst. Histopathological examination revealed neoplastic cells were arranged in well-defined cell clusters separated by fibrous septa (Figure 3). Mitotic figures were not conspicuous. Immunohistochemical analysis showed that tumor cells were strongly positive for chromogranin A (cga), synaptophysin (syn) and...
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Figure 1. CT scan of paraganglioma. CT scan of paraganglioma showing a round homodensity lesion in the sellar region, at front part of the lesion a high density mass which was confirmed to be bony like mass in the operation.

neuron-specific enolase (NSE), and were negative for endomysial antibody (EMA) (Figure 4). The tumor cells were also immunonegative for all pituitary hormones. Combination histological features with immunohistochemical findings confirmed the diagnosis of a paraganglioma.

Discussion

Since the first description in 1967 [1], only 22 cases of paraganglioma in sellar region have been reported in English literature (Table 1). The average age at diagnosis for them is 47 years (14-84) and predominance in male was seen in these reported cases (16 males and 6 males), in contrast to paragangliomas of the glomus-jugular region, whose predominance in female was reported by large case series (33 males and 198 females) [2]. More cases and further study were needed to explain this discrepancy.

Paragangliomas are assumed to originate from ectodermic cells derived from neuron crest progenitor cells [3], Extra-adrenal paragangliomas can be found in the intercarotidium tissue, tympanojugular and vagal nerve [4], in central nervous system paraganglioma most frequently affect glomus jugular [2] and cauda equine [5]. The vast majority of paragangliomas in central nervous system presented as tumors at cauda equine and glomus jugular, however, rare examples of purely intracranial presentation, including sellar region like the present case, cerebellar region, and skull base, have been reported. There is no definite explanation for the development of intracranial paragangliomas. The possible hypotheses include (1) anomaly of migration of neural crest cells; (2) anomaly of involution of fetal paraganglia (persistence of vestigialtissue); (3) intracerebral metastasis of an undiagnosed paraganglioma in another location [6, 7]. Further studies are needed to clarify the tumorigenesis of intracranial paraganglioma.

Since intracranial paragangliomas show no characteristic clinical features, such as endocrinological symptoms seen in adrenal paragangliogliomas, pre-operative diagnosis of paragangliomas in sellar region can be obtained only by radiographic examinations. Radiographic findings of paragangliomas in the sellar region were summarized in Table 1. Because majority of them appeared as well-enhanced hypo-isointense mass on T1-weighted images, which led to the preoperative diagnosis of pituitary adenoma or meningioma due to their radiographic similarities. Conversely, the tumor
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in the present case showed hyperintense in T1 weighted image, which induced presurgical diagnosis of craniopharyngioma. During the operation, the yellow colloid mass inside the cyst and highly vascularized reddish cyst wall were observed. To the best of our knowledge this is the first report of paraganglioma in sellar with such kind of radiological and intraoperative manifestation.

Pathological features of paraganglioma resembles normal paraganglia, composed of chief cells in well-defined cell nests (Zellballen), surrounded by layer of sustentacular cells. In our case, the typical zellballen architecture was evident, however, the S-100 positive sustentacular cells were scarce. The same phenomena were also reported by other author, these authors regarded the tumor with few sustentacular cells as more aggressive forms of paragangliomas. The correlation between the presence of sustentacular cells and clinical behavior should be further confirmed.

We treated the present case with postoperative radiotherapy. During the 3-year follow-up, the residual tumor remains radiographically stable. Most of the reported paragangliomas at sellar region underwent less than subtotal remove in order to avoid damaging adjacent structures, subsequently received radiotherapy (Table 1). There was no clear evidence of radiation effects for intracranial paragangliomas, Postoperative radiation for re-

Figure 2. MR image of paraganglioma. Left, T1-weighted without gadolinium; right, T1-weighted with gadolinium. MRI image showing a hyper intensive lesion of sellar region in T1, and showing enhancement of the wall after gadolinium administration.
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Figure 3. HE stain of the tumor. Nest of neoplastic cells separated fibrous septa. Tumor cells showing pleomorphism (×400).

Figure 4. Immunohistochemical stain of the tumor. Positive to chromogranin A (A), neuron-specific enolase (B), and synaptophysin (C), confirmed the neuroendocrine native of the tumor (×400).

radiologic manifestation of sellar region paraganglioma was complex. Histopathological and immunohistochemical diagnosis was essential for the final confirmation. When diagnosed of sellar region tumors, especially the cystic lesion in this region, paraganglioma should also be considered.

Disclosure of conflict of interest
None.

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References
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Table 1: Over review of reported sellar paraganglioma

<table>
<thead>
<tr>
<th>Author</th>
<th>Age/sex</th>
<th>Location</th>
<th>Radiology</th>
<th>Operation finding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chaudhry NS [8]</td>
<td>44/male</td>
<td>s/ss</td>
<td>Hyperdense in CT, enhancing, well-defined heterogeneous in consistency (MRI)</td>
<td>Very bloody, freshly/subtotal</td>
</tr>
<tr>
<td>Yokoo H [9]</td>
<td>52/female</td>
<td>ss</td>
<td>T1 low signal T2 high signal homogenously Enhanced (MRI)</td>
<td>Extremely hemorrhagic/biopsy</td>
</tr>
<tr>
<td>Ünal Özüm [10]</td>
<td>70/male</td>
<td>s/ps</td>
<td>Heterogeneous sellar mass (CT)</td>
<td>Soft, small grayish-brown tissue fragments/subtotal</td>
</tr>
<tr>
<td>Nagpara O [11]</td>
<td>47/male</td>
<td>s/ss</td>
<td>Isointense T1, hyperintense T2, enhanced (MRI)</td>
<td>Extravascular greyish-to-pinkish soft tumor/subtotal</td>
</tr>
<tr>
<td>Boari N [12]</td>
<td>52/male</td>
<td>s/ss</td>
<td>Homogenously enhanced (MRI)</td>
<td>Grey-pink fleshy tissue/subtotal</td>
</tr>
<tr>
<td>F. Hertel [13]</td>
<td>51/ambel</td>
<td>s/ps</td>
<td>Isointense T1 hypointense T2 enhanced (MRI)</td>
<td>Inhomogeneous, not extremely vascular tumor</td>
</tr>
<tr>
<td>Zorlu F [14]</td>
<td>37/male</td>
<td>s/ps</td>
<td>Contrast enhancing parasellar mass (MRI)</td>
<td>Subtotal</td>
</tr>
<tr>
<td>Haresh KP [15]</td>
<td>17/male</td>
<td>s/ps</td>
<td>Isointense on T1 hypointense T2, well defined mass (MRI)</td>
<td>Highly vascular firm to hard tumor/biopsy (bone metasis)</td>
</tr>
<tr>
<td>Sinha S [16]</td>
<td>18/male</td>
<td>s/ss</td>
<td>Isointense on T1 hypointense T2, enhanced (MRI)</td>
<td>Subtotal (malignant)</td>
</tr>
<tr>
<td>Lu QJ [17]</td>
<td>81/male</td>
<td>s/ss</td>
<td>Well-circumscribed heterogeneous enhanced (MRI)</td>
<td>Variable consistency, both soft and firm/subtotal/metastasis from esophageal</td>
</tr>
<tr>
<td>Voulgaris SG [18]</td>
<td>48/male</td>
<td>s/ss</td>
<td>Solid and cystic hypointense T1 hyperintense T2 enhanced (MRI)</td>
<td>Encapsulated, with reddish-brown mass, in consistency/subtotal</td>
</tr>
<tr>
<td>Del Basso De Caro ML [19]</td>
<td>84/male</td>
<td>s/ss/ps</td>
<td>Enlarged and eroded sellar turcica occupied by isodense mass (CT)</td>
<td>Soft yellow-reddish very hemorrhagic tissue</td>
</tr>
<tr>
<td>Mokry M [20]</td>
<td>76/female</td>
<td>s/ss</td>
<td>Lower signal T1 inhomogeneous T2 Enhanced (MRI)</td>
<td>Soft, highly vascular, friable tumor/subtotal</td>
</tr>
<tr>
<td>Flint EW [21]</td>
<td>17/female</td>
<td>s/ss</td>
<td>Mass with some calcification (CT) vessel in the mass (MRI)</td>
<td>Not mention</td>
</tr>
<tr>
<td>Do Nascimento [22]</td>
<td>32/female</td>
<td>s/ss</td>
<td>Mass suggestive of meningioma or pituitary adenoma (CT)</td>
<td>Not mention</td>
</tr>
<tr>
<td>Sambaziotis D [24]</td>
<td>54/male</td>
<td>s/ss</td>
<td>Clepsydra-shaped mass (MRI)</td>
<td>Highly fibrous and vascular tumor with extremely firm consistency</td>
</tr>
<tr>
<td>Salame K [25]</td>
<td>48/female</td>
<td>s/ss</td>
<td>Homogenous, isointense T1, hypertense T2 diffuse enhanced (MRI)</td>
<td>Red-gray somewhat fibrous mass, highly vascular/subtotal;</td>
</tr>
<tr>
<td>Scheithauer BW [26]</td>
<td>14/male</td>
<td>s/ss</td>
<td>Contrast-enhancing mass (CT)</td>
<td></td>
</tr>
<tr>
<td>Steel TR [27]</td>
<td>44/female</td>
<td>s/ss</td>
<td>Lobulated lesion isointonic T1 uniformly enhancement (MRI)</td>
<td>Soft white to slightly pink encased tumor/subtotal</td>
</tr>
<tr>
<td></td>
<td>41/female</td>
<td>s/ss</td>
<td>Uniformly enhancement (MRI)</td>
<td>Soft pinkish-white tumor/subtotal</td>
</tr>
</tbody>
</table>

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