A Case of Coincidental Intrasellar Chordoma and Pituitary Adenoma

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INTRODUCTION

Chordomas are midline tumors that arise from ectopic cells of notochord remnants, though primarily intrasellar chordomas are extremely rare. In this report, the authors describe the case of a 68-year-old female with partial abducens nerve palsy in the right eye due to the intrasellar cystic tumor. After endonasal trans-sphenoidal surgery, intraoperative and histopathological findings confirmed the co-occurrence of an entirely intrasellar chordoma and pituitary adenoma. To our knowledge, the present case is the third reported case of an intrasellar chordoma with a pituitary adenoma.

Key Words  Chordoma; Pituitary adenoma; Sella turcica; Pathology.

CASE REPORT

A 68-year-old female presented with slowly progressive diplopia for 1 month when she stared to the right. During the neurological examination, the patient exhibited partial abducens nerve palsy in the right eye, but there was no other neurologic deficit. And, the finding of visual field examination was unremarkable. A computed tomography (CT) scan revealed an intrasellar mass extending to right cavernous sinus. Although the sellar floor was thinned, there was no bony destruction of the skull and no calcification (Fig. 1). Magnetic resonance imaging (MRI) demonstrated a 1.5×2×1.5 cm-sized cystic lesion with a well-enhanced margin and a slightly enhanced inside. The mass compressed the medial wall of the cavernous sinus without invasion (Fig. 2). Pituitary hormone function test revealed normal levels.

Based on the clinical and radiological findings, either a pituitary macroadenoma or a craniopharyngioma were suspected. The patient underwent an endonasal trans-sphenoidal resection. However, an intraoperative finding showed two different tumors in the sella turcica. The first mass was located in the left and anterior side of the sella turcica, and revealed a small and friable tissue with a yellowish color. Whereas, the second cystic mass including a mucoid substance was located in the right side of the sella turcica (Fig. 3). A grossly-total resection was performed.

Histopathological findings confirmed two different tumors. First, the yellowish and friable tumor demonstrated as a pituitary adenoma, according to the typical findings of the hematoxylin and eosin (H&E) stain (Fig. 4A). Immunohisto-
Fig. 1. Preoperative CT. A: Enhanced brain CT showing a thinned sellar floor and an intrasellar cystic mass compressing the right cavernous sinus without calcification. B: Sagittal images revealing the normal parasellar bony abnormalities.

Fig. 2. Preoperative MRI. Brain MRI revealing the intrasellar cystic tumor with a well-enhanced rim and without invasion to the cavernous sinus. A: T2 WI coronal image. B: T1 WI coronal image. C: Enhanced T1 WI coronal image. D: Enhanced T1 WI sagittal image.
Chemical stains were negative for prolactin, adrenocorticotrophic, and growth hormones. Second, the cystic tumor demonstrated as a chordoma, according to the typical findings of the H&E and immunohistochemical stains (cytokeratin-20 and epithelial membrane antigen) (Fig. 4B, C, and D). Therefore, the present case suggested a coincidental chordoma and pituitary adenoma.

Immediately postoperative MRI demonstrated complete tumor removal. After surgery, the diplopia disappeared, and she did not complain of any symptoms. Pituitary hormone function test also revealed normal levels. Additional treatment, including radiosurgery or proton beam radiotherapy for chordoma, was considered.

**DISCUSSION**

Chordomas are slow-growing malignant bone tumors, and are thought to arise from notochordal cell rests. Their incidence is rare, and they represent approximately 1% of all malignant bone tumors and 0.2% of intracranial tumors [1]. Their location is usually related to the midline, and approximately 50% develop in the sacrococcygeal region, while 35% develop in the sphenoid region and 15% in vertebrae [9].

Most chordomas involving the sellar region likely originate from parasellar or suprasellar bones [2]. Although the incidence of intrasellar chordoma is extremely rare, several studies to date have reported intrasellar chordomas, such as chordomas mimicking pituitary adenomas [1, 3-6]. In addition, a chordoma associated with a pituitary adenoma has been twice reported [7, 8]. The first case was a parasellar chordoma combined with a non-functioning pituitary adenoma, and the second case was a clival chordoma associated with a prolactinoma. Therefore, the present case may be the third case of an intrasellar chordoma associated with a coincidental pituitary adenoma.

Radiologically, chordomas are characterized by a solid type, and appear hyperintense on T2-weighted MRI. And, a CT scan may show a destructive and lytic lesion [2, 7]. However, several researchers have reported chordomas during their early development stage and intrasellar chordomas unassociated with sphenoid bone involvement [3, 6, 7]. Similarly, the present case revealed an intrasellar chordoma without sphenoid bone involvement.

In conclusion, we report the third case of an intrasellar chordoma with a pituitary adenoma. Although it may be difficult to distinguish a cystic chordoma without bone involve-
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ment from other intrasellar tumors, such as a pituitary adenoma, a craniopharyngioma, or a Rathke’s cleft cyst, the present case illustrates that the co-occurrence of two tumors in the sellar region is possible.

Conflicts of Interest

The authors have no financial conflicts of interest.

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