A Primary Sellar Neuroblastoma Mimicking a Pituitary Adenoma: A Case Report
뇌하수체 선종으로 오인되었던 원발성 터키안 신경모세포종: 증례 보고

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INTRODUCTION

Neuroblastomas are embryonal tumors that arise from the migrating neuroectodermal cells derived from the neural crest (1). Intracranial neuroblastomas are uncommon and are generally located in the periventricular or supratentorial parenchyma in contrast to olfactory neuroblastomas that originate from the olfactory receptor cells in the nasal cavity (1, 2). To our knowledge, primary sellar neuroblastoma is extremely rare and only 11 cases have been reported since 1979 (1, 3).

We report a case of primary sellar neuroblastoma that showed a solitary sellar mass with supra- and parasellar extension; the tumor mimicked a non-functioning pituitary adenoma or meningioma. We describe the clinical, radiologic, and histopathologic features of our case; additionally, we discuss the current method of treatment for primary sellar neuroblastoma reported in the literature.

CASE REPORT

A 76-year-old man was admitted to our institution for evaluation and treatment of a sellar mass detected after he experienced progressive visual disturbance for 2 months. His medical history was unremarkable except for hypertension. Neuroimaging studies revealed a solitary sellar mass with suprasellar and parasellar extension that mimicked a non-functioning pituitary adenoma or meningioma. The tumor was excised by transsphenoidal resection. Histopathologic analysis revealed small cells surrounded by a dense fibrillary stroma as well as strong expression of neural markers. Hence, the patient was diagnosed with sellar neuroblastoma. Progesterone levels normalized in the immediate postoperative period, although visual disturbances persisted. Herein, we describe the clinical manifestations, MRI characteristics, and histopathologic findings of this case.

Index terms
Neuroblastoma
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Pituitary MRI at admission revealed a large sellar mass (3.2 × 2.5 × 3.8 cm) with supra- and parasellar extension that caused left posterolateral displacement of the pituitary stalk. The lesion showed homogeneous iso-signal intensity on T1-weighted images, high signal intensity on fluid attenuation inversion recovery images, and relatively homogeneous contrast enhancement. There was no evidence of an abnormality in the floor of the sella turcica or any nasal/ethmoidal bone lesions (Fig. 1).

The patient underwent transsphenoidal resection of the tumor. A friable yellow mass was identified upon performing a linear incision over the dura; the mass adhered to adjacent structures. A near total resection was performed, and an intraoperative frozen biopsy indicated a pituitary adenoma. However, histopathologic examination revealed that tumor lobules were separated by dense fibromuscular tissue, and neoplastic cells were small and round.
with scant cytoplasm, dispersed coarse-to-fine nuclear chromatin, and inconspicuous nuclei. Immunohistochemical results showed strong expression of neural markers including synaptophysin, neural cell-adhesion molecule (NCAM/CD56), and neuron-specific enolase (NSE) (Fig. 2). Immunohistochemical staining was negative for pituitary hormones.

The patient's prolactin levels normalized; however, there was no improvement in his visual disturbance symptoms during the immediate postoperative period. Several imaging examinations were performed to detect any primary or distant lesions; these imaging examinations included neck, abdominopelvic, and chest CT, as well as whole-body positron emission tomography-CT. No evidence of additional pathologic lesions was detected with these modalities.

**DISCUSSION**

Intracranial neuroblastomas are rare, and they commonly occur in the supratentorial parenchyma or a paraventricular region (4); primary neuroblastomas are extremely rare (5). The origin of this tumor still remains unclear, but Sarwar (5) and Roy et al. (6) suggested that these tumors arise from the ganglion of Locy which grows between the olfactory fossa and the telencephalic vesicle.
Histopathologic examination of sellar neuroblastoma typically reveals homogeneous small cells with round nuclei, obvious fibrillary stroma, and rosette or pseudorosette formations representing a “packed arrangement” of well-differentiated tumor cells. Tumors with a more undifferentiated status show anaplastic hyperchromic small cells with high mitotic activity (7). Immunohistochemical analysis of sellar neuroblastomas typically shows strong expression of neural markers such as synaptophysin, chromogranin, NCAM/CD56, and S-100 protein (7). In the present case, tumor lobules were surrounded by a dense fibrillary stroma, leading to a diagnosis of sellar neuroblastoma. Furthermore, strong expression of synaptophysin, NSE, and NCAM/CD56 was detected on immunohistochemistry.

Dupuy et al. (3) reviewed 9 patients with primary sellar neuroblastomas; their report described that most of these tumors arise primarily in women (77%) with a mean age of 40 years at diagnosis (range, 31 to 57 years). They described that the presenting symptoms were similar to those of a non-functioning pituitary adenoma (98%, bilateral hemianopsia; 66%, hyperprolactinemia; 44%, gonadotrophic insufficiency; 11%, hypopituitarism; and 22%, syndrome of inappropriate secretion of anti-diuretic hormone). Our patient was a 76-year-old man; hence, he was the oldest patient among all of the previously reported cases of primary sellar neuroblastomas and one of the infrequent cases of primary sellar neuroblastomas in males.

Radiologic findings of primary sellar neuroblastomas are typically nonspecific on CT and MRI; a sellar mass with or without supra- or parasellar extension can show variable signal intensities on T1- and T2-weighted sequences as well as inconsistent enhancement patterns after gadolinium injection. These findings mimic those of non-secretory pituitary adenoma, tuberculum sellae, or diaphragma sellae meningioma (1, 3). Intraoperative diagnosis can also be equivocal because of the extremely rare occurrence of sellar neuroblastoma and non-specificity of intraoperative findings (1).

Although surgical resection via a transphenoidal or transcranial approach is the first-line treatment, a uniform postoperative therapy has not yet been established. Dupuy et al. (3) reported the case of a patient who underwent subtotal tumor resection without adjuvant treatment, but was healthy after 3 years. Adjuvant radiotherapy was recommended in the previous studies because of the radiosensitivity of tumors (1, 5). In the present case, a transphenoidal approach was used for surgical resection; although the tumor was totally excised as per the intraoperative assessment, a follow-up brain MRI performed 2 months later revealed a residual tumor in the posterosuperior aspect of the sella turcica. Therefore, adjuvant radiotherapy was scheduled to prevent tumor progression or metastasis.

In conclusion, primary neuroblastomas located in the sella turcica are extremely rare and they can be difficult to diagnose via radiologic and pathologic examinations. However, primary sellar neuroblastoma should be included in the differential diagnoses of masses located in the sella turcica. Furthermore, understanding primary sellar neuroblastomas in greater detail may assist the radiologists in devising more appropriate multidisciplinary treatment strategies.

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뇌하수체 선종으로 오인되었던 원발성 터키안 신경모세포종: 증례 보고

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두개 내 신경모세포종은 드문 악성 종양으로 대개는 천막상부의 뇌실질 혹은 뇌실주변에 발생한다. 터키안에서 발생하는 원발성 신경모세포종은 극히 드물다. 저자들은 진행성 양이측반응을 호소한 76세 남자 환자의 증례를 보고하고자 한다. 프로락틴의 경미한 상승을 제외한 환자의 뇌하수체 호르몬 수치는 정상이었다. 뇌하수체 자기공명영상에서 터키안위와 터키안 주변부로의 확장을 동반한 고립성 터키안 내의 종양으로 관찰되었고 이는 비기능성 뇌하수체 선종 혹은 뇌수막종과 유사하게 보였다. 종양은 경질형동 접근으로 제거되었고 조직 검사에서 신경세포 표지자에 강한 양상반응을 보이는 치밀한 원섬유성 간질에 돌려싸인 작은 세포들로 관찰되어 터키안 내 신경모세포종으로 진단되었다. 수술 직후 프로락틴 수치는 정상화되었지만 시야장애는 지속되었다. 저자들은 본 증례의 임상적 소견, 자기공명영상의 특징과 조직병리학적 소견을 보고하고자 한다.

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