Atypical Location of Intracranial Germinoma: A Case Report

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Introduction

Intracranial germinomas are rare and account for only 0.4–3.4% of primary intracranial tumors. They predominantly develop around the onset of puberty and during the second and third decades of life and seem to be more frequent in Asia, where their incidence in children with a brain tumor reaches 2.1–9.4% [1].

These tumors usually develop in the midline structures, i.e., the pineal gland, hypothalamic region, and basal ganglia.

Herein, we report an atypical radiological presentation of intracranial germinoma, and we discuss the magnetic resonance imaging (MRI) characteristics and the diagnostic challenges of this entity.

Case Report

A previously fit and well 13-year-old boy of Russian origin presented with a 3-week history of vomiting and headaches and a 1-week history of diplopia. Physical examination disclosed he was alert and oriented. Ophthalmologic examination showed a bilateral papilledema associated with bilateral sixth cranial nerve palsy. He had no other neurological deficit.

Cerebral MRI showed bilateral frontal periventricular lesions infiltrating the corpus callosum, the floor of the frontal horns, and the anterior wall of the left frontal horn (Fig. 1). These lesions were heterogeneous both in T1- and T2-weighted sequences and presented a slight diffusion restriction. Small cystic changes were visualized. Two other lesions were also visualized in the floor of the third ventricle and in the Sylvian aqueduct (Fig. 2), generating a supratentorial hydrocephalus associated with transependymal resorption signs. A complementary MRI centered on the hypothalamic–pituitary axis showed invasion of the posterior (mamillary) region of the hypothalamus and absence of invasion of the anterior (supraoptic) and middle (tuberal) regions of the hypothalamus, the pituitary gland, and the pineal gland. MRI of the spine showed no metastatic dissemination.

Biological results showed undetectable serum and cerebrospinal fluid (CSF) levels of alpha-fetoprotein and human chorionic gonadotropin. There were no tumoral cells in the CSF. Hormonal assessment showed decreased adrenocorticotropic hormone levels (2.4 pg/ml; normal values: 10–60 pg/ml), reduced serum cortisol value (0.4 µg/dl; normal values: 6.2–19.4 µg/dl), and low urinary cortisol excretion (4 µg/24 h; normal values: 10–85 µg/24 h). There was no deficiency among the other pituitary hormones.

Endoscopic third ventriculostomy was performed and confirmed evidence of involvement of the floor of the third ventricle. Simultaneous biopsy sampling was obtained and revealed a pure germinoma (Fig. 3).
Computed tomography scan of the chest, abdomen, and pelvis and ultrasonography of the scrotum did not show any primary or metastatic lesion.

The patient was treated with radiotherapy alone. Radiation therapy consisted of 24 Gy to the craniospinal axis, followed by a 16 Gy boost to the tumor bed and the intracranial metastasis, thereby increasing the tumor bed and metastasis dose to 40 Gy. The MRI performed 1 month after the end of radiotherapy showed complete resolution, and the patient remained stable without evidence of recurrent disease 14 months after completion of radiotherapy.

**Discussion**

The main distinguishing radiological features of pure germinomas relate to their typical sites of involvement. They are indeed most often located in the midline structures, i.e., the pineal gland, hypothalamic region, and basal ganglia. Approximately 30% of them are bifocal, involving both the suprasellar and pineal region.

Other sites have rarely been described, including the corpus callosum [2], the corona radiata [3], and the frontal lobe [4]. The incidence of metastatic intracranial germinomas is not well established and varies largely among series. In a series of 23 patients [5], 34% were found to have CSF seed-
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Tions illustrate the importance of a comprehensive clinical and biological assessment of the hypothalamic–pituitary axis in complex cases.

In the presented case, most parts of the lesions were intraparenchymal, and there was no involvement of the pituitary gland and the pineal gland, making the radiological findings nonspecific. The differential diagnosis of periventricular tumors in children and adolescents comprises ependymoma, lymphoma, supratentorial primitive neuroectodermal tumor (PNET), and glioblastoma. Typically, ependymoma shows heterogeneous enhancement with cystic areas; lymphoma in immunocompetent patients presents as a homogeneously enhancing parenchymal mass; supratentorial PNET is heterogeneous with moderate gadolinium enhancement; and glioblastomas often contain areas of necrosis. Nevertheless, these criteria are not specific. Furthermore, advanced imaging such as diffusion and spectroscopy does not reliably differentiate these different tumor types.

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Conflict of Interest  On behalf of all authors, the corresponding author states that there is no conflict of interest.

References


Fig. 3 Photomicrographs of histological findings (original magnification × 400). Hematoxylin and eosin stain (a) showing two-cell pattern of germinoma with small lymphoid elements and large poorly differentiated cells with abundant cytoplasm and round nuclei. Immunohistochemical stainings of CD45 (b) revealing positive lymphocytes and of OCT4 (c) revealing positive tumor cell nuclei.