Lymphoplasmacyte-Rich Meningioma of the Third Ventricle: Case Report

Meningioma linfoplasmocítico do terceiro ventrículo: relato de caso

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Abstract

Introduction Meningiomas are the most common primary intracranial tumors, accounting for up to 35% of the neoplasms in this category. Approximately 10–20% of these neoplasms are histologically atypical, and the lymphoplasmacyte-rich meningioma (LPM) corresponds to a very rare subtype of meningioma that is characterized histopathologically by massive infiltrates of inflammatory cells. The case described in the present study is the sixth case of an intraventricular LPM found in the literature and the first case considering the location in the third ventricle.

Case Description A 21-year-old male without previous comorbidities sought medical attention due to visual impairment (complaining of intermittent visual blur) for 2 months. A magnetic resonance imaging of the brain confirmed the presence of a well-delimited solid mass in the third ventricle of 3.0 × 2.3 cm with a cystic component that extended itself inferiorly and distorted the visual pathway anatomy. Neurosurgeons decided to access the lesion using an interhemispheric transcallosal approach with a transfornaminal access, and the lesion was resected completely. The patient has an ambulatorial endocrinological follow-up and is neurologically stable 6 months after the procedure. No new visual deficits were noted.

Conclusion Lymphoplasmacyte-rich meningioma is a very rare intracranial tumor, and the involvement of the third ventricle make this case unique.

Keywords
► intracranial tumor
► meningioma
► lymphoplasmacyte-rich meningioma
► intraventricular tumor
► neuro-oncology

Resumo

Introdução Os meningiomas são os tumores intracranianos primários mais comuns, responsáveis por até 35% das neoplasias nessa categoria. Aproximadamente 10–20% desses tumores são histologicamente atípicos, e o meningioma linfoplasmocítico (LPM) corresponde a um subtipo muito raro que é caracterizado histopatologicamente por infiltrados maciços de células inflamatórias. Este é o sexto caso encontrado na literatura de uma LPM intraventricular e o primeiro caso considerando a localização no terceiro ventrículo.

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Introduction

Meningiomas are the most common primary intracranial tumors, accounting for up to 35% of the neoplasms in this category.\(^1\)\(^2\) Their origin is from arachnoid caput cells and, of the 16 subtypes recognized by World Health Organization (WHO) classification of tumors of the nervous system, the meningothelial, fibrous and transitional meningiomas are the most common.\(^3\) Approximately 10–20% of these neoplasms are histologically atypical, and the lymphoplasmacyte-rich meningioma (LPM) corresponds to a very rare subtype of meningioma that is characterized histopathologically by massive infiltrates of inflammatory cells, such as lymphocytes and plasma cells.\(^3\)\(^4\) Fewer than 110 cases of LPMs have been reported in the literature, most of which were isolated case reports.\(^5\)\(^6\)

Regarding topography, most meningiomas develop in the cerebral convexity, parasagittal, or in the sphenoid ridge, and the LPMs follow this pattern of distribution. Intraventricular cases are considered very rare and occur in 0.5 to 3% of all meningiomas.\(^7\)

The authors report the sixth case found in the literature of an intraventricular LPM (first case considering the third ventricle location) and describe the clinical and histopathological features of this rare pathology.

Case Report

A 21-year-old male, without previous comorbidities, sought medical attention due to visual impairment (complaining of intermittent visual blur) for 2 months. He was referred to an ophthalmologist, who found bilateral papilledema in the fundoscopy. The patient was sent to an emergency hospital, where he performed a computed tomography (CT) of the head that showed an expansion lesion in the third ventricle.

At neurological examination, there were no other deficits, such as weakness, paresthesias, loss of the visual field or other cranial nerves disturbances. The endocrinological evaluation did not indicate any disturbance of the hormonal axes. The investigation was complemented with a magnetic resonance imaging (MRI) of the brain, which confirmed the presence of a well-delimited solid mass in the 3rd ventricle of 3.0 × 2.3 cm, with a cystic component that extended itself inferiorly and distorted the visual pathway anatomy (Fig. 1).

Due to the lesion characteristics, the hypothesis of craniohypophyseal was formulated. The patient was evaluated by the neurosurgery department, and surgical treatment was indicated due to the high risk of permanent visual deficit and deterioration by ventricular obstruction with the lesion growth.

Neurosurgeons decided to access the lesion using an interhemispheric transcallosal approach with a transforaminal access. It was noted during the intraoperative period that, in fact, the cystic component presented in the MRI was a septation of the third ventricle and was not part of the tumor itself. All the visualized tumor was resected using microsurgical technique, and tumoral material was collected and sent for histopathological analysis. At this point, the cyst was opened and its contents (motor oil aspect) were drained, decompressing the optic nerve.

The immunohistochemical and histological analysis confirmed the diagnosis of LPM, showing positive immunexpression for epithelial membrane antigen (EMA) and cluster of differentiation 68 (CD68) (Fig. 2).

After the surgery, the patient developed panhypopituitarism and diabetes insipidus, probably due to manipulation of the hypothalamus-pituitary pathways. In addition, he developed a right hemiparesis and paralysis of the left oculomotor nerve, probably due to surgical manipulation, that resolved almost completely in the postoperative period.

The postoperative MRI showed completed resection of the tumor (Fig. 3). The patient has an ambulatorial endocrinological follow-up and is neurologically stable 6 months after the procedure. No new visual deficits were noted.

Discussion

The LPM was first described by Banerjee and Blackwood, in 1971, as a “subfrontal tumor with the features of plasmacytoma and meningioma”\(^8\) and has been classified as grade I by the WHO since 1993.\(^5\) According to a more recent series of
case, PML is diagnosed more frequently between the third and fifth decade of life (earlier than in meningiomas in general). Also, in contrast with regular meningiomas, there was no gender prevalence in LPMs.

The epicenter of growth of the LPMs is similar to that found in meningiomas in general, with the cerebral convexity being the most common topography. The symptoms are associated with increased intracranial pressure (headache, nausea, vomiting, papilledema, and altered level of consciousness), and focal deficits are related to local brain compression caused by the tumor.

Due to the classification in the WHO class I, LPMs are considered benign tumors with low rates of recurrence and aggressiveness. Our case corresponded to this benign profile by presenting a low Ki67 index (less than 0.1% of the cells). Some reports in the literature point to a more malignant character of LPM, and these are associated with higher indices of the Ki67 and earlier neurological deterioration due to the infiltration of inflammatory cells in the cerebral parenchyma, but this characteristic was not noticed in the neuroimaging studies or in the transoperative scenario.

Regarding the histological characteristics, this tumor is marked by dense lymphocyte and plasma cell infiltration that occasionally overlies the meningeal epithelial cells. Immunohistochemically, expressions of certain markers, like EMA, vimentin, and CD68, are fundamental to confirm and distinguish the origin of the tumor. The differential diagnosis is important due to similar lesions that can mimic the LPM. Plasma cell granulomas (PCGs) are morphologically difficult to differentiate. Plasma cell granulomas are characterized as non-neoplastic hyperplasia, which have massive infiltration of the lymphocytes, plasmaocytes, and histiocytes. However, the typical meningioma morphology of a fibrous, spiral, or storiform shape is absent, and there is no positivity on the EMA and vimentin markers.

Examples of other neoplasms that can mimic LPM include chordoid meningioma, multiple myeloma, solitary plasmacytoma, plasma cell granuloma, giant lymph node hyperplasia, and sinus histiocytosis. Lymphoplasmacyte-rich meningiomas of the ventricular system are extremely rare, and there are only 5 cases reported in the literature. Most cases occurred in the lateral ventricle, and there is no record of this type of meningioma.

**Fig. 1** Preoperative magnetic resonance imaging of the brain. T1 weighted with contrast injection images. (A) coronal view. (B) sagittal view. (C) axial view of the solid component. (D) axial view of the cystic component. An expansive lesion enhanced by contrast is noted on the floor of the third ventricle with a cystic component that exerted mass effect and distorted the visual pathway.
affecting the third ventricle. Meningiomas of this location are a great therapeutic challenge due to the difficult surgical access as well as the risk of damaging important midline structures, such as the diencephalon.

Usually, these lesions are indicated for surgical treatment due to the rapid neurological deterioration and the risk of developing acute hydrocephalus by obstruction of the foramen of Monro. The approach method should be planned and individualized for the best optimization and effectiveness of the surgical procedure. Our case had a complex lesion involving the third ventricle and the suprachiasmatic cistern as well as an important relation with the optic pathway. Due to the location of the tumor, an interhemispheric transcallosal approach was chosen, which allowed a great visualization of the tumor and, consequently, complete macroscopic resection of the tumor. Despite the effort to preserve adjacent structures, the development of deficits due to surgical manipulation cannot be avoided.

**Conclusion**

Lymphoplasmacyte-rich meningioma is a very rare intracranial tumor, and the involvement of the third ventricle makes this case unique. The definitive diagnosis of LPM relies on
histopathological findings and total surgical excision remains as the standard treatment for this type of tumor.

Conflict of Interests
The authors have no conflict of interests to declare.

References