Lymphoplasmacyte rich meningioma: A rare variant of meningioma mimicking diffuse granulomatous disease with leptomeningeal spread

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Date of submission: 16th August 2020 Date of acceptance: 13th November 2020 Date of publication: 1st December 2020

Abstract

Lymphoplasmacyte-rich meningioma is a rare variant of World Health Organization (WHO) Grade I meningioma, which may present as diffuse granulomatous disease with leptomeningeal spread. Here, we report a case of 61 years old male who presented with gradual onset of progressive quadriperesis. Patient was initially treated with antitubercular medications as imaging studies and serum markers were suggestive of granulomatous lesion. Surgical intervention was advised as patient did not respond to medical management. Histopathology and immunohistochemistry confirmed the diagnosis of lymphoplasmacyte-rich meningioma. This meningioma can present as diffuse leptomeningeal infiltrative lesion involving both the hemispheres and all the tentorial surfaces mimicking inflammatory lesion. Tuberculosis is very common in our part and the diffuse infiltrative nature of lymphoplasmacyte-rich meningioma with similar radiological features of these two pathologies may confuse and delay the diagnosis. Recognition of this entity is of utmost importance as it guides proper management.

Key words: meningioma, lymphoplasmacyte rich meningioma, granulomatous disease

Introduction

Meningiomas are divided into three grades and 15 subtypes according to the 2007 WHO classification of tumors of CNS.¹,² Lymphoplasmacyte Meningioma (LPM) was first reported by Banerjee and Blackwood in 1971.³,⁴ It is a Grade I subtype of intracranial meningiomas which has been adapted to the WHO classification since 1993, LPM is characterized by exuberant lymphoplasmacytic inflammatory cell infiltrate and is one of the rarest variants of meningioma. Lymphoplasmacytic meningiomas occur most commonly over cerebral convexities. Other sites include sphenoid ridges, olfactory grooves, parasellar regions, petrous ridges, tentorium and posterior fossa.⁵ Because of its atypical histology and clinical presentation, some investigators have argued against diagnosis of this particular type of tumor as a meningioma.⁶,⁷ Meningiomas presenting as tumors with the features of a LPM are very unusual.⁸-¹² The pathophysiology associated with this form of tumor is poorly understood. We report a case of LPM in a 61 years old male patient, surgical resection of the lesion was performed and the histopathological diagnosis was made.
Case report

A 61 years old male presented with gradual onset weakness of bilateral upper and lower limbs. Initially the weakness started from right lower limb then gradually progressed to involve all the four limbs over the period of 3 months. The weakness was associated with paresthesia with inability to perform his daily activities on his own with Modified Rankin Scale (MRS) of 4. His upper limb power was right 2/5, left 3/5 and lower limb bilateral 3/5 according to Medical Research Council (MRC) power grading system. He was a known case of hypertension for 30 years and Type 2 Diabetes Mellitus for 10 years under medication. Nerve conduction studies ruled out neuropathy. Lumbar puncture was done which showed lymphocyte predominance and high protein levels. Therefore, a provisional diagnosis of tuberculosis was made and patient was started on Anti Tubercular Therapy (ATT) along with steroid. However, no clinical improvements were seen and repeat CSF examinations showed no significant findings. The MRI brain showed T2 and FLAIR high signal in bilateral frontal lobes (Figure 1A, 1.B) and post contrast MRI showed patchy enhancing homogenous plastering lesion along the skull base meninges, falx cerebri, tentorium and encasing bilateral carotid arteries (Figure 1.C, 1.D, 1.E, 1.F). However, there was no edema surrounding the lesion.

The tissue biopsy was carried out from suprasellar area extending to the skull base and dural surface including medulla. Grossly the lesion was highly vascular, pinkish, diffuse, plastered to the surface with abnormal tissue attached to the dura and pial surface involving the optic nerve as well as carotid artery (arrow) (Figure 2A). The lesion was dissected meticulously from brain surface using dissector (arrow) (Figure 2B, 2C) and part of it was cut with micro-scissor for biopsy (arrow) (Figure 2D).

The Hematoxylin Staining (H.E) x200 (Figure 3A) showed scattered small clusters of meningo-geothelial cells with characteristic whirling and indistinct cells outlines on a fibrous stroma. Prominent lymphoplasmacytic infiltrate is seen with lymphoid aggregates, increased vascularity and sheets of histiocytes. The histopathology showed lymphoplasmacytic-rich (inflammation-rich) meningioma, WHO Grade I (Figure 3). The diagnosis was further confirmed by IHC where the tumor cells were positive for vimentin (Figure 3B), EMA (Figure 3C), PR (Figure 3D), CD20 (Figure 3E), Ki67 (Figure 3F), CD138 (Figure 3G), CD68 (Figure 3H) and negative for GFAP (Figure 3I).

In our case the lesion was diffuse infiltrative and surgically inaccessible for total resection. Radiotherapy was suggested after biopsy.

![Figure 1A](image1.png)
**Figure 1A:** The T2 Weighted Coronal brain MRI showing bilateral frontal lobe signal changes. 1.B. The Axial Fluid-attenuated inversion recovery (FLAIR) image showing bilateral frontal lobe hyperintensity patches. 1.C, D, E, and F. The post Gadolinium contrast MRI showing patchy enhancement of adjacent meninges along skull base, tentorium, diffuse homogenous enhancing supresellar lesion extending to skull base and dural surface including medulla, base of bifrontal lobe, along the falx cerebri and superior to bitemporal region and encasement of bilateral carotid artery.
Figure 2A: The arrow showing grossly highly vascular, pinkish white lesion intraoperatively. B. The arrow showing plastered lesion over pial surface of brain being dissected intraoperatively. C. The arrow showing plastered lesion over pial surface and its dissection along with vessels intraoperatively. D. The arrow showing biopsy of the lesion being taken intraoperatively.

Figure 3A: The Hematoxylin Staining (H.E) x200 showing scattered small clusters of meningo-geothelial cells with characteristic whorling and indistinct cells outlines on a fibrous stroma. Prominent lymphoplasmacytic infiltrate is seen with lymphoid aggregates, increased vascularity and sheets of histiocytes. B, C, D, E, F, G, H: The Immunohistochemistry (IHC) Study showing positive for vimentin (V9), Epithelial membrane antigen (EMA), progesterone receptor (PR), CD20 staining B cells, Ki67, CD138 and CD68. I. The Immunohistochemistry (IHC) study showing negative for Glial fibrillary acidic protein (GFAP).
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Discussion

Meningiomas are the most common primary CNS tumors and accounts for approximately one-third of all primary brain and spinal tumors. Meningiomas are divided into 3 grades and 15 sub-types according to the 2007 WHO classification of tumors of central nervous system (CNS). LPM is a rare WHO Grade I subtype of meningioma.

LPM, was first reported by Banerjee and Blackwood in 1971. It has been adapted to the WHO classification since 1993. The patients with LPM have variable clinical manifestations according to the location of the tumor. The common presentations include headache, hemiparesis, seizure, vomiting, dizziness, visual disturbance, dyscalculia, dysgraphia and slurred speech. LPM is characterized by exuberant lymphoplasmacytic inflammatory cell infiltrate. It occurs most commonly over cerebral convexities. Other sites include sphenoid ridges, olfactory grooves, parasellar regions, petrous ridges, tentorium and posterior fossa. In the new WHO classification of tumors of the meninges, lymphoplasmacyte-rich meningioma, is characterized by the dense infiltration of lymphocytes and plasma cells. Banerjee and Blackwood regarded it as a collision tumor with ordinary meningioma and plasmacytoma components. On the other hand, Russell and Rubinstein expressed the opinion that the infiltration of plasma cells was of a secondary character rather than that of collision tumors.

Actually, the origin (neoplastic or inflammatory) of this tumor is unclear; its biological behavior and clinical course are anomalous so it is considered closer to intracranial inflammatory masses rather than typical meningioma. Like in our case the LPM does not have typical imaging features of a meningioma so it can mimic intracranial inflammatory condition or brain neoplasm. Intracranial plasma cell granulomas and dural plasmacytoma are often confused with meningiomas at the radiographic level. Moreover, inflammatory pseudo tumors, inflammatory fibrous histiocytomas, lymphomatoid granulomatosis and sinus histiocytosis with massive lymphadenopathy are also considered in the differential diagnosis. The tumor exhibits a positive immune-expression for EMA in a membranous pattern, a diffuse immunoreactivity for vimentin, which are similar to our case IHC findings.

Total excision is the treatment of choice in LMP but in our case we could only do biopsy of the lesion due to its unusual pattern. As the lesion was diffuse, infiltrative and surgically inaccessible radiotherapy was suggested after biopsy. It is reported that in complex skull base meningiomas, a high tumour control rate in the range of 85-100% at 5 years with a low risk of significant incidence of long-term toxicity after radiotherapy.

Our case had initial clinical and laboratory diagnosis of tubercular meningitis but on further investigations like histopathology and immunohistochemistry, we could come out with final diagnosis of LPM. LPM is rare and usually associated with a favorable outcome. Surgical resection is the treatment of choice. However, whether this entity has an inflammatory origin or a neoplastic one is still uncertain and requires research on the subject.

Conclusion

Lymphoplasmacyte–rich meningioma is rare and usually associated with a favourable outcome. However, it can be confused with inflammatory lesion which delays the proper treatment. Hence, recognition of this entity is of utmost importance as it guides proper management.

Conflict of Interest: None
Source(s) of support: None

References


