Cystic Meningioma in the Inter-Hemisferic Space Location

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Abstract

Objective: The presentation of cystic meningioma in the inter-hemispheric near the falx cerebri is uncommon. It is difficult to differentiate it from intra-axial tumors, such as gliomas. Therefore, it is likely that it is misdiagnosed as other types of brain tumors.

Methods: In this study, we reported a cystic meningioma case in the inter-hemispheric location, showing an intramural nodule on magnetic resonance imaging scans.

Results: Patient underwent surgical treatment and pathological section confirmation revealing meningioma. The patient was a middle-age woman and had been misdiagnosed as suffering from glioma followed by slight hemiparesis on the right extremities.

Conclusions: Although this is a rare case, it will be good if we always consider cystic meningioma in inter-hemispheric space when diagnosing this type of cystic lesion if the magnetic resonance imaging (MRI) shows a cystic lesion mimicking glioma image presentation.

Keywords: Cystic meningioma, interhemispheric space, magnetic resonance imaging, diagnosis

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Case

A case of 57 years-old female with weakness on the right extremities for one month followed by intermittent headache for the past three months is presented. Neurological examinations revealed hemiparesis of right motor with increased physiology reflex and also pathologic reflex. General examination showed no abnormality. CT-scan was then performed, identifying the presence of an isohypodense mass in the left frontal, with peritumoral edema. This tumor was compressing sulcy and giry. Ventric and sylvian fissures were also compressed. Cistern was opened and the midline had shifted more than 5 mm to the right (Fig. 1). The magnetic
Isohypodense Mass in Left Frontal, with Peritumoral Edema. Sulcy, Giry, Ventricle, and Sylvian Fissure was Compressed. Cistern was Opened and Midline had Shifted more than 5 Mm to the Right

Fig. 1 Isohypodense Mass in Left Frontal on T1 (*). Inhomogenously Enhanced with Positif Ring Enhanced on T1 with Contrast (**). Hyperintense Mass in Left Frontal on T2 (***) with Peritumoral Edema (****)

Fig. 2 Isohypoointense Mass in Left Frontal on T1 (*). Inhomogenously Enhanced with Positif Ring Enhanced on T1 with Contrast (**). Hyperintense Mass in Left Frontal on T2 (***) with Peritumoral Edema (****)
resonance imaging (MRI) confirmed this condition (Fig. 2). With these results, the tumor was then diagnosed as a space occupying lesion (SOL) supratentorial in left frontal due to suspected high grade glioma. Craniotomy was performed under general anesthesia. Additionally, at the time of surgery, a mass located in interhemispheric space (Fig. 3) was completely resected (Fig. 4).

The pathologic specimen was submitted for microscopic analysis. The macroscopic features showed a cystic lesion containing a dense proteinaceous liquid (Fig. 5). The confirmation of pathological section revealed that it was meningioma (Fig. 6).

Discussion

Meningiomas are generally solid tumors, and their classical appearance on CT and MRI usually leads to a correct diagnosis. Mittal et al. mentioned that meningiomas generally have the characteristic imaging appearance of a well circumscribed, solid, homogeneously enhancing extra-axial mass on both CT and MRI.

A problem with the diagnosis arises when meningiomas have cystic components that can be confused with other tumors, including glial or metastatic tumors with cystic or necrotic
changes.8 In solid meningiomas, the sensitivity of a CT-scan is virtually 100% and the specificity is 90%; however, with cystic meningiomas, the diagnosis is made pre-operatively in less than 38% of cases.2 At MRI, cystic meningiomas can be difficult to differentiate from gliomas which partially show enhancement after the injection of contrast media or metastases, due to the presence of cyst which does not enhance and also due to the focal edema. Contrast enhanced MRI can distinguish cystic walls infiltrated by tumor cells from those formed by gliotic tissue.4 Cyst formation in meningiomas is thought to be a resultant of ischemic necrosis, cystic degeneration, intratumoral hemorrhage, trapping of CSF, peritumoral edema into cyst, active secretion by tumor cells, or due to glial reaction and transudation.1 The most common location of cystic meningiomas is in the cerebral convexity, particularly in the

Fig. 5 Macroscopic Features Showed a Cystic Lesion Containing a Dense Proteinaceous Liquid

Fig. 6 Pathological Anatomy Revealed a Meningioma
frontal and parietal regions. The cerebral falx is the second most frequent location. In this case, location of cystic meningiomas in interhemispheric space make this case very rare.

Cystic meningiomas have been classified into types 1 to 4. Type I is a central intratumoral cyst, type II is a peripheral intratumoral cyst, type III is a peritumoral cyst in the adjacent brain parenchyma, and type IV is a cyst between the tumor and adjacent brain parenchyma. Based on the fact that the cysts were intratumoral, peritumoral with/without tumor lining cyst wall, and that the cyst is at the brain-tumor interface, the present case was of type 2.

In terms of therapy for cystic meningiomas, previous studies have shown that total removal of tumor is the key for preventing recurrence. In this present case, the tumor was completely resected. The aim of this report was to highlight an extremely rare location of cystic meningioma for future differential diagnosis.

Imaging studies, including computed tomography and magnetic resonance imaging, had demonstrated a round, enhancing anterior interhemispheric mass, appearing to arise from the falx and caused mild mass effect. The patient underwent a craniotomy and removal of the anterior mass without complication, and her headache and hemiparesis resolved. A pathological examination of the specimen confirmed the diagnosis of meningioma, not glioma.

References