Original Article

Meningioma - A common tumor with diagnostic challenge

Authors

Pranita Mohanty¹, Anima Hota², Sunil Patra³, Shirin Dasgupta⁴

¹Prof. Pathology, Dept. of Pathology, IMS & SUM Hospital
²Asst. Prof. Pathology, Dept. of Pathology, IMS & SUM Hospital
³Prof. Neurosurgery, IMS & SUM Hospital
⁴Tutor, Dept. of Pathology, IMS & SUM Hospital

IMS & SUM Hospital, S’O’A (Deemed to be University), Kalinga Nagar, Bhubaneswar, Odisha

Bhubaneswar

*Corresponding Author

Dr Anima Hota

Asst. Prof. Pathology, Dept. of Pathology, IMS & SUM Hospital, K8, Kalinga Nagar, Bhubaneswar, Odisha

Abstract

Background: Meningioma are generally solid tumor. Rarely they are associated with cysts and mistaken as glial or metastatic tumor with cystic or necrotic changes. It is easily recognized with modern imaging techniques.

Method: A prospective study of 21 meningioma cases was conducted out of total 98 intracranial CNS tumors within a period of 2& ½ yr i.e., from Sept 2015 to Feb 2018. Cases were studied with clinical findings, radiological appearance and reviewed with histopathological examination using routine haemotoxylene and eosin stain, also immunohistochemical whenever necessary.

Results: Of the 21 cases of meningioma; were presented mostly in 5-6 decade with female predominance. Histopathological diagnoses were found to be meningothelial meningioma, transitional meningioma with extremely rare forms like intraparenchymal cystic meningioma, chordoid meningioma, microcystic meningioma & NF2 associated Schwannoma and meningioma.

Keywords: Intraparenchymal, cystic Meningioma, Intratumoural Cyst, MRI.

Introduction

Meningiomas are common tumours of the central nervous system which counts for 15 % of all intracranial tumours and are most common extra-axial benign neoplasms¹. Malignant meningiomas are relatively infrequent. Typical imaging characteristics include a well circumscribed, homogenously enhancing extra-axial mass on both CT and MRI¹. However intraparenchymal cystic meningiomas are rare, accounting 4-7% of all meningiomas and are challenging to diagnose both radiologically and intraoperatively and may be mistaken as a glial / metastatic tumour with cystic and necrotic change/hemangioblastoma/ neuroblastoma on imaging²,³,⁴. But chordoid meningioma, an atypical meningioma though have conventional imaging picture, is found to be clinically more aggressive with higher rate of recurrence.

We studied clinical presentations, radiological appearances and pathological characteristics of 21
cases of intracranial meningiomas out of total 98 cases of intracranial central nervous system tumors over a period of two and a half years (i.e., September 2015-February 2018). One extremely rare form encountered: that is an intraparenchymal cystic meningioma (21.4%). & Three rare types: one microcystic meningioma, a chordoid meningioma and other is meningioma with schwannoma and Neurofibromatosis 2.

Material and Methods
A prospective review of 21 cases of intracranial meningiomas that are operated in between September 2015 to February 2018 was undertaken. The clinical details, radiological appearances, operative findings & histopathological details with both routine haematoxylin and eosin and immunostained sections are elaborately studied.

Results
The male:female ratio was 1:1.3 and mean age was 51 years (25-70 years). The chief complaints were headache, hemiparesis, dizziness, seizure and slurring of speech.

Table 1: Age, sex, location and histological type distribution of meningiomas

| No. | Histological Subtype                  | No. Of Cases | Age(years) / Sex | Location                      |
|-----|--------------------------------------|--------------|------------------|-------------------------------|
| 1   | Meningothelial                       | 4            | 70 / Female      | Left Frontal Lobe             |
|     |                                       |              | 62 / Male        | Parasagittal region           |
|     |                                       |              | 45 / Female      | Frontal Region                |
|     |                                       |              | 58 / Female      | Sphenoid ridge                |
| 2   | Chordoid                             | 1            | 63 / Male        | Lt. Temporo parietal region   |
| 3   | Microcystic                          | 1            | 45 / Male        | Left Frontal lobe             |
| 4   | Transitional                         | 3            | 43 / Male        | Frontal lobe                  |
|     |                                       |              | 55 / Female      | Parietal lobe                 |
|     |                                       |              | 50 / Female      | Lt Frontal lobe               |
| 5   | Psammomatous                         | 3            | 48/Female        | Parasagittal region           |
|     |                                       |              | 51/Male          | Front parietal Lobe           |
|     |                                       |              | 58/Female        | Frontal lobe                  |
| 6   | Cystic (Transitional)                | 1            | 41/Female        | Fronto parietal lobe          |
| 7   | Anaplastic                           | 1            | 44/Female        | Frontal lobe                  |
| 8   | Clear cell                           | 1            | 54/Female        | Parietal lobe                 |
| 9   | Fibrous                              | 2            | 56/Male          | Cerebellar cortex             |
|     |                                       |              | 25/Female        | Parietal lobe                 |
| 10  | Angiomatous                          | 1            | 61/Female        | Tentorial region              |
| 11  | Papillary                            | 2            | 49/Male          | Supratentorial region         |
|     |                                       |              | 62/Male          | Left temporal region          |
| 12  | NF2 associated Meningioma (Transitional type) and Schwannoma | 1 | 33/Male | Left Frontal and Temporal lobe |

Out of total 21 meningiomas 17 cases were the classical histological variants (Fig. 1). There were 3 rare histological forms of meningioma which has been extensively studied along with an extremely rare form of intraparenchymal cystic meningioma which has been elaborately reviewed with the literature.

The sites were in different parts of the extra axial locations as cited (Table 1), except cystic meningioma which was intra axial in location. The mean duration of complain was 1.8 months (2 days-6 months). Preoperatively all were diagnosed as meningioma on imaging (CT/MRI) as per its classical description whereas the cystic type were misdiagnosed as ganglioglioma/DNET.

Illustrative Cases
Case 1
A 50 year old female presented with right hemiparesis for 1 month. In the MRI it was noted as a well defined intra axial complex solid cystic mass lesion in left frontal lobe closely abutting...
falx and SGS having enhancing solid component as well as cystic areas within. The lesion was hypointense in T1W and hyperintense in T2W. It measured 6x6x5.5 cm. The lesion was having significant mass effect causing contralateral shift to midline of about 2 cm, subfalcine herniation and compression of ipsilateral ventricles. There was mild surrounding edema seen as well [Fig. 2].

Operative finding - Left frontal SOL craniotomy and tumour exenteration was done. Greyish white intraaxial soft to firm moderately vascular lesion excised in piece meal and submitted for histopathological study [Fig. 3]. Routine histopathology with Hematoxyline and eosin revealed meningothelial cells with complex cystic pattern arranged both in whorls and fascicles which was diagnosed as Transitional meningioma with intra tumoral cysts [Fig 4]. Which showed immunoreactivity towards EMA and PR [Fig. 5].

Case 2
A 63 year old male presented with Right hemiparesis with features of raised intracranial pressure for last 10 days. His CECT brain revealed rim of extradural haemorrhage (1.4 cm thickness) noted in left temperoparietal convexity. Edema and gyral swelling in the left temperoparietal lobe with significant mass effect & midline shift, infarct. MRI showed Left temperoparietal convexity Meningioma with perilesional edema, mass effect and midline shift. Operative finding and postoperative follow up revealed Craniotomy with evidence of pneumocephalus/small extradural hematoma and soft tissue edema in left side without any residual enhancing soft mass seen in post CT.

The tumour mass was submitted, which revealed chordoid meningioma (grade 2) in histopathology with features reminiscent of chordomas. Tumor was composed of cords & nests of eosinophilic vacuolated cells with an abundant myxoid matrix and focal areas of typical meningothelial pattern [Fig 6A]. Immunohistochemistry showed EMA and Vimentin positivity. Patient was followed up for next 6 months without any recurrence.

Case 3
A 45 year old male presented with tonic clonic seizure attack with mild dizziness within 3 month whose CT showed low density with intense homogenous enhancement in left frontal area and MRI revealed low intensity with solid component & T2 with very high signal. Craniotomy followed by Histopathological diagnosis was Microcystic variant of meningioma. (Grade 1) [Fig 6B]. This variant is an unusual histological subtype of meningioma & was challenging for its morphology revealing extensive microcyst formation though the clinical and neuro-imaging features were identical with those of meningioma in general. Diagnosis was confirmed by PR & EMA immune positivity.

Case 4
A 33 year old man admitted to the hospital for an attack of generalized seizure with features of NF2 with similar history in other 2 siblings. There was no associated vomiting, visual difficulty. CT-revealed two mass lesions; one at temporal, other one at frontal location of left side. Surgical removal of both the mass lesions & then histopathological diagnosis-revealed Transitional type of meningioma (frontal lobe) & Schwannoma (temporal lobe). Patient being followed of for last 10 months without any neurological deficit.
Figure 1: Distribution of Histological types of meningioma cases

Figure 2: Image of Intraparenchymal cystic meningioma showing cystic structures (Arrow marked) within the brain tissue

Figure 3: Gross picture of intraparenchymal cystic meningioma showing cystic structures (Arrow marked) within the brain tissue
Figure 4: Photomicrograph of Transitional Meningioma (100x, H & E)

EMA POSITIVITY               PR POSITIVITY

Figure 5: Photomicrograph showing Immunohistochemistry of Fibroblastic Meningioma (100x)

Figure 6: A: Photomicrograph of choroid Meningioma showing abundant extracellular mucin (arrow marked) (400x, H & E). B: Photomicrograph of microcystic Meningioma showing numerous microcysts (arrow marked) with vacuolated cells (400x, H & E)
Discussion

Meningiomas are most common extraaxial neoplasms, accounting approximately 15% of all intracranial tumours\(^1\). Affects mostly adults, women almost twice as that of men. Most meningiomas are benign and grow slowly; arise from arachnoidal cells located anywhere in the brain and spinal cord and about half of them arise over the cerebral convexities and one fifth at the sphenoid ridge.

WHO classifies and grades meningiomas into many different subtypes based on histological parameters. Of these subtypes meningothelial, transitional, fibroblastic ones are most common. Some histological types such as papillary, chordoid, rhabdoid and clear cell meningioma have a more aggressive behavior and high recurrence rate.

The etiology of meningioma is not entirely known though these tumours are occasionally seen at locations of prior trauma/radiation\(^5\). Genetic alterations, including inherited mutation of NF 2 in the setting of neurofibromatosis type 2 and loss of chromosome 22q in sporadic cases are associated with a higher incidence\(^5\). Patients with neurofibromatosis type 2 have an association with increased incidence of other CNS tumours, including schwannoma and ependymomas. Our case 4 a bright example of it.

Meningiomas are uncommonly associated with cysts and their occurrence has been reported at 4-7% of all intracranial meningiomas\(^2,3\). However high incidence rate of >10% has been in record in case of children\(^2,3\). Cysts are more commonly found in males\(^6\) and are of peritumoral cyst types whereas females predominates in tumours with intratumoral cysts.

Nauta et al\(^7\) described four types of cysts associated with meningiomas 1.a centally located intratumoral cyst that is surrounded by microscopic tumour throughout, 2. A peripherally situated intratumoral cyst, 3. A peritumoral cyst that actually lies within the adjacent brain and 4. Peritumoral cyst at the interface of tumor and brain.

Cysts in meningioma is divided into intratumoral or peritumoral cysts depending on whether the cyst walls are lined by meningothelial cells\(^8\). Fortuna et al believe that intratumoral cysts develop due to microcystic degeneration, ischemic necrosis and / haemorrhage in addition to secretory changes within the tumour whereas peritumoural cysts are large, contain xanthochromic fluid with a high protein content and are lined by fibrillary astrocytes and their processes – a glial response to the presence of a meningioma\(^9,10\). From surgical point of view the Nauta type 2 cysts are important as neoplastic cells are found in the distal cyst wall. So scrupulous search should be made so as to remove all the neoplastic tissue component to prevent recurrence.

Szilwowski and Cummings studied the chemical nature an enzyme levels of 214 cerebral cysts. The protein concentration from their two meningiomas was within the range for cysts associated with malignant tumours. SGOT and LDH levels tended to be lower in the meningioma cysts and were considered to be no diagnostic\(^11\).

Our cystic meningioma (case1) though diagnosed histopathologically as Transitional Meningioma with intratumoral cyst its radiological and surgical findings were misleading and mimicked with that of glial tumor hence misdiagnosed preoperatively with a differential of Glioma/DNET. Because of its unusual appearance of no dural attachment, intraparenchymal location, complex solid cystic mass lesion (intratumoral type), it was quite challenging to be diagnosed preoperatively and established the fact that not all meningiomas are dura based.

Conclusion

Meningiomas though easily diagnosed usually by conventional radiological (CT and MRI), operative findings yet some misdiagnosis occurs due to its atypical intraparenchymal location and intracystic lesion. Which is a crucial fact that has to be always kept in mind as a potential surgically curable tumor can be prevented from recurrence
and other pattern of treatment modalities. So the role of pathology is ultimate in establishing a diagnosis by submitting a specimen for histopathological and immunohistochemical study.

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