INTRODUCTION

Myxopapillary ependymoma (MPE), a rare pathological subtype of ependymoma which shows predilection for middle-aged adults between third and fifth decades of life. It is a benign tumor that primarily occurs at filum terminale of spinal cord with a rare possibility to metastasize to the brain. Primary intracranial MPEs have been reported only rarely. Two-year-old child presented with a space-occupying lesion in fourth ventricle. Based on clinical and radiological findings, particularly owing to patient’s age differential diagnosis of medulloblastoma was considered. Biopsy followed by histopathological and immunohistochemical analysis yielded diagnosis of primary intracranial myxopapillary ependymoma.

In conclusion, MPEs can occur anywhere in intracranial space and in any age group. Despite their indolent behavior, their metastatic potential necessitates thorough investigation to rule out former before establishing diagnosis of primary intracranial MPE.

KEY WORDS: Ependymoma (MeSH); Ependymoma, Myxopapillary (MeSH); Brain Tumor (MeSH); Fourth Ventricle (MeSH); Ependymogial Cells (MeSH); Brain Neoplasms (MeSH); Pediatric brain tumor (Non-MeSH); WHO grade I (Non-MeSH); Neoplasm Grading (MeSH); Immunohistochemistry (MeSH); Children, Preschool (MeSH).


CASE REPORT

Two-year-old child presented with a space-occupying lesion in fourth ventricle. On radiological examination, MRI revealed an intraventricular mass lesion in fourth ventricle resulting in obstructive hydrocephalus. No calcifications or hemorrhage were seen. Based on clinical findings, differential diagnosis of medulloblastoma was considered (Figure 1). Biopsy was performed, and specimen sent for histopathological examination for establishment of diagnosis.

Grossly, the biopsy material comprised of multiple pale white soft tissue pieces measuring 4×3×2 cm, collectively. Microscopically, sections examined revealed neoplastic lesion exhibiting characteristic papillary architecture and cuboidal to elongated neoplastic cells with fibrillar background. Individual neoplastic cells show hyperchromatic, vesicular nuclei, small nucleoli and pale cytoplasm. Perivascular pseudo-rosette formation was also noted. Occasional mitotic figures were seen with mitotic count of 1-2 per 10 high power fields. No evidence of increased cellularity, increased mitoses or necrosis was noted. Abundant perivascular mucin with microcyst formation was noted, highlighted on special stain alcian blue (Figure 2). Focally reactive glial tissue is noted.

Immunohistochemical staining was performed and tumor cells were reactive against glial fibrillary acidic protein (GFAP), vimentin and S100. No reactivity was noted against pan-cytokeratin (Figure 3).

Based on radiological and histopathological features, diagnosis of primary intracranial myxopapillary ependymoma was established. Spinal cord MRI excluded possibility of metastasis.

DISCUSSION

Ependymomas are relatively uncommon neoplasms of ependymal cell origin constituting 6% of all gliomas, 60% of spinal gliomas, and up to 90% of primary tumors of cauda equina and filum terminale. MPE was first described as a distinct morphological variant of former by Kernohan in 1932 based on histological findings of papillary architecture, perivascular pseudorosettes and abundance of mucin.

MPEs are slow-growing, WHO grade I tumors manifesting almost exclusively as intraspinal lesions in 3rd or 4th decade of life. Slight male predilection with male-to-female ratio of 1:7:1 has been reported. In children, the neoplasm tends to behave more aggressively with greater propensity for CNS metastasis. Metastases may develop, albeit rarely, through dissemination via cerebrospinal fluid (CSF) circulation. Before diagnosis of primary intracranial MPE can be established, radiological imaging of both brain and spinal cord are mandatory.
Depression among individuals having lower limb physical disability

Figure 1: MRI scans showing intraventricular mass resulting in obstructive hydrocephalus. No evidence of calcification or hemorrhage is seen. No lesion was noted in Spinal cord.

Figure 2: Hematoxylin and Eosin (H&E) stained sections showing: A: occasional mitotic figures and fibrillary background, B: Perivascular pseudorosette formation, C: Microcyst formation, D: Acidic mucin in cystic spaces (PAS Alcian blue), and E - F: Adjacent reactive brain tissue showing foci of hemorrhage. No evidence of necrosis was seen.

Only 17 cases of primary intracranial MPEs have been reported prior to this case. Only three cases have been reported in fourth ventricle, all of which have been females with youngest age at diagnosis recorded at 7 years, which is also the youngest age in which primary intracranial MPEs have been reported up until now. Our case is first to report involvement of fourth ventricle in male patient and youngest age, i.e. 2 years.

In conclusion, MPEs can occur anywhere in intracranial space and in any age group. Despite their indolent behavior, their metastatic potential necessitates thorough investigation to rule out former before establishing diagnosis of primary intracranial MPE.

REFERENCES

AUTHORS’ CONTRIBUTIONS

Following authors have made substantial contributions to the manuscript as under:

UB & DeSK: Identification, diagnosis & management of the case, critical review, final approval of the version to be published.

HS: Diagnosis, drafting the manuscript, final approval of the version to be published.

Authors agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

CONFLICT OF INTEREST

Authors declared no conflict of interest

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