A 4-year-old boy with a ventricular mass

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1 | CLINICAL HISTORY

A 4-year-old boy was admitted to our hospital because of fever and convulsions, with a body temperature of 39.4°C. No specific abnormalities were found in immunochemistry, urine, or fecal analyzes, except for elevated serum neuron-specific enolase (21.45 ng/ml, normal range: 0–16.3). Computed tomography showed what might be an enlarged frontal boundary of the right ventricle with calcified foci. On magnetic resonance imaging, this seemed to correspond to a T1 low intensity mass, measuring 2.5 × 2.0 × 2.1 cm, with a few darker strands and no enhancement (Figure 1). A preliminary diagnosis of a low grade glial/glioneuronal neoplasm was made. The tumor was completely removed, with no complications.

2 | FINDINGS

Histopathological examination showed a low-grade neoplasm composed of oligodendroglial-like tumor cells with round to ovoid nuclei and scant cytoplasm immersed in a mucus-rich extracellular matrix. Focal microcalcifications were also observed. No mitoses, microvascular proliferation or necrosis were observed (Figure 2A, Box 1). Immunohistochemical staining demonstrated Olig2 (Figure 2B), Sox10 (Figure 2C), GFAP (Figure 2D), ATRX (Figure 2E), and nestin (Figure 2F) expression.

Jun Zhou and Kexuan Qu are contributed equally to this work.

Access the whole slide scan at http://image.upmc.edu:8080/NeuroPathology/BPA/BPA-22-02-047.svs/view.apml?

FIGURE 1 Axial T1-weighted contrast enhanced magnetic resonance imaging shows the low intensity mass was located on the front edge of the right brain ventricle, with a few darker strands and no enhancement (arrowhead).
Myxoid glioneuronal tumor (CNS WHO grade 1).

Targeted next-generation sequencing analysis using a 599 gene panel revealed PDGFRA p.K385L (rate: 38.76%) and NSD2 p.L581H (rate: 2.02%) mutations.

Myxoid glioneuronal tumor (MGNT) is a newly defined glioneuronal tumor in the 2021 WHO Classification of Tumors of the central nervous system, usually found in the septal area (nucleus accumbens or cavum septum pellucidum) and sometimes in the surrounding white matter or corpus callosum. MGNT can occur across a wide age range (6–65 years) [1], our patient, 4 years old, being the youngest patient reported at present. Clinical manifestations are variable, including headaches, vomiting, seizures, and behavioral disorders. In terms of its histological appearance, MGNT usually exhibits oligodendrocyte-like tumor cells immersed in a prominent myxoid matrix. Floating neurons, neurocytic rosettes, and/or perivascular neuropil resembling findings typical of dysembrioplastic neuroepithelial tumor (DNET) or rosette-forming glioneuronal tumor (RGNT) have also been observed. Mitotic activity is low or absent (<1/10 under high-magnification view). Eosinophilic granular bodies, Rosenthal fibers, and microcalcifications typical of other low grade glioneuronal tumor are generally absent. It is considered a CNS WHO grade 1 tumor [2]. Examples of this tumors were previously reported under the term “dysembryoplastic neuroepithelial tumor of the septum” [3]. In 2018, Solomon et al. described mutations at codon p.K385 of PDGFRA, typically a dinucleotide mutation at codon 385 of the PDGFRA oncogene replacing lysine with either leucine or isoleucine (p.K385L/I), which appear to be highly characteristic for this tumor [3]. MGNT lacks PIK3CA/PIK3R1 alterations or BRAF/FGFR1 mutations which are characteristic of the majority of RGNT and DNET, respectively. Copy number profile is generally balanced. MGNT has a distinct DNA methylation profile, which together with the mutational pattern may be critical in some cases in distinguishing it from other central nervous system tumors.

In our case, the diagnosis of MGNT was suspected based on its histological appearance and confirmed by the molecular analysis. Targeted next-generation sequencing analysis revealed PDGFRA p.K385L mutation. An additional gene mutation was observed in this case, namely, NSD2 p.L581H (NSD2:NM_133330: exon10:c.1742T>A:p.L581H), also referred to as the MMSET or WHSC1. NSD2 is located on chromosome pair 4 and encodes nuclear receptor-binding SET domain protein 2. Specifically, NSD2 catalyzes the methylation of the lysine site on the histone protein, thereby promoting the occurrence and progression of tumors through interactions with other proteins or regulation of target genes. However, as NSD2 rarely been reported in CNS tumors, its significance in the occurrence and progression of MGNT is unclear.

**AUTHOR CONTRIBUTIONS**

Jun Zhou and Kexuan Qu designed the study and drafted the manuscript; Yucheng Xie made the diagnosis; Yan Gao and Mengxing Lv reviewed the specimen; Ling Duan and Zhixiang A conducted the IHC and molecular
analysis; Hong-Fang Wu provided clinical data; Lin Zhang provided imaging data; all authors took part in writing the manuscript and approved the final, submitted version.

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**KEYWORDS**
corpus callosum, molecular neuro-oncology, myxoid glioneuronal tumor, *PDGFRα*

**CONFLICT OF INTEREST**
The authors declare that they have no conflicts of interest.

**ETHICAL STANDARDS AND PATIENT CONSENT**
Ethical approval was obtained from the ethical committee from the Kunming Children’s Hospital, Kunming, China.

**PATIENT CONSENT**
Written informed consent was obtained from the patient’s parents.

**DATA AVAILABILITY STATEMENT**
The data on investigation results of this case are available from the corresponding author upon reasonable request.

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**REFERENCES**
1. Lucas CG, Villanueva-Meyer JE, Whipple N, Bush NAO, Cooney T, Chang S, et al. Myxoid glioneuronal tumor, *PDGFRα* p.K385-mutant: clinical, radiologic, and histopathologic features. Brain Pathol. 2020;30(3):479–94.
2. Central Nervous System Tumours. WHO classification of tumours. 5th ed. Lyons, France: International Agency for Research on Cancer; 2021.
3. Baisden BL, Brat DJ, Melhem ER, Rosenblum MK, King AP, Burger PC. Dysembryoplastic neuroepithelial tumor-like neoplasm of the septum pellucidum: a lesion often misdiagnosed as glioma: report of 10 cases. Am J Surg Pathol. 2001;25(4):494–9.

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