Case Report

The Multifaceted Appearance of Supratentorial Ependymoma with ZFTA-MAML2 Fusion

Ming Liang Oon¹, Lutfi Hendriansyah², Patricia Diana Pratiseyo³, Eka J Wahjoepramono², Jian Yuan Goh⁴, Chik Hong Kuick⁴, Kenneth TE Chang^{4,5}, Arie Perry⁶, Char Loo Tan^{1,4}

- ¹ Department of Pathology, National University Health System, Singapore, Singapore
- ² Department of Neurosurgery, Siloam Hospital Lippo Village Karawaci, Tangerang, Indonesia
- ³ Department of Pathology, Siloam Hospital Lippo Village Karawaci, Tangerang, Indonesia
- ⁴ Department of Pathology and Laboratory Medicine, KK Women's and Children's Hospital, Singapore, Singapore
- ⁵ Duke-NUS Medical School, Singapore, Singapore
- ⁶ Departments of Pathology and Neurological Surgery, University of California, San Francisco, California, USA

Corresponding author:

 $Char \ Loo \ Tan \cdot Department \ of \ Pathology \cdot National \ University \ Hospital \cdot 5 \ Lower \ Kent \ Ridge \ Road \cdot Singapore \ {\tt 119074} \ \underline{char \ loo \ tan@nuhs.edu.sg}$

Submitted: 12 June 2021 · Accepted: 11 September 2021 · Copyedited by: Jerry J. Lou · Published: 20 September 2021

Abstract

Ependymomas are glial neoplasms with a wide morphological spectrum. The majority of supratentorial ependymomas are known to harbor *ZFTA* fusions, most commonly to *RELA*. We present an unusual case of a 9-year-old boy with a supratentorial ependymoma harboring a noncanonical *ZFTA-MAML2* fusion. This case had unusual histomorphological features lacking typical findings of ependymoma and bearing resemblance to a primitive neoplasm with focal, previously undescribed myogenic differentiation. We discuss the diagnostic pitfalls in this case and briefly review the histological features of ependymoma with noncanonical gene fusions. Our report underscores the importance of molecular testing in such cases to arrive at the correct diagnosis. Supratentorial ependymomas with noncanonical fusions are rare, and more studies are necessary for better risk stratification and identification of potential treatment targets.

Keywords: Ependymoma, ZFTA, C11orf95, MAML2, Myogenic

Introduction

Ependymomas (EPN) form a diverse group of morphologically and molecularly heterogeneous glial neoplasms [5, 14]. While the classical histological appearance of EPN is well-described, the morphological spectrum is broad and may confound diagnosis. Unusual features encountered in ependymomas include chondro-osseous elements [13, 17], neuropil-like islands [6], melanin-containing cells [4], tumor giant cells [8], neuronal differentiation [16], granular cell features [18], et cetera. Recent advances in genomic and methylome profiling have



identified distinct molecular subgroups characterized by recurrent genetic or epigenetic alterations that may better inform diagnosis, clinical outcomes and risk stratification of EPN as compared to conventional light microscopic diagnosis and histological grading [5]. The majority of supratentorial (ST) EPNs have gene fusions involving either ZFTA (zinc finger translocation associated; previously known as C11orf95) or YAP1. These gene fusions characterize the two molecular subgroups of ST-EPN: ST-EPN-ZFTA and ST-EPN-YAP1, respectively [1, 14]. Here, we present an unusual case of a ST-EPN with ZFTA-MAML2 fusion demonstrating diagnostic challenges due to primitive morphology, suggesting an embryonal neoplasm and divergent differentiation, including hitherto undescribed myogenic differentiation. Our case underscores the importance of molecular testing in such cases to arrive at the correct diagnosis.

Clinical summary

A 9-year-old boy presented with a visible, enlarging right forehead lump for 2 months. The child complained of an occasional headache but was otherwise well without any focal neurological deficits. Brain magnetic resonance imaging (MRI) revealed a well-demarcated, right frontal cystic tumor with a mural nodule, surrounding vasogenic edema and midline shift (Fig. 1A-C). The tumor showed a high choline peak and high choline:N-acetylaspartate (NAA) ratio. The child underwent craniotomy, revealing a well-demarcated bosselated tumor with a clear interface between tumor and brain (Fig. 1D-E). Gross total resection of the tumor was achieved (Fig. 1F), and the postoperative course was uneventful. He received adjuvant radiation therapy followed by temozolomide. The child remained well at follow up 9 months post-resection, with no radiological evidence of recurrence.

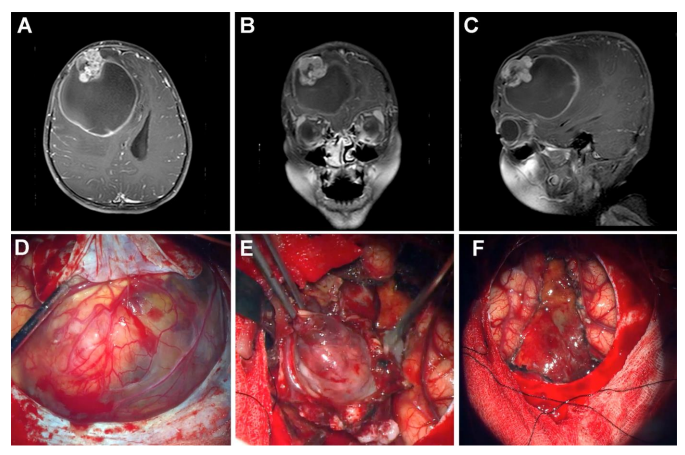


Figure 1. MRI findings of the tumor. A) Axial, B) coronal and C) sagittal views showing a cystic tumor with an enhancing multilobulated mural nodule in the right frontal lobe with mass effect. D) Intraoperative view after lifting of dural flap, prior to resection E) Dissection of a well-demarcated multinodular cystic tumor. F) Final intraoperative view after gross total resection of the solid tumor showing part of cyst lining with ventricular wall.

Material and methods

The specimen was routinely processed for paraffin embedding and staining with hematoxylin and eosin (H&E). Immunohistochemistry (IHC) with commercially available primary antibodies, including GFAP, OLIG2, synaptophysin, desmin, myogenin, MyoD1, CAM 5.2, EMA, INI1, BRG1, LIN28A, BCOR, HMB45 and Ki67, was performed following locally validated technical protocols.

Gene fusion detection was performed using a next-generation sequencing-based anchored multiplex polymerase chain reaction (PCR) assay (Archer® FusionPlex, Boulder, CO, USA) that can detect and identify gene fusions involving any of 101 covered genes, including *RELA, YAP1, MAML2, NCOA1* and *NCOA2*. Briefly, total ribonucleic acid (RNA) was extracted from formalin-fixed paraffin-embedded tissue sections of tumor. 150 ng of RNA was used for library preparation utilizing the Archer® FusionPlex

kit, according to the manufacturer's protocol (ArcherDX, Boulder, CO, USA). The prepared library was sequenced using an Illumina MiniSeq sequencer. The data obtained was analyzed by the Archer Data Analysis (version 6.2.3) portal. A confirmatory reverse transcription-PCR was designed flanking the breakpoint of the two genes and the PCR product was Sanger-sequenced.

DNA methylation analysis was performed using the Illumina Infinium Human Methylation EPIC BeadChip Kit according to published protocols [2, 7]. The results of the methylation profiling and copy number variation analysis were obtained using an automated web-based deoxyribonucleic acid (DNA) methylation profiling program (molecularneuropathology.org) [12]. The case was also integrated onto the t-distributed Stochastic Neighbor Embedding (t-SNE) analysis in the German Cancer Research Center (DKFZ) database.

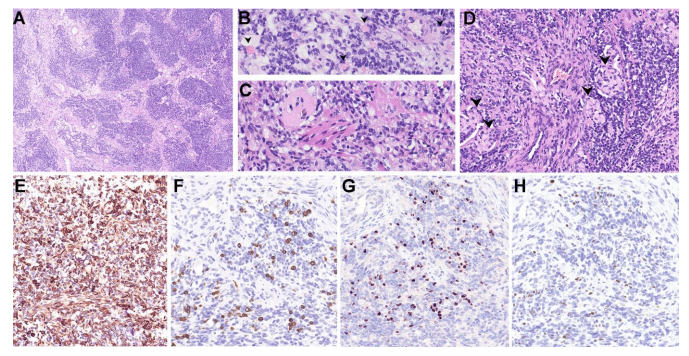


Figure 2. Light microscopy and immunohistochemical findings of the tumor. A) Low-power appearance of the tumor consisting of nests of tumor cells separated by fibrous septa, imparting a desmoplastic appearance. The bulk of the tumor cells show primitive cytomorphology with minimal cytoplasm, imparting a nondescript "small blue cell" appearance. No perivascular pseudorosettes are seen, although tumor cells aggregate along blood vessels. B) A minor subset of tumor cells showed prominent intracytoplasmic fine eosinophilic granules (arrowheads). C) Rare tumor cells showed spindled morphology, resembling skeletal muscle fibers. D) Aggregates of tumor cells with eccentric nuclei and mucinous cytoplasm (arrowheads). The tumor cells were positive for E) GFAP. Very focal areas showed myogenic differentiation, being positive for F) desmin, G) myogenin and H) MyoD1 (F-H were images taken from same area of the tumor.)

Clicking the figure will lead you to the full virtual slide (H&E).

Results

Light microscopy

Sections showed a discrete tumor composed of hypercellular sheets of small, round to ovoid cells with hyperchromatic nuclei and indistinct cell boundaries, imparting a primitive appearance to the tumor. In other areas, the tumor cells formed nodular aggregates, short fascicles and palisades/ trabeculae, separated by fibrous septa imparting a desmoplastic appearance (Fig. 2A). A minor subset of tumor cells showed prominent intracytoplasmic fine eosinophilic granules, some with a more globular appearance (Fig. 2B). A minor component of the tumor was composed of spindled cells with eosinophilic cytoplasm reminiscent of skeletal muscle fibers (Fig. 2C). Focally, aggregates of tumor cells with eccentric nuclei and mucinous cytoplasm were also identified (Fig. 2D). No other heterologous differentiation was seen. There was a distinct lack of a fibrillary background in the tumor. While some tumor cells appeared to congregate around large thickwalled vessels, no distinctive perivascular pseudorosettes were present. Geographic areas of necrosis accompanied by dystrophic calcification were present. The mitotic count reached up to 8 per 10 high power fields (3 per mm²).

The tumor cells showed patchy positivity for GFAP (Fig. 2E), and focal reactivity for synaptophysin, CAM 5.2 and EMA (cytoplasmic; no dot/ring-like pattern was found). Focally, some cells expressed desmin (Fig. 2F) and nuclear reactivity for myogenin (Fig. 2G) and MyoD1 (Fig. 2H). No neurofilament-positive axons were identified, in keeping with the non-infiltrative nature of this tumor. There was retained expression for INI1 and BRG1. OLIG2, LIN28A, BCOR and HMB45 stains were negative. The Ki67 labeling index reached up to 40%.

Gene fusion detection and confirmation

The Archer FusionPlex assay detected a ZFTA (exon 5)- MAML2 (exon 2) fusion (Fig. 3A). This gene fusion was confirmed by RT-PCR with Sanger sequencing of the amplicon product (Fig. 3B).

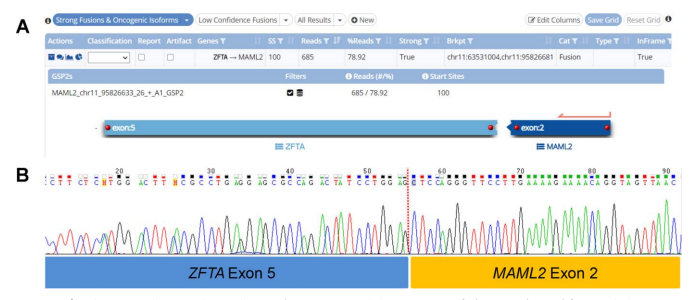


Figure 3. A) Archer FusionPlex assay detected a ZFTA (exon 5; previously known as C11orf95)-MAML2 (exon 2) fusion, which was confirmed by B) Sanger sequencing.

Methylation analysis

DNA-based methylation analysis of the tumor revealed a methylation class corresponding most closely to "Ependymoma, RELA fusion", with a calibrated score of 0.65. Concurrent copy number analysis revealed 1q gain and CDKN2A/B loss (Fig. 4A). No chromothripsis of chromosome 11 was present. On the t-SNE plot, our case localized to the methylation cluster of ST-EPN-RELA, consisting of 444 cases including our case, out of a total of 3272 cases of ependymomas of different anatomical locations and molecular groups, lending further diagnostic confidence (Fig. 4B).

A final diagnosis of ST-EPN with *ZFTA-MAML2* fusion was rendered after synthesizing the pathological and molecular findings of this case.

Discussion

ZFTA, now recognized as a key player in the oncogenesis of ST-EPN, confers transforming capabilities upon oncogenic fusion proteins by increasing their translocation to the nucleus and permitting access to the transcriptional machinery [9]. Most commonly, ZFTA is fused to RELA [9, 14, 19]. Being promiscuous, ZFTA may also rarely fuse with alternative partners such as MAML2, MAML3, NCOA1, NCOA2 and CTNNA2 [14, 18, 19]. Of note, the ZFTA-MAML2 fusion has only been identified in 18 cases of ST-EPN in the literature thus far, including the current case [14, 18, 19]. Tumors with such alternative fusion partners form satellite clusters around the previously known ST-EPN-RELA cluster in the t-SNE plot of methylation profiles and show distinct transcriptional profiles [19]. While most ST-EPN-ZFTA demonstrate classical features of EPN such as perivascular pseudorosettes and fibrillary matrix, a recent paper by Zheng et al found that EPNs with ZFTA fusions with alternative partners tend to demonstrate unusual histological features. These include sarcomatous features, areas mimicking high grade central nervous system (CNS) neoplasms like diffuse high-grade glioma, CNS embryonal tumors and other primitive tumors [19]. Such atypical histological findings are in keeping with the case we present herein.

Similar to the case by Tamai et al, our case also demonstrated granular cell features [18]. In their case, the granular cells showed eccentrically placed nuclei and cytoplasm filled with eosinophilic granules which were positive for periodic acid-Schiff (PAS) and alpha-1-antitrypsin stains. In addition, our case showed a small population of tumor cells with unequivocal expression for desmin, myogenin and myoD1, consistent with myogenic differentiation, a finding which has not been previously described. Initial diagnosis on the basis of light microscopy was difficult because of the atypical morphologic features of the tumor including the conspicuous absence of a fibrillary stroma, the absence of classical EPN morphology, and the presence of primitive-appearing cells in a desmoplastic background. Coupled with the polyphenotypic immunoprofile, which included desmin reactivity, the differential of desmoplastic small round cell tumor was initially entertained [10]. However, the subsequent molecular finding of the ZFTA-MAML2 gene fusion and the absence of an EWSR1-WT1 fusion clinched the diagnosis of ST-EPN with ZFTA-MAML2 fusion and led to revision of the histologic diagnosis. Recurrent fusions involving ZFTA are characteristic of EPN and fusions involving this gene have only been described previously in chondroid lipoma [15]. No primitive tumor has been described in the literature with this characteristic fusion to date. While lacking in histological elements of classical EPN, the circumscribed, non-infiltrative nature of this tumor and the presence of the signature fusion and methylation profile were all in keeping with a diagnosis of ST-EPN. The novel finding of myogenic differentiation, likely attributable to metaplasia of the neoplastic glial component or the mesenchymal component of the tumor cells, broadens the spectrum of heterologous differentiation in EPN [3]. The protean morphological manifestations of EPN has the potential to obfuscate morphological diagnosis, particularly when classical EPN features are not present. This underscores the limitations of morphological evaluation alone and the necessity of molecular testing.

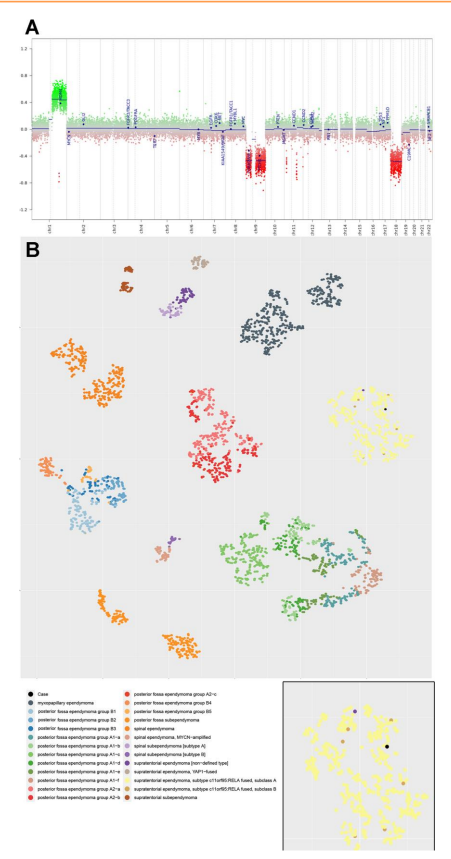


Figure 4. A) Copy number variation profile of the tumor, with 1q gain and CDKN2A/B losses. **B)** t-distributed Stochastic Neighbor Embedding (t-SNE) plot containing a total of 3272 cases of EPNs for cluster analysis. Our case localized to methylation cluster of supratentorial EPN with RELA fusion (n= 444, including our case) (Inset, case indicated in black).

There has been longstanding controversy surrounding the utility of histological grading of EPN. While ST-EPN-RELA was found to be associated with adverse patient outcomes in one retrospective study [14], this was not reproduced in another trialbased study [11]. Interestingly, mice with EPN induced by ZFTA-MAML2 fusion demonstrated a reduced survival compared to those with ZFTA-RELA fusion [19]. Out of the 18 cases of ST-EPN with ZFTA-MAML2 fusion, follow up data was only available in two cases. Both cases showed anaplastic features, corresponded to WHO grade 3. The case reported by Tamai et al. showed no recurrence for 30 months without chemoradiotherapy, which may indicate a favorable prognosis [18]. Our patient remained disease free at 9 months follow up. Owing to the paucity of survival data, it remains to be seen if differences in survival exist in ST-EPN with alternative fusions. Similarly, the utility of assignment of a WHO grade to molecularly defined EPN remains insufficiently characterized [5]. Additional studies are necessary to better delineate the behavior of these tumors, which may inform treatment decisions and risk stratification.

In conclusion, we present an unusual case of ST-EPN with *ZFTA-MAML2* fusion exhibiting primitive morphology and a focal area of previously undescribed myogenic differentiation. Awareness of the multifaceted appearance of EPN is critical and an integrated clinico-pathological-molecular diagnosis is essential for the contemporary diagnosis of this entity.

Acknowledgements

We would like to thank Drs. Felix Hinz and Felix Sahm from the German Cancer Research Center (DKFZ) for their invaluable help in getting the t-SNE data of our case.

References

- Andreiuolo F, Varlet P, Tauziède-Espariat A, Jünger ST, Dörner E, Dreschmann V, et al (2019) Childhood supratentorial ependymomas with YAP1-MAMLD1 fusion: an entity with characteristic clinical, radiological, cytogenetic and histopathological features. Brain Pathol 29: 205-216 https://doi.org/10.1111/bpa.12659
- 2 Capper D, Jones DTW, Sill M, Hovestadt V, Schrimpf D, Sturm D, et al (2018) DNA methylation-based classification of central nervous system tumours. Nature 555: 469-474 https://doi.org/10.1038/nature26000
- 3 Chakraborti S, Govindan A, Alapatt JP, Radhakrishnan M, Santosh V (2012) Primary myxopapillary ependymoma of the fourth ventricle with cartilaginous metaplasia: a case report and review of the literature. Brain Tumor Pathology 29: 25-30 https://doi.org/10.1007/s10014-011-0059-8
- 4 Chan AC, Ho LC, Yip WW, Cheung FC (2003) Pigmented ependymoma with lipofuscin and neuromelanin production. Archives of pathology & laboratory medicine 127: 872-875 https://doi.org/10.5858/2003-127-872-PEWLAN
- 5 Ellison DW, Aldape KD, Capper D, Fouladi M, Gilbert MR, Gilbertson RJ, et al (2020) cIMPACT-NOW update 7: advancing the molecular classification of ependymal tumors. Brain Pathol 30: 863-866 https://doi.org/10.1111/bpa.12866
- 6 Gessi M, Marani C, Geddes J, Arcella A, Cenacchi G, Giangaspero F (2005) Ependymoma with neuropil-like islands: a case report with diagnostic and histogenetic implications. Acta neuropathologica 109: 231-234 https://doi.org/10.1007/s00401-004-0927-y
- 7 Illumina (2019) Infinium HD Methylation AssayReference Guide.
- 8 Jeon YK, Jung HW, Park SH (2004) Infratentorial giant cell ependymoma: a rare variant of ependymoma. Pathology, research and practice 200: 717-725 https://doi.org/10.1016/j.prp.2004.08.003

- 9 Kupp R, Ruff L, Terranova S, Nathan E, Ballereau S, Stark R, et al (2021) ZFTA-translocations constitute ependymoma chromatin remodeling and transcription factors. Cancer Discovery: candisc.1052.2020 https://doi.org/10.1158/2159-8290.cd-20-1052
- 10 Lee JC, Villanueva-Meyer JE, Ferris SP, Cham EM, Zucker J, Cooney T, et al (2020) Clinicopathologic and molecular features of intracranial desmoplastic small round cell tumors. Brain Pathol 30: 213-225 https://doi.org/10.1111/bpa.12809
- 11 Merchant TE, Bendel AE, Sabin ND, Burger PC, Shaw DW, Chang E, et al (2019) Conformal Radiation Therapy for Pediatric Ependymoma, Chemotherapy for Incompletely Resected Ependymoma, and Observation for Completely Resected, Supratentorial Ependymoma. J Clin Oncol 37: 974-983 https://doi.org/10.1200/JCO.18.01765
- 12 MolecularNeuropathology.org (2020) MNP Welcome https://www.molecularneuropathology.org/mnp. Accessed 10 August 2021.
- 13 Mridha AR, Sharma MC, Sarkar C, Garg A, Singh MM, Suri V (2007) Anaplastic ependymoma with cartilaginous and osseous metaplasia: report of a rare case and review of literature. Journal of neurooncology 82: 75-80 https://doi.org/10.1007/s11060-006-9239-5
- 14 Pajtler KW, Witt H, Sill M, Jones DT, Hovestadt V, Kratochwil F, et al (2015) Molecular Classification of Ependymal Tumors across All CNS Compartments, Histopathological Grades, and Age Groups. Cancer Cell 27: 728-743 https://doi.org/10.1016/j.ccell.2015.04.002
- 15 Parker M, Mohankumar KM, Punchihewa C, Weinlich R, Dalton JD, Li Y, et al (2014) C11orf95-RELA fusions drive oncogenic NF-κB signalling in ependymoma. Nature 506: 451-455 https://doi.org/10.1038/nature13109

- 16 Rodriguez FJ, Scheithauer BW, Robbins PD, Burger PC, Hessler RB, Perry A, et al (2007) Ependymomas with neuronal differentiation: a morphologic and immunohistochemical spectrum. Acta neuropathologica 113: 313-324 https://doi.org/10.1007/s00401-006-0153-x
- 17 Salazar MF, Tena-Suck ML, Ortiz-Plata A, Salinas-Lara C, Rembao-Bojórquez D (2017) Lipomatous/Extensively Vacuolated Ependymoma with Signet-Ring Cell-Like Appearance: Analysis of a Case with Extensive Literature Review. Case Reports in Pathology 2017: 8617050 https://doi.org/10.1155/2017/8617050
- 18 Tamai S, Nakano Y, Kinoshita M, Sabit H, Nobusawa S, Arai Y, et al (2021) Ependymoma with C11orf95-MAML2 fusion: presenting with granular cell and ganglion cell features. Brain Tumor Pathol 38: 64-70 https://doi.org/10.1007/s10014-020-00388-6
- 19 Zheng T, Ghasemi DR, Okonechnikov K, Korshunov A, Sill M, Maass KK, et al (2021) Cross-species genomics reveals oncogenic dependencies in ZFTA/C11orf95 fusion-positive supratentorial ependymomas. Cancer Discov: https://doi.org/10.1158/2159-8290.CD-20-0963