Novel KRAS mutation in an unusual tectal low-grade glioma

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DESCRIPTION

An 8-year-old boy with no significant medical history or family history presented with 1–2 weeks of decreased visual acuity and blurry vision. The patient denied any headaches, vomiting, dizziness, seizures or weakness. Physical examination was significant for poorly reactive pupils, constricted bilateral visual fields and bilateral papilledema. MRI (figure 1A,B) revealed a non-enhancing tectal mass with associated obstructive hydrocephalus and a second lesion in the left middle cerebellar peduncle. MRI of the spine was normal. The patient underwent a robotic-guided biopsy of the tectal mass and endoscopic third ventriculostomy. Neuroradiological features of tectal neoplasm (figure 2A,B) demonstrated an infiltrative mildly hypercellular glial neoplasm composed of elongated glial cells with mild nuclear pleomorphism. Small collections of eosinophilic granular bodies were identified. Vascular proliferation, necrosis or increased mitotic activity were not identified on higher power. There were several neurons present throughout the biopsy indicative of infiltration. Immunohistochemistry for glial fibrillary acidic protein (GFAP) was diffusely positive. Antigen Ki-67 (Ki-67) had a focal positivity of approximately 3%. These findings were consistent with a tectal low-grade astrocytoma with histological features of pilocytic astrocytoma and evidence of infiltration (WHO Grade I–II). Next generation sequencing was significant for a Kirsten rat sarcoma viral oncogene homolog (KRAS) c181C>A variant. Variants of unknown significance in Cbl Proto-Oncogene B (CBLB) (c1877A>G) adenomatous polyposis coli gene (APC) (c5026A>G) were also detected. Following surgery, the patient had mild improvement on his bilateral papilledema. However, the patient continues with stable visual deficits and no progression of disease following 3 years of routine neuroimaging in the absence of therapy.

Tectal gliomas are rare low-grade tumours that are more often seen in the paediatric population. The most frequent clinical presentation are symptoms related to an increased intracranial pressure. Most tectal gliomas have similar histopathological features similar to pilocytic astrocytoma and diffuse astrocytoma. Immunohistochemically, tectal gliomas are characteristically diffusely and strongly positive for GFAP and Olig2, while having minimal Ki67 staining. However, tectal gliomas have a unique DNA methylation profile. Additionally, tectal gliomas have a high frequency of v-raf murine sarcoma viral oncogene homolog B1 (BRAF) mutations (KIAA1549-BRAF, BRAF V600E and BRAF T599dup), although at lower rates (especially the KIAA1549-BRAF fusion) compared with pilocytic astrocytoma. Moreover, high frequency of spot KRAS G12R mutations has also been identified and tectal gliomas with both BRAF and KRAS mutations have also been reported. Other rare KRAS mutations like E63K have also been identified. KRAS is involved in many signalling cascades (ie, Phosphoinositide 3-Kinase (PI3K)/AKT axis, mitogen-activated protein kinase (MAPK) kinase

![Figure 1](image1.png)  Neuroradiological features of tectal neoplasm with associated obstructive hydrocephalus. (A) Axial fluid-attenuated inversion recovery magnetic resonance sequencing reveals a non-enhancing tectal neoplasm (arrow) with associated acute obstructive hydrocephalus as evidenced by transependymal oedema. (B) An additional non-contiguous, non-enhancing lesion is noted in the left middle cerebellar peduncle (arrow).

![Figure 2](image2.png)  Neuropathological features of tectal glioma. (A) H&E staining demonstrates an infiltrative mildly hypercellular glial neoplasm composed of elongated glial cells with mild nuclear pleomorphism. Small collections of eosinophilic granular bodies were identified (not pictured). (B) Vascular proliferation, necrosis or increased mitotic activity were not identified on higher power (40× magnification).

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cascade and Janus kinase-signal transducers and activators of transcription (JAK/STAT3)) and is therefore involved in many cellular processes. When mutated it acquires oncogenic properties that allow for perpetual activation of the RAS molecules, leading to unregulated cellular proliferation. Although KRAS mutations have been implicated in numerous cancers and is the most frequently involved MAPK oncogene across all cancer types, KRAS mutations are infrequent in paediatric central nervous system (CNS) tumours.

While the ideal approach to KRAS mutant targeted therapy for glioma is not clear, consideration of downstream MAPK inhibition may be worthwhile.

The KRAS c181C>A mutation found in our patient has not been reported to the best of our knowledge and it is unclear what role this mutation or the other mutations identified in CBLB and APC play in tumour growth. Our findings add to the genetic diversity of paediatric tectal glioma.

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**REFERENCES**


