High-Grade Gliomas in Children with Neurofibromatosis Type 1: Literature Review and Illustrative Cases

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High-Grade Gliomas in Children with Neurofibromatosis Type 1: Literature Review and Illustrative Cases

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ABSTRACT

SUMMARY: High-grade gliomas in patients with neurofibromatosis type 1 are rare and may therefore not be considered in the differential of brain lesions. Here, we describe 5 children with neurofibromatosis type 1; four of them developed various types of high-grade gliomas. The fifth patient had imaging features concerning for a high-grade lesion, but tissue diagnosis showed a low-grade glioma. The cases and literature summary provided here are to raise awareness for the occurrence of high-grade gliomas in children with neurofibromatosis type 1 and the limited ability of imaging features alone to predict a high-grade malignancy.

ABBREVIATIONS: HGG = high-grade glioma; LGG = low-grade glioma; NF1 = neurofibromatosis type 1

Brain tumors associated with neurofibromatosis type 1 (NF1) are typically low-grade gliomas (LGGs), such as pilocytic astrocytomas.1 The mutation of neurofibromin, a protein that regulates growth and the proto-oncogene Ras, causes unregulated cell growth and can result in formation of either LGGs or high-grade gliomas (HGGs).2 The incidence of HGG in children with NF1 is low and ranges from 0.28% to 5%.3-5

Using MR imaging for differentiating LGG from HGG can be problematic. While typical imaging features of LGG include lack of contrast enhancement, lack of restricted diffusion, and lack of hyperperfusion,6-9 the lack of MR imaging contrast enhancement in supratentorial brain tumors does not always equate with low-grade malignancy.10-12 In addition, contrast enhancement is a typical imaging feature of pilocytic astrocytoma, a common low-grade tumor in NF1.10,13 Thus, tissue sampling may become necessary for some patients.14

Here, we present several pediatric patients with NF1 with suspicion for HGG based on imaging, along with clinical features and neuropathology outcomes. Our goal was to raise awareness of HGG in children with NF1 and to facilitate appropriate management, including tissue sampling.

MATERIALS AND METHODS

The study received an exempt status by the institutional review board at Children’s Healthcare of Atlanta and was Health Insurance Portability and Accountability Act–compliant. mPower™ (Nuance, Burlington, Massachusetts) was used to identify brain imaging studies performed during an 18-month period (January 1, 2015, to June 30, 2016) at our institution. Of 176 unique patients with NF1, five had concerning imaging findings.

Case Series

Patient 1. A 7-year-old boy with no known medical history presented to an outside institution with a 1-month history of roving eye movements with upgaze palsy, ataxia, and headaches. An outside MR imaging at baseline demonstrated a 2.6 × 2.2 × 2.5 cm pineal region mass (Fig 1) with restricted diffusion (Fig 1C, -D), loss of the NAA peak, and elevated choline on MR spectroscopy. Tissue sampling at an outside facility showed a low-grade pineal parenchymal tumor. A 6-week MR imaging follow-up showed lesion growth to 2.9 × 2.5 × 2.9 cm, which was localized to the tectal plate rather than the pineal gland, and interval development of an irregular and centrally necrotic–appearing area of contrast enhancement (Fig 1B). The patient was diagnosed with NF1 on admission in accordance with diagnostic criteria.15 Subtotal resection of the mass was performed, and pathology was now read as consistent with a midline, H3 K27M–mutant glioblastoma multi-forme (On-line Table). When we compared it with the initial
pathology, the current specimen displayed a different histomorphology with necrosis and vascular proliferation, which could have been related to intratumoral heterogeneity or possibly secondary to malignant transformation of the tumor. After a 6-week course of focal radiation therapy, the patient is currently stable clinically and by imaging surveillance.

Patient 2. An 11-year-old girl with no known medical history presented with a 1-month history of left upper extremity weakness, left foot drop, facial droop, mild dysarthria, headache, and intermittent nausea and vomiting. MR imaging demonstrated a peripherally enhancing heterogeneous mass containing blood products centered within the right thalamus (Fig 2A). The patient was diagnosed with NF1 on admission in accordance with diagnostic criteria.15 Tissue sampling showed histomorphology consistent with a grade II–III astrocytoma (On-line Table). However, the amount of Ki-67/MIB1 staining was elevated at 10%, and H3 K27M immunostaining was strongly positive, consistent with a diffuse midline, H3 K27M–mutated glioma (World Health Organization grade IV). After chemotherapy induction, follow-up MR imaging demonstrated tumor progression and aggressive features consistent with a high-grade malignancy (Fig 2B). Despite aggressive treatment, the patient’s tumor progressed and the patient died 17 months after the initial diagnosis.

Patient 3. An asymptomatic 10-year-old boy with a diagnosis of NF1 in accordance with diagnostic criteria15 and previously treated optic glioma underwent a routine brain MR imaging surveillance scan, and a tiny T2 hyperintense focus along the posterior aspect of the right hippocampus was found (Fig 3A). Follow-up MRIs performed 6 and 12 months later demonstrated a progressive increase in size, new heterogeneous contrast enhancement (B, arrow), and new restricted diffusion (Fig 3C, -D). After gross total tumor resection, neuropathology revealed a high-grade pleomorphic glioma, with imaging and histologic features consistent with a pleomorphic xanthoastrocytoma (On-line Table). Due to the high-grade features of this tumor, he was treated with
focal proton radiation therapy. The patient is currently active and asymptomatic 18 months after completion of proton radiation.

**Patient 4.** A 4-year-old boy with a history of “skin spots” presented to our institution with headache, nausea and vomiting, weight loss, fatigue, and forehead swelling for 1 month. A brain MR imaging revealed a heterogeneous and partially enhancing mass measuring $3.5 \times 2.3 \times 2.1$ cm in the midbrain tectum with compression of the cerebral aqueduct leading to severe obstructive hydrocephalus (Fig 4A). The patient was diagnosed with NF1 on admission in accordance with diagnostic criteria. Tissue biopsies revealed a diffuse, midline World Health Organization grade IV glioma with immunohistochemistry positive for nuclear $H3 K27M$–mutant protein, $\alpha$-thalassemia/mental retardation syndrome X-linked (ATRX), and wild type $p53$, with an MIB-1/Ki-67 index of 10%. However, OncoScan (https://www.thermofisher.com/order/catalog/product/902695) copy-number microarray only revealed a $PIK3CA$ p.11047R point mutation, but no copy number changes. Given that the molecular findings were inconsistent with a high-grade malignancy, the initial plan was to follow the patient clinically and by MR imaging. On follow-up MR imaging at 3 and 6 months, the lesion remained relatively stable in size but developed multiple peripherally enhancing components within the mass (Fig 4B). The patient experienced intratumoral hemorrhage, and MR imaging at that time showed evidence of tumor dissemination. The patient was started on chemotherapy with carboplatin and vincristine with a plan for re-evaluation for evidence of tumor progression by MR imaging after the induction phase of chemotherapy.

**Patient 5.** A 7-year-old boy with a medical history of migraines and slowed speech presented to an outside institution with a 2-month history of clumsiness, truncal ataxia, worsening headache, nausea and vomiting, somnolence, and worsened slowed/slurred speech. MR imaging demonstrated a robustly enhancing suprasellar mass and a peripherally enhancing mass centered within the left globus pallidus (Fig 5A), an infiltrating T2 hyperintense signal with punctuate foci of enhancement in the right basal ganglia, and enlargement of the optic nerves, chiasm, and optic radiations. The constellation of findings was most compatible with an infiltrating optic pathway/hypothalamic glioma. The patient was diagnosed with NF1 on admission in accordance with diagnostic criteria. Tissue sampling showed a high-grade infiltrating astrocytoma, but without morphologic criteria diagnostic for glioblastoma multiforme, with staining negative for $p53$ and a mutation of the $BRAF$ gene (V600E) mutant protein (On-line Table). A second biopsy of both the suprasellar mass and the left globus pallidus revealed findings more compatible with a pilocytic astrocytoma with low $Ki-67/MIB-1$ immunostaining. The patient was intolerant of initial chemotherapy with carboplatin and vincristine, so he was treated with vinblastine for 8 months. Follow-up MR imaging 2 months later showed an enlarging, enhancing mass lesion in the inferior right basal ganglia (Fig 5B). He began treatment with $5'$-Inosinic acid, polymers, complex with $5'$-cytidylic acid, polymers and L-lysine, homopolymer, compound with cellulose carboxymethyl ether (cancer drug) (Poly-ICLC) at that time and has experienced improvement of neurologic symptoms; his MRI findings have remained stable on immunotherapy with Poly-ICLC for 24 months.

**DISCUSSION**

In our case series, new enhancement on MR imaging or a change in the contrast-enhancement pattern in a pre-existing presumed low-grade lesion occurred in 3 of our patients (patients 1–3) and correctly indicated a higher grade malignancy. Only 2 patients (patients 1 and 4) had restricted diffusion on conventional DWI and also had a high-grade glioma. The use of kurtosis diffusion imaging may have demonstrated superiority to conventional diffusion metrics in distinguishing various cerebral glioma grades. Most interesting, patient 5 had concerning clinical and MR imaging signs but turned out to have a low-grade histopathology.

There are 2 scenarios by which an HGG may occur in patients with NF1: 1) Lesions presumed to be pilocytic astrocytoma convert to HGG at that site years later; or 2) patients with presumed low-grade lesions subsequently develop a high-grade glioma at a different location. In our study, 1 patient (patient 3) developed a new lesion at a different site that grew slowly during a 16-month period; the other 4 patients did not have prior neuroimaging.
Recent retrospective analyses found that pediatric LGGs rarely transform to HGGs. In 1 retrospective analysis of 886 pediatric LGGs, only 2.9% of pediatric LGGs transformed into HGGs.\(^{20}\)

Approximately half of biopsied intracranial tumors in patients with NF1 are pilocytic or pilomyxoid astrocytomas, and the remainder are mostly diffuse/infiltrating gliomas, of which some are high-grade (anaplastic astrocytoma or glioblastoma).\(^1\) Lesions of the 3 patients with HGGs in our series were diffuse and contained an H3 K27M point mutation. NF1 mutations have been demonstrated in a subset of H3 K27M–mutant HGGs.\(^{21}\) Another HGG in our series was an anaplastic pleomorphic xanthoastrocytoma, a rare tumor that has been reported several times in the literature in the context of NF1.\(^{22-25}\) Recently, the molecular genetic definition of HGG and apparent LGG has gained more importance, emphasizing the need to obtain tissue surgically.\(^{26}\) The 2-year survival in patients with NF1 with HGG is higher than in patients without NF1.\(^{27}\)

Despite the small number of cases, we believe that raising awareness of an HGG in patients with NF1 among neuroradiologists is important. We were unable to evaluate the relative specificity of MR spectroscopy, perfusion, and kurtosis imaging, which may have prevented the tissue biopsy in the patient without HGG.

In conclusion, HGG in children with NF1 is rare, but concerning imaging signs should be taken seriously. We pursue tissue sampling when imaging findings are highly concerning for an HGG, though one of our patients in this series had a low-grade tumor on surgical pathology.

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