Pediatric low-grade gliomas and the need for new options for therapy

Why and how?

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Abbreviations: LGG, low-grade glioma; PFS, progression-free survival; OPG, optic pathway glioma; NF1, neurofibromatosis 1; PA, pilocytic astrocytoma; NF1-PA, neurofibromatosis 1-associated pilocytic astrocytoma; ERK, extracellular signal-regulated kinase; MAPK, mitogen-activated protein kinase; MEK, mitogen-activated protein kinase kinase; PI3K, phosphatidylinositol-3 kinase; mTOR, mammalian target of rapamycin; GG, ganglioglioma; GTP, guanosine-5’-triphosphate; GDP, guanosine-5’-diphosphate; PTEN, phosphatase and tensin homolog; Rheb, Ras homolog enriched in brain

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Pediatric low-grade gliomas are the most common tumors of the central nervous system in children, accounting for almost 50% of all childhood brain tumors. They are a heterogeneous group of tumors with different histologic subtypes. Most treatment studies address low-grade gliomas as a single entity, depriving us of histology-specific treatment outcomes. This is mostly due to a lack of understanding of tumor biology at the molecular level. Pediatric low-grade gliomas are not benign, and most incompletely resected tumors will progress and negatively affect quality of life. The advancements made in understanding sporadic pilocytic astrocytoma and neurofibromatosis 1-associated pilocytic astrocytoma in particular have paved the way for potential targeted therapy and biological stratification. Such progress in pilocytic astrocytoma needs to be consolidated and expanded to other histologic varieties of pediatric low-grade gliomas.

Introduction

Low-grade gliomas (LGGs) include a wide spectrum of grade I and grade II tumors, according to the World Health Organization 2007 classification.1 Table 1 lists the characteristics of these tumors, but this review will focus on the tumors most common in children.

If completely surgically resected, LGGs do not require further therapy. Unfortunately, total resection is not attainable in many of these tumors that are centrally located, so adjuvant treatments such as radiation therapy and chemotherapy were introduced. The initial study of chemotherapy in pediatric LGG in 1985 utilized vincristine and actinomycin D. Various regimens have emerged since then, with variable progression-free survival (PFS) curves that have not plateaued.3 Such regimens have been used for all LGG and did not reflect a biological understanding of LGG. Radiotherapy is an effective treatment with better PFS results than chemotherapy alone; 5-year PFS rates are 38% for chemotherapy alone and 68% for chemotherapy plus radiotherapy.4

Better targeted therapy is needed, and this can be achieved through a better understanding of the genetics and biology of the histologic subtypes that constitute LGG.

Why We Need New Options for Therapy

LGGs are the most common brain tumors in children, and pilocytic astrocytoma (PA) alone accounts for 21–23% of all pediatric brain tumors.5,6 Although complete surgical resection is the preferred therapeutic option, it is not always possible to achieve gross total resection in eloquent areas such as the dienecphalon. In a multicenter study of 198 low-grade chiasmatic hypotalamic tumors, gross total resection was achieved in only 5 cases.7 Radiotherapy is another effective treatment but with significant morbidities, such as neuroendocrine-cognitive deficits, vasculopathy and second tumors, especially in patients with neurofibromatosis 1 (NF1).8–10 Chemotherapy is widely used after surgery in an effort to avoid or defer radiation therapy, especially in young children. Some of the most common regimens used are carboplatin/vincristine, thioguanine/procarbazine/lomustine or CCNU/vincristine (TPC), and temozolomide. In three studies utilizing these regimens, only 4 patients (2.6%) achieved complete radiologic response: 4/78 (5%) on carboplatin/vincristine, 0/42 (0%) on TPC, and 0/30 (0%) on temozolomide.11–13 A further concern is that chemotherapy is not without complications. Carboplatin allergy, which can affect as many as 42% of patients, significantly limits delivery of an effective chemotherapy regimen.14 The randomized Children’s Oncology Group A 9,952 trial showed 5-year PFS of 35% using carboplatin/vincristine and 48% using TPC (p = 0.11), thus demonstrating no clear superiority between the 2 commonly used regimens.15

The choice of therapy for patients whose disease progresses on chemotherapy is an area of ongoing research. The preference of most treating physicians is to try to delay the use of definitive radiation
therapy until the child is past the first decade of life. Recent studies have identified vinblastine as an effective second-line treatment for recurrent or refractory LGG.\textsuperscript{16,17} Current studies are seeking to determine the feasibility of delivering vinblastine in combination with carboplatin.

LGGs are not benign tumors. They can negatively affect quality of life significantly, even in cerebellar PA treated with surgery alone: these patients are still at risk for cognitive and adaptive impairment, in addition to language, memory, attention and spatial function problems.\textsuperscript{18,19} In patients with optical pathway gliomas (OPGs), the quality of life can be compromised further due to effects on vision and the endocrine system.\textsuperscript{20,21} Visual outcomes in patients with OPGs have not been well documented, with most reports just focusing on limited aspects such as visual acuity or visual field alone.\textsuperscript{21,22} More detailed outcome measures such as contrast sensitivity and color vision are rarely addressed, and when such outcomes are documented, it is likely that the visual outcome will be much worse.\textsuperscript{20}

LGGs can also, although rarely, metastasize or transform into high-grade gliomas.\textsuperscript{23,24} Most important, LGGs can recur even if resected completely.\textsuperscript{25} In one study, the PFS in 278 patients was 55\% at 5 years and 42\% at 10 years.\textsuperscript{26} In the Children’s Oncology Group A 9,952 trial, the 5-year PFS for children without NF1 was 41.7\%. This high rate of progression exposes these children to different lines of treatment and more side effects.

**How can We Achieve New Options for Therapy for LGG?**

**Improved pathologic classification.** The pathologic classification of LGG can be challenging. In the Children’s Cancer Group CCG-945 high-grade glioma study, 70/250 patients (28\%) were reclassified as having LGG on further central pathologic review.\textsuperscript{4} These patients were exposed to unnecessarily aggressive therapy that could have been avoided. In other studies, unclassified or "not otherwise specified" LGGs were predominant, constituting 42\% and 32\% of patients enrolled in the protocol, respectively, in two series.\textsuperscript{26,27}

To better understand some of the rare types of LGGs, it is pivotal that we investigate them as such. Launching international studies may facilitate accrual of larger numbers of different types, allowing better conclusions to be reached. The “Montevideo initiative” for childhood leukemia launched in 1995 in an attempt to understand the biology and heterogeneity of rare leukemia subgroups is a good model to follow for rare LGGs.\textsuperscript{28} A similar example at the national level in pediatric brain tumors is the NF1 OPG task force.\textsuperscript{29} Such an initiative is responsible for many of the advancements seen in NF1-associated PA (NF1-PA).

**Better understanding of the behavior of LGG.** There is still much we do not understand about the behavior of LGG. Metastasis is a real phenomenon estimated to afflict 3–5\% of patients at diagnosis and 7–10\% at progression.\textsuperscript{21,24,30,31} Most of the literature consists of case reports, with the largest series reporting only 13, 11, 8 and 6 cases.\textsuperscript{24,31-33} Even the etiology of metastasis in LGG is poorly understood. Tabori et al.,\textsuperscript{32} identified amplification of the epidermal growth factor receptor gene in 6/6 (100\%) cases of metastatic LGG.

Another poorly understood and understudied event in LGG is transformation to high-grade glioma. Malignant transformation is a controversial issue, with some investigators doubting that it occurs independent of radiation therapy in PA.\textsuperscript{34} Our group documented that malignant transformation can occur in patients who have not been treated with radiotherapy.\textsuperscript{23} Five of the 11 patients in this study did not receive prior radiotherapy. Three of those 5 patients had undergone gross total resection before the malignant transformation. In 9 patients (82\%), tissue was available for molecular studies before and after malignant transformation. The overexpression of p53 and deletion of phosphatase and tensin homolog (PTEN) were more frequently seen after malignant transformation. Our data confirmed that malignant transformation is exceedingly rare in children with PA. On the other hand, the 15-year risk of malignant transformation among children with World Health Organization grade II LGG was 6.7\% in our study.

At the other extreme is spontaneous regression in pediatric LGG with and without associated NF1.\textsuperscript{35,36} Radiologically, the regression presents as a decrease in tumor size or a change in signal and clinically can be associated with improved symptoms. The biology of this phenomenon is poorly understood. An interesting study by Tabori et al.,\textsuperscript{37} identified shortening telomeres as an explanation for the growth arrest seen in some pediatric LGG. In their series, they did not find any telomerase activity or alternative lengthening of telomeres in 56 LGG samples.

Many studies have been launched in an attempt to identify clinical and biological risk factors for progression in LGG. Age of less than 5 years at diagnosis was found by most researchers to be associated with poorer 5-year PFS.\textsuperscript{15,27} Others found that infants and young children, less than 2 years, had worse PFS.\textsuperscript{7,21,38,39} On the other hand, one study found that children younger than 5 years had a better prognosis.\textsuperscript{12} Other factors such as NF1 presence, location, quality of response to chemotherapy, extent of resection and pathologic grade have been suggested as risk factors by different studies with contradicting findings.\textsuperscript{26,27,58-40}
are among the most common chromosomal abnormalities identified focused on PAs, which are the most common type. astrocytoma.

3 growth, proliferation and differentiation. An interesting finding in matrix is involved in creating the microenvironment that influences aggressive tumors. involved in subgrouping sporadic PA into more aggressive and less risk of progression in LGG. To date, the importance of the MIB-1 labeling index is controversial, with one study showing it to be a good predictor of tumor progression and others showing no such correlation. 

In an interesting study, tumor vascularity and angiogenesis were found to predict progression in OPGs. In addition, Wong et al., found that PA can be divided into two subgroups based on the differential expression of genes involved in cell adhesion, cell motility, angiogenesis and nerve ensheathment. The group overexpressing gene products associated with cell motility, adhesion and angiogenesis but underexpressing gene products associated with nerve ensheathment such as proteolipid protein and myelin basic protein was associated with more aggressive behavior and incomplete resection.

Chromosomal abnormalities were the focus of the earliest genetic studies done on LGGs, particularly PA. Many numerical and structural abnormalities were identified, with chromosomes 5, 7, 8 and 17 being the most frequently affected. In one study, the deletion of 17p (p53 site) was associated with rapid recurrence in 4 cases of PA regardless of aggressive treatment. 

Individualized understanding for each tumor type. Pilocytic astrocytoma. Most of the literature regarding the biology of LGGs is focused on PAs, which are the most common type. PAs are a heterogeneous group with a wide range of patterns that vary by location and sometimes even within the same lesion. Sporadic PA differs from NF1-PA in clinical behavior and natural history.

Sporadic PA. We have addressed some of genetic aberrations involved in subgrouping sporadic PA into more aggressive and less aggressive tumors. As mentioned earlier, chromosomes 7 and 8 are among the most common chromosomal abnormalities identified in sporadic PA. The importance of such chromosomal abnormalities was demonstrated by identifying the matrilin-2 gene (MATN2) and band 7q34 on chromosomes 8 and 7, respectively. Matrilin-2 is an extracellular matrix protein. The extracellular matrix is involved in creating the microenvironment that influences growth, proliferation and differentiation. An interesting finding in regard to matrilin-2 was the differential in the three-tier expression of the matrilin-2 gene and protein. First, matrilin-2 was overexpressed in sporadic PA compared with NF1-PA. Second, matrilin-2 was overexpressed in 14 of 15 (93%) supratentorial PA compared with 10 of 19 (53%) infratentorial PA. Third, matrilin-2 was overexpressed in tumors with aggressive behavior, whereas all 4 recurrent and 5 fatal tumors showed increased levels of matrilin-2.

On chromosome 7, band 7q34 had been intensively studied. Band 7q34 gain in these tumors included part of the BRAF gene locus that encodes the kinase domain. BRAF is a downstream effector in the mitogen-activated protein kinase (MAPK) pathway that is activated in some cases of sporadic PA. The mechanism of BRAF activation was found to be duplication of 7q34 in 53% or an activating mutation in 6% of sporadic PA. Such activation was proved by showing overexpression of CCND1, a downstream target of the MAPK pathway, and increased phosphorylation of extracellular signal-regulated kinase (ERK) 1/2, an immediate downstream target of BRAF when phosphorylated. Another interesting finding in this study was that the duplication at the 7q34 band was associated more with supratentorial location, recurrence and incomplete resection. Moreover, 7q34 duplication was specific to PA and was negative in all 28 ganglioglioma (GG) samples and 10 pleomorphic xanthoastrocytomas.

Similar findings were confirmed by Bar et al., who found 7q34 gain in 17/25 (68%) cases of sporadic PA. In this study, there were 20 infratentorial and 5 supratentorial tumors. The gain was again associated with location, but all 17 tumors with the 7q34 gain were in the infratentorial group, which contrasts with what Pfister et al., found. This inconsistency between studies may be due to sample size.

Another possible genetic aberration involved in the pathogenesis of sporadic PA is activation of the phosphatidylinositol-3 kinase (PI3K) pathway. In one study, Ras was found to be activated in all 21 sporadic PA cases tested by demonstrating phosphorylation of its downstream effector, AKT kinase. However, the mechanism of activation was found to be an activating mutation in K-RAS in only one tumor (5%). The same group confirmed this finding in other tumor (7%) out of 15 sporadic PA samples. A summary of putative genetic abnormalities that may be responsible for the tumorigenesis of sporadic PA is provided in Figure 1. Understanding such genetic aberrations in sporadic PA may allow us to better biologically target these tumors.

Decoding mechanisms of the MAPK and PI3K pathways in sporadic PA and NF1-PA may help guide us in selecting potential targeted therapies (Fig. 2). This can include RAF inhibitors, such as urea derivatives. One such compound, BAY 43-9006 (sorafenib), which inhibits C-RAF and B-RAF, is being investigated in clinical trials and has been approved for renal cell cancer. Among other therapeutic targets under investigation in vitro are the following:

- Inhibiting cyclin D1 by neomycin, which has shown efficacy in glioma cell lines.
- Inhibiting ERK and mitogen-activated protein kinase kinase (MEK) using compounds such as AZD6244 and perifosine, which also inhibits Akt in the PI3K pathway.
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- Inhibiting AKT1/2 using AKT1/2, which is effective against cancer cell lines with activated AKT.

- Inhibiting PI3K and mammalian target of rapamycin (mTOR) with the dual inhibitor PI-103, which has shown efficacy in preclinical trials on malignant glioma cell lines. It also can be a putative targeted therapy in PA when the PI3K pathway is activated.

NF1-PA. NF1 is an autosomal dominant disease that affects 1 in 3500 individuals. Those affected are at risk for many central nervous system tumors, with OPG being the most common, occurring in 15–20% of NF1 patients. NF1-PA differs from sporadic PA in clinical behavior and genetic signature.

The loss of NF1 on 17q is the main genetic difference between the two types. The NF1 gene product, neurofibromin, is a negative regulator of RAS by exerting guanosine-5'-triphosphate (GTP)ase activity on the GTP-bound RAS and converting it to guanosine-5'-diphosphate (GDP)-bound RAS. As seen in Figure 2, the constitutively activated Ras will trigger both the MAPK (RAF/MEK/ERK) and the PI3K (RAS/PI3K/AKT/mTOR) pathways. Normally, RAS can be activated by a receptor tyrosine kinase such as epidermal growth factor receptor (Fig. 2). For RAS to perform its functions, it must localize to the cell membrane. This is done by adding hydrophobic molecules through the process of prenylation.

Using Nf1-deficient mice, Dasgupta et al., found specific activation of K-RAS responsible for glioma formation in vitro and in vivo.

One of the downstream targets of the PI3K pathway is mTOR, a serine/threonine kinase involved in protein synthesis and growth that can be activated by AKT, as in Figure 2. Neurofibromin is a negative regulator of mTOR via RAS, and its loss causes hyperactivation of mTOR. In an interesting study, rapamycin, an mTOR inhibitor, was administered in an NF1 mouse model and yielded a positive response in a dose-dependent manner. In addition to mTOR inhibitors, other inhibitors of the RAS/PI3K/AKT/mTOR pathway can be putative targeted therapies, as mentioned earlier in connection with sporadic PA (Fig. 2).

Subependymal giant cell astrocytoma and PA in tuberous sclerosis complex. Another mechanism to activate mTOR is through RAS homolog enriched in brain (Rheb). As shown in Figure 2, the TSC1/2 (tuberin/hamartin) complex inactivates Rheb. The loss of this complex, as in tuberous sclerosis complex, results in hyperactivation of mTOR.

In 5 patients with a clinical diagnosis of tuberous sclerosis complex who developed subependymal giant cell astrocytoma (n = 4) or PA (n = 1) with documented progressive lesions, rapamycin was administered. In all 5 patients, lesions and PA decreased in size.

Pilomyxoid astrocytoma. First reported as a separate entity in 1999, pilomyxoid astrocytoma is closely related to PA, with bipolar cells in a myxoid matrix but no Rosenthal fibers or eosinophilic granular bodies. It occurs in the very young (less than 1 year old) with hypothalamic/chiasmatic predilection and carries a worse prognosis with a 1-year PFS of only 37.8%. Reports about the genetics of pilomyxoid astrocytoma are scarce, consisting of only one case report of an NF1 patient and another report of disruption of the BCR gene in a child.

Dysembryoplastic neuroepithelial tumor. First reported by Daumas-Dupont in 1988 on a combined series of 39 patients from the United States and France, dysembryoplastic neuroepithelial tumors are cortical lesions that occur in young individuals with longstanding refractory seizures. Pathologically, these tumors are composed of glial nodules, cortical dysplasia and specific glioneuronal elements. Initial reports showed no recurrence even after incomplete resection. However, later reports showed one case of recurrence after gross total resection and another case of malignant...
transformation 14 years after resection with no chemotherapy or radiotherapy. Genetically, we know very little about this tumor so far. Case reports have shown these tumors in association with NF1 and XYY syndromes. Other studies showed a lack of 1p and 19q deletions and p53 mutations. A unique study by Prayson’s group investigated some of the apoptosis-associated proteins such as Bcl-2, bcl-x, and bax in 18 cases. They found these genes differentially expressed between the oligodendroglial and neuronal components. The significance of such findings is not yet known.

Ganglioglioma. GG is a well differentiated, indolent, mixed neuronal-glial tumor composed of mature ganglion cells and abnormal glial cells. It is also associated with a long history of seizure. One of the initial studies on the genetics of GG investigated 13 tumors (9 temporal and 4 at other sites) from 120 total cases of pediatric brain tumors. Only 2 non-temporal cases showed abnormalities, with both involving inversion 1q. Later publications included an unusual case report on spinal GG in an NF2 patient and 2 reports of metastatic Ggs. Genetic analysis on both metastatic cases was performed and showed inversion of chromosome 7 and loss of 17p. A recent large study investigated 61 cases of GG, 19 of which were in patients 18 years old or younger. In the study, 2 groups of GG were identified. The first group had a complete gain of chromosome 7 with additional gains in 5, 8 and 12. The second group had mostly normal chromosomes, with occasional losses on chromosome 9 and 22q. It should be noted that the second group was similar to the 11 of 13 GG cases with normal cytogenetics mentioned above. Another interesting finding in this study was the anaplastic recurrence of 2/61 (3%) as GG grade III. These two tumors had genetic aberrations similar to those of high-grade gliomas. Moreover, these genetic aberrations were present on the primary tumor as GG grade I at diagnosis.

There is a distinct entity called dysplastic gangliocytoma of the cerebellum or Lhermitte-Duclos disease that is associated with Cowden disease. It can occur in infancy and adulthood. A PTEN inactivating mutation is the genetic basis for this syndrome. PTEN is a tumor suppressor that inhibits the phosphorylation of PI3K. Such inactivation causes hyperactivation in the PI3K/Akt/mTOR pathway (Fig. 2). Interestingly, PTEN mutations were identified only in adult-onset Lhermitte-Duclos disease and not in the pediatric cases.

Conclusion

In the last decade, we have come to know somewhat more about PAs. We still know very little about other types of LGGs. We believe that the future of research in pediatric LGGs will provide a better understanding of the biology and genetics of each type. Furthermore, within each type, there will be additional stratification based on genetic signatures that provide clues to clinical behavior, such as recurrence, metastasis, or a more indolent natural history. Such understanding may allow us to stratify treatment accordingly. For this to occur, multicenter and multination studies are needed to provide enough numbers for basic and clinical research for each group of LGGs.

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