Cerebral Neoplasms in L-2 Hydroxyglutaric Aciduria: 3 New Cases and Meta-Analysis of Literature Data


doi: https://doi.org/10.3174/ajnr.A2869
http://www.ajnr.org/content/33/5/940
Cerebral Neoplasms in L-2 Hydroxyglutaric Aciduria: 3 New Cases and Meta-Analysis of Literature Data

SUMMARY: Increasing evidence suggests that patients with L2-HGA have a predisposition to cerebral neoplasms. This may be related to the pathologic accumulation of L2-HG because high amounts of 2-HG have been found in brain neoplasms that have IDH1 mutations. Our experience, on the basis of 11 previously unreported cases of L2-HGA, 3 of which developed cerebral neoplasms during the course of the disease, also supports an association between L2-HGA and cerebral neoplasms. We conducted a meta-analysis of published data, and we identified 295 patients (including our 11 patients) with L2-HGA. In 14 patients, the metabolic disorder was associated with cerebral neoplasms, suggesting an approximately 5% prevalence rate of CNS neoplasms in patients with L2-HGA; nonetheless, it may still be an underestimate. L2-HGA is an important disease “model” that provides further evidence to support the recently proposed pathogenetic role of 2-HG in the development of cerebral neoplasms.

Abbreviations: 2-HG = 2-hydroxyglutarate; IDH = isocitrate dehydrogenase; L2-HG = L-2-hydroxyglutarate; L2-HGA = L-2-hydroxyglutaric aciduria; L2HGDH = L-2-hydroxyglutarate dehydrogenase; MDH = malate dehydrogenase; PNET = primitive neuroectodermal tumor; WHO = World Health Organization

L2-HGA is a rare organic acidopathy. Compelling evidence suggests that the disease predisposes to the development of cerebral neoplasms. On the basis of 4 cerebral neoplasms observed in 7 patients with confirmed L2-HGA, the most comprehensive description to suggest an association between L2-HGA and malignant brain tumors is that of Moroni et al. Our experience with patients affected by L2-HGA supports this association, which we report here alongside a meta-analysis of pertinent published data on the prevalence of brain tumors in patients with L2-HGA.

Case Series and Review of Literature Data

We have encountered 11 patients (male/female: 9:2) with L2-HGA in our clinical practice. Three of these patients developed cerebral neoplasms during the course of their disease (Fig 1). In fact, 2 of the patients were known to have L2-HGA at the time of tumor diagnosis (cases 1 and 2, presenting with a right hippocampal anaplastic astrocytoma [Fig 2] and a right frontal oligodendroglioma, respectively), whereas in the third case, the diagnosis of underlying metabolic disorder was made at the preoperative radiologic work-up of the tumor (case 3, with a left thalamic low-grade glioma). The metabolic diagnosis was subsequently confirmed by laboratory tests. The demographic, clinical and histopathologic data on these patients together with all published data on patients with L2-HGA is summarized in the Table.

We found 295 cases of L2-HGA (including our 11 patients) in the literature, in which the diagnosis of the metabolic disorder was established with the necessary degree of confidence. A cerebral neoplasm developed in 14 patients. Seven of the cases were described in case reports; 1 report described a series of 4 patients; and as mentioned above, our 3 patients were taken from a cohort of 11.

Based on these data, the overall prevalence of brain tumors in patients with L2-HGA would be estimated at <5%. There is no apparent sex predisposition to the carcinogenic effect.

Analysis of patient age at diagnosis reveals some interesting additional phenomena. Tumors occurred between the ages of 3 and 36 years but were not evenly distributed across this age range; indeed, 3 putative subgroups may be identified within the cohort. Tumors in the youngest patients probably correspond to medulloblastomas (2 patients; the age of 1 of these patients is not known). Three of the 4 patients with low-grade supratentorial tumors (astrocytoma or low-grade astrocytoma) were children between 9 and 12 years of age. Only 1 low-grade glioma was found in an adult. Tumors encountered in older teenagers or young adults tended to be high-grade gliomas, though PNETs were also represented. In 1 patient, neither age at tumor diagnosis nor pathologic diagnosis was available. Supratentorial tumors occurred in patients older than 9 years, were located in the frontal or temporal lobe or within the thalamus, and were gliomas.

Discussion

L2-HGA is a rare panethnic organic aciduria. A finding of urinary excretion of 2-hydroxyglutaric acid is required for a formal diagnosis and is accomplished by using chiral gas chromatography and mass spectrometry. All 295 patients reviewed here had the L-isomer variant of 2-hydroxyglutaric aciduria (D-2-hydroxyglutaric aciduria is a very rare inborn error of metabolism, with only 75 cases in the literature). To the best of our knowledge, there have been no reports linking neoplasms of any kind to D-2-hydroxyglutaric aciduria, and this also applies to the extremely rare dual disease, L- and D-2-hydroxyglutaric aciduria.

L2-HGA is a peculiar organic acidopathy in many ways.
Unlike most other organic acidopathies, it develops insidiously, without episodes of metabolic decompensation. Clinically, it presents with slowly progressive encephalopathy, pyramidal and cerebellar signs, and intellectual deterioration. Thus, patients, though affected to varying degrees neurologically, typically receive a diagnosis late in childhood and often live into adulthood. The disease is associated with the optical isomer of 2-HG (D-2-hydroxyglutaric aciduria) and may have a less benign clinical course.

Abnormalities on MR imaging in L2-HGA are characteristic and, at later stages of the disease, even pathognomonic; hence, the diagnosis is quite often suggested by the radiologist, to be confirmed by laboratory tests. Our 3 cases illustrate this well. Despite the presence of an obvious neoplastic process and its secondary effects (eg, peritumoral vasogenic edema), the background abnormalities of L2-HGA were easy to recognize. Remarkably, a neurometabolic disease was unknown in patient 3 until a left thalamic tumor and L2-HGA were diagnosed, both through the use of MR imaging, when she was 36 years old.

The typical imaging presentation of L2-HGA includes an extensive leukoencephalopathy exhibiting a centripetal and anteroposterior gradient, as well as bilateral symmetric involvement of the globus pallidus, caudate nucleus and putamen, and (characteristically) the cerebellar dentate nuclei. The leukodystrophy-like appearance of the disease by imaging is unique among organic acidopathies, with the exception of Canavan disease, which also presents with white matter disease and characteristic involvement of the globus pallidus and thalamus (but sparing of the striatum).

From our meta-analysis, an association between L-2-hydroglutaric aciduria and primary cerebral tumors is obvious. Regarding the estimated prevalence of about 5%, it may be either an overestimate or an underestimate. On one hand, it is true that not all L2-HGA cases are necessarily reported, especially in recent years; therefore, the actual prevalence may be lower. However, if one looks at 2 specific comprehensive series (ours and that of Moroni et al1), 7 cerebral neoplasms were found among 18 patients, and the actual prevalence may be closer to 40%. It is, of course, possible that an inadvertent accrual bias (ie, patients had more severe disease and hence were more prone to neoplasms) could account for the apparent higher prevalence in these 2 cohorts compared with the rest of the reported cases. Another explanation for the relative under-reporting of an association between L2-HGA and brain tumors is that the metabolic disorder is typically encountered in childhood; therefore, most published cases are likely to have been reported by pediatricians or metabolic specialists. Because the disease is untreatable and is not complicated by episodes of metabolic decompensation later in life, it is possible that discontinuity of care by the time a brain tumor develops in a patient prevents the association being made by the neurosurgeon or radiation oncologist who manages the new disease episode. In this context, it is perhaps relevant that the 2 largest series and the most compelling data come from large referral centers where pediatric neurologists, neurosurgeons, neuropathologists, and neuroradiologists work closely in multidisciplinary teams. In any case, a more systematic follow-up (perhaps including periodic brain MR imaging screening) of patients with L2-HGA may allow us in the future to determine the true prevalence of cerebral neoplasms associated with this metabolic disorder.

Although various theories have been proposed for the association between L2-HGA and brain tumors, the mechanism
of tumorigenesis has only recently been established. Inborn errors of metabolism lead to an accumulation of some metabolites upstream of the defect in the metabolic pathway; in L2-HGA, other metabolites in excess but without obvious tumor-igenic effect, include glycolate, glycerate, pyruvate, laurate, hydroxybutyrate, citrate, isocitrate, palmitate, myristate, oleate, and stearate. However, recent studies involving novel sequencing strategies, which were directed first at glioblastoma, uncovered a high frequency of \( IDH \) mutations in diffuse gliomas of WHO grade II/III and secondary glioblastomas. More than 80% of these tumors harbor an \( IDH1 \) mutation, which mainly involves 1 codon (R132H) and induces a gain of function that promotes conversion of isocitrate to excessive (nonphysiologic) quantities of potentially carcinogenic \( 2\)-HG (IDH normally converts isocitrate to alpha-ketoglutarate in the citric acid cycle).

In addition to its primary function of converting malate to oxaloacetate in the citric acid cycle, MDH converts small amounts of alpha-ketoglutarate to \( 2\)-HG under normal physiologic conditions. This conversion of alpha-ketoglutarate to \( 2\)-HG is insignificant under physiologic conditions because \( 2\)-HG is reconverted to alpha-ketoglutarate by L2HGDH as a metabolic “repair” or “clean-up” mechanism (Fig 3). However, in L2-HGA, \( 2\)-HG accumulates and may reach pathologic and, hence, potentially carcinogenic concentrations because of a mutation in the \( L2HGDH \) gene at 14q22 that causes loss of function of \( L2HGDH \). The significance of these physiologic intracellular “housecleaning” processes is being increasingly recognized, and L2-HGA is a prime example of the failure of one such mechanism with subsequent deleterious consequences. In patients with L2-HGA, the mechanism of the \( 2\)-HG increase is, therefore, very different from that seen in \( IDH1 \) mutants, but the end result is the same.

As described earlier, of the 14 brain tumors encountered in patients with L2-HGA, 9 were diffuse gliomas (WHO grades II, III, IV) of a type found to harbor \( IDH1 \) mutations in the sporadic disease. Medulloblastomas and PNETs seem to occur more frequently in patients with L2-HGA; however, \( IDH1 \) mutations have not been reported in medulloblastomas or other PNETs. Finally, although other tumors (bone tumors and Wilms tumor) have also been reported in L2-HGA, these tumors are so rare that these occurrences may be merely coincidental.
tricular white matter, which constitutes the basis of the diagnosis of the disease by MR imaging. This selective vulnerability also suggests that myelin produced early in life (before the onset of the metabolic derangement and the myelinotoxic effect) may be somewhat resistant to the myelinotoxic pathogenic process but not necessarily to the oncogenetic effect. It is interesting to speculate that in L2-HGA, the short-term effect of excess 2-HG in brain parenchyma may be essentially myelinotoxic in vulnerable areas, whereas long-term exposure of the brain to 2-HG induces neoplastic transformation of glial progenitors or less committed neural stem cells, but to date, no data exist to support this hypothesis.

This review of the clinicopathologic features of patients with L2-HGA who occasionally have brain tumors highlights associations between the metabolic consequences of this disorder and IDH1 mutations in diffuse gliomas, though we must await a clearer understanding of the pathogenesis.

References