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### ► To cite this version:

Alberto Picca, Giulia Berzero, Franck Bielle, Mehdi Touat, Julien Savatovsky, et al.. FGFR1 actionable mutations, molecular specificities, and outcome of adult midline gliomas. *Neurology*, 2018, 90 (23), pp.e2086-e2094. 10.1212/WNL.0000000000005658 . hal-01971530

**HAL Id: hal-01971530**

**<https://hal.sorbonne-universite.fr/hal-01971530>**

Submitted on 7 Jan 2019

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# *FGFR1* actionable mutations, molecular specificities, and outcome of adult midline gliomas

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## Abstract

### Objective

To characterize the prevalence and prognostic significance of major driver molecular alterations in adult midline diffuse gliomas (MLG).

### Methods

Adults with histologically proven MLG diagnosed between 1996 and 2017 were identified from our tumor bank, systematically reviewed, and reclassified according to WHO 2016. Targeted sequencing was performed, including determination of *H3F3A*, *HIST1H3B*, *TERT**p*, *IDH1/2*, *FGFR1*, *p16/CDKN2A*, and *EGFR* status.

### Results

A total of 116 adult patients (M/F 71/45, median age 46.5 years) with MLG (17 cerebellar, 8 spinal, 30 brainstem, 57 thalamic, and 4 diencephalic nonthalamic) were identified. Most patients had high-grade disease at presentation (grade II: 11%, grade III: 15%, grade IV: 75%). Median overall survival was 17.3 months (14.5–23.8 months). Main molecular alterations observed were *TERT* promoter, *H3F3A*, and hotspot *FGFR1* (N546 and K656) mutations, in 37%, 34%, and 18% of patients, respectively. *IDH1* mutations only affected brainstem gliomas (6/24 vs 0/78;  $p = 7.5 \times 10^{-5}$ ), were mostly non-R132H (contrasting with hemispheric gliomas,  $p = 0.0001$ ), and were associated with longer survival (54 vs 12 months). *TERT* promoter mutation (9.1 vs 24.2 months), *CDKN2A* deletion (9.9 vs 23.8 months), and *EGFR* amplification (4.3 vs 23.8 months) were associated with shorter survival. Of interest, in contrast with pediatric MLG, H3K27M mutations were not associated with worse prognosis (23 vs 15 months).

### Conclusions

Patients with adult MLG present with unique clinical and molecular characteristics, differing from their pediatric counterparts. The identification of potentially actionable *FGFR1* mutations in a subset of adult MLG highlights the importance of comprehensive genomic analysis in this rare affection.

## Glossary

CI = confidence interval; MLG = midline gliomas; OS = overall survival.

Diffuse midline gliomas (MLG) are rare entities occurring predominantly in children and young adults.<sup>1</sup> Genomic studies in pediatric MLG showed that these tumors frequently harbor recurrent K27M missense mutations in *H3F3A* and *HIST1H3B* genes, which both encode variants of the histone H3 protein.<sup>2–4</sup> While *HIST1H3B* mutations are anatomically restricted to brainstem,<sup>4,5</sup> *H3F3A* mutants have been reported in MLG of various locations.<sup>2,5–8</sup> In pediatric MLG, these mutations associate with more aggressive clinical phenotype and poorer prognosis.<sup>6,8,9</sup> Based on these observations, the WHO 2016 classification introduced “diffuse midline glioma, H3K27M-mutant” as a new grade IV entity, even in the absence of histopathologic markers of anaplasia.<sup>10</sup>

In adults, fewer than 10% of diffuse gliomas arise in midline structures.<sup>1,11</sup> Recent works suggested that they may differ from both pediatric MLG and adult supratentorial gliomas. As an illustration, although *H3F3A* mutations have been reported in adult MLG,<sup>7,12–15</sup> their prognostic significance in adults remains unclear.<sup>12,13</sup> While less frequent, actionable<sup>16,17</sup> *IDH1* and *FGFR1* hotspot mutations have been described in this population.<sup>18,19</sup>

Overall, the genomic landscape of adult MLG remains poorly understood. Indeed, the need for histologic confirmation in presence of a typical radiologic presentation is debated, because of the surgical risks and assumption of limited benefit on overall survival (OS).<sup>20</sup>

We assumed that unravelling the genomic landscape of these tumors will enable better defining the prognostic value of molecular biomarkers in this disease and identify new therapeutic strategies. In this study, we report the clinical, histologic, and molecular characteristics of 116 adult patients with MLG, and assess the prevalence and prognostic significance of major driver molecular alterations.

## Methods

### Patients and tumor samples

Patients (≥15 years) with diagnosis of MLG were retrospectively identified from our database (OncoNeuroTek, GH Pitié-Salpêtrière, Paris, 1996–2017) according to the following criteria: 1) tumor arising in midbrain location (thalamus, hypothalamus, pineal region, brainstem, cerebellum, or spinal cord), verified on neuroimaging at diagnosis; 2) histopathologic confirmation of diffuse glioma according to WHO 2016 classification; and 3) available follow-up data.

All tumor samples underwent histopathologic review (F.B. and K.M.) and, if required, additional diagnostic immunohistochemical stainings were performed.

Radiologic analysis was performed by A.P. and J.S. and scored contrast enhancement (absent, mild/moderate, marked/avid) and radiologic necrosis (none, <5%, 6%–33%, 34%–67%, >67% of the bulk), according to Visually Accessible Rembrandt Images (VASARI) criteria.<sup>21</sup>

As a control, we used a set of hemispheric diffuse gliomas with available tumor DNA.

### Standard protocol approvals, registrations, and patient consents

All tumor samples and clinical data were collected with written informed consent in accordance with the tenets of the Declaration of Helsinki.

### DNA extraction

DNA was extracted from formalin-fixed paraffin-embedded or frozen samples, as previously described.<sup>15</sup>

### Sanger sequencing

Genes were sequenced using the Sanger method for the codons 132 of *IDH1*, 172 of *IDH2*, 27 of *H3F3A*, 27 of *HIST1H3B*, 546 and 656 of *FGFR1*, 600 of *BRAF* and *TERT* promoter at position –250 and –228 after standard PCR amplification, as previously described.<sup>15,22</sup>

The primers used were as follows: for *IDH1*, F-TGTGTTGATGATGGAGCGCTATTTG, R-TGCCACCAACGACCAAGTCA; for *IDH2*, F-5-GC CCGGTCTGCCACAAAGTC, R-5-TTGGCAGACTCCAGAGCCCA; for *H3F3A*, F-GTGATCGTGCCAGGAAAAGT, R-CAAGAGAGACTTTGTCCATTTT; for *HIST1H3B*, F-GTTTTGCCATGGCTCGTACT, R-AAGCGAAGATCGGTCTTGAA; for *TERT* promoter region, F-GGCCGATTCGACCTCTCT, R-AGCACCTCGCGGTAGTGG; for *FGFR1*, F-CCTCCTCCCTCCCAAGTAA, R-GGACTGATACCCAGCTCAG; for codon 546 hotspot (exon 12), F-CTTCCAGCTCCCTCACC TC, R-CCCACTCCTTGCTTCTCAGAT for codon 656 hotspot (exon 14); and for *BRAF* codon 600, F-TGCTTGCTCTGATAGGAAAATG, R-CAGGATCTCAGGGCCAAAAAT.

### Array comparative genomic hybridization

When enough material was available, array comparative genomic hybridization analysis was performed as previously described,<sup>23</sup> to determine *EGFR* amplification, *p16/CDKN2A* deletion, and chromosomes 1p, 19q, 9p, 10p, and 10q losses.

### MGMT methylation

DNA methylation status of the *MGMT* promoter was determined by bisulfite conversion and subsequent nested methylation-specific PCR, a 2-stage PCR approach.

## Immunohistochemical analyses

Detection of IDH1 R132H mutant enzyme, Ki67, and ATRX nuclear expression were performed as previously described.<sup>24,25</sup> In particular, the loss of nuclear ATRX expression was considered specific when tumor cell nuclei were unstained while the nuclei of non-neoplastic cells were ATRX-positive. Antibody antihistone H3.3 K27M mutant (ABE419; Millipore, Burlington, MA) was used at dilution 1/1,000. Samples were considered histone H3 K27M or IDH1 R132H mutated when immunohistochemistry was positive for the presence of the altered protein.<sup>26,27</sup>

## Statistical analysis

OS was defined as the time between diagnosis and death. Patients who were still alive at the last follow-up were censored for the analysis. Survival curves were calculated using the Kaplan-Meier method. Statistically significant differences between survival were assessed using the log-rank test. A log-rank test  $p$  value  $\leq 0.05$  (2-sided) was considered to be statistically significant. Cox model was used to evaluate the effect of quantitative variables on survival and for multivariate survival analysis.  $\chi^2$  and Fisher exact test were used to compare qualitative variables, with  $p < 0.05$  considered as significant. Hierarchical clustering and multidimensional representations were used using the Ward method. The analyses were performed using R software packages.

## Data availability

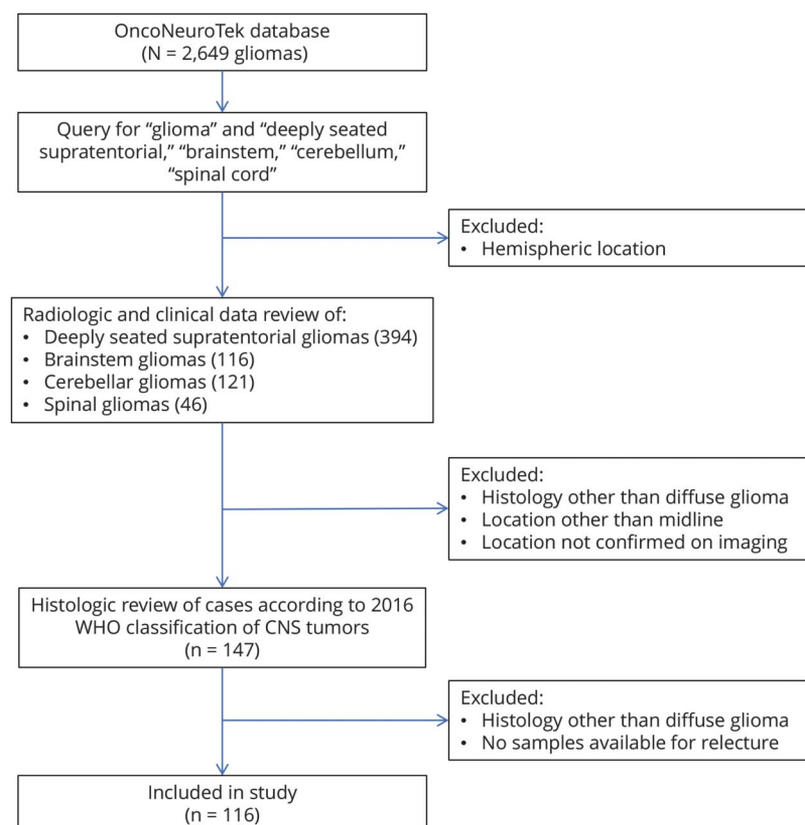
The complete individual data and supplementary material are available at doi: 10.5281/zenodo.1183373.

## Results

### Clinical and histologic features of adult MLG

We identified 116 patients (M/F 71/45; median age 46.5 years, range 15–75) fulfilling the inclusion criteria, out of 2,649 patients with diffuse glioma of any location from our database (4.4%), and harboring cerebellar (17), spinal (8), brainstem (30), thalamic (57), and diencephalic non-thalamic (4) midline gliomas (table e-1, [links.lww.com/WNL/A511](https://links.lww.com/WNL/A511); figure 1). All the cases have been reviewed by a neuropathologist (F.B.) according to the 2016 WHO classification (table e-1). The predominant histologic aspect was astrocytic diffuse gliomas, but other patterns were also observed: oligodendroglial, piloid, pseudo-ependymal, pleomorphic xanthoastrocytoma-like, or embryonal tumor-like. A total of 84/109 (77%) patients had a biopsy only; 25 (23%) had partial or subtotal surgical resection. Sixty-eight percent (68/101) of patients received combined chemoradiation with temozolomide as first-line therapy, 17% received chemotherapy, and 12% received radiation therapy alone. Four patients did not receive any adjuvant treatment after initial diagnosis.

**Figure 1** Inclusion flowchart



## Molecular characterization of adult MLG reveals the presence of recurrent hotspot mutations in FGFR1

The frequency and type of the main molecular alterations is detailed in figure 2. As diagnosis was based on tumor biopsy in most of the patients and tissue availability was limited, molecular analyses were focused on the alterations hypothesized to be the most relevant to the disease. Comparative genomic hybridization array could be performed in 56 cases.

Histone *H3K27M* mutations were found in 38/110 patients (34%) and were associated with a younger age at diagnosis (median 33 years vs 53 years,  $p = 6 \times 10^{-5}$ , figure e-1e, links. lww.com/WNL/A510). Histone H3 mutations were mutually exclusive with *IDH1*, and inversely associated with *TERT* mutation, *p16/CDKN2A* loss, and *EGFR* amplification (table e-1, links.lww.com/WNL/A511). *H3K27M* was found in all the locations but was associated with thalamic and spinal locations (table 1).

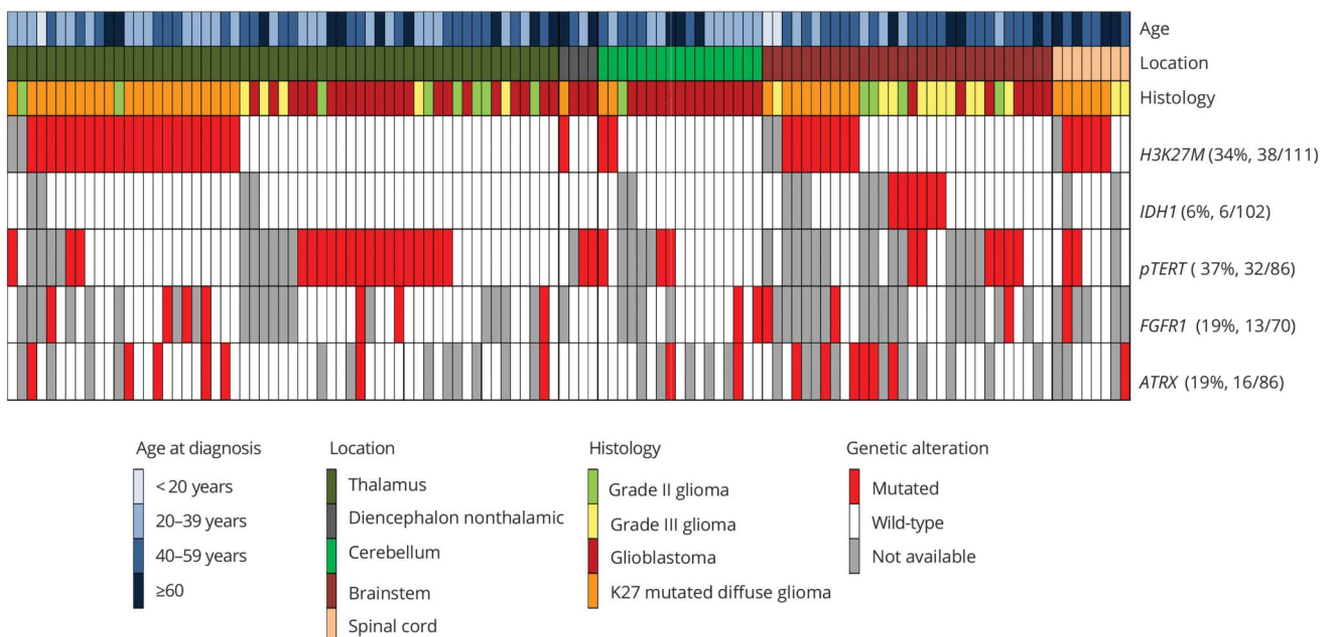
*IDH1* mutations were found in 6 patients (1 grade II, 4 grade III, 1 grade IV) out of 102: strikingly, all 6 *IDH1* mutated tumors were located in brainstem (6/24 vs 0/78 for MLG not involving the brainstem;  $p = 7.5 \cdot 10^{-5}$ ). Only 1 out of the 5 *IDH1* mutations identified by sequencing was *IDH1* R132H; the other mutations were p.Arg132Gly (R132G) (2), p.Arg132Cys (R132C), and p.Arg132Leu (R132L), which contrasted with non-midline *IDH1*-mutant gliomas: we compared these results to the data from non-midline gliomas recorded in our tumor base (OncoNeuroTek) and found a difference in the pattern of *IDH1* mutations between midline and non-midline gliomas ( $p = 0.0001$ ): out of 2,234 *IDH1* mutated supratentorial gliomas, 2,083 (93.2%) were R132H,

55 (2.5%) were R132G, 54 (2.4%) were R132C, 27 (1.2%) were R132S, and 15 (0.7%) were R132L. These data indicate that *IDH*-mutated brainstem gliomas differ from supratentorial *IDH*-mutated gliomas, and that immunohistochemistry anti-IDH1R132H is not the valid method for *IDH* screening in these tumors.

Fifty-nine patients were found *IDH* wild-type and K27 wild-type: this group—*IDH*wt-K27wt—of MLG had more frequent *TERT* mutation (24/48,  $p < 1 \times 10^{-5}$ ), *CDKN2A* deletion (7/24,  $p = 0.028$ ), and *EGFR* amplification (5/24,  $p = 0.016$ ) (figure 2). No patient had *BRAF* V600 mutation (0/69).

Recurrent *FGFR1* mutations affecting 2 hotspots were identified in 18% of patients (13/73): p.Asn546Lys (9), p.Asn546Asp (2), and p.Lys656Glu (2) with diverse locations: 7/37 thalamus (19%), 2/12 brainstem (17%), 2/13 cerebellar (15%), and 1/3 spinal cord (33%) ( $p = 0.9$ ), in both K27M MLG (6/23) and K27wt MLG (6/45),  $p = 0.3$ ; and 6/41 *IDH* wild-type vs 0/4 *IDH* mutated,  $p = 1$  (figure 2). In order to investigate whether *FGFR1* mutations were specific to midline located diffuse gliomas, we sequenced from our OncoNeuroTek tumor bank 479 DNA samples extracted from hemispheric gliomas (170 GBM, 151 grade III, 157 grade II, of which 197 were *IDH* mutated, 212 *IDH* wild-type, and 70 undetermined) for the presence of *FGFR1* mutations: we found only one mutation in a patient with an *IDH* wild-type GBM involving the corpus callosum, suggesting that *FGFR1* mutations are restricted to midline diffuse gliomas (13/73 vs 1/479;  $p = 1.2 \cdot 10^{-10}$ ). Our data were further confirmed by public data (cbiportal.org): out of 1,722 patients with diffuse gliomas, only 2 patients had *FGFR1* activating mutation (13/73 vs 2/1,722;  $p = 10^{-16}$ ): 1 GBM with K656E

**Figure 2** Age, tumor location, tumor grade, and molecular alterations of 116 patients with midline gliomas



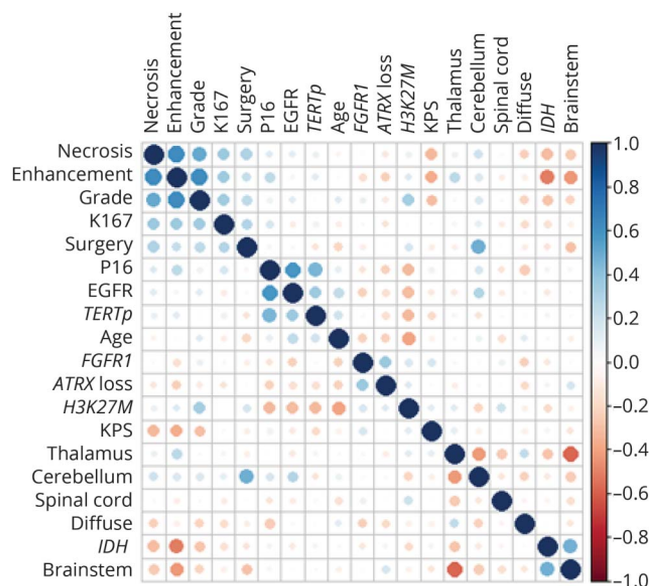
**Table 1** Clinical, histologic, and molecular characteristics in patients with H3K27M mutation vs H3K27 wild-type midline gliomas (MLG)

	K27M wild-type	K27M mutated	p Value
<b>N</b>	73	38	NA
<b>Sex ratio (M/F)</b>	1.52 (44/29)	1.53 (23/15)	0.98
<b>Age at surgery, y, median (range)</b>	53 (23–75)	33 (15–65)	<0.0001
<b>Location, % (n)</b>			0.04
<b>Nonthalamic diencephalic</b>	4 (3/73)	3 (1/38)	
<b>Thalamic</b>	45 (33/73)	58 (22/38)	
<b>Brainstem</b>	27 (20/73)	21 (8/38)	
<b>Cerebellar</b>	21 (15/73)	5 (2/38)	
<b>Spinal cord</b>	3 (2/73)	13 (5/38)	
<b>Contrast enhancement, % (n)</b>	72 (34/47)	88 (21/24)	0.16
<b>First-line treatment, % (n)</b>			
<b>Surgery + RT</b>	14 (9/63)	9 (3/34)	
<b>Surgery + CHT</b>	19 (12/63)	15 (5/34)	
<b>Surgery + RT-CHT</b>	63 (40/63)	74 (25/34)	
<b>Surgery only</b>	3 (2/63)	3 (1/34)	
<b>Median OS, mo (95% CI)</b>	15.0 months (10.4–19.7 months)	18.6 months (14.7–33.6 months)	0.65
<b>IDH mutation, % (n)</b>	17 (5/30)	0 (0/14)	0.16
<b>ATRX loss, % (n)</b>	15 (8/54)	26 (8/31)	0.25
<b>pTERT mutation, % (n)</b>	52 (27/52)	19 (5/27)	0.007
<b>C228T</b>	20	4	
<b>C250T</b>	7	1	
<b>MGMT methylation, % (n)</b>	38 (3/8)	0 (0/5)	0.23
<b>FGFR1 mutation, % (n)</b>	13 (6/46)	26 (6/23)	0.19
<b>EGFR amplification, % (n)</b>	23 (7/31)	0 (0/22)	0.03
<b>P16 loss, % (n)</b>	32 (10/31)	5 (1/22)	0.02
<b>BRAF mutation, % (n)</b>	0 (0/43)	0 (0/24)	NA

Abbreviations: CHT = chemotherapy; CI = confidence interval; OS = overall survival; RT = radiotherapy.

mutation but no clinical information on location<sup>28</sup> and 1 1p/19q-codeleted IDH1 mutant frontal oligodendroglioma with NS46K mutation.<sup>29</sup>

**Figure 3** Correlation matrix of the major variables



The variables are indicated in blue when associated, in purple when inversely associated. Circle diameter reflects the strength of the correlation ( $r$  value). KPS = Karnofsky performance status.

We then performed a hierarchical clustering on the clinical, histologic, radiologic, and molecular alterations (figure e-2, links. lww.com/WNL/A510). This table shows that *IDH1* mutations cluster with the brainstem location, and *FGFR1* mutations cluster with extinction of *ATRX* expression ( $p = 0.02$ ), while *CDKN2A* loss clusters with *EGFR* ( $p = 0.0004$ ) and *TERTp* ( $p = 0.01$ ) mutations. Grade, radiologic necrosis, and contrast enhancement clustered together. All the variables are reported in a correlation matrix (figure 3) showing the positive (blue) and negative (red) associations. *FGFR1* mutations tended to be associated with younger age ( $p = 0.06$ ), as *ATRX* loss ( $p = 0.05$ ), while *TERTp* mutation ( $p = 0.1$ ) and *EGFR* amplification ( $p = 0.05$ ) tended to be associated with older age.

### Survival analysis

OS was 17.3 months (95% confidence interval [CI] 14.7–23.0 months) for the whole group, 17.3 months (95% CI 14.5–27.5 months) for the thalamic gliomas, 16.3 months (95% CI 11.6–52.2 months) for brainstem, 19.7 months (95% CI 8.3–NA months) for cerebellar, and 23.0 months (95% CI 9.3–NA months) for spinal gliomas. Out of the main clinical prognostic factors validated in gliomas,<sup>30–32</sup> only poor Karnofsky performance status ( $p = 0.0007$ ), high tumor grade ( $p = 0.01$ ), % of Ki67 labeling ( $p = 0.003$ ), and presence of contrast enhancement ( $p = 0.007$ ), but not age ( $p = 0.11$ ), were associated with poor survival.

According to the 2016 WHO classification, all K27M-mutated diffuse MLG were considered as grade IV tumors. However, the presence of histone H3K27M mutation did not correlate with a worse prognosis in our patients (table 2, figure 4A): 23 (95% CI 16.4–45.0) vs 15 months (95% CI 11.3–21.3),  $p = 0.7$ .

**Table 2** Median survival according to the main genetic alterations, with log-rank test *p* value (2-sided)

Genetic alteration	Median OS (95% CI), mo, alteration present	Median OS (95% CI), mo, alteration absent	<i>p</i> Value
H3K27M mutation	23 (16.4–45.0)	15 (11.3–21.3)	0.7
FGFR1 mutation	45 (14.5–NA)	13.8 (8.3–18.6)	0.01
TERTp mutation	9.1 (5.7–14.8)	24.2 (18.6–45.0)	<0.0001
p16/CDKN2A deletion	9.9 (4.3–NA)	23.8 (16.9–46.1)	0.0001
EGFR amplification	4.3 (1.3–NA)	23.8 (16.2–40.4)	<0.0001
IDH1 mutation	54.1 (51.3–NA)	15.9 (12.2–19.3)	0.05
ATRX loss	19.7 (14.5–NA)	15.0 (11.6–18.5)	0.05
Loss of chromosome 10p	9.15 (2.0–NA)	32.0 (18.6–54.1)	0.016
Loss of chromosome 10q	9.8 (3.8–NA)	29.2 (18.0–52.2)	<0.0001
Loss of chromosome 9p	14.8 (9.1–NA)	32.0 (18.5–53.5)	0.03

Abbreviations: CI = confidence interval; OS = overall survival.

We therefore investigated whether grading, according to the WHO 2007 classification, had a prognostic relevance in H3K27M mutant gliomas and found no difference of survival between WHO 2007 grade II to grade IV tumors, suggesting that histone H3 mutant MLG represents a separated entity in adult patients (figure e-3a, [links.lww.com/WNL/A510](#)). We hypothesized that prognostic effect of histone H3K27M could have been masked by the association with younger age, which is a favorable prognostic factor. We therefore entered the 2 variables in a Cox model: however, neither K27M ( $p = 0.2$ ) nor age ( $p = 0.07$ ) was associated with poor survival.

Finally, by stratifying for single locations, we found that K27M mutations were associated with longer OS in thalamic MLG as compared to wild-type tumors: 29.2 (95% CI 24.2–NA) vs 14.4 months (95% CI 9.5–33.6),  $p = 0.04$  (figure e-3b, [links.lww.com/WNL/A510](#)). In brainstem tumors, after exclusion of IDH1-mutated cases, we observed a similar trend: 16.4 (9.8–NA) vs 9.1 months (0.2–13),  $p = 0.24$ . These data together suggest that K27M mutation in adults is not associated with poorer outcome, showing a different effect on survival compared to what is observed in the pediatric populations so far.

IDH1 mutation, loss of ATRX, and FGFR1 mutations were associated with longer survival (table 2 and figures 4, B and D, and e-4a, [links.lww.com/WNL/A510](#)). As the IDH1 mutation was present only in brainstem gliomas, we considered only this group and found it associated with a much longer OS: 54.1 (51.3–NA) vs 11.9 months (9.1–19.3),  $p = 0.03$  (figure e-5a). None of the IDH1 mutant brainstem MLG showed

contrast enhancement (0/5) vs 6/10 in IDH1 wild-type MLG ( $p < 0.05$ ) (figure e-5, b–d).

IDH wild-type–K27 wild-type patients showed the worst outcome (figure 4C). This group had frequent TERTp mutation, CDKN2A deletion, and EGFR amplification, all associated with shorter survival (table 2 and figure e-4, b–d, [links.lww.com/WNL/A510](#)). Loss of chromosomes 10q and 9p were also associated with poorer outcome (table 2 and figure e-4, e and f).

Finally, to identify independent prognostic factors, we performed multivariate survival analysis. Due to missing data, we limited the Cox model analysis to the most relevant (based on our data or current knowledge) and most documented measures: age, grade, K27M, TERTp, IDH, and FGFR1 status: only TERTp, IDH, and FGFR1 status were independent prognostic markers (table 3).

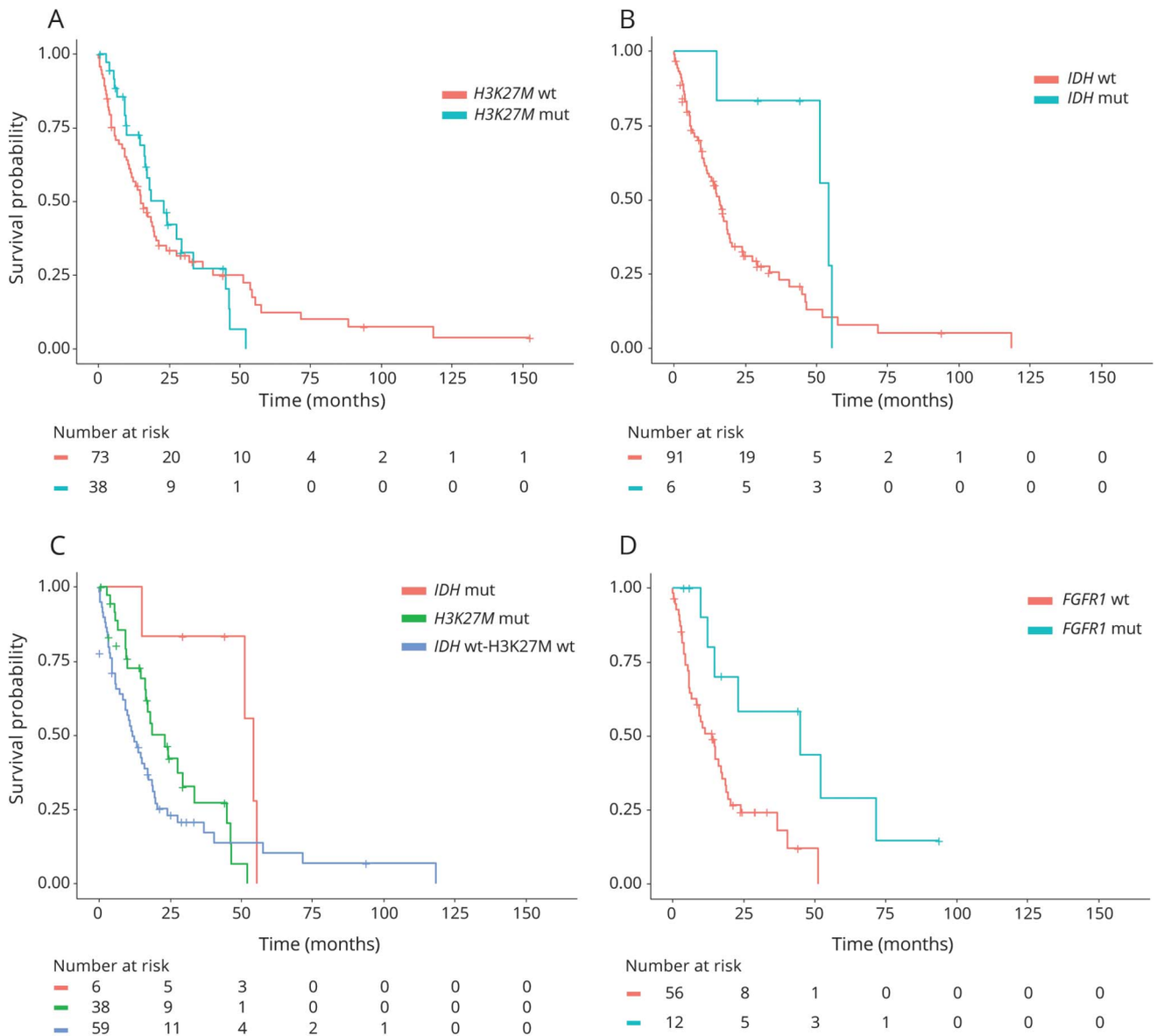
## Discussion

This retrospective study shows that adult MLG constitute a heterogeneous group with mostly high-grade neoplasms and clinical and histomolecular characteristics differing from both pediatric MLG and adult supratentorial high-grade gliomas. We identified 2 main subgroups based on the presence of histone H3 or IDH1 mutations, and identified hotspot FGFR1 mutations as the main potential target in this population.

The most prominent molecular subgroup consisted of histone H3 mutant MLG, in line with previous studies showing a strong association between histone H3 mutations and gliomagenesis of midline tumors.<sup>2–6,8,12–15,18,33</sup> However, we observed a lower prevalence of histone H3 mutations compared to pediatric high-grade thalamic and brainstem gliomas (70%–80%).<sup>2,4–6</sup> In addition, clinical and molecular characteristics of patients from our cohort with histone H3 mutant MLG differed from pediatric patients.<sup>2,4,6,33</sup> First, although histone H3 mutations were associated with younger age at diagnosis (median 33 years vs 53 years,  $p = 6 \times 10^{-5}$ ), we found H3 mutation in patients over 60 years, indicating that screening for histone H3 mutants should be considered in all MLG cases, regardless of age. Moreover, in contrast to pediatric gliomas, we found that histone H3 mutations were not associated with a worse prognosis compared to the other IDH wild-type gliomas<sup>12,14</sup>: in the largest group, i.e., the thalamic MLG, H3 mutations were even associated with better outcome. While H3 G34 and to a lesser extent K27 mutations have been associated with alternative lengthening of telomeres in pediatric tumors, as reflected by frequent ATRX inactivation,<sup>3,6,8,12,34</sup> we also found H3 mutation associated with TERTp mutation in 5 patients. Interestingly, either ATRX loss or TERTp mutation tended to be associated with poorer outcome in the subset of H3 mutated gliomas (OS 9.8 vs 24.2 months;  $p = 0.08$ ; data not shown).

The other main driver of alteration is IDH mutation, which occurs specifically in brainstem (6/24 vs 0/78 in other midline

**Figure 4** Kaplan-Meier survival curves



(A) Kaplan-Meier survival curves by H3K27M status (black = wild-type, blue = mutated). Median survival for mutated patients (blue): 23 months (95% confidence interval [CI] 16.4–45.0 months) vs median survival for wild-type patients (black): 15 months (95% CI 11.3–21.3 months),  $p = 0.7$ . Vertical bars indicate censored events. (B) Kaplan-Meier survival curves by IDH status (black = wild-type, red = mutated). Median survival for mutated patients (red): 54.1 months (95% CI 51.3–NA months) vs median survival for wild-type patients (black): 15.9 months (95% CI 12.2–19.3 months),  $p = 0.05$ . Vertical bars indicate censored events. (C) Kaplan-Meier survival curves by H3K27M and IDH status (black = both wild-type, blue = H3K27M mutated, red = IDH mutated). Median survival for H3K27M mutated patients (blue): 23.0 months (95% CI 16.4–45.0 months) vs median survival for IDH mutated patients (red): 54.1 months (95% CI 51.3–NA months) vs median survival for wild-type patients (black): 11.6 months (95% CI 9.1–18.5 months),  $p = 0.05$ . Vertical bars indicate censored events. (D) Kaplan-Meier survival curves by FGFR1 status (black = wild-type, red = mutated). Median survival for mutated patients (red): 45 months (95% CI 14.5–NA months) vs median survival for wild-type patients (black): 13.8 months (95% CI 8.3–18.6 months),  $p = 0.01$ . Vertical bars indicate censored events.

localizations). The majority of *IDH1* mutations in this cohort were non-R132H, indicating that the widely used anti-R132H immunohistochemistry is not appropriate in brainstem gliomas and has probably underestimated the proportion of *IDH1*-mutant tumors.<sup>35</sup> Systematic *IDH1* sequencing should be performed whenever feasible in brainstem gliomas. Indeed, as observed in hemispheric gliomas, *IDH1* mutation is associated with better outcome (OS 54.1 vs 15.9 months in our cohort), and may in part explain why adults with brainstem gliomas have better survival than children.<sup>19,35,36</sup> In addition to its prognostic

value, the recognition of *IDH1* mutations may open therapeutic opportunities, as inhibitors of *IDH1*-mutant enzymes have demonstrated antitumor potential and entered clinical trials for patients with *IDH1* mutations.<sup>37</sup>

In addition to *IDH1* mutation, we identified other potentially targetable hotspot mutations affecting N546 and K656 residues of *FGFR1* in 19% of our patients. In contrast to *IDH1* mutation, *FGFR1* mutations were not associated with any specific location or any molecular subgroup. Somatic *FGFR1*

**Table 3** Cox model analysis (due to missing data, we limited the analysis to the most relevant and most documented measures: age, grade, K27M, TERTp, IDH, and FGFR1 status)

	Coeff	OR (exp [coeff])	SE (coeff)	Z	p
Age	0.003	1.003	0.013	0.234	0.8
Grade	-0.191	0.826	0.288	-0.667	0.5
H3K27M	-0.159	0.853	0.484	-0.329	0.7
TERTp	1.367	3.926	0.347	3.936	8.3e-05
IDH	-1.756	0.173	0.652	-2.693	0.007
FGFR1	-2.117	0.120	0.799	-2.649	0.008

N546 and K656 mutations were previously reported in several other cancer types,<sup>36,38</sup> comprising a wide range of CNS tumors: rosette-forming glioneuronal tumors,<sup>39</sup> dysembryoplastic neuroepithelial tumors,<sup>40</sup> pilocytic astrocytomas,<sup>41,42</sup> diffuse leptomeningeal tumor with glial and neuronal markers,<sup>43</sup> and pediatric thalamic gliomas.<sup>3,18,19,43</sup> These studies analyzed isolated cases or small cohorts, which made difficult the assessment of the incidence of *FGFR1* mutations in this population. *FGFR1* N546 and K656 mutants lead to constitutive activation of *FGFR1* tyrosine kinase and downstream RAS-MAPK signaling, contributing to transformation,<sup>44</sup> and can be targeted by a number of oral highly specific *FGFR* inhibitors, currently under clinical evaluation,<sup>16</sup> such as BGJ398, AZD4547, JNJ-42756493, and TAS-120 (a novel irreversible *FGFR* inhibitor that demonstrated potent activity against N546K mutants),<sup>45</sup> opening new therapeutic opportunities in this population. *BRAF* V600E is another actionable target, which has been rarely associated with H3K27M in pediatric gangliogliomas and diffuse midline gliomas.<sup>7,46–48</sup> However, we did not find any *BRAF* V600 mutation in all the 68 sequenced patients.

Our study was restricted by the scarcity of available tumor material limiting both histopathologic and genomic analyses. Therefore, our genomic analyses, when possible, were restricted to a panel of selected molecular alterations, while several samples were exhausted at the time of this retrospective analysis. However, the majority of the patients could be assessed for *TERT* promoter status, which was not covered by previous whole-exome sequencing studies or was technically limited in whole-genome sequencing studies by the very low coverage due to the high GC content of the *TERT* promoter region.<sup>4,19</sup> As observed in adult supratentorial glioblastomas, *TERTp* mutation, which affected 37% of our patients, was associated with older age, grade IV, poor survival, *EGFR* amplifications, loss of chromosome 10, and deletion of *CDKN2A*. Survival of these patients was particularly poor (OS 9.1 months vs 24.2 months for patients without *TERTp* mutations), even compared to adult patients with hemispheric glioblastomas followed in our institution (OS 13.5 months<sup>49</sup>), which may be related to the inability to resect these tumors. *TERT* promoter appears

therefore a prognosticator biomarker of poor survival in patients with adult MLG.

Our findings reinforce the need for histologic confirmation and molecular analyses in adult patients with MLG, given the significant heterogeneity among midline tumors and the presence of recurrent potentially targetable molecular alterations in these often-fatal diseases. While *IDH1* and *TERT* promoter mutations may assist treatment stratification in adult patients with MLG, we show here that histone H3 mutations do not confer worse prognosis in adult patients with MLG. Our finding of frequent and potentially targetable *FGFR1* mutations has important clinical implications in the current context of anti-*FGFR* trials, and further reinforces the need for molecular analyses. In this population characterized by a risk for biopsies, the development of liquid biopsy allowing a noninvasive assessment of these few mutational hotspots therefore would be particularly relevant.

### Author contributions

A.P., A.L.D.S., and M.S. designed the study. A.P., M.T., A.L.D.S., and M.S. wrote the manuscript. G.B., E.T., S.M., and Y.S. performed the gene analysis. F.B. performed the histologic analysis and immunohistochemistry. A.P. and J.S. analyzed the MRI. Clinical data were collected and analyzed by A.P., M.S., and A.L.D.S. All authors contributed to the data analysis and interpretation and read and approved the manuscript.

### Acknowledgment

The authors thank Anne-Marie Lekieffre, Muriel Brandel, Armelle Rametti, Ines Detrait, and Amithys Rahimian for database management and technical support.

### Study funding

Supported by the Institut National du Cancer (INCa; PRT-K15-061), the Ligue Nationale Contre le Cancer, Comité de Paris, and the “Princesse Margot” association. A.L.D.S. received support from “Premio Riquier” and a donation in the memory of Olivier Ribes.

### Disclosure

The authors report no disclosures relevant to the manuscript. Go to Neurology.org/N for full disclosures.

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