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Characteristics of diffuse hemispheric gliomas, H3 G34-mutant in adults

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- TP, DM and FD designed the study.
- TP, MB, DP, CI, ET, VB, FBV, ECJM, JG substantially contributed to data acquisition.
- TP, MB, LPB, CI, DFB, AI, FB, EUC, JG, MG, DM, FD interpreted the data.
- TP and FD drafted the manuscript.
- All authors revised the manuscript critically for important intellectual content and approved the final version to be published.
ABSTRACT

Background: Diffuse hemispheric gliomas, H3 G34-mutant (DHG H3G34-mutant) constitute a distinct type of aggressive brain tumors. Although initially described in children, they can also affect adults. The aims were to describe the characteristics of DHG H3G34-mutant in adults and to compare them to those of established types of adult WHO grade IV gliomas.

Methods: The characteristics of 17 adult DHG H3G34-mutant, 32 H3.3 K27M-mutant diffuse midline gliomas (DMG), 100 IDH-wildtype and 36 IDH-mutant glioblastomas were retrospectively analyzed.

Results: Median age at diagnosis in adult DHG H3G34-mutant was 25 years (range: 19-33). All tumors were hemispheric. For 9 patients (56%), absent or faint contrast enhancement initially suggested another diagnosis than a high-grade glioma, and diffusion-weighted imaging seemed retrospectively more helpful to suspect an aggressive tumor than MR-spectroscopy and perfusion MRI. All cases were IDH-wildtype. Most cases were immunonegative for ATRX (93%) and Olig2 (100%) and exhibited MGMT promoter methylation (82%). The clinical and radiological presentations of adult DHG H3G34-mutant were different from those of established types of adult grade IV gliomas. Median overall survival of adult DHG H3G34-mutant was 12.4 months compared to 19.6 months (p=.56), 11.7 months (p=.45) and 50.5 months (p=0.006) in H3.3 K27M-mutant DMG, IDH-wildtype and IDH-mutant glioblastomas, respectively.
Conclusions: Adult DHG H3G34-mutant are associated with distinct characteristics compared to those of established types of adult WHO grade IV gliomas. The present study supports considering these tumors as a new type of WHO grade IV glioma in future classifications.

KEYWORDS
Diffuse Hemispheric Glioma; H3.3 G34R/V mutation; H3.3 K27M mutation; PNET; Survival
KEYPOINTS

1. Adult diffuse gliomas G34-mutant display a misleading radiological presentation.
2. Adult diffuse gliomas G34-mutant differ from other grade IV gliomas.
3. Adult diffuse gliomas G34-mutant have a poor outcome.

IMPORTANCE OF THE STUDY

In future classifications, diffuse gliomas H3 G34-mutant will be considered as a new WHO grade IV tumor type. Since the median age at diagnosis in diffuse hemispheric gliomas, H3 G34-mutant is around 20 years, half of the patients are followed by adult neuro-oncologists. The present study provides a detailed analysis of clinical, radiological and survival of a series of 17 adult diffuse hemispheric gliomas, H3 G34-mutant. The results highlight that adult diffuse hemispheric gliomas, H3 G34-mutant frequently display a misleading radiological presentation that can lead to a potentially detrimental diagnostic delay. Although these gliomas have a poor prognosis, similar to that of H3.3 K27M-mutant diffuse midline gliomas and IDH-wildtype glioblastomas, they display distinct characteristics compared to established types of WHO grade IV adult gliomas.
Abbreviations: ADC = Apparent Diffusion Coefficient, ALT = Alternative Lengthening of Telomeres, ATRX/DAXX = Alpha-Thalassemia X-linked mental Retardation/Death-domain Associated protein, DMG = Diffuse Midline Glioma, EGFR = Epidermal Growth Factor Receptor, FLAIR = FLuid-Attenuated Inversion Recovery, H3.3 = Histone protein H3.3, IDH = Isocitrate DeHydrogenase, MGMT = O6-MethylGunanine MethylTransferase, PNET = Primitive NeuroEctodermal Tumor, TERT = TElomerase Reverse Transcriptase
INTRODUCTION

Recurrent somatic gain-of-function mutations in the genes encoding the histone H3 variants H3.3 and H3.1 are encountered in and characterize two different types of high-grade gliomas that have initially been described in the pediatric population but can also affect adults.\textsuperscript{1–4} Lysine to methionine substitution at codon 27 in the \textit{H3F3A} or in the \textit{HIST1H3B/C} genes (H3.3 K27M mutation) are associated with aggressive diffuse midline gliomas and define “Diffuse midline glioma, H3.3 K27M-mutant” which corresponds to grade IV in the revised 2016 World Health Organization (WHO) classification.\textsuperscript{5} In contrast, glycine to arginine or valine substitutions at codon 34 in the \textit{H3F3A} gene (H3.3 G34 mutation) are associated with non-midline hemispheric high-grade gliomas.\textsuperscript{6–8} H3.3 K27M and H3.3 G34 mutations are mutually exclusive and exclusive with IDH mutations.\textsuperscript{1,9,10} H3.3 G34 mutations are about twice less frequent than H3.3 K27M mutations. They are encountered in less than 1% of all gliomas but in 15% of high-grade gliomas in adolescents and young adults.\textsuperscript{7,11,12}

In the 2016 WHO classification, H3.3 G34-mutant diffuse gliomas were not considered as a distinct tumor type and most cases are currently classified as IDH-wildtype glioblastomas.\textsuperscript{5} However, in future classifications, H3.3 G34-mutant diffuse gliomas will be individualized. In its 6\textsuperscript{th} update, cIMPACT-NOW (the consortium to Inform Molecular and Practical Approaches to CNS Tumor Taxonomy – Not Official WHO) indeed recommended to consider H3.3 G34-mutant diffuse gliomas as a new WHO grade IV tumor type that should be referred to as “Diffuse hemispheric glioma, H3.3 G34-mutant” and the grade will be given in arabic.\textsuperscript{13} In the fifth edition of the CNS WHO classification, which is in progress, the suggested name is consequently “Diffuse hemispheric glioma, H3G34-mutant”, grade 4. Since the median age at diagnosis in diffuse hemispheric gliomas, H3 G34-mutant is around 18 to 19 years, half of the patients are followed by adult neuro-oncologists.\textsuperscript{3,4,6,10} Yet, the characteristics and the outcome of adult diffuse hemispheric gliomas, H3 G34-mutant remain
to be fully described\textsuperscript{14-16}. Consequently, the aim of the present study was to report the characteristics of diffuse hemispheric gliomas, H3 G34-mutant in adults and compare them to those of established types of adult WHO grade IV gliomas.

**MATERIAL AND METHODS**

**Patient identification**

Adult patients (≥18-year-old) diagnosed with diffuse hemispheric glioma, H3 G34-mutant were retrospectively identified in 5 French University Hospitals (Lyon, Marseille, Nice, Paris and Toulouse) between January 2012 and May 2018. Their clinical, radiological, histological, and molecular characteristics were retrospectively reviewed.

The characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant were compared to those of retrospective contemporaneous series of H3.3 K27M-mutant diffuse midline gliomas (DMG), IDH-wildtype and IDH-mutant glioblastomas diagnosed in Lyon University Hospital according to the revised WHO 2016 classification. The inclusion criteria were: (1) age ≥ 18 years, (2) available MRI for radiological review, (3) available TERT promoter mutation status for IDH-wildtype glioblastomas.

Brain MRI scans were independently reviewed by 3 investigators (TP, CI and LPB) for the location of T2/fluid-attenuated inversion recovery (FLAIR) hyperintensity, the presence and type (fainted, nodular, or ring-like) of parenchymal contrast enhancement, the presence of Apparent Diffusion Coefficient (ADC) restriction, as well as abnormalities on proton magnetic resonance spectroscopy (1H-MRS) and, dynamic contrast-enhanced perfusion MRI. For patients with available MRI follow-up, tumor progression was defined according to RANO criteria\textsuperscript{17}. 

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Molecular analysis

DNA was extracted from formalin-fixed paraffin-embedded samples using a standard protocol (QIAmp DNA mini Kit, Qiagen, Courtaboeuf, France). After PCR amplification, the mutations of codon 132 of IDH1, codon 172 of IDH2, codons 27 and 34 of H3F3A, codon 28 of HIST1H3B, codon 600 of BRAF, as well as mutational hotspots C228 and C250 of the TERT promoter were investigated by single-nucleotide primer extension with the ABI PRISM SNaPshot Multiplex kit, Applied Biosystems (Foster City, California, USA).18

MGMT promoter methylation was assessed by pyrosequencing, which was performed using the PyroMark Q96 MGMT kit (Qiagen) on a PSQTM96 MA system (Biotage, Uppsala, Sweden)19. EGFR gene amplification was detected by using SYBR Green real-time quantitative polymerase chain reaction (qPCR) analysis (Absolute SYBR Green Rox Mix; Abgene, Paris, France) with the following set of primers EGFR-F: GTGCAGATCGCAAAGGTAATCAG, EGFR-R: GCAGACCGCATGTGAGGAT; hydrolysis probe: FAM CCCCTCCCCGTATCTC.20

Statistical analysis

Categorical comparisons were performed using the Fisher’s exact test or the Chi² test when the conditions of application were met. The nonparametric Wilcoxon test was used for quantitative variable as the hypothesis of normality of distribution was not verified. Overall survival and progression-free survival were represented with Kaplan–Meier plots and compared using the log-rank test.

The statistical tests were bilateral and the level of significance was set at 5% (p <0.05). Statistical analyses were conducted using SAS version 9.4 (SAS Institute Inc, Cary, NC, US).
Standard protocol approvals and registrations

Study design and manuscript organization were guided by the STROBE statement on cohort studies. The study was approved by the Institutional Review Board (local number 19-113). Informed consents were obtained from alive patients.

RESULTS

Patients and tumors characteristics

A total of 17 adult patients diagnosed with a diffuse hemispheric glioma, H3 G34-mutant were retrospectively identified. Their characteristics are summarized in Table 1. Median age at diagnosis was 25 years (range: 19-33). Most patients were aged ≤ 30 years (n=13, 76%). Clinical presentation consisted of epileptic seizure (n=9, 53%), focal deficit (n=5, 29%) and acute intracranial hypertension (n=3, 18%) that was imputable to a massive tumor hemorrhage in two patients (12%).

Histologically, all cases presented as a diffusely infiltrating high-grade glioma. Mitotic activity was observed in all cases, and microvascular proliferation and/or necrosis in all but one case (94%). All cases were IDH-wildtype (17/17). According to the 2016 WHO classification, one case was diagnosed as an IDH-wildtype anaplastic astrocytoma and 16 as IDH-wildtype glioblastomas. Among the latter cases, eight tumors displayed a PNET-like morphology (50%), four displayed an undifferentiated morphology (25%), three contained multinucleated giant cells (19%) and one displayed an oligodendroglial appearance (6%). These features are illustrated in Figure 1. Median Ki67 labeling was 42% (range: 15-95%). Olig2 immunostaining was completely negative in 13/16 cases (81%) and focally positive in 3/16 cases (19%). In these cases, however, Olig2 positivity corresponded to reactive glial cells and after review they were finally considered as negative. TP53 staining was positive...
(＞10%) in 14/16 cases (88%) and ATRX expression was lost in 13/14 cases (93%). Finally, all cases displayed ATRX loss and/or TP53 positive immunostaining. The H3F3A mutation was a G34R mutation in all cases and was conserved at recurrence in the two patients from the series who underwent a second surgery. All studied cases were BRAF wildtype (n=15/15) and lacked TERT promoter mutations (n=10/10). An EGFR amplification was present in 2/14 patients (14%) and MGMT promoter was methylated in 9/11 patients (82%).

**Radiological presentation**

Initial MRI scans were available for review for 16/17 patients. Radiological characteristics are summarized in Table 1. One patient (patient 6) underwent a brain MRI which was normal, three years before diagnosis. At diagnosis, all tumors had a hemispheric location and were monocentric. Midline involvement was observed in 4 patients but always as an extension of a primarily hemispheric tumor. Most cases demonstrated no or only a faint contrast-enhancement (n=11/15, 73%) and presented as cortico-subcortical (n=13/16, 81%), poorly delimited (n=9/16, 69%), infiltrative lesions that were mostly located in the fronto-parietal (n=11/16, 69%) or in the temporo-insular lobes (n=3/16, 19%). All but one tumor (n=12/13, 92%) harboured areas of ADC restriction on Diffusion Weighted Imaging. Only half of the cases in which perfusion MRI and/or MR spectroscopy were performed presented an elevated (＞1.7) intratumoral relative cerebral blood flow (50%, n=4/8) or a metabolic profile suggestive of an aggressive tumor (elevated Choline/N-Acetyl-Aspartate ratio ＞ 2 and/or of lipid/lactate peaks, 44% n=4/9).

For 6 patients, the initial radiological presentation led to suspect a high-grade glioma. For 10 cases, however, the young age of the patient, absent or faint contrast enhancement (n=9) or the presence of an isolated haemorrhage (n=1) led to suspect another diagnosis (Figure 2). In these patients, the initially suspected diagnoses were: low-grade glioma (n=3),
primary CNS lymphoma (n=2), multiple sclerosis (n=2), limbic encephalitis (n=1), metabolic disease (n=1) and ruptured arterio-venous malformation (n=1). Subsequently, in 8 patients the decision was made to perform an MRI follow-up leading to a median delay of 2.3 months before histological diagnosis. Retrospectively, in all but one of these patients, the initial MRI demonstrated focal ADC restriction that could have led to suspect an aggressive glioma. In contrast, an elevated intra-tumoral relative cerebral blood flow and an aggressive metabolic profile on MR spectroscopy were both present in only 2/8 patients (25%).

Among the 11 patients with initially absent or faint contrast-enhancement, 8 underwent a repeated MRI before (n=4) or after (n=4) histological diagnosis that demonstrated the apparition of a ring-like or nodular contrast enhancement in all of the patients after a median interval of 2.6 months (range: 0.8-5.3 months).

Treatment and outcome

A total of 12 patients underwent a biopsy (70%), one underwent a partial (6%) and four a complete resection (24%). One patient presented with a brain hematoma leading to neurological worsening after biopsy and one patient presented a lethal refractory raised intracranial pressure after biopsy. In the 15 remaining patients, post-surgical course was uneventful. Post-operative oncological treatment consisted of temozolomide radio-chemotherapy (n=13, 82%), radiotherapy alone (n=1, 6%), radiotherapy followed by PCV (procarbazine, CCNU, and vincristine) chemotherapy (n=1, 6%) and temozolomide chemotherapy alone (n=1, 6%). No severe adverse effect was reported during the oncological treatment and no patient developed drug-resistant epilepsy.

At last news, after a median follow-up of 12.7 months, tumor progression was diagnosed in 17 patients and 15 patients had died. The median progression-free survival was 8.8 months and the median overall survival was 12.4 months, in the entire series. For the 13
patients who were treated with temozolomide radio-chemotherapy, the median overall survival was 25 months. The pattern of tumor progression was analyzed for 15 patients and consisted of gliomatosis cerebri (n=9, 60%) with contralateral hemispheric (n=4, 27%) or infratentorial (n=5, 33%) infiltration, local recurrence (n=3, 20%), distant progression (n=3, 20%) and leptomeningeal dissemination (n=4, 26%). At progression, treatment consisted of chemotherapy alone (n=8, 53%), chemotherapy combined with cyberknife radiation (n=1, 8%) or conformational radiotherapy (n=1, 8%), surgery followed by chemotherapy (n=2), radiotherapy at a remote site (n=1, 8%) or palliative cares (n=3, 23%).

Comparison of adult diffuse hemispheric gliomas, H3 G34-mutant with established types of adult WHO grade IV gliomas

As shown in Table 2, several characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant differed significantly from those of established types of adult WHO grade IV gliomas. Compared to H3.3 K27M-mutant DMG, IDH-mutant and IDH-wildtype glioblastomas, diffuse hemispheric gliomas, H3 G34-mutant occurred in younger patients, more frequently involved the parietal lobe, more frequently presented as a haemorrhagic tumor at diagnosis, less frequently demonstrated contrast-enhancement and necrosis, more frequently displayed ADC restriction and less frequently expressed Olig2. In addition, TERT promoter mutations were lacking in diffuse hemispheric gliomas in contrast to IDH-wildtype glioblastomas while MGMT promoter methylation was more frequent in diffuse hemispheric gliomas, H3 G34-mutant than in H3.3 K27M-mutant DMG and IDH-wildtype glioblastomas. Median overall survival of adult diffuse hemispheric gliomas, H3 G34-mutant was 12.4 months compared to 19.6 months (p=.56), 11.7 months (p=.45) and 50.5 months (p=0.006) in H3.3 K27M-mutant DMG, IDH-wildtype glioblastomas and IDH-mutant glioblastomas, respectively (Figure 3). The comparison yielded similar results when the analysis was
restricted to H3.3 K27M-mutant DMG and IDH-mutant glioblastomas with comparable age (i.e. aged < 35 years, data not shown). In IDH-wildtype glioblastomas, the number of patients aged < 35 years was too limited to allow this analysis.

DISCUSSION

Diffuse hemispheric gliomas, H3 G34-mutant are rare brain tumors that affect children and adolescents but also adults. In the present study, that is the first dedicated to adult cases, we show that adult diffuse hemispheric gliomas, H3 G34-mutant frequently displayed a misleading radiological presentation, had distinct characteristics compared to established types of WHO grade IV adult gliomas and a poor prognosis.

Radiological presentation

The radiological characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant in the present series are consistent with those reported in previous studies, although these studies were not dedicated to adults (Table 3). In the present series, adult diffuse hemispheric gliomas, H3 G34-mutant were constantly located within cerebral hemispheres, frequently affected the parietal and temporal lobes, frequently demonstrated no or faint contrast-enhancement, not rarely presented a massive initial hemorrhage and, frequently presented an unusual presentation for a high-grade glioma. Actually, less than half of cases fulfilled the imaging criteria of high-grade glioma which led in some cases to a potentially detrimental diagnostic delay. Patients for whom the radiological presentation was misleading typically presented with an MRI aspect that has been previously described as “gliomatosis cerebri-like”, consisting of a non-contrast enhancing ill-defined infiltrating lesion that led to suspect another diagnosis than a high-grade glioma because of the young age of the patients. Non-tumoral diagnoses, including inflammatory, infectious, vascular
and metabolic diseases, could be initially suspected. Retrospectively, a striking feature in these patients that could have raised suspicions towards an aggressive hypercellular tumor was the presence of areas of profound focal ADC restriction. In addition to Diffusion-Weighted-Imaging, dynamic $^{18}$F-Fluoro-Ethyl-L-Tyrosine positron emission tomography (FET-PET) may be useful to pre-operatively identify non-contrast enhancing diffuse hemispheric gliomas, H3 G34-mutant as aggressive gliomas.

**Histo-molecular presentation**

At the histo-molecular level, it was evidenced herein that adult diffuse hemispheric gliomas, H3 G34-mutant could present a PNET-like or a GBM-like morphology, lacked Olig2 expression, were most frequently immunonegative for ATRX, immunopositive for TP53, frequently demonstrated a methylated MGMT promoter, and lacked TERT promoter mutations. These findings are consistent with those reported in previous series that included both pediatric and adult cases (Table 3). The absence of Olig2 expression is a hallmark of diffuse hemispheric gliomas, H3 G34-mutant. Yet, as observed in our series, reactive glial cells may wrongly suggest positivity in some cases. As highlighted in children, due to a frequent PNET-like morphology, adult diffuse hemispheric gliomas, H3 G34-mutant could have previously been misdiagnosed as CNS PNET, a now obsolete entity. Actually, in a young adult presenting with an IDH-wildtype hemispheric PNET-like tumor, the most likely diagnosis is a diffuse hemispheric glioma, H3 G34-mutant since other entities that can present as PNET-like tumors occur in younger patients. Importantly, in the present series, despite the young age of the patients and a high rate of MGMT promoter methylation, adult diffuse hemispheric gliomas, H3 G34-mutant had a poor prognosis. Despite many similarities, it has been shown that pediatric and adult H3.3 K27M-mutant DMG present significant differences. Compared to pediatric cases, adult H3.3 K27M-mutant DMG more
frequently affect the thalamus and less frequently the pons, almost exclusively harbour
H3F3A mutations and have a better prognosis. It remains to be determined whether this is
also the case in diffuse hemispheric gliomas, H3 G34-mutant, however, these gliomas seem
to be a more homogeneous entity than H3.3 K27M-mutant DMG. Age at diagnosis is
narrower in H3 G34-mutant gliomas than in H3.3 K27M-mutant DMG and in a previous
study neither histological appearance nor molecular characteristics were associated with age.6
Also consistent with this hypothesis, as detailed in Table 3, the characteristics of adult diffuse
hemispheric gliomas, H3 G34-mutant in the series herein seemed very similar to those
reported in previous studies that included both adult and pediatric cases3,4,6,21,22,27 or only
pediatric cases.3,4,6,21,22,27 However, analysis of larger series will be necessary to assess
whether there are age-related differences in H3 G34-mutant gliomas.

Outcomes and therapeutic opportunities

The median overall survival of adult diffuse hemispheric gliomas, H3 G34-mutant
herein is consistent with the median overall survival ranging from 12 to 36 months reported
in previous series (Table 3).3,4,6,12,21,22,26 Although several studies suggested that diffuse
hemispheric gliomas, H3 G34-mutant had a better prognosis than H3.3 K27M-mutant
DMG4,10,30,31 the prognosis of these two tumour types was similar in the present series and
was also similar to that of IDH-wildtype glioblastomas. However, the clinical, radiological
and histo-molecular characteristics of diffuse hemispheric gliomas, H3 G34-mutant were
very different from those of H3.3 K27M-mutant DMG, IDH-mutant and IDH-wildtype
glioblastomas. These findings support that diffuse hemispheric gliomas, H3 G34-mutant
should be considered as a new WHO grade IV tumor type in future classifications.13

Beside gliomas, H3.3 G34 mutations are found in the majority of giant cell tumors of
the bone (92%) and some osteosarcomas (5%).32,33 Although the oncogenesis of diffuse
hemispheric gliomas, H3 G34-mutant remains to be fully understood, some therapeutic opportunities have already been identified. Histones pack DNA and are involved in transcriptional regulation and telomeres stability through post-translational modifications. Mutations of some histone residues, such as the H3.3 K27 and K36 residue, generate conformational changes or steric inhibition that prevent post-translational modifications and trigger deep epigenetic reprogramming. In contrast to the H3.3 K27 residue, the H3.3 G34 residue does not undergo post-translational modifications. However, the H3.3 G34 mutations alter accessibility of H3 K36 to histone methyltransferases affecting H3 K36 methylation and subsequently gene expression. H3.3 G34 mutations have been shown to result in the upregulation of transcription factor MYCN and to impair H3 K36 methylation by SETD2 which promotes aberrant PRC2 activity and drives tumor progression. H3.3 G34 mutations also promote genomic instability by blocking the interaction between H3 K36 and the mismatch repair protein MutSα. In addition, it has been shown that diffuse hemispheric gliomas, H3 G34-mutant are characterized by a global DNA hypomethylation and ATRX/DAXX mutations, resulting in alternative lengthening of telomeres (ALT). Recently, it has been shown that although considered gliomas, H3 G34-mutant tumors arise in specific interneuron progenitors, in which G34 mutations prevent neuronal differentiation and that PDGFRA signalling plays a major role in these tumors. PDGFRA activating mutations (mostly in the extracellular domain) which are present in 40% of the cases at diagnosis and, in 80% of the cases at recurrence, promote astroglial features and are potent oncogenic drivers. In the future, targeting MYCN, DNA hypomethylation, ALT and PDGFRA signaling could constitute new treatment strategies in diffuse hemispheric gliomas, H3 G34-mutant. Targeting MYCN upregulation could be achieved by inhibiting kinases such as CHK1 and AURKA which stabilize MYCN or by using BRD3 and BRD4 inhibitors which play essential roles in the transcription of
Regarding PDGFRA signaling inhibition, given the high rate of extracellular mutations, it remains to be determined whether it could be best achieved through PDGFRA inhibitors or through inhibitors targeting downstream MAPK activation. In H3.3 K27M-mutant DMG histone deacetylase inhibitors and the selective antagonist of dopamine receptor DRD2/3 ONC201 are promising therapeutic strategies that are currently being evaluated in clinical trials. Whether these strategies may benefit patients with H3 G34-mutant gliomas remains to be determined.

Study limitations

Other than the limited sample-size and its retrospective design, the present study is limited by the absence of a comprehensive molecular analysis, the absence of multi-modal imaging in all of the patients, and the absence of comparative pediatric series. In addition, the results will need to be validated in an independent and larger series. Nevertheless, the study provides for the first time a picture of adult diffuse hemispheric gliomas, H3 G34-mutant characteristics.

CONCLUSIONS

Adult diffuse hemispheric gliomas, H3 G34-mutant are associated with distinct characteristics compared to those of established types of adult WHO grade IV gliomas. The present study supports considering these tumors as a new type of WHO grade IV glioma in future classifications. If further studies confirm that adult and pediatric diffuse hemispheric gliomas, H3 G34-mutant share similar characteristics, it may be reasonable to include patients in the same clinical trials, irrespective of age.
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REFERENCES

1. Wu G, Broniscer A, McEachron TA, et al. Somatic histone H3 alterations in pediatric diffuse intrinsic pontine gliomas and non-brainstem glioblastomas. *Nat Genet*. 2012;44(3):251-253. doi:10.1038/ng.1102

2. Wu G, Diaz AK, Paugh BS, et al. The genomic landscape of diffuse intrinsic pontine glioma and pediatric non-brainstem high-grade glioma. *Nat Genet*. 2014;46(5):444-450. doi:10.1038/ng.2938

3. Schwartzentruber J, Korshunov A, Liu X-Y, et al. Driver mutations in histone H3.3 and chromatin remodelling genes in paediatric glioblastoma. *Nature*. 2012;482(7384):226-231. doi:10.1038/nature10833

4. Sturm D, Witt H, Hovestadt V, et al. Hotspot mutations in H3F3A and IDH1 define distinct epigenetic and biological subgroups of glioblastoma. *Cancer Cell*. 2012;22(4):425-437. doi:10.1016/j.ccr.2012.08.024

5. Louis DN, Perry A, Reifenberger G, et al. The 2016 World Health Organization Classification of Tumors of the Central Nervous System: a summary. *Acta Neuropathol (Berl)*. 2016;131(6):803-820. doi:10.1007/s00401-016-1545-1

6. Korshunov A, Capper D, Reuss D, et al. Histologically distinct neuroepithelial tumors with histone 3 G34 mutation are molecularly similar and comprise a single nosologic entity. *Acta Neuropathol (Berl)*. 2016;131(1):137-146. doi:10.1007/s00401-015-1493-1

7. Yoshimoto K, Hatae R, Sangatsuda Y, et al. Prevalence and clinicopathological features of H3.3 G34-mutant high-grade gliomas: a retrospective study of 411 consecutive glioma cases in a single institution. *Brain Tumor Pathol*. 2017;34(3):103-112. doi:10.1007/s10014-017-0287-7

8. Korshunov A, Ryzhova M, Hovestadt V, et al. Integrated analysis of pediatric glioblastoma reveals a subset of biologically favorable tumors with associated molecular prognostic markers. *Acta Neuropathol (Berl)*. 2015;129(5):669-678. doi:10.1007/s00401-015-1405-4

9. Pathak P, Jha P, Purkait S, et al. Altered global histone-trimethylation code and H3F3A-ATRX mutation in pediatric GBM. *J Neurooncol*. 2015;121(3):489-497. doi:10.1007/s11060-014-1675-z

10. Fontebasso AM, Liu X-Y, Sturm D, Jabado N. Chromatin remodeling defects in pediatric and young adult glioblastoma: a tale of a variant histone 3 tail. *Brain Pathol Zurich Switz*. 2013;23(2):210-216. doi:10.1111/bpa.12023

11. Neumann JE, Dorostkar MM, Korshunov A, et al. Distinct Histomorphology in Molecular Subgroups of Glioblastomas in Young Patients. *J Neuropathol Exp Neurol*. 2016;75(5):408-414. doi:10.1093/jnen/nlw015
12. Roux A, Pallud J, Saffroy R, et al. High-grade gliomas in adolescents and young adults highlight histomolecular differences from their adult and pediatric counterparts. *Neuro-Oncol*. 2020;22(8):1190-1202. doi:10.1093/neuonc/noaa024

13. Louis DN, Wesseling P, Aldape K, et al. cIMPACT-NOW update 6: new entity and diagnostic principle recommendations of the cIMPACT-Utrecht meeting on future CNS tumor classification and grading. *Brain Pathol Zurich Switz*. 2020;30(4):844-856. doi:10.1111/bpa.12832

14. Lee J, Solomon DA, Tihan T. The role of histone modifications and telomere alterations in the pathogenesis of diffuse gliomas in adults and children. *J Neurooncol*. 2017;132(1):1-11. doi:10.1007/s11060-016-2349-9

15. Puntonet J, Dangouloff-Ros V, Saffroy R, et al. Histорadiological correlations in high-grade glioma with the histone 3.3 G34R mutation. *J Neuroradiol J Neuroradiol*. Published online March 2, 2018. doi:10.1016/j.neurad.2018.02.006

16. Zhang R-Q, Shi Z, Chen H, et al. Biomarker-based prognostic stratification of young adult glioblastoma. *Oncotarget*. 2016;7(4):5030-5041. doi:10.18632/oncotarget.5456

17. Wen PY, Macdonald DR, Reardon DA, et al. Updated response assessment criteria for high-grade gliomas: response assessment in neuro-oncology working group. *J Clin Oncol Off J Am Soc Clin Oncol*. 2010;28(11):1963-1972. doi:10.1200/JCO.2009.26.3541

18. Meyronet D, Esteban-Mader M, Bonnet C, et al. Characteristics of H3 K27M-mutant gliomas in adults. *Neuro-Oncol*. 2017;19(8):1127-1134. doi:10.1093/neuonc/now274

19. Walter T, van Brakel B, Vercherat C, et al. O6-Methylguanine-DNA methyltransferase status in neuroendocrine tumours: prognostic relevance and association with response to alkylating agents. *Br J Cancer*. 2015;112(3):523-531. doi:10.1038/bjc.2014.660

20. Idbaih A, Crinière E, Marie Y, et al. Gene amplification is a poor prognostic factor in anaplastic oligodendrogliomas. *Neuro-Oncol*. 2008;10(4):540-547. doi:10.1215/15228517-2008-022

21. Mackay A, Burford A, Carvalho D, et al. Integrated Molecular Meta-Analysis of 1,000 Pediatric High-Grade and Diffuse Intrinsic Pontine Glioma. *Cancer Cell*. 2017;32(4):520-537.e5. doi:10.1016/j.ccell.2017.08.017

22. Mackay A, Burford A, Molinari V, et al. Molecular, Pathological, Radiological, and Immune Profiling of Non-brainstem Pediatric High-Grade Glioma from the HERBY Phase II Randomized Trial. *Cancer Cell*. 2018;33(5):829-842.e5. doi:10.1016/j.ccell.2018.04.004

23. Vettermann FJ, Felsberg J, Reifenberger G, et al. Characterization of Diffuse Gliomas With Histone H3-G34 Mutation by MRI and Dynamic 18F-FET PET. *Clin Nucl Med*. 2018;43(12):895-898. doi:10.1097/RLU.0000000000002300

24. Pramanik PP, Parmar HA, Mammoser AG, et al. Hypercellularity Components of Glioblastoma Identified by High b-Value Diffusion-Weighted Imaging. *Int J Radiat Oncol Biol Phys*. 2015;92(4):811-819. doi:10.1016/j.ijrobp.2015.02.058
25. Ebrahimi A, Skardelly M, Bonzheim I, et al. ATRX immunostaining predicts IDH and H3F3A status in gliomas. Acta Neuropathol Commun. 2016;4(1):60. doi:10.1186/s40478-016-0331-6

26. Chen CCL, Deshmukh S, Jessa S, et al. Histone H3.3G34-Mutant Interneuron Progenitors Co-opt PDGFRA for Gliomagenesis. Cell. 2020;183(6):1617-1633.e22. doi:10.1016/j.cell.2020.11.012

27. Gessi M, Gielen GH, Hammes J, et al. H3.3 G34R mutations in pediatric primitive neuroectodermal tumors of central nervous system (CNS-PNET) and pediatric glioblastomas: possible diagnostic and therapeutic implications? J Neurooncol. 2013;112(1):67-72. doi:10.1007/s11060-012-1040-z

28. Sturm D, Orr BA, Toprak UH, et al. New Brain Tumor Entities Emerge from Molecular Classification of CNS-PNETs. Cell. 2016;164(5):1060-1072. doi:10.1016/j.cell.2016.01.015

29. Schulte JD, Buerki RA, Lapointe S, et al. Clinical, radiologic, and genetic characteristics of histone H3 K27M-mutant diffuse midline gliomas in adults. Neuro-Oncol Adv. 2020;2(1):vdaa142. doi:10.1093/noajnl/vdaa142

30. Sturm D, Bender S, Jones DTW, et al. Paediatric and adult glioblastoma: multiform (epi)genomic culprits emerge. Nat Rev Cancer. 2014;14(2):92-107. doi:10.1038/nrc3655

31. Khuong-Quang D-A, Buczkowicz P, Rakopoulos P, et al. K27M mutation in histone H3.3 defines clinically and biologically distinct subgroups of pediatric diffuse intrinsic pontine gliomas. Acta Neuropathol (Berl). 2012;124(3):439-447. doi:10.1007/s00401-012-0998-0

32. Behjati S, Tarpey PS, Presneau N, et al. Distinct H3F3A and H3F3B driver mutations define chondroblastoma and giant cell tumor of bone. Nat Genet. 2013;45(12):1479-1482. doi:10.1038/ng.2814

33. Koelsche C, Schrumpf D, Tharun L, et al. Histone 3.3 hotspot mutations in conventional osteosarcomas: a comprehensive clinical and molecular characterization of six H3F3A mutated cases. Clin Sarcoma Res. 2017;7: doi:10.1186/s13569-017-0075-5

34. Kallappagoudar S, Yadav RK, Lowe BR, Partridge JF. Histone H3 mutations--a special role for H3.3 in tumorigenesis? Chromosoma. 2015;124(2):177-189. doi:10.1007/s00412-015-0510-4

35. Martin C, Zhang Y. The diverse functions of histone lysine methylation. Nat Rev Mol Cell Biol. 2005;6(11):838-849. doi:10.1038/nrm1761

36. Vanan MI, Underhill DA, Eisenstat DD. Targeting Epigenetic Pathways in the Treatment of Pediatric Diffuse (High Grade) Gliomas. Neurother J Am Soc Exp Neurother. 2017;14(2):274-283. doi:10.1007/s13311-017-0514-2

37. Shi L, Shi J, Shi X, Li W, Wen H. Histone H3.3 G34 Mutations Alter Histone H3K36 and H3K27 Methylation In Cis. J Mol Biol. 2018;430(11):1562-1565. doi:10.1016/j.jmb.2018.04.014
38. Bjerke L, Mackay A, Nandhabalan M, et al. Histone H3.3 mutations drive pediatric glioblastoma through upregulation of MYCN. Cancer Discov. 2013;3(5):512-519. doi:10.1158/2159-8290.CD-12-0426

39. Huang M, Weiss WA. G34, another connection between MYCN and a pediatric tumor. Cancer Discov. 2013;3(5):484-486. doi:10.1158/2159-8290.CD-13-0126

40. Jain SU, Khazaei S, Marchione DM, et al. Histone H3.3 G34 mutations promote aberrant PRC2 activity and drive tumor progression. Proc Natl Acad Sci. 2020;117(44):27354-27364. doi:10.1073/pnas.2006076117

41. Fang J, Huang Y, Mao G, et al. Cancer-driving H3G34V/R/D mutations block H3K36 methylation and H3K36me3-MutSa interaction. Proc Natl Acad Sci U S A. 2018;115(38):9598-9603. doi:10.1073/pnas.1806355115

42. Puissant A, Frumm SM, Alexe G, et al. Targeting MYCN in neuroblastoma by BET bromodomain inhibition. Cancer Discov. 2013;3(3):308-323. doi:10.1158/2159-8290.CD-12-0418

43. Shahbazi J, Liu PY, Atmadibrata B, et al. The Bromodomain Inhibitor JQ1 and the Histone Deacetylase Inhibitor Panobinostat Synergistically Reduce N-Myc Expression and Induce Anticancer Effects. Clin Cancer Res. 2016;22(10):2534-2544. doi:10.1158/1078-0432.CCR-15-1666

44. Grasso CS, Tang Y, Truffaux N, et al. Functionally defined therapeutic targets in diffuse intrinsic pontine glioma. Nat Med. 2015;21(6):555-559. doi:10.1038/nm.3855

45. Chi AS, Tarapore RS, Hall MD, et al. Pediatric and adult H3 K27M-mutant diffuse midline glioma treated with the selective DRD2 antagonist ONC201. J Neurooncol. 2019;145(1):97-105. doi:10.1007/s11060-019-03271-3
FIGURE CAPTIONS

Figure 1: Histological characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant (Scale bar = 100 µm)

A. PNET-like morphology, characterized by small round cells, with anaplastic features including endothelial cell proliferation, negative Olig2 immunostaining, positive TP53 staining and ATRX loss

B. Undifferentiated glial morphology, characterized by irregularly-shaped nuclei and poorly-delimited cytoplasms, with anaplastic features including endothelial cell proliferation, positive Olig2 (reactive glial cells), ATRX and TP53 stainings

C. Monstrocellular morphology, characterized by multinucleated cells, with anaplastic features including endothelial cell proliferation, necrosis, negative Olig2 immunostaining, positive TP53 staining and ATRX loss

D. Oligoid morphology, characterized by small cells with a clear peri-nuclear halo, with anaplastic features including endothelial cell proliferation and 2 mitoses, positive Olig2 (reactive glial cells) and TP53 immunostaining and ATRX loss

Figure 2: Radiological characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant

For each case, T1-weighted contrast-enhanced MRI (left), T2-weighted FLAIR MRI (middle) and ADC map (right) are shown.

A. Frontal cortico-subcortical, poorly delimited, diffuse hemispheric glioma, H3 G34-mutant without contrast enhancement, displaying focal ADC restriction and a metabolic profile on MR spectroscopy demonstrating no Choline/Creatinine ratio
increase but an elevated Choline/N-Acetyl-Aspartate ratio (2.35) and a lipid peak (patient 10). In this patient, a low-grade glioma was initially suspected.

B, C, D. Three other cases of diffuse hemispheric gliomas, H3 G34-mutant with a misleading presentation, including absent or faint contrast enhancement and a focal ADC restriction. These characteristics led to suspect at first diagnosis multiple sclerosis (B -Patient 5, D – Patient 3), limbic encephalitis (C – Patient 4) or low-grade glioma (D – Patient 3)

E. Fronto-callosal diffuse hemispheric glioma, H3 G34-mutant with a presentation suggestive of high-grade glioma, including necrosis area and an evident contrast enhancement (patient 6). Three years earlier, this patient underwent an MRI which was normal.

**Figure 3: Kaplan-Meier survival analysis of diffuse hemispheric gliomas, H3 G34-mutant, H3.3 K27M-mutant diffuse midline gliomas, IDH-mutant glioblastomas, and IDH-wildtype glioblastomas patients**

Patients at risk are indicated below Figure 3.
Table 1: Characteristics of the 17 included adult patients diagnosed with diffuse hemispheric glioma, H3 G34-mutant

| Patients | Sex | Age (years) | Suspected diagnosis | Tumor location | Gadolinium enhancement | Presence of necrotic areas | Restriction of diffusion | Surgical manipulation | Microscopic Ki67 index % | Olig2 staining | TP53 staining | IDH mutation | ATRX loss | EGFR amplification | MGMT methylation | Progression-free survival (months) | Overall survival (months) |
|----------|-----|-------------|---------------------|----------------|------------------------|---------------------------|--------------------------|------------------------|--------------------------|----------------|---------------|--------------|------------|-----------------|----------------|-----------------------------|--------------------------|
| 1        | M   | 9           | HG                  | P              | Ye                     | Y                        | N                        | T                      | Monomorphic            | 3              | 0             | 80           | N          | N               | Y             | 9                           | 7                         |
| 2        | F   | 5           | HG                  | F              | Ye                     | N                        | N                        | T                      | Polymorphic            | 6              | 0             | 80           | N          | Y               | Y             | 4                           | 0                         |
| 3        | F   | 3           | MS                  | F              | No                     | N                        | N                        | P                      | Polymorphic            | 4              | 0             | 60           | N          | N               | Y             | 9                           | 3                         |
| 4        | M   | 3           | LE                  | T              | No                     | N                        | N                        | P                      | Polymorphic            | 7              | 0             | 10           | N          | Y               | Y             | 4                           | 1                         |
| 5        | F   | 9           | LG                  | F              | No                     | N                        | N                        | B                      | Polymorphic            | 2              | 0             | 10           | N          | N               | -             | -                           | -                         |
| 6        | F   | 0           | HG                  | F              | Ye                     | Y                        | Ye s(F)                  | Undiff                | Polymorphic            | 2              | 0             | 80           | N          | N               | Y             | 9                           | 7                         |
| 7        | M   | 2           | H/H                 | T              | Ye s(F)                | Ye s(F)                  | Ye s(F)                 | Undiff                | Polymorphic            | 4              | 0             | 10           | N          | Y               | Y             | 2                           | 3                         |
| 8        | M   | 0           | PCN                 | F              | Ye s(F)                | Ye s(F)                  | Ye s(F)                 | Olig2oid              | Polymorphic            | 6              | 0             | 10           | N          | Y               | Y             | 9                           | 3                         |
| 9        | M   | 1           | VM                  | T              | Monomorphic            | Ye s(F)                  | Ye s(F)                 | Undiff                | Polymorphic            | 9              | 0             | 10           | N          | Y               | Y             | 2                           | 3                         |
| 10       | M   | 1           | LG                  | F              | No                     | N                        | N                        | B                      | Anaplastic             | 3              | 0             | 10           | N          | N               | Y             | 2                           | 5                         |
| 11       | F   | 7           | LG                  | F              | No                     | N                        | No                      | B                      | Anaplastic             | 3              | 0             | 10           | N          | N               | Y             | 2                           | 5                         |
| 12       | M   | 3           | HG                  | T              | Ye s(F)                | Ye s(F)                  | Ye s(F)                 | Undiff                | Polymorphic            | 4              | 0             | 90           | N          | N               | N             | 7                           | 4                         |
| 13       | M   | 2           | A                   | F              | Ye                     | N                        | N                        | B                      | Polymorphic            | 5              | 0             | 90           | N          | Y               | -             | 8                           | 1                         |
|   |   |   | Sex | F: Female, M: Male; | Initially suspected diagnosis. A: Abscess, AVM: Arterio-venous Malformation, H: Hematoma, HGG: High-grade glioma, LE: Limbic encephalitis, LGG: Low-grade glioma, MD: Metabolic disease, MS: Multiple sclerosis, PCNSL: Primary central nervous system lymphoma; | Tumor location. FC: fronto-callosal, FP: fronto-parietal, PO: parieto-occipital, TI: temporo-insular, a indicates tumors that were only cortical at diagnosis, b indicates midline invasion; | Gadolinium enhancement. (F): Faint; | Restriction of Diffusion. (F): Focal; | Surgical management. B: Biopsy, PR: Partial Resection, TR: Total Resection; | Microscopic aspect. A-III: Grade III Astrocytoma, Monstro: Monstrocellular glioblastoma, Oligoid: Glioblastoma with oligoid morphology, PNET: Glioblastoma with Primitive Neuro-Ectodermic Tumor-like foci, Undiff: Glioblastoma with undiffernciated morphology; | Olig2 staining. * indicates cases containing Olig2-imunopositive reactive glial cells | Overall survival. c indicates patients who are still alive. |}

|   |   |   |   |   |   |   |   |   |   |   |   |
|---|---|---|---|---|---|---|---|---|---|---|---|
| 3 | 9 | P | s(F) | o | o | s(F) | ET | 0 | % | % | o | es | 8 | 2.5 |
| 1 | M | 1 | MD | P | Ye | s(F) | N | N | Ye | B | Und | iff | 2 | 5 | 0 | 40 | N | Y | - | - | 3 | 4. |
| 4 | 8 | H/H | PP | Ye | s(F) | N | N | Ye | B | PN | ET | 2 | 5 | 0 | 5 | N | Y | - | - | 0 | 0. |
| 5 | 2 | GG | F | Ye | s(F) | N | N | Ye | B | Mo | nstr | o | 1 | 5 | 0 | 90 | N | Y | N | - | 6 | 8. |
| 6 | 9 | PCN | SL | Ye | s(F) | N | N | Ye | B | Und | iff | - | - | - | N | N | o | o | 1 | 1. |
| 7 | 8 | - | - | - | - | - | B | Und | iff | - | - | N | N | o | o | 1 | 1. |

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Table 2: Comparison of the characteristics of adult diffuse hemispheric gliomas, H3 G34-mutant with those of H3.3 K27-mutant DMG, IDH1/2-mutant glioblastomas and pTERT-mutant glioblastomas

|                  | H3.3 G34-mutant gliomas (n=17) | H3.3 K27-mutant DMG (n=33) | IDH-mutant glioblastomas (n=36) | IDH-wt glioblastomas (n=100) |
|------------------|-------------------------------|-----------------------------|---------------------------------|-------------------------------|
| Median age at diagnosis (years) | 25.8 (SD =5.9) | 34.8 (SD=16.5) | 38.5 (SD=13.7) | <0.00 | 66.4 (SD=11.8) | <0.00 |
| Sex              |                               |                             |                                 |                               |                                 |
| Male             | 11/17 (65%)                   | 15/33 (45%)                 | 13/36 (36%)                     | 49/100 (49%)                  |
| Female           | 6/17 (35%)                    | 18/33 (55%)                 | 23/36 (64%)                     | 51/100 (51%)                  |
| Location         |                               |                             |                                 |                               |                                 |
| Frontal          | 11/16 (69%)                   | 0/33 (0%)                   | 24/36 (67%)                     | 48/100 (48%)                  |
| Parietal         | 11/16 (69%)                   | 0/33 (0%)                   | 7/36 (19%)                      | 22/100 (22%)                  |
| Temporal         | 3/16 (19%)                    | 0/33 (0%)                   | 11/36 (31%)                     | 32/100 (32%)                  |
| Occipital        | 1/16 (6%)                     | 0/33 (0%)                   | 1/36 (3%)                       | 14/100 (14%)                  |
| Corpus callosum  | 1/16 (6%)                     | 0/33 (0%)                   | 11/36 (31%)                     | 33/100 (33%)                  |
| Midline          | 3/16 (19%)                    | 33/33 (100%)                | 10/36 (28%)                     | 33/100 (33%)                  |
| Initial haemorrhage |                             |                             |                                 |                               |
| Yes              | 3/16 (19%)                    | 0/33 (0%)                   | 2/36 (6%)                       | 2/100 (2%)                    |
| No               | 13/16 (81%)                   | 33/33 (100%)                | 34/36 (94%)                     | 98/100 (98%)                  |
| Radiological features |                     |                             |                                 |                               |
| CE  | 4/15 | 18/29 | 0.05 | 27/36 (75%) | **0.002** | 93/100 (93%) | <0.00 |
|-----|------|-------|------|-------------|-----------|---------------|-------|
| Cyst | (27%) | (62%) | 0.60 | 7/36 (19%)  | 0.70      | 8/100 (8%)    | 1     |
| Cyst | 4/15 | 3/28  | 0.03 | 14/36 (39%) | **0.02**  | 70/100 (70%) | 0.60  |
| ADC restriction | 1/16 | 10/28 | 1    | 4/30 (13%)  | <0.00     | 35/68 (51%)  | <0.00 |
|     | (6%)  | (36%) | | (92%)       | (13%)     | (92%)        |      |

**IHC features**

| OLIG2 positivity | 3/16 | 29/29 | <0.00 | 36/36 (100%) | 0.03 | 89/91 (98%) | <0.00 |
|------------------|------|-------|-------|---------------|------|---------------|-------|
|                  | (19%)| (100%)|       |               | 1    |               | 1     |

**Molecular characteristics**

| IDH mut. | 0/17 | 2/34 (6%) | 34/96 (35%) |
|-----------|------|-----------|-------------|
| (0%)      |      | 1/28 (4%) |             |

| EGFR amplificacion | 2/14 | 0/33 (0%) | 88/100 (88%) |
|-------------------|------|----------|-------------|
| (14%)             |      | 2/30 (7%)|             |

| pTERT mut. | 0/15 | 0/33 (0%) | 46/93 (49%) |
|------------|------|----------|-------------|
| (14%)      |      | 0/18 (0%)|             |

| MGMT meth. | 0/15 | 2/30 (7%) | 1/59 (2%) |
|------------|------|----------|----------|
| (14%)      |      | 0/18 (0%)| <0.00    |

| BrafV600E mut. | 9/11 | 4/31 | 0.001 |
|----------------|------|------|-------|
| (82%)          |      |      |       |

| Median Overall Survival | 12.4 | 19.6 | 50.5 | 11.7 |
|-------------------------|------|------|------|------|
| (months)                | [6.3-27.5] | [6.1-29.1] | [45.9-79.2] | [9.3-17.5] |

\[a\] H3.3 K27-mutant DMG compared to diffuse hemispheric gliomas, H3 G34-mutant, \[b\] IDH-mutant glioblastomas compared to diffuse hemispheric gliomas, H3 G34-mutant, \[c\] IDH-wt glioblastomas compared to diffuse hemispheric gliomas, H3 G34-mutant

ADC: Apparent Diffusion Coefficient, CE: Contrast Enhancement, SD: Standard Deviation
Table 3: Summary of H3 G34-mutant gliomas characteristics in the present and in previously reported series

| Populations | Present series | Chen et al. 28 | Roux et al. 2012 | Mackay et al. 2018 | Vetter et al. 2018 | Mackay et al. 2017 | Puntoni et al. 2017 | Neumann et al. 2016 | Korshunov et al. 2015 | Sturm, Schwartzenbruck et al. 2012 |
|-------------|----------------|----------------|------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
| N           | Adult          | 17             | 95               | 11                | 7                 | 8                 | 67                | 12                | 12                | 81                |
| Median age  | Adult          | 25             | -                | 17                | 12                | (9-52)            | 15 (7-31)         | 16 (6-31)         | 19 (9-51)         | 18 (9-42)         |
|             | Pediatric      | 95             | 17               | 8                 | 6                 | 27%               | 100%              | 100%              | 100%              | 100%              |
| Radiological characteristics | FP (%) | 27% | 100 | (50%) | 37% | (80%) | 27% |
|             | TP (%)         | 92%            | %                | 62%               | %                 | 100%             | -                 | -                 | -                 | -                 |
| Main location | GBM-like | 53% | 72 | 57% | - | 83% | 0% | 50% | - | 50% |
| ADC restriction (%) | -    | - | - | - | - | - | - | - | - | - |
| Histologic and molecular characteristics | GBM-like | 53% | 72 | 57% | - | 83% | 0% | 50% | - | 50% |
|            | PNET-like | 53% | 72 | 57% | - | 83% | 0% | 50% | - | 50% |
|            | Olig2 | 14% | 44% | 8% | - | 13% | - | 27% | 10% | 75% |
|            | Immunone | 1% | 71% | - | 65% | - | 74% | - | - | - |
|            | ATRX loss or mutant | 82% | 50% | - | - | - | - | - | - | - |
|            | TERTp mutation | 82% | 50% | - | - | - | - | - | - | - |
|            | TP53 expression or mutant | 82% | 50% | - | - | - | - | - | - | - |
|            | EGFR amplification | 82% | 50% | - | - | - | - | - | - | - |
|            | PDGFRA alteration | 82% | 50% | - | - | - | - | - | - | - |
|            | MGMT methylation | 82% | 50% | - | - | - | - | - | - | - |

Note: The data in the table includes the following:
- **N**: Number of patients
- **Median age**: Median age in years (range)
- **Radiological characteristics**: FP and TP percentages
- **Histologic and molecular characteristics**: GBM-like, PNET-like, Olig2, Immunone, ATRX loss, TERTp mutation, TP53 expression, EGFR amplification, and PDGFRA alteration
- **MGMT methylation**: Percentage of patients with methylated MGMT

*The data is compiled from various studies, with references provided for each entry.*
| Median survival (months) | 12.5 | 36.2 | 12 | 18 | 22 | 24 | 12 to 36.2 |
|-------------------------|------|------|----|----|----|----|------------|

* only series with > 4 cases were included, y: years, FP: fronto-parietal, TP: temporo-parietal, T: temporal, ** PDGFRA mutations diagnosis/recurrence, *** PDGFRA mutations and amplifications, $: PDGFRA amplification, $$: excluding Puntonet et al study that focused on cases with a PNET-like presentation.
Figure 1

|    | HPS | Olig2 | TP53 | ATRX |
|----|-----|-------|------|------|
| A  | ![Image A](image1) | ![Image B](image2) | ![Image C](image3) | ![Image D](image4) |
| B  | ![Image A](image1) | ![Image B](image2) | ![Image C](image3) | ![Image D](image4) |
| C  | ![Image A](image1) | ![Image B](image2) | ![Image C](image3) | ![Image D](image4) |
| D  | ![Image A](image1) | ![Image B](image2) | ![Image C](image3) | ![Image D](image4) |
Figure 3

The graph shows the probability of survival over overall survival (months) for different subtypes of glioblastoma:

- H3.3 G34-mt: 17, 4, 0
- H3K27M-mt: 33, 5, 0
- GBM IDH-mt: 36, 18, 6
- GBM IDH-wt: 100, 3, 0