Combinations of Single-Gene Biomarkers Can Precisely Stratify 1,028 Adult Gliomas for Prognostication

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Advanced genomic techniques have now been incorporated into diagnostic practice in neuro-oncology in the literature. However, these assays are expensive and time-consuming and demand bioinformatics expertise for data interpretation. In contrast, single-gene tests can be run much more cheaply, with a short turnaround time, and are available in general pathology laboratories. The objective of this study was to establish a molecular grading scheme for adult gliomas using combinations of commonly available single-gene tests. We retrospectively evaluated molecular diagnostic data of 1,275 cases of adult diffuse gliomas from three institutions where we were testing for IDH1/2 mutation, TERTp mutation, 1p19q codeletion, EGFR amplification, 10q deletion, BRAF V600E, and H3 mutations liberally in our regular diagnostic workup. We found that a molecular grading scheme of Group 1 (1p19q codeleted, IDH mutant), Group 2 (IDH mutant, 1p19q non-deleted, TERT mutant), Group 3 (IDH mutant, 1p19q non-deleted, TERT wild type), Group 4 (IDH wild type, BRAF mutant), Group 5 (IDH wild type, BRAF wild type and not possessing the criteria of Group 6), and Group 6 (IDH wild type, and any one of TERT mutant, EGFR amplification, 10q deletion, or H3 mutant) could significantly stratify this large cohort of gliomas for risk. A total of 1,028 (80.6%) cases were thus classifiable with sufficient molecular data. There were 270 cases of molecular Group 1, 59 cases of molecular Group 2, 248 cases of molecular Group 3, 27 cases of molecular Group 4, 117 cases of molecular Group 5, and 307 cases of molecular Group 6. The molecular groups were independent prognosticators by multivariate analyses and in specific instances, superseded conventional histological grades. We were also able to validate the usefulness
INTRODUCTION

There has been a vast increase in our knowledge of the molecular pathology of central nervous system (CNS) tumors, and a new WHO classification will soon be published (1). The WHO 2016 Classification already introduced for the first time molecular criteria in the classification of gliomas by IDH status and the molecular groups of medulloblastomas (2). In many advanced centers, brain tumor samples are now routinely tested with genomic techniques, such as targeted sequencing, RNAsseq, and genome-wide DNA methylation profiling (3–6). The latter has become a regular genomic diagnostic tool in many centers (7–9), and many tumor entities in the upcoming WHO classification will have the methylation groups listed as desirable diagnostic criteria (1). However, genomic tests are still expensive, especially in countries with a national health system, and also typically, their results only come back several weeks later. Interpretation of genomic data also requires expertise and reporting of genomic tests, especially those conducted by commercial companies, is variable. In contrast, single-gene tests are faster for results, often done in-house, and are much cheaper. Results of many single-gene tests, e.g., 1p19q, IDH genotyping, are needed soon after the surgical resection so that planning for further treatment can be made and patients can receive appropriate counseling as soon as possible after the operation. There is therefore a need for precise molecular diagnosis and grading of brain tumors using single-gene tests.

Isocitrate dehydrogenase (IDH) genotype has been shown to be a major classifier and prognosticator for gliomas (10, 11) and WHO 2016 classifies astrocytic tumors as to whether they are IDH mutant or IDH wild type. Together with another group, we were the first to show the usefulness of mutations of TERT promoter (TERTp), when combined with IDH genotype, in the precise prognostication of low-grade gliomas (12, 13). The Cancer Genome Atlas (TCGA) showed that a combination of IDH genotype and 1p19q status stratified survivals well for the low-grade gliomas (14), but Eckel-Passow et al. in the same issue of New England Journal of Medicine, in addition to using IDH and 1p19q, introduced also TERT mutation to stratify low-grade gliomas (15). BRAF is also a gene that can be easily tested in most laboratories without too much difficulty with a mutation hotspot at V600E. We showed that in the younger adults, it is associated with good survival, and similarly, good survival with BRAF V600E mutated gliomas was shown in smaller series (16–18). Phillips et al. showed that among BRAF mutated anaplastic pleomorphic xanthoastrocytoma, the regular molecular hallmarks of glioblastomas, e.g., EGFR amplification, 10q loss, were uncommon (19). Mutations of the chromatin genes, H3.3 and H3.1, were also found to be poor prognosticators for gliomas, and the poor prognosis is not limited to tumors located in mid-line, nor were mutations restricted to pediatric high-grade gliomas (20–24). We hypothesized that combinations of genes IDH1/2, 1p19q, TERTp, EGFR, BRAF, TERTp, 10q, and H3 status can stratify gliomas for risk and may even be superior to conventional histological grading for prognostication. We recognize that this gene list is not exhaustive as regards the vast amount of literature concerning the molecular pathology of gliomas. However, since there is a limit as to how many single-gene tests an in-house facility can possibly carry out with a reasonable turnover time, we hypothesize that such a list would adequately molecularly grade a glioma and that it could be superior to conventional histological grading in some instances.

MATERIALS AND METHODS

Patients

A total of 1,275 adult diffuse gliomas diagnosed as astrocytic or oligodendrogial from patients aged 18 years or above were unselectively and retrospectively retrieved from the pathology archives of three institutions: Prince of Wales Hospital, Chinese University of Hong Kong, Huashan Hospital, Fudan University, Shanghai and First Affiliated Hospital of Zhengzhou University, Zhengzhou. Pediatric gliomas were excluded due to the vastly different molecular pathology between adult and pediatric gliomas (25, 26). Some cases were part of published studies (27–29), and the others were diagnosed between 2011 and 2017, and in the latter cases, molecular tests were done as part of the diagnostic workup. Cases lacking enough clinical information or survival data were excluded. Histological sections were centrally reviewed by two experienced neuropathologists before the present analyses (H-KN and HC). Histological grading from Grade II to Grade IV was made also according to the WHO 2016 Classification. The diagnostic criteria for Grade III tumors were increased mitotic activity, nuclear atypia, and high cellularity as described in WHO 2016, and the defining criteria for Grade IV tumors (glioblastomas) were either the presence of necrosis or endothelial proliferation. Ethics approvals were obtained from The Joint Chinese University of Hong Kong–New Territories East Cluster Clinical Research Ethics Committee; the Ethics Committees of Huashan Hospital, Shanghai; and The First Affiliated Hospital of Zhengzhou University, Zhengzhou. The study was performed in accordance with the Declaration of Helsinki.
Demographics and survival data were obtained from the hospital information systems of the three hospitals. The protocol for treatment for adult gliomas of Grades II–IV in all three hospitals was maximal safe resection in the first instance. Some but not all patients received adjuvant radio-chemotherapy due to individual clinical situations (Table 1), and this information was obtained from the hospital information systems. Overall survival (OS) was obtained either with the hospital information systems or by phone calls as practiced in our previous studies (30–32). OS was defined as the period of time between operation and death or the last follow-up.

Results of molecular tests for this cohort were all retrieved. We had a long-term interest in TERTp mutations in adult gliomas (12, 27–29), and therefore adult gliomas diagnosed between the years 2011 and 2017 at our three institutions had been examined extensively by this test. Cases diagnosed prior to this period of time were part of the previous studies (27–29). EGFR amplification, H3.3 and H3.1 mutations, and 10q deletion were also examined as much as tissues would allow in IDH wild-type gliomas because of our previous findings of their prognosticative value and also findings from the literature (14, 33, 34). BRAF V600E mutations were also evaluated extensively by this test. Case diagnosis prior to this period of time was routinely obtained with the hospital information systems or by phone calls as practiced in our previous studies (30–32). The methodology used for the single-gene test was the same for the three institutions.

Mutation Analysis

The hotspots for IDH1/2, TERTp, BRAF, H3.3, and H3.1 mutations were evaluated by Sanger sequencing as described (18, 30). In brief, tissue sections obtained from either macrodissection or direct scraping off the slides were placed into Tris–HCl buffer (pH 8.5) containing proteinase K. The mixture was then incubated at 56°C for overnight followed by 98°C for 10 min. The crude lysate was then mixed with primers, KAPA Robust HotStart ReadyMix (Sigma, St. Louis, MO, USA)/KAPA HiFi HotStart ReadyMix (Sigma), for PCR amplification. PCR products were visualized on an electrophoresis gel, cleaned with a spin column-based PCR product purification kit, and sequenced with BigDye Terminator Cycle Sequencing kit v1.1 (Life Technologies, Carlsbad, CA, USA). The PCR primers can be found in Supplementary Table 1.

Fluorescence In Situ Hybridization Analysis

EGFR amplification, 10q deletion, 1p19q codeletion, and CDKN2A homozygous deletion were evaluated by FISH. The BAC clone (CTD-2199A14) containing the genomic sequences of the 7p11.2 and a centromere probe (CEP7, Vysis, Downers Grove, IL, USA) was used for EGFR amplification detection. Vysis LSI PTEN/CEP 10 FISH Probe Kit was applied for detection of 10q loss. Vysis Lp36/1q25 and 19q13/19p13 FISH Probe Kit (Vysis) was used to examine 1p19q codeletion. Vysis LSI CDKN2A SpectrumOrange/CEP 9 SpectrumGreen Probes (Vysis) was employed to study CDKN2A. At least 100 non-overlapping signals were counted and analyzed in each case. EGFR amplifications were considered when >5% recorded cells displayed many tight clusters or a ratio of the target (red) to reference (green) signal >2 (33, 36). Deletion for 1p,
Molecular Group 4: IDH wild type, positive for TERTp mutation, EGFR amplification, 10q loss, or H3 mutations

The numbers of different tests carried out on this cohort are shown in Supplementary Table 2.

Overall, there were 270 cases of molecular Group 1, 59 cases of molecular grade 2, 248 molecular grade 3, 27 cases of molecular grade 4, 117 cases of molecular Group 5, and 307 cases of molecular grade 6 (Table 1). A total of 247 cases were unclassifiable for the six molecular groups, all due to a lack of data for some biomarkers. We found that molecular grading was significantly associated with histological grade (p < 0.001). Molecular Group 1 was enriched for WHO Grade II tumors and absent in WHO Grade IV tumors. Nearly 80% (76.3%) of molecular Group 1 tumors were of WHO grade II. Molecular Group 6 was highly enriched for WHO Grade IV, and nearly 70% of molecular grade 6 tumors were of WHO grade IV. Molecular groups were not associated with sex. The 27 cases of molecularly Group 4 were not epithelioid, PXA-like, or gangliogliomas on histological review.

As shown in Figure 1, molecular grades defined by molecular signatures were strongly associated with prognostication across the whole cohort (p < 0.001). Univariate analyses showed that all molecular grades were significantly associated with OS (Supplementary Figure 1 and Figure 1). Multivariable analysis showed that molecular groups had independent prognostic value (p < 0.001) across the cohort by adjusting for age, gender, histological grades, tumor resection, radiotherapy, and chemotherapy. Notably, molecular Groups 3, 5, and 6 tumors demonstrated significantly worse prognosis, and molecular Groups 2 and 4 tumors showed trends of unfavorable prognosis as compared to the reference group (molecular Group 1) (Figure 2).

We also tried to compare the power of prognostication of the molecular groups against histological grades in different scenarios. In three scenarios, the molecular groups were better than histological grades in prognostication. Figure 3A shows that WHO Grade III molecular Group 1 tumor demonstrated significantly longer OS than WHO Grade II molecular Group 6 tumors (p < 0.001). Figure 3B also illustrates that WHO Grade IV, molecular Group 4 tumors in fact had a remarkably longer OS as compared to the WHO Grade III, molecular Group 6 tumors (p = 0.004). Interestingly, WHO Grade IV molecular Group 4 tumors had similar OS as compared to the WHO Grade II molecular Group 6 tumors (Figure 3C). A total of 93 cases of IDH mutant, 1p19q non-codeleted gliomas were evaluated for CDKN2A; 11.8% of cases (11/93) were positive for CDKN2A homozygous deletion, and this was not associated with OS (p = 0.235).

We next retrieved an unselected adult glioma cohort with relevant clinical and molecular data from TCGA (10). The proportion of available biomarkers relevant to this study in this TCGA validation cohort is listed in Supplementary Table 3. When the molecular grading scheme of this study was used in TCGA cohort, there were 169 cases of Group 1, 8 cases of Group 2, 146 cases of Group 3, 7 cases of Group 4, 140 cases of Group 5, and 303 cases of Group 6. When tested against OS, the six molecular groups remain statistically significant (p < 0.001) (Supplementary Figure 2).
DISCUSSION

The diagnostic protocols for brain tumors have been vastly changed in the advanced centers due to the advent of next-generation sequencing, for both DNA and RNA and also genome-wide methylation profiling. The former can evaluate gene mutations or gene fusion on a panel of a large number of genes, while the latter can map all brain tumors according to their methylation profiling into 91 methylation classes (7). Copy number variations (CNVs) can be obtained from both methods. Both have been widely used in the day-to-day fine diagnosis of brain tumors as well as research (8, 30, 38, 39). The WHO 2016 Classification of CNS Tumors for the first time introduced molecular criteria in the diagnosis of some brain tumors, and many centers in the world use an integrated, multi-layered approach in the pathological diagnoses of brain tumors.
integrating histological features, WHO grades, and molecular characteristics. Adult diffuse astrocytic tumors are classified into IDH-mutant and IDH wild type, and oligodendrogliomas are defined by both IDH mutation and 1p19q codeletion by the WHO 2016 Classification. However, grading of adult diffuse gliomas remains entirely histological depending mainly on mitoses, cellular anaplasia, necrosis, and microvascular proliferation.

Most general pathology laboratories in the world are not in a position to carry out next-generation sequencing or genome-wide methylation profiling routinely on all their neurosurgical specimens, as these tests remain expensive and demanding on expertise for interpretation, e.g., t-distributed stochastic neighbor embedding (t-SNE) plots. Most national types of health finance systems are unable to support such extensive genomic tests regularly. However, most pathology departments in the more advanced societies will have the capacity for single-gene studies and molecular tests like Sanger sequencing, for example, for BRAF V600E, IDH1 R132, and IDH2 R172H. Similarly, FISH is also widely available in pathology departments, especially in the context of Her-2 testing for breast cancers. We therefore believe that there is a need to develop a relatively simple molecular grading algorithm for the gliomas based on single-gene testing based on current literature on the molecular pathology of gliomas (15, 32, 33, 40–42). There are many excellent studies on the grading of special groups of gliomas by a panel of single genes. For example, TCGA concluded that IDH and 1p19q status satisfactorily grades diffuse lower-grade gliomas, while Eckel-Passow et al. further fine-graded lower-grade gliomas with the addition of TERT status (14, 15).

Mutations of the promoter region of TERT have been suggested to be a key terminal event in the evolution of glioblastomas, but it has an intriguing property of being mutated also in oligodendrogliomas, which are mostly low-grade gliomas (43). The reason for its dual and seemingly contradictory properties is unknown. The dual impact of TERT can be seen in that other than being present in IDH mutant, 1p19q codeleted tumors, it was also a prognosticating factor among IDH mutant 1p19q non-deleted tumors (44, 45) and, in this series, the difference between molecular grade 2 (TERT mutant, IDH mutant, 1p19q non-codeleted) and molecular grade 3 (TERT wild type, IDH mutant, 1p19q non-codeleted). However, molecular group 2 remains a small group in our cohort (n = 59).

Other than lower-grade gliomas, there have also been a few studies in the literature devoted to molecular-grade special groups of gliomas by single-gene studies (32, 33), but we are only aware of one previous attempt to use a panel of single-gene status to grade holistically all adult gliomas (46). In this study, Jiao et al. showed that gliomas can be satisfactorily graded for prognosis by complete sequencing of IDH1/2, ATRX, CIC, and FUBP1. However, the latter genes are large, and sequencing of entire genes is laborious and expensive.

The biomarkers used in this study were known in the literature on their own. Group 1, IDH mutant, 1p19q codeleted is the regular oligodendroglioma as currently defined by WHO Classification, and Group 6 constitutes the bulk of the regular IDH wild-type glioblastomas as defined by cIMPACT-NOW update 3 (47). Group 5 would constitute most of the IDH wild-type lower-grade gliomas that do not fulfill the molecular criteria of glioblastoma (33, 48, 49). It is also interesting that in our molecular Group 4, BRAF V600E was able to impart a better prognosis among the IDH wild-type gliomas. This is consistent with the few papers concerning the potentially better prognosis of BRAF V600E in adult gliomas (16–18). The overall number in this IDH wild type, BRAF mutant group is small (27/1,028 cases), consistent with the overall low percentage of BRAF V600E in...
glioblastoma is the homozygous deletion of CDKN2A/B (51). In lower-grade gliomas, the molecular diagnostic criterion for methylation and the critical reference for this recommendation used events that are usually not tested in a general pathology laboratory, this study, we have used extensively EGFR amplification to test for whole chromosomes (7). The molecular grading scheme we propose can easily be made available in general pathology departments. Adoption of this system provides a fine prognostication for gliomas and in, some instances, is superior to the histological grading, as shown in Figure 3. The use of this scheme also fulfills many of the recommended markers of the WHO Classification. While most of the biomarkers used in this grading scheme including 1p19q codeletion, IDH2 mutations, 10q loss, and TERT mutation do not have any immunohistochemical tests, IDH1 and BRAF V600E mutations are readily tested by immunohistochemistry, and these can be used in busy diagnostic laboratories. We recognize that not all single-gene biomarkers that were described in the new WHO Classification and cIMPACT (e.g., CDKN2A/B) have been included in the grading scheme. In our cohort of molecular Grade 2 (IDH mutant, 1p19q non-codeleted, TERT mutant) and Grade 3 (IDH1 mutant, 1p19q non-codeleted, TERT wild type) tumors, we were unable to demonstrate a statistical significance between cases with or without homozygous CDKN2A deletion. Marker and Pearce also showed that CDKN2A deletion was not associated with prognostication in Grades II and III IDH-mutant astrocytomas (54). Methylated MGMT has been known for a long time to be a favorable prognostic marker in glioblastomas (55) and predicts their response to alkylating chemotherapy (56, 57). However, it is not in regular use for the lower-grade glioma as a prognosticator, and most of the low histological grade IDH mutant gliomas are likely to be MGMT methylated.

We also recognize the limitations of our study, as we used retrospective analyses of existing data, and it would be impossible for us to re-do a new panel of single-gene markers for 1,275 cases according to the recommendation of the new WHO Classification. Ours is not a proposal to replace the new WHO Classification (1) but merely a proof of principle that a set of well-placed single-gene biomarkers can fully cover risk stratification in adult gliomas. Even in the new WHO Classification (1), there are cases that do not fit and are classified as NEC (Not Elsewhere Classified), e.g., IDH wt gliomas that do not exhibit EGFR amplification or TERTp or combined whole chromosomes +7/−10q. It is possible that other algorithms may have developed that use some biomarkers might be shown to be equally useful in molecular grading of gliomas in the future. But our study showed that a simple molecular grading scheme such as the one proposed can satisfactorily fine-grade adult gliomas, and these tests can be performed relatively quickly and are highly achievable in practically all pathology laboratories. This grading scheme will provide critical and timely information for immediate clinical management of glioma patients.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by The Joint Chinese University of Hong Kong–New Territories East Cluster Clinical Research Ethics Committee; the Ethics Committees of Huashan Hospital, Shanghai; and The First Affiliated Hospital of Zhengzhou University, Zhengzhou. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

The authors confirm contribution to the paper, as follows: study conception and design: AC, W-SP, HL, and H-KN. Data acquisition: Z-FS, KL, W-WW, HC, NC, DC, X-ZL, Z-YZ, and YM. Analysis and interpretation of data: AC, Z-FS, KL, NC, and H-KN. Writing—original draft: AC and H-KN. Writing—review and editing: AC, W-SP, HL, and H-KN. All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.839302/full#supplementary-material
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