## Treatment Response Assessment in IDH-Mutant Glioma Patients by Noninvasive 3D Functional Spectroscopic Mapping of 2-Hydroxyglutarate

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Treatment response assessment in IDH-mutant glioma patients by non-invasive 3D functional Spectroscopic Mapping of 2-Hydroxyglutarate

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Abstract
**Purpose**—Measurements of objective response rates are critical to evaluate new glioma therapies. The hallmark metabolic alteration in gliomas with mutant isocitrate dehydrogenase (IDH) is the overproduction of oncometabolite 2-hydroxyglutarate (2HG), which plays a key role in malignant transformation. 2HG represents an ideal biomarker to probe treatment response in IDH-mutant glioma patients, and we hypothesized a decrease in 2HG levels would be measureable by *in vivo* magnetic resonance spectroscopy (MRS) as a result of antitumor therapy.

**Experimental Design**—We report a prospective longitudinal imaging study performed in 25 IDH-mutant glioma patients receiving adjuvant radiation and chemotherapy. A newly developed 3D MRS imaging was used to non-invasively image 2HG. Paired Student’s T-test was used to compare pre- and post-treatment tumor 2HG values. Test-retest measurements were performed to determine the threshold for 2HG functional spectroscopic maps (fSM). Univariate and multivariate regression were performed to correlate 2HG changes with Karnofsky performance score (KPS).

**Results**—We found that mean 2HG (2HG/Cre) levels decreased significantly (median = 48.1%, CI = 27.3–56.5%, P = 0.007) in the post-treatment scan. The volume of decreased 2HG correlates ($R^2 = 0.88$, $P = 0.002$) with clinical status evaluated by KPS.

**Conclusion**—We demonstrate that dynamic measurements of 2HG are feasible by 3D fSM, and the decrease of 2HG levels can monitor treatment response in patients with IDH-mutant gliomas. Our results indicate that quantitative *in vivo* 2HG imaging may be employed for precision medicine and early response assessment in clinical trials of therapies targeting IDH-mutant gliomas.

**Keywords**
Isocitrate Dehydrogenase (*IDH*) mutation; Glioma; 2-Hydroxyglutarate (2HG); Magnetic Resonance Spectroscopic Imaging (MRSI); functional Spectroscopic Maps (fSM); Radiochemotherapy

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**Introduction**

Intracranial malignant diffuse gliomas are a leading cause of cancer death in people under age 45. After initial diagnosis and therapy with surgery, radiation (1) and chemotherapy (2, 3) these diffusely infiltrative gliomas often regrow as malignant lesions, and then lack effective treatment, representing a substantial unmet clinical need. Indeed, the longitudinal treatment monitoring of diffuse gliomas with conventional magnetic resonance imaging (MRI) has been a substantial challenge, as radio-chemotherapy induced alterations and treatment with anti-angiogenic agents often result in treatment-related imaging changes on MRI that are difficult to distinguish from true disease progression (4, 5). Methods to discern tumor progression from treatment associated changes (true/pseudo-progression/response) are urgently needed, particularly to measure objective response rates in clinical trials of novel glioma treatments.

The majority (50–86%) of gliomas diagnosed in younger adults (<45 yo) have recurrent somatic mutations in the genes encoding NADP+-dependent isocitrate dehydrogenase 1 and 2 (*IDH1/IDH2*) (6, 7). These heterozygous mutations, characteristically non-synonymous point-mutations of Arginine at the 132 residue (R132) of the *IDH1* gene or the 172 residue...
(R172) of IDH2, result in a neomorphic gain of enzymatic function, leading to excess turnover of \( \alpha \)-ketoglutarate (\( \alpha \)KG, known as 2-oxoglutarate) and accumulation of the oncometabolite 2-hydroxyglutarate (2HG) (8). Increased levels of 2HG inhibit \( \alpha \)KG-dependent dioxygenases (9) and chromatin modifiers, promoting tumorigenesis through DNA-/histone- hypermethylation and genome-wide alterations in the epigenetic control of gene expression (10, 11).

Hence, 2HG may serve as a unique biomarker to probe IDH-mutant tumor burden, cancer mechanisms and therapeutic opportunity. This possibility is based on a powerful combination of mutually exclusive events: i) IDH mutation is an early oncogenic event in gliomas (12), ii) it is not found outside the tumor volume in patients harboring this mutation (13), iii) IDH-mutant cancer cells produce very high levels of 2HG (8), iv) only trace levels of 2HG are found in IDH-wild-type cells where it is cleared rapidly by 2-HG dehydrogenases (8).

The markedly increased intracellular concentration of 2HG, up to 5–35 mM (8), exceeds the sensitivity threshold for in vivo magnetic resonance spectroscopy (MRS). Early work by our group (14) and others (15, 16) has reported on the feasibility of single-voxel or single slice MRS detection of 2HG in IDH mutant glioma patients. Unambiguous in vivo detection of 2HG by conventional MRS is challenging, and special editing spectroscopic methods were designed (14, 15) to disentangle the 2HG signal from overlapping signals from metabolites with similar structure.

While initial studies (14–18) focused on 2HG imaging for diagnosing IDH mutant gliomas, currently the interest has shifted towards longitudinal 2HG imaging for monitoring treatment response in IDH-mutant patients. Nevertheless, the existing data suggest that in addition to its diagnostic/prognostic value (7, 19), 2HG may be also predictive of treatment outcome (20). It is expected that reliable longitudinal quantitative measurement of 2-HG levels in response to treatment would accelerate clinical trials, either as a pharmacodynamic marker for small molecule inhibitors (20) that block the production of 2-HG by mutant IDH, or as an efficacy marker for drugs that selectively kill IDH mutant tumors based on synthetic lethality (21, 22) This is highly relevant for intracranial gliomas where multiple patient biopsies to test activity of novel therapeutics in a longitudinal study are often not feasible. On the other hand, it has been shown in previous studies that MRS can be safely performed to monitor treatment response in glioma patients (23, 24).

Due to tumor heterogeneity and the physical remodeling of tumor-brain structure during treatment course regional changes of 2HG levels need to be measured longitudinally over the entire tumor volume. The imaging requirements for monitoring tumor treatment response are more stringent than for diagnosis, and this has not been demonstrated yet for 2HG. By comparison, for diagnosing IDH mutation the one time presence or absence of 2HG signal is sufficient (14, 15), while for tracking response accurate changes in 2HG levels have to be reliably measured in the same anatomical location over time.

In the current work we hypothesized that a decrease in 2HG levels would be measureable by in-vivo MRS in mutant IDH glioma patients as a result of antitumor therapy. Since only the
mutant IDH glioma cells produce 2HG, a reduction in the number of tumor cells or impairment of their metabolism would result in decreased 2HG levels. To test our hypothesis we employed a recently developed 3D MRS imaging (MRSI) sequence (25) that can selectively edit metabolites of interest, such as 2HG, with increased temporal stability and spatial coverage. Existing data suggest that IDH mutant gliomas are more radiosensitive compared to their wild-type counterparts (26, 27), hence a 2HG reduction may be observed in patients treated with radiotherapy. Therefore, we tested our hypothesis in a prospective longitudinal study of a cohort of IDH-mutant gliomas patients undergoing radiation, and evaluated the potential of non-invasive 3D 2HG quantification for response assessment.

For this study, we further propose a novel analytic framework, which we label functional spectroscopic map (fSM), as a metric to quantify spatiotemporal changes of metabolite levels. Importantly, 3D fSM provides comprehensive tumor metabolic information that cannot be captured by a single mean value of metabolite levels. In this respect, fSM of 2HG is similar to functional diffusion map (fDM) (28) based on apparent diffusion coefficient (ADC) that showed increased value over mean ADC values for treatment monitoring of glioblastoma patients (29). It is expected that tumor regions with decreased 2HG/Cre and/or increased ADC would correlate with treatment response. Results from 3D 2HG fSM were compared against other imaging modalities for determining treatment response such as RANO criteria (4, 5) and diffusion MRI (29).

In our study we found a statistically significant decrease of mean 2HG levels and tumor volume with quantifiable 2HG between pre- and post-treatment scans, early after completion of radiotherapy. In addition, correlation between changes of 2HG levels and clinical improvement supports the hypothesis that 2HG could be used as an endpoint to determine objective response rates complementary to traditional clinical outcome criteria in IDH mutant patients.

Materials and Methods

Patient population

Glioma patients (WHO grades II–IV) seeking clinical care at the Pappas Center of Neuro-Oncology from Massachusetts General Hospital were selected for our study over the last year. Inclusion criteria specified: i) independent confirmation of mutant IDH and ii) clinical indication for radiation and/or chemotherapy. Standard external beam radiotherapy (RT, photons) was administered focal in 2 Gy/day fractions for 5 days over a 6-week period. Focal RT included a 2–3 cm margin beyond tumor or resection margin. Chemotherapy (CT) was administered using the oral methylating drug temozolomide, at a dose 75 mg/m$^2$/day, 7-days per week for the same duration as radiotherapy. Temozolomide was also continued after the completion of radiation (150–200 mg/m$^2$/day, days 1–5 of a 28-day cycle, 6 cycles). All patients had surgical procedures (i.e. diagnostic biopsy, subtotal or gross-total resection) but none received prior adjuvant treatment. Exclusion criteria were patients who could not be safely imaged with MRI. A number of twenty five patients (N=25) were consented with an approved IRB protocol and enrolled in the imaging study. Patients are listed in Supplementary Tables 1S and 2S.
Image acquisition

All images were acquired on a clinical 3T MR scanner (Tim Trio, Siemens, Erlangen) with a 32-channel head coil. A robust 3D MRSI sequence for 2HG imaging was newly developed (25) by integrating three highly optimized modules: i) adiabatic J-difference spectral editing MEGA-LASER, ii) spiral spectroscopic imaging, and iii) real-time prospective motion correction, shim update with reacquisition. This sequence efficiently edits 2HG signal while minimizing subtraction errors caused by subject motion or scanner instability. Details of the pulse sequence are provided in Supplementary Figure 1S.

The acquisition parameters of the 3D MRSI sequence were: TR=1600 ms, TE=68 ms, FOV=200×200×200 mm$^3$, VOI=100×80×50 mm$^3$, 20 mm isotropic voxels, acquisition matrix 10×10×10 zero-filled to 16×16×16, NA=20, acquisition time 9:55 min:s. Our choice of spatial resolution favored the SNR, but higher resolutions are technically possible for the same acquisition time. The timing of the MEGA-LASER excitation was optimized by simulations and phantom measurements for the maximum 2HG signal. MEGA pulses were applied in an interleaved fashion at 1.9 ppm (ON) and 7.5 ppm (OFF) to edit the 2HG signal (Hα) at 4.02 ppm. To assess repeatability, 3D MRSI was acquired twice at the baseline in six patients. For test-retest, patients were taken out from the scanner and repositioned between first and second 3D MRSI.

In addition, 3D T2 weighted (TR/TE = 3200/420 ms, FOV = 256×256×176 mm$^3$, matrix = 256×256×176) and 2D fluid attenuated inversion recovery (FLAIR, TR/TI/TE = 10000/2500/70 ms, FOV = 220×186×138 mm$^3$, matrix = 256×192×23) images were acquired. In twenty patients diffusion tensor imaging (DTI) was also acquired (TR/TE = 7980/84 ms, b-values = 0/700 s/mm$^2$, gradient directions = 12, FOV = 236×236×118 mm$^3$, matrix = 128×128×64).

To reproduce positioning across visits for each patient the on-line automatic slice positioning (AA scout, (30)) was used during acquisition of all imaging data.

Image processing

The difference (ON-OFF) and OFF spectra from each voxel of the VOI (approx. 200 voxels per VOI) were fitted with LCModel (31) software using the corresponding simulated basis sets for the MEGA-LASER sequence. Difference spectra were used to quantify 2HG and Glx (glutamate and glutamine), and OFF spectra were used to quantify total choline (tCho), creatine (Cre), lactate (Lac), and N-acetyl-aspartate (NAA). Cramer-Rao lower bound (CRLB) less than 25% (relative CRLB) as calculated by LCModel was considered acceptable for goodness of fits (see the note and references at the end of Supplementary Data for rationale of CRLB threshold). 3D metabolic maps were further constructed using a combination of MINC (Montreal Neurological Institute), FSL (FMRIB Software Library, Oxford, UK) and MATLAB (Mathworks, Natick, MA) software tools. Relative metabolic maps were obtained for 2HG, Glx, Lac, tCho and NAA as ratio to Cre. Ratios to Cre were preferred to water because of edema, cysts and surgical cavities that need to be accurately segmented, while Cre excludes these confounding. All metabolic maps were coregistered to anatomical T2w, FLAIR and ADC images for each visit.
Coregistration of T2w, FLAIR and ADC images obtained at different time points was performed using the robust register method of Freesurfer (32) to a common middle space. The same transformations were applied to coregister the corresponding 3D metabolic maps. In addition, AAscout ensured prospectively that images and spectral maps were identically positioned at each visit.

Difference maps were calculated by voxel-wise subtraction between coregistered post- and pre-treatment scans for metabolic and ADC maps. Functional spectroscopic (fSM) and diffusion (fDM) maps were obtained by thresholding the 2HG/Cre and ADC difference maps, respectively. Functional mapping classifies voxels in three categories: i) decrease, ii) stable, and iii) increase according to a statistically significant functional threshold. The functional threshold for difference 2HG/Cre (Δ2HG/Cre) was taken as the 95% confidence interval (CI) of the test-retest distribution. The normality of test-retest Δ2HG/Cre distribution was verified by Kolmogorov-Smirnov test. The 95% CI of test-retest Δ2HG/Cre distribution was found to be 0.11, hence voxels with Δ2HG/Cre < −0.11 were considered to have decreased 2HG/Cre, voxels with Δ2HG/Cre ∈ [−0.11, 0.11] were considered stable, and voxels with Δ2HG/Cre > 0.11 were considered to have increased 2HG/Cre. For difference ADC (ΔADC) a functional threshold of 0.4 µm²/ms representing the 95% CI of gray and white matter distribution was used, as proposed by Ellingson et al (33).

Tumor regions of interest (ROI) were manually outlined on the FLAIR images to include tumor tissue and infiltrating peritumoral edema, with exclusion of surgical cavities, cysts or necrosis areas. For each ROI multiple imaging biomarkers were calculated: a) FLAIR tumor volumes, b) mean, median and CI for 2HG/Cre, Glx/Cre, Lac/Cre, tCho/Cre, NAA/Cre, c) mean, median and CI for ADC, d) volume of voxels with quantifiable 2HG/Cre > 0.1, and e) volume of voxels with decreased, increased and stable 2HG/Cre and ADC, respectively.

**Statistical Analysis**

Differences in imaging biomarkers between post- and pre-treatment scans were deemed significant at a level of P<0.05 using two-sided Student’s T-test. Correlations between 2HG and other imaging biomarkers were calculated using Pearson product-moment correlation coefficient. Multivariate linear regression (general linear model) was performed using KPS as dependent variable and 2HG/Cre, age and grade as covariates. All statistical analysis was performed with the statistical package of MATLAB.

**Genomic analysis**

IDH-mutational status was confirmed by immunohistochemistry (IHC) analysis using an anti-human R132H antibody (DIANOVA) (34), or by genetic sequencing (SNapshot) (35). In addition, molecular markers such 1p/19q codeletion (36) and MGMT promoter methylation (37) were tested using fluorescence in-situ hybridization (FISH) and polymerase chain reaction (PCR), respectively (35).

**Clinical and radiologic evaluation**

Patients were evaluated clinically and radiologically by the treating neuro-oncologist and radiographic images were centrally reviewed by A.S.C and D.P.C. Karnofsky Performance
Status (KPS) (38) was used to assess functional improvement or decline. Primary clinical endpoints such as overall survival (OS) and progression free survival (PFS) were limited due to length of follow up time. RANO (Response Assessment in Neuro-Oncology) (4, 5) criteria were used to determine whether patients had complete response (CR), partial response (PR), minor response (MR), stable disease (SD) or progressive disease (PD).

**Results**

**Optimization of 3D MRSI editing of 2HG**

The MEGA-LASER editing was optimized before the start of the longitudinal study. Detailed results pertaining to this phase are shown in the Supplementary Data (Figs. S2–S4). In Supplementary Figure 2S the efficiency of 2HG editing of MEGA-LASER (25) is compared to MEGA-PRESS and PRESS-97 (PRESS, TE=97ms), the other two sequences which were previously demonstrated (15) for 2HG detection in patients. MEGA-LASER was found to provide 60% signal increase compared to MEGA-PRESS, and 90% of the signal of PRESS-97. When considering the range of B1 field inhomogeneity at 3T the MEGA-LASER has improved performance, yielding 94% and 115% of the PRESS-97 signal for a variation of −15% to +15% B1 amplitude, respectively (Supplementary Figure 3S). Real-time correction (25) of subtraction errors due to patient motion or scanner instability is shown in Supplementary Figure 4S.

**Longitudinal imaging**

A number of twenty five patients (N=25) were enrolled and imaged in this study. Thirteen (N=13) patients were imaged twice: i) a pre-treatment baseline scan 1–14 days before starting adjuvant treatment, and ii) a post-treatment follow-up scan acquired in the time interval of 1–3 months after the end of radio-chemotherapy. The remaining twelve patients had only baseline scans either due to change of the treatment plan or lack of compliance. Out of the thirteen patients imaged longitudinally, four had extensive surgical resection and 2HG levels could not be detected on the postoperative scan. The baseline metabolite levels from all 25 patients are given in Supplementary Table 1S. The nine patients that had only biopsy or subtotal resection with 2HG present at baseline and follow-up are listed in Supplementary Table 2S with their demographic, genomic, molecular, histological, radiological and clinical data. Note that any surgical procedure was done before the baseline scan, and no tumor was removed between baseline and follow-up scans.

Representative metabolic maps obtained from the pre- and post-treatment scans are shown in Figure 1. Comparing the intra-scan maps (either pre- or post-treatment), regions of high 2HG correspond to regions of high lactate and choline, and to regions of reduced Glx (glutamate and glutamine) and NAA, respectively. Comparing the inter-scan maps (pre- vs. post-treatment) it can also be seen that 2HG has the largest change among all metabolites with treatment. The metabolic maps corresponding to the acquired 3D MRSI matrix are shown before image interpolation in Supplementary Figure 5S.

Difference and OFF spectra from voxels selected in the tumor and contralateral normal appearing white matter before and after treatment, respectively, are shown in Supplementary...
Figure 6S. The edited 2HG signal at 4.02 ppm can be clearly distinguished only in the difference spectra in the tumor, with reduced amplitude after treatment. In addition, other metabolites can be identified in the OFF spectra (NAA, choline, creatine and lactate), or are co-edited in the difference spectra (GABA+ and Glx).

In Figure 2 three examples of functional spectroscopic maps (fSM) for 2HG/Cre are shown from a WHO grade II, WHO grade III, and WHO grade IV patient, respectively. Lower grade (WHO-II, WHO-III) and smaller tumors seem to have a more uniform response (decrease) of 2HG/Cre to treatment. Higher grades (WHO-IV) typically present with a larger tumor volume and show a more heterogeneous treatment response with regions of decreased, stable, and increased 2HG/Cre within the tumor after treatment. The distribution of Δ2HG/Cre values for test-retest and longitudinal scans are shown in Supplementary Figure 7S. For test-retest the Δ2HG/Cre values follow a single normal distribution with a mean of −0.022 and 95% CI of ±0.115. For the longitudinal scans the Δ2HG/Cre values may be arranged on two normal distributions with a mean/CI of −0.45/±0.19 (voxels of large response), and a mean/CI of −0.13/±0.15 (voxels of low response). The two distributions showed that there were approximately three times less voxels of large response compared to voxels of low response. Comparison of fSM and functional diffusion maps (fDM) maps is shown in Supplementary Figure 8S. It was noticed that regions of decreased 2HG/Cre partially overlap with regions of increased apparent diffusion coefficient (ADC), suggestive of reduction in cellular density associated with treatment response.

Quantification of biomarkers

In Figure 3 histograms of 2HG/Cre values within tumor ROI are shown for all patients for both pre- and post-treatment scans. In general, a shift towards lower and a narrower range of 2HG/Cre values are seen in post-treatment scans as compared to pre-treatment scans. Bar plots are shown in the enclosed bottom panel for pre- and post- median values of 2HG/Cre (Fig. 3A), pre- and post-fractional volumes of 2HG/Cre (Fig. 3B) and fractional volumes of 2HG/Cre changes from fSM (Fig. 3C). Typically, larger differences between pre- and post-treatment values were noticed for lower grade patients. In particular, the effect of tumor heterogeneity on changes of 2HG levels can be noticed in Fig. 3C, showing tumor volumes where 2HG levels were decreased, stable and increased, respectively. Fractional changes for all 2HG/Cre parameters are listed in Table 1. Fractional changes for all imaging biomarkers are provided in Supplementary Table 3S.

Group mean values and 95% CI of all imaging biomarkers over tumor ROIs are listed in Table 2. The only biomarkers that demonstrated a statistically significant change between pre- and post-treatment scans were 2HG/Cre (P = 0.0069) and the volume of 2HG/Cre (P = 0.0406). Across the patient group the mean pre-treatment 2HG/Cre was 0.36 (CI = ±0.09) and the mean post-treatment 2HG/Cre was 0.2 (CI = ±0.05). The mean pre-treatment volume of 2HG/Cre was 7.65 cm$^3$ (CI = ±6.55 cm$^3$), while the mean post-treatment volume of 2HG/Cre was 4.5 cm$^3$ (CI = ±4.43 cm$^3$).

Box plots of tumor mean fractional changes of metabolites are shown in Figure 4A. The median fractional change of 2HG/Cre −41.87% (CI = ±14.62%, median = 48.17%), while the median fractional change of 2HG/Cre volume was −39.07% (CI = ±9.3%). The box plots
of fractional volumes of decreased, increased and stable 2HG/Cre and ADC are plotted in Figure 4B. A reduction in the number of tumor cells due to treatment is expected to correlate with the fractional volume of decreased 2HG/Cre and the fractional volume of increased ADC. We found that the mean fractional volume of decreased 2HG/Cre was 58.2% (CI = ±15.8%) and the mean fractional volume of increased ADC was 18.2% (CI = ±13.6%). The relationship between the quality of 2HG data and the apparent (FLAIR) tumor size is presented in Supplementary Figure 9S, showing a trend for higher SNR and lower CRLB of 2HG in tumors that have larger FLAIR volumes, albeit not statistically significant. Note that within the mask of 2HG goodness of fit (e.g. Fig. 1, last column) a significant number of voxels had relative CRLB lower than 20%, as indicated in Suppl. Fig. 9S.

Correlation of clinical and imaging data

Univariate linear regression was first employed to establish if serial changes of 2HG correlated with serial changes in the Karnofsky Performance Status (KPS) scores. We separately tested all metrics calculated for 2HG/Cre (mean difference, volume, and fractional decreased volume), and we found that the fractional volume of decreased 2HG/Cre correlated strongest with the increase of KPS ($R^2 = 0.88$, $P = 0.002$). The scatter plot of KPS change versus the fractional volume of decreased 2HG/Cre is shown in Figure 4C. Interestingly, the patient with RANO progressive disease has also the lowest fractional volume of decreased 2HG/Cre and the largest decline of KPS, while one of the patients with minor response had the largest fractional volume of decreased 2HG/Cre and largest improvement of KPS. The change in FLAIR volume and mean ADC did not correlate with KPS scores. Results from a general linear regression model including tumor grade and age besides the fractional volume of decreased 2HG/Cre are given in Supplementary Table 4S, with the caveat that the number of patients was not enough for the number of independent variables.

Discussion

In vivo MRS has the unique ability to probe specifically and non-invasively IDH mutant gliomas by measuring the IDH mutant specific oncometabolite 2HG. Compared to single voxel or single slice methods (14, 15), the 3D MRSI imaging technique reported here provides complete spatial coverage of tumors and exploits the full potential of advanced image analysis. 3D mapping of 2HG and other metabolites has the advantage of accounting for tumor heterogeneity and may be especially important for longitudinal studies, because the change in size, shape and molecular composition of tumors during treatment makes positioning of a single voxel or a single slice difficult to match precisely over time. Indeed, 3D imaging has been shown to reduce variance in longitudinal MRS studies (39), likely because it is not affected by the sampling bias of single voxel methods, or for that matter, of invasive biopsies. In addition to obtaining a single mean or median 2HG/Cre value for a tumor, several more useful metrics can be computed from 3D MRSI data, such as histogram and volumetric analysis. Similar to functional diffusion maps (fDM) (28, 29) which showed increased clinical utility over mean ADC value in longitudinal studies of glioma patients, functional spectroscopic maps (fSM) can be constructed to obviate regional significant changes of 2HG. While we noticed a trend between the decrease of 2HG and the increase of...
ADC in response to therapy, we did not find a statistical significant correlation between these two imaging biomarkers. This limitation of our study may require further investigation to demonstrate whether 2HG is a biomarker that presents with an earlier treatment response as compared to ADC.

The observed 2HG levels are linked to NADPH (40) depletion and the impaired reductive capacity of IDH-mutant cells. Therefore 2HG quantification could probe directly the potential mechanisms of radiation damage (26, 27) in IDH-mutant cells and decrease of tumor burden. In our study the fractional volume of decreased 2HG/Cre derived from 3D fSM has been found to correlate stronger than the median 2HG/Cre with KPS, used as a standard measure of clinical status. Though KPS is often a secondary endpoint in clinical studies and not a primary endpoint such as the gold-standard endpoints of overall survival (OS) or progression-free survival (PFS), KPS is a well-established prognostic factor in gliomas and has been used in conjunction with imaging to predict survival (41). Although clinical scoring systems such as KPS are known to be subjective with large inter-observer variability, clinical assessments are an integral part of the criteria used to establish imaging endpoints (4, 5). Ultimately, 2HG imaging correlation with OS and PFS in this subset of patients will be informative, however these endpoints will require of many more years of follow up, as retrospective studies (7, 42) indicate these patients have a median PFS of >50 months and median OS of >100 months.

Multiple metabolites can be quantified from the same 3D MRSI data set allowing further exploratory studies of aberrant metabolic networks and pathways in tumors. In our data the tumor ROI median value of 2HG/Cre was found positively correlated with that of Lac/Cre in pre-treatment scans ($R^2 = 0.84$, $P = 0.0047$), but correlation was not statistically significant in post-treatment scans ($R^2 = 0.13$, $P = 0.744$). It’s possible that the increased lactate levels in IDH-mutant gliomas may be the result of Warburg effect due to silencing of lactate dehydrogenase A (43). In addition, monocarboxylate transporters (MCTs) are responsible for lactate transport and modulation of MCT has been suggested as potential therapeutic in glioma (44).

Conversely, the tumor ROI median value of 2HG/Cre was negatively correlated with that of Glx/Cre in post-treatment scans ($R^2 = -0.86$, $P = 0.0026$), but correlation was not statistically significant in pre-treatment scans ($R^2 = -0.36$, $P = 0.34$). Negative correlation between 2HG and glutamate/glutamine levels may be possible in mutant IDH through a shunting effect. Glutamine via glutamate is used to produce $\alpha$-ketoglutarate that is further a substrate for 2HG synthesis. Glutamine represents the major source of 2HG production (8) in mutant IDH cells and is also a major source for tricarboxylic acid cycle (TCA) anaplerosis, suggesting glutaminolysis (21) as a plausible therapeutic intervention. Lower glutamate and glutamine levels have been measured in mutant versus wild-type IDH cell cultures (45), while results from animal models have yielded a more complex fate of glutamate and glutamine (46, 47). To our knowledge our findings are the first corroborating in vivo evidence in patients suggesting that changes in glutamate and glutamine may accompany the loss of 2HG production with glioma treatment.
Except of 2HG no other metabolite showed a statistical significant effect in the post-treatment scan. In particular, choline that is the most measured metabolite for monitoring cancer progression and response by MRS was not sensitive to treatment changes within 1–3 months post-treatment, highlighting the potential of 2HG as an earlier imaging marker sensitive to treatment effects. In an earlier study (23) of temozolomide in glioma patients choline MRS at three months has been shown to not significantly different from baseline, while a more recent study (24) has shown that Cho/Cre at 3 months predicts tumor response. The difference between our results and of previous studies may be explained by both different patient population (all IDH mutant) and treatment (radiotherapy). In addition, arguably choline is not a specific marker for tumor cells, and the levels of choline can change also due to inflammation, gliosis and other biological responses of non-tumor cells, while 2HG accumulates only in the mutant cells.

Special attention and careful analysis needs to be exercised when computing functional maps. The difference method that is at the core of the functional mapping requires accurate longitudinal coregistration of baselines and follow up images, especially of the tumor ROI, which is not a trivial problem. Besides rigid misalignment of head position which is easier to correct, more challenging situations can arise from deformation within the brain due to the tumor growth or treatment. To minimize rigid displacement we have used automatic alignment of baseline and follow up MRSI during the acquisition coupled with real-time motion correction, and followed by post-processing head coregistration. On the other hand the fact that IDH tumors grow very slow and the time between two scans is relatively short the internal brain deformation is minimal, and the influence at the spatial resolution of MRSI may be minimal too.

Additionally, the quality of MRSI is critical for the end results. While our methodology has improved robustness designed to increase the number of patients from which good quality MRSI can be obtained, there will be a limited number of cases, in particular with tumor location in areas of large B0 inhomogeneity close to air cavities (sinuses, nasal) which will be challenging for MRSI. For these situations further development is necessary.

Several limitations exist in our study. There are few patients of different glioma grades and histologic subtypes, limiting our ability to assess correlation between 2HG imaging and histologic subgroups. The utility of 2HG imaging to predict OS and PFS cannot yet be investigated, as this determination will require at least 5–10 years of follow-up after baseline imaging, due to the prolonged clinical course of mutant IDH gliomas. The value of 2HG imaging for stratifying patients according to their RANO (4, 5) criteria treatment response (CR, PR, MR, SD, PD) could only be partially compared, since RANO criteria are based upon mixed glioma cohorts (IDH wild-type and mutant) and differ across WHO histological grades (high-grade, WHO grades III and IV - Ref. (4), versus low-grade, WHO grade II - Ref. (5)). We did nevertheless observed a correlation of 2HG imaging with high-grade RANO criteria for patients that show progressive disease, but other patient categories are under-represented at this interface between the two paradigms of glioma classification (histologic versus molecular). Our methods will only apply to the molecularly-homogeneous population of IDH mutant gliomas, a cohort for which objective clinical response criteria are not yet established. Existing data suggest that mutant IDH glioma of different WHO grades
may be more similar, than for example wild-type and mutant glioma of the same WHO grade (48). In addition, it is known that the peak “best response” of traditional radiographic response criteria are often delayed (49) with respect to metabolic activity, and IDH-mutant tumors have typically less contrast enhancement (48, 50) compared to wild-type IDH gliomas.

A 2HG/Cre threshold of 0.115 was found to be statistically significant for change of 2HG levels. Assuming a total Cre level between 8–10 mM (Suppl. Fig. 7S and Suppl. Ref. 7), the 2HG/Cre threshold of 0.115 would correspond to a 2HG level around 1 mM, which corresponds to the minimum concentration that can be detected by in-vivo MRS. Interestingly, the 0.115 threshold compared to the range [0.15–0.45] of 2HG/Cre values is similar to what was observed for ADC (threshold of $0.4 \times 10^{-3}$ mm$^2$/s for a range of $[0.5–1.5] \times 10^{-3}$ mm$^2$/s, (33)). However a longer study and more subjects will be required to establish the optimal imaging time point and the threshold of 2HG as an endpoint that can be used to determine or predict the objective response rate in this patient population.

Nevertheless, we demonstrate that longitudinal 3D functional imaging of 2HG is clinically feasible for non-invasive treatment monitoring. Our methodology combines imaging and genomics, and 2HG imaging is a clear example for the value of imaging genomics (radiogenomics) paradigm (50) in IDH mutant gliomas. As an imaging technique 3D fSM has the benefit of posing minimal risks to patients with the possibility to repeat examinations as many times as necessary. Compared to other molecular imaging modalities (PET, SPECT) the 3D MRSI is cost effective, easy to perform, and not confounded by the permeability of blood-brain barrier. Our novel method represents a major advance over prior point-diagnostic methods (14–16), and has immediate practical implications for patients with mutant IDH gliomas, who can now enter ISM surveillance after initial treatment. In contrast to other MRI modalities that can be used to assess treatment response (27, 50) our methods provide quantitative and mechanistic information about IDH enzymatic activity, metabolic interactions and networks involved in this subtype of glioma. In vivo longitudinal measurements of intratumoral 2HG levels will be highly valuable to probe target modulation in combination with pharmaco-kinetic/dynamic modeling of novel drugs acting on IDH-mutant glioma. We anticipate that it will prove critical to understand the complex and ongoing relationship between mutant IDH1 enzyme activity, 2-HG levels, and tumor maintenance to successfully impact the course of this disease through targeted treatment or other selective agents such as metabolic inhibitors, demethylating agents and synthetic lethality (20–22). Our measurements demonstrate the potential to probe pharmacological modulation by in vivo 2HG measurements, however at the moment our study does not provide evidence for an early therapy termination based on the 2HG levels. Our methods are therefore expected to play an important role as precision medicine tools to accelerate clinical trials of IDH mutant specific glioma therapies.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.
Acknowledgements

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References


Statement of Translational Relevance

IDH mutations are at the center of genetic, epigenetic and molecular alterations that define a distinct subtype of glioma patients. As a direct product of mutant IDH enzyme, 2HG can probe enzymatic activity and target modulation by inhibitory drugs. Hence, measurements of 2HG levels can accelerate clinical trials and translation of IDH mutant specific glioma therapies. However, repeated invasive sampling of tumor tissue is generally not feasible in glioma patients and non-invasive measurements of 2HG are necessary. In vivo imaging of 2HG levels using magnetic resonance spectroscopy (MRS) provides a safe and effective way to probe mutant IDH glioma patients. Our 3D MR spectroscopic imaging of 2HG has the temporal stability and spatial coverage required to monitor treatment response and tumor heterogeneity in IDH-mutant glioma patients, and can be easily incorporated into clinical trials of targeted therapies as precision medicine tool to determine objective response rates.
Figure 1.
3D metabolic maps in a mutant IDH glioma patient (grade II, astrocytoma, R132C) before and after radiotherapy. The pre- and post-treatment maps of each metabolite are scaled to the same intensity scale. In the last column the mask of CRLB < 25% is shown for 2HG goodness of fit by LCModel.
Figure 2.
Functional spectroscopic maps (fSM) of 2HG/Cre. fSM from three patients are shown: upper patient grade II (WHO) astrocytoma, middle patient grade III (WHO) anaplastic astrocytoma, and lower patient grade IV (WHO) glioblastoma. A threshold $\Delta 2HG/\text{Cre} = 0.11$ was used to classify voxels as decreased (blue), increased (red) or stable (green) 2HG/Cre. The fSM maps are masked by the CRLB < 25% mask.
Figure 3.
Histograms for 2HG/Cre values within tumor ROI for all patients: post-treatment histograms (blue line) compared to pre-treatment histograms (red line). Bottom panel includes bar plots for: A) median 2HG/Cre, B) fractional tumor volume of 2HG/Cre (>0.1), and C) fractional volumes of decreased, stable and increased 2HG/Cre (Δ2HG/Cre = 0.11). Fractional volumes are calculated as the ratio of 2HG/Cre or Δ2HG/Cre volume within specified thresholds over the total volume of tumor outlined on FLAIR images and masked by CRLB < 25%.

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Figure 4.
Box plots of imaging biomarkers and clinical correlation. A) Fractional changes ((post-pre)/pre*100, %) of metabolite ratios, ADC mean tumor ROI values, and tumor volumes of 2HG/Cre (>0.1) and FLAIR. Vertical red lines indicate the variability of test-retest measurements. Stars indicate statistical significant change (P < 0.05) between pre- and post-treatment scans. B) Fractional volumes of decreased, increased and stable 2HG/Cre and ADC, respectively. The hash symbol (#) indicates statistical significant correlation with KPS clinical outcome. C) Univariate linear regression of Karnofsky Performance Status (post-pre change) and the fractional tumor volume of decreased 2HG/Cre (PR = partial response, MR = minor response, SD = stable disease, PD = progressive disease by RANO criteria).
Fractional values for several 2HG metrics: median 2HG/Cre, volume of 2HG/Cre>0.1, volume of decreased 2HG/Cre (fVd), volume of increased 2HG/Cre (fVi), and volume of stable 2HG/Cre (fVs).

<table>
<thead>
<tr>
<th>Patient #</th>
<th>2HG/Cre</th>
<th>Volume (2HG/Cre&gt;0.1)</th>
<th>fVd (Δ&lt;−0.11)</th>
<th>fVi (Δ&gt;0.11)</th>
<th>fVs (Δ∈[−0.11,0.11])</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>−37.37</td>
<td>−50.09</td>
<td>60.01</td>
<td>0.00</td>
<td>39.99</td>
</tr>
<tr>
<td>2</td>
<td>−69.32</td>
<td>−51.87</td>
<td>58.23</td>
<td>24.58</td>
<td>17.18</td>
</tr>
<tr>
<td>3</td>
<td>−54.01</td>
<td>−61.55</td>
<td>76.74</td>
<td>0.86</td>
<td>22.41</td>
</tr>
<tr>
<td>4</td>
<td>−65.39</td>
<td>−34.61</td>
<td>99.12</td>
<td>0.00</td>
<td>0.88</td>
</tr>
<tr>
<td>5</td>
<td>2.88</td>
<td>−32.15</td>
<td>24.82</td>
<td>8.75</td>
<td>66.43</td>
</tr>
<tr>
<td>6</td>
<td>−34.34</td>
<td>−14.86</td>
<td>60.76</td>
<td>1.09</td>
<td>38.15</td>
</tr>
<tr>
<td>7</td>
<td>−48.09</td>
<td>−42.53</td>
<td>49.11</td>
<td>3.29</td>
<td>47.59</td>
</tr>
<tr>
<td>8</td>
<td>−22.30</td>
<td>−26.29</td>
<td>19.78</td>
<td>7.95</td>
<td>72.27</td>
</tr>
<tr>
<td>9</td>
<td>−48.89</td>
<td>−37.70</td>
<td>58.01</td>
<td>0.00</td>
<td>41.99</td>
</tr>
</tbody>
</table>

All values are given as percentage change (%). Fractional changes of all imaging biomarkers are given in Supplementary Table 3S.
Table 2

Mean and confidence interval (95% CI) of imaging biomarkers across patient group. The median over the tumor ROI was considered for the biomarkers of each patient.

<table>
<thead>
<tr>
<th>Imaging Biomarker</th>
<th>Pre</th>
<th>Post</th>
<th>Difference</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2HG/Cre</td>
<td>0.36 ± 0.09</td>
<td>0.20 ± 0.05</td>
<td>−0.16 ± 0.09</td>
<td>0.0069</td>
</tr>
<tr>
<td>GLX/Cre</td>
<td>1.06 ± 0.18</td>
<td>0.99 ± 0.18</td>
<td>−0.07 ± 0.10</td>
<td>0.2278</td>
</tr>
<tr>
<td>Lac/Cre</td>
<td>0.35 ± 0.16</td>
<td>0.30 ± 0.12</td>
<td>−0.05 ± 0.04</td>
<td>0.0576</td>
</tr>
<tr>
<td>tCho/Cre</td>
<td>0.33 ± 0.09</td>
<td>0.30 ± 0.09</td>
<td>−0.03 ± 0.04</td>
<td>0.1455</td>
</tr>
<tr>
<td>NAA/Cre</td>
<td>0.83 ± 0.10</td>
<td>0.80 ± 0.11</td>
<td>−0.02 ± 0.06</td>
<td>0.4686</td>
</tr>
<tr>
<td>tCho/NAA</td>
<td>0.47 ± 0.19</td>
<td>0.41 ± 0.15</td>
<td>−0.06 ± 0.11</td>
<td>0.2543</td>
</tr>
<tr>
<td>ADC (µm²/ms)</td>
<td>1.29 ± 0.09</td>
<td>1.32 ± 0.11</td>
<td>0.03 ± 0.04</td>
<td>0.2075</td>
</tr>
<tr>
<td>Volume 2HG/Cre (cm³)</td>
<td>7.65 ± 6.55</td>
<td>4.50 ± 4.43</td>
<td>−3.15 ± 2.53</td>
<td>0.0406</td>
</tr>
<tr>
<td>Volume FLAIR (cm³)</td>
<td>35.90±17.36</td>
<td>38.59 ± 20.57</td>
<td>2.69 ± 7.61</td>
<td>0.5075</td>
</tr>
</tbody>
</table>

Values are given for pre-, post-treatment, and the difference (post-pre)-treatment.