Longitudinal MR spectroscopy to detect progression in patients with lower-grade glioma in the surveillance phase

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Abstract

Background. Monitoring lower-grade gliomas (LrGGs) for disease progression is made difficult by the limits of anatomical MRI to distinguish treatment related tissue changes from tumor progression. MR spectroscopic imaging (MRSI) offers additional metabolic information that can help address these challenges. The goal of this study was to compare longitudinal changes in multiparametric MRI, including diffusion weighted imaging, perfusion imaging, and 3D MRSI, for LrGG patients who progressed at the final time-point and those who remained clinically stable.

Methods. Forty-one patients with LrGG who were clinically stable were longitudinally assessed for progression. Changes in anatomical, diffusion, perfusion and MRSI data were acquired and compared between patients who remained clinically stable and those who progressed.

Results. Thirty-one patients remained stable, and 10 patients progressed. Over the study period, progressed patients had a significantly greater increase in normalized choline, choline-to- N-acetylaspartic acid index (CNI), normalized creatine, and creatine-to- N-acetylaspartic acid index (CRNI), than stable patients. CRNI was significantly associated with progression status and WHO type. Progressed astrocytoma patients had greater increases in CRNI than stable astrocytoma patients.

Conclusions. LrGG patients in surveillance with tumors that progressed had significantly increasing choline and creatine metabolite signals on MRSI, with a trend of increasing T2 FLAIR volumes, compared to LrGG patients who remained stable. These data show that MRSI can be used in conjunction with anatomical imaging studies to gain a clearer picture of LrGG progression, especially in the setting of clinical ambiguity.

Key Points

• Progressed LrGG patients had greater increases in multiple metabolic markers than stable patients.
• MRSI can be used in conjunction with anatomical imaging studies to gain a clearer picture of LrGG progression, especially in the setting of clinical ambiguity.

Lower-grade gliomas (LrGGs) are either WHO grade 2 and 3 astrocytomas (isocitrate dehydrogenase (IDH)-mutant) or WHO grade 2 oligodendrogliomas (IDH-mutant with chromosome 1p/19q co-deletion), that can be differentiated from higher-grade gliomas (WHO grade 4, astrocytoma, IDH-mutant and the more common glioblastoma; WHO grade 3 oligodendrogliomas) by their unique histological features. Advances in molecular characterization of these tumors have
Importance of the Study

The goal of this study was to compare longitudinal changes in multiparametric MRI, including diffusion weighted imaging, perfusion imaging, and 3D MRSI, for LrGG patients in surveillance with tumors that progressed and those who remained clinically stable. LrGG patients that progressed had significantly increasing choline and creatine metabolite signals on MRSI, with a trend of increasing T2 FLAIR volumes, compared to LrGG patients who remained stable. Together, these data show that MRSI can be used in conjunction with anatomical imaging to gain a clearer picture of lower-grade glioma progression, especially in the setting of clinical ambiguity.

clarified classification opening up new avenues for potential targeted treatments that could lead to improvements in treatment paradigms, increased survival, and reduced morbidity for long term survivors. While the ability to delay radiation and chemotherapy is important for reducing side effects and optimizing quality of life, the relatively slow rate of change in lesion volume and lack of specificity from standard imaging methods, such as the Brain Tumor Imaging Protocol, hinders the identification of an optimal time point for initiating a new treatment. Further, standard imaging methods are not able to predict which patients will experience stable disease, progression, or a favorable response to therapy. Determination of tumor growth in LrGG is highly dependent on imaging technique, which is likely to vary given the long-time course over which LrGG patients are followed and the heterogeneity of treatment. LrGGs are also often irregular in shape due to their infiltrative growth pattern and can be further distorted by treatment effects (eg, post-surgical gliosis, radiotherapy related changes, edema), which can impair accurate measurements of tumor size from linear 2D measurements with T2 FLAIR being the standard surveillance modality for LrGG typically showing non-enhancing tumor growth. Further, 2D measurements of tumor size over serial MR exams for slow growing tumors can be affected by technical factors such as variations in slice orientation, acquisition parameters, or scanner type. These characteristics of LrGGs and their imaging can lead to significant inter-observer and intra-observer variability in tumor measures when using a 2D product method to estimate tumor size. Thus, there is a need for more sensitive imaging predictors of tumor growth and alternative measures of treatment response.

Multiparametric MR examinations that integrate conventional anatomic imaging with more advanced imaging techniques such as diffusion weighted imaging (DWI), perfusion imaging, and magnetic resonance spectroscopic imaging (MRSI) have been developed to provide a more comprehensive evaluation of the structural, physiologic, and metabolic properties of LrGGs. To date, these advanced imaging techniques have been primarily studied in the evaluation of suspected newly diagnosed LrGG to assist with targeting for tissue sampling and in non-invasive diagnosis. The discovery of IDH mutation status as an important differentiator of glioma classification has afforded detection of its metabolite 2-hydroxyglutarate (2HG) using MRSI. While recent studies have supported its use as a biomarker for metabolic activity in lower-grade glioma, the technical and tumor volume requirements for 2HG measurement limit its current clinical utility, and emphasize the continued need for multiparametric MRI biomarkers of progression in LrGG.

The goal of this study was to compare longitudinal changes in multiparametric MRI, including DWI, perfusion imaging, and 3D MRSI, for LrGG patients who progressed at the final timepoint and those who remained both clinically and radiographically stable. We hypothesized that these techniques would be valuable in the serial evaluation of LrGGs undergoing surveillance and would offer additional information beyond conventional anatomic imaging in identifying tumor progression.

Methods

Patients

This study included histologically confirmed LrGG patients who were clinically stable and off treatment (ie, surgery, radiation, chemotherapy). There was no upper limit on the time between completed treatment and enrollment, or the number and types of previous treatments. Clinical stability was determined by the referring neuro-oncologist which included patients that recently received surgery and no further therapies. Eligible patients needed to be over 18 years old, have a Karnofsky performance status (KPS) ≥ 70, and be fluent in English. All participants gave written informed consent, and ethical approval was granted by the UCSF Institutional Review Board, in compliance with the Helsinki Declaration.

Clinical and Demographic Variables

Clinical and demographic data were collected by review of patients’ electronic medical records. This included age, education, time since diagnosis/surgery, previous treatment, tumor grade, extent of resection (biopsy, subtotal, gross total), KPS and use of anti-epileptic drugs (AEDs). Patients were classified according to the 2021 WHO central nervous system classification integrated diagnoses: astrocytoma, IDH-mutant; oligodendroglioma, IDH-mutant and 1p19q co-deleted; and NOS (not otherwise specified: molecular status unknown). IDH mutation status and co-deletion of
1p19q were determined by immunohistochemistry and FISH, respectively. MRI was performed at intervals according to the treating neuro-oncologist recommendations ranging from every 3–6 months depending on point in disease course.1,2 Serial MRIs were performed for each patient until the time of progression or until the time the dataset was locked for analysis.

**Clinical Definition of Progression**

Progression was determined by the treating neuro-oncologist with or without consensus of multidisciplinary tumor board, based on change in the T2 FLAIR or T1 contrast-enhancing lesions, or new or worsening neurological symptoms, with a recommendation for a change in treatment.

**MRI Acquisition**

All scans were performed on a GE Discovery 750 3T scanner (GE Healthcare, Waukesha, WI) with a 32-channel head coil (Nova Medical, Wilmington, MA). Standard anatomical imaging included 3D T2-weighted Fluid Attenuated Inversion Recovery (FLAIR), 2D T2-weighted fast spin echo, and 3D T1-weighted pre- and post-gadolinium sequences. DWI was obtained in the axial plane with 24 gradient directions [repetition time (TR)/echo time (TE) = 1000/108 ms, voxel size = 1.7 x 1.7 x 3 mm, b = 1000 s/mm]. Dynamic susceptibility contrast-enhanced (DSC) perfusion images were acquired before, during, and following a 5 ml/s bolus injection of 0.1 mmol/kg body weight gadolinium diethylenetriamine pentaacetic acid (Gd-DTPA) using a series of T2*-weighted echo-planar images [TR/TE/Flip-angle = 1500/35 ms/30–35°, 128 x 128 matrix, slice thickness = 3–4 mm, 20–25 slices with 80 time points].

**MRSI Acquisition Using Atlas-Based Prescription**

Our MRSI method predefines the prescription on a standard image atlas and transforms both the selected volume and outer volume suppression bands to individual subject space during each scan session.13 This allows a fully automated prescription that covers the desired anatomy consistently for different time points in a single subject.13 In each scan session, the atlas T1-weighted image was aligned to the 3D T1 image volume of the subject, and the predefined MRSI volume and outer volume suppression bands were transformed by applying the transformation matrix resulting from the alignment. Higher order shimming was performed to reduce the degree of magnetic field in homogeneity. Immediately after the field shimming, the 3D multi-voxel H-1 MRSI was acquired with point resolved spectroscopy (PRESS) volume selection, chemical shift selective (CHESS) water suppression, phase encoding in two dimensions and flyback encoding in the superior-inferior direction.13 The sequence parameters were as follows: TE/TR 144/1250 ms, phase encoding grid 18 x 18 x 16, nominal resolution 1cm, total acquisition time 11 min. An over excitation factor of PRESS volume 1.2 was used to reduce chemical shift misregistration.14

**MR Data Processing**

The anatomic, diffusion, perfusion and spectroscopic images were aligned to the T1 post-contrast image using FMRIB’s Linear Image Registration Tool.15,16 Each region of interest (ROI) was defined by a single investigator with the guidance of UCSF neuroradiologists. The T2 lesion ROI was defined using semi-automated software (3DSlicer) to include all T2 FLAIR hyperintensity relative to the surrounding normal tissue. These T2 lesion ROIs are 3D volumetric ROIs that include all T2 FLAIR hyperintensity on all slices. A contrast-enhancing lesion ROI was defined when contrast-enhancement was present.

A previously published algorithm was applied to estimate relevant DWI parameters and normalize between field strengths using estimates from normal appearing brain tissue.17 To provide metrics that described regions with abnormal intensities, the voxel values for ADC and FA maps were first normalized to the mode of intensities in normal appearing brain tissue (calculated across the entire cerebrum, excluding the T2 lesion ROI). Percentiles (10th, 50th, 90th) were then calculated from histograms of normalized intensities within T2 lesion ROIs.

Perfusion imaging data, including cerebral blood volume (CBV) and peak height (PH), were calculated for each voxel using software developed in our research group.18-20 The T2* signal-intensity time curves acquired during the first-pass of the gadolinium bolus were converted to change in the relaxation rate (ΔR2*). PH was defined as the maximum DR2* value of the first-pass curve. CBV maps were calculated on a voxel-by-voxel basis utilizing a modified gamma-variate function that considers leakage of the contrast agent.19 The CBV and PH values from the T2 FLAIR lesions were normalized by the median value of the histogram derived from normal brain tissue.21,22

MRSI data acquisition was conducted using fully automated reconstruction and post-processing described in previous publications.13,14,23 In brief, the start point was the raw data file obtained from the lactate-edited MRSI sequence. This comprised interleaved acquisitions (or cycles) obtained with radiofrequency pulses that modulated the phase of the lactate peak, with each cycle having 8-channels of data corresponding to the multiple receiver coils. The k-space time domain data for each cycle and channel were first filtered with a 4-Hz exponential function in the time domain, zero filled to 1024 points, and Fourier transformed to produce a k-space array of spectra. The next step was to apply the k-space Fourier transforms to produce 3D spatial arrays of spectral data, followed by combination using in-house developed software that weights the data by coil sensitivities estimated from low resolution proton density weighted images.13 Additional phase corrections were applied in the SI dimension to account for the flyback echo-planar readout gradient.24 The cycles were summed to produce an array of spectra containing choline (Cho), creatine (Cr), NAA (N-Acetylaspartic acid), and lipid and subtracted to produce an array of spectra containing lactate. Spectra were baseline subtracted and phase and frequency corrected using parameters estimated from the summed array.

Peak heights and areas were determined from baseline subtracted, frequency and phase corrected spectra on a
voxel-by-voxel basis. Metabolite PHs were normalized by median PHs in normal brain for choline and creatine (nCho and nCr), and normalized lactate and lipid (nLac, nLip) were estimated by metabolite PHs divided by the median peak NAA intensity in normal brain. The choline-to-NAA index (CNI), a z-score that reflects changes in the relative levels of these two metabolites compared to normal brain voxels, and the creatine-to-NAA (CRNI) index and choline-to-creatine (CCRI) index, defined in a similar manner, were computed automatically for each voxel using in-house software.

### Statistical Analysis

The relationships among the within-subjects effect of time (days to progression/latest scan), the between-subjects effect of WHO type (histological and molecular integrated diagnosis), and MRI parameters were assessed with repeated-measures analyses of variance (using JMP Pro 13.0, SAS Institute Inc). The cut-off for defining a significant result was a $P$ value of .05, with a Benjamini-Hochberg False Discovery Rate correction for multiple comparisons. Significant main effects for three-way interactions were followed up with pairwise Tukey-Kramer tests.

### Results

#### Patient Characteristics

Our study cohort included 41 total patients with LrGG, 31 who remained stable throughout the study and 10 who had disease progression. The median age of the population was 45 (range 24–66 years) and 17 (41%) were female. There were 26 patients with astrocytoma, IDH-mutant, 12 patients with oligodendroglioma, IDH-mutant, and

| Table 1. Patient demographics |
|--------------------------------|
|                              | Progressed | Stable | Progressed vs stable |
| Age Median years              | 45         | 44     | $F = .86, P = .36$    |
| Grade                         |            |        |                      |
| 2                             | 7          | 19     | Chisq = 25, $P = .62$ |
| 3                             | 3          | 12     |                      |
| WHO type                      |            |        |                      |
| Astrocytoma IDH-mutant        | 6          | 20     | Chisq = 3.55, $P = .06$ |
| Oligodendroglioma              | 4          | 8      |                      |
| NOS                           | 0          | 3      |                      |
| Upgraded at progression       |            |        |                      |
| Yes                           | 2          | 9      | Chisq = 5.28, $P = .022$ |
| No                            | 5          | 22     |                      |
| No surgery                    | 3          | 1      |                      |
| Prior chemotherapy            |            |        |                      |
| No                            | 7          | 9      | Chisq = 3.93, $P = .048$ |
| Yes                           | 3          | 14     |                      |
| Prior radiation therapy       |            |        |                      |
| No                            | 8          | 14     | Chisq = 3.93, $P = .048$ |
| Yes                           | 2          | 17     |                      |
| Number of prior surgeries     |            |        |                      |
| Median                        | 1          | 1      | $F = .27, P = .61$    |
| Mean                          | 1.5        | 1.4    |                      |
| Time from diagnosis           |            |        |                      |
| Median days                   | 1184       | 1856   | $F = .5, P = .48$     |
| Mean days                     | 1978       | 2483   |                      |
| Number of MRI scans           |            |        |                      |
| Median                        | 6          | 4      | $F = 3.75, P = .06$   |
| Mean                          | 5.4        | 4.3    |                      |
| Presence of T1 contrast-enhancement |        |        |                      |
| Baseline scan                 | 1          | 2      | ns                   |
| Progression/final scan        | 2          | 2      |                      |
1p/19q-co-deleted, and 3 NOS patients for whom the molecular status was unknown (Table 1). Additional clinical data can be found in Supplementary Table 1.

Volumetrics

The trend towards increasing T2 FLAIR volume over time in progressed patients compared to stable patients did not reach significance ($F = 3.64, P = .065$) (Table 2, Figure 1). Progressed patients had a greater increase in contrast-enhancing volume over time than stable patients ($F = 4.55, P = .039$), although only two stable patients and two progressed patients had any contrast-enhancement. Regarding the two patients with contrast-enhancement, in one patient the enhancement was attributed to post-radiation change and was stable over the course of the study and the other patient the enhancement was attributed to post-surgical change and resolved over the course of the study. The remaining 37 patients had no contrast-enhancement at any time-point.

Diffusion Weighted and Perfusion Imaging

There were no significant differences between progressed and stable patients in change over time for any diffusion or perfusion measures. Across all patients, normalized CBV (nCBV) decreased over time ($F = 5.19, P = .028$). There was also a significant interaction between progression status and WHO diagnosis on nCBV that did not vary over time ($F = 5.5, P = .025$) Although post-hoc assessments in oligodendroglioma patients did not reach significance, those who progressed had higher nCBV than stable patients ($F = 3.86, P = .076$), whereas astrocytoma progressed patients had no difference in nCBV from stable astrocytoma patients ($F = 1.6, P = .22$).

MRSI

Progressed patients had a greater increase in CNI, nCho, CRNI, and nCre over time than stable patients ($F = 5.13, P = .03$; $F = 4.12, P = .05$; $F = 5.8, P = .023$; and $F = 5.57, P = .025$, respectively) (Figure 1). There was also a significant three-way interaction between change in CRNI over time, progression status and WHO type (oligodendrogliomas vs astrocytomas vs Not Otherwise Specified tumors (NOS) patients excluded) ($F = 9.72, P = .005$). In astrocytomas, progressed patients had a greater increase in CRNI over time than stable patients ($F = 14.2, P = .0012$), while in oligodendrogliomas, the progress...
difference between progressed and stable patients was not significant ($F = .45$, $P = .51$). Although nLip and nLac decreased over time across all patients ($F = 5.6$, $P = .023$; $F = 14.73$, $P = .0005$), progressed patients had significantly higher levels of nLip ($F = 9.87$, $P = .004$) and nLac ($F = 5.18$, $P = .03$) compared to stable patients, regardless of time-point. Figures 2 and 3 show example metabolite spectra for a patient that progressed and a patient that remained stable, respectively.

**Discussion**

In this study, we found 3D MRSI characteristics of post-treatment LrGG in surveillance that significantly increased over time in patients who progressed, compared to patients who remained stable. Overall, our findings build upon a growing body of evidence that routine MRSI in LrGG provides prognostic value through its unique capability to measure local metabolic activity, indicating tumor changes that can guide management. T2 FLAIR volume increase, the primary indicator of clinical progression in LrGG, can be subtle and non-specific in LrGG patients undergoing post-treatment surveillance. In this study, both progressed and stable LrGGs showed T2 FLAIR volumes increasing over time, and the difference between the groups did not reach significance. Given this ambiguity, MRSI provides unique and complementary information to understand tissue metabolism and characterize subtle morphologic changes. This is particularly relevant in light of recent evidence that highlights the importance of delaying chemotherapy or radiation in LrGG to reduce the risk of malignant transformation through induced hypermutation as well as a preventative measure against lasting effects of chemotherapy and radiation related neurocognitive changes.

Our findings demonstrate the utility of multiple metabolic indicators that showed significant changes leading up to progression. Significantly larger increases in the levels of nCho, choline-to-NAA index, nCre, and creatine-to-NAA index were observed in patients who progressed, compared to those who remained stable. Choline is associated with cell proliferation and altered membrane phospholipid metabolism. The MRSI creatine peak includes both creatine and phosphocreatine, and is a marker of cellular metabolism, while NAA is a marker of normal brain tissue and actively functioning neurons helpful in most brain tumors classes.

Despite the small number of patients studied, CRNI showed a significant increase over time in progressed
astrocytoma compared to stable astrocytoma, an effect that was not significant in oligodendroglioma, suggesting that changes in creatine levels may offer information specific to WHO glioma types. Howe et al. showed that glioblastomas had lower levels of creatine than grade 2 and 3 astrocytomas. Lupo et al. found that within abnormal perfusion PH areas of grade 4 gliomas (indicative of leaky blood vessels), reductions in creatine and increases in lactate were observed. Lower creatine within areas of abnormal PH was also observed in grade 3 LrGGs, while areas of elevated CBV had higher creatine, potentially signifying that more energy is needed in the initial recruitment and formation of new vessels. Ozturk-Isik et al. reported creatine was reduced in grade 3 LrGG lesions compared to normal appearing white matter, but in areas of high metabolic activity (CNI > 4) within the lesion, creatine was higher. Increased creatine may be an early biomarker of reduced energy reserves from anaerobic respiration before lactate accumulation. We hypothesize that once transformation to a more malignant phenotype occurs, the creatine declines and lactate levels begin to rise, especially in regions with leaky vessels. Previous cross-sectional studies have shown metabolic differences between oligodendroglioma and astrocytoma using MRSI. Stadlbauer et al. found that grade 3 oligodendrogliomas had a significantly lower maximum creatine concentration than grade 3 astrocytomas. Vuori et al. reported that grade 2 astrocytomas had lower normalized creatine, and grade 2 oligodendrogliomas had higher normalized creatine, compared with matched controls subjects. Our longitudinal results may offer further insight into creatine's role in glioma metabolism.

The utility of MRSI is exemplified in its ability to depict areas of active tumor metabolism even in the setting of non-specific or inconclusive structural MRI findings. MRSI may direct clinicians to follow suspicious areas of interest or label MRI changes as clinically silent with greater confidence. Caveats of MRSI include the required additional acquisition time per scan, the necessity for an experienced technologist in protocols beyond the typical MRI scan procedures, and the added complexity of post-acquisition processing. To combat this, we have organized a streamlined acquisition process, including atlas-based automatic prescription and fast MRSI acquisitions using flyback echoplanar trajectory in addition to previously published institutional processes.

Higher-grade gliomas show DWI alterations, indicating increased cellularity, and increased perfusion, indicating neoangiogenesis and hypervascularity, while LrGGs often do not demonstrate these imaging changes. We did not find any significant differences in diffusion or perfusion changes over time between the progressed and stable cases.
LrGGs. Across all patients, diffusion metrics remained consistent over time, and nCBV decreased over time. These results suggest that longitudinal monitoring of diffusion and perfusion are not likely to aid in identifying LrGG progression in the setting of inconclusive structural MRI findings.

The limitations of this study include the relatively low numbers of patients with progression, as well as our patients being recruited from a single center. Including more patients will improve our ability to look at the role of MRSI and other advanced imaging parameters to determine progression in particular clinical subgroups of LrGG (eg, astrocytomas with previous radiotherapy and/or chemotherapy vs. those treated solely with surgery). Nonetheless, as treating physicians and clinical scientists, our impact is often measured in incremental improvements and while our data show small, though significant, differences the results provide support for the use of MRSI as an adjunct to often indeterminant imaging and clinical findings surrounding glioma progression.

Research connecting metabolic findings to the cellular and molecular underpinnings of higher-grade glioma biology has previously been done with glioblastoma cells lines, which showed high variability in metabolites as measured by MRSI and cell culture metabolites. A similar study has not been done for LrGGs. Several studies have shown considerable upregulation of genomic and transcriptomic components of gliomas pertaining to metabolic pathway enzymes such as lactate dehydrogenase and creatine kinase with levels increasing with glioma grade. Similarly, our results support these data from a metabolic imaging perspective with increased metabolic indexes of creatine and choline. Though we did not conduct a comprehensive enough study to validate these findings, future studies could combine pathology, transcriptomic, and MRSI data to create robust metabolic phenotyping of LrGGs.

**Conclusion**

This study supports the use of MRSI as an imaging complement to the clinical evaluation of lower-grade glioma progression. LrGG patients in surveillance with tumors that progressed had significantly increasing choline and creatine metabolite signals on MRSI, with a trend of increasing T2 FLAIR volumes, compared to LrGG patients who remained stable. Together, these data show that MRSI can be used in conjunction with anatomical imaging to gain a clearer picture of lower-grade glioma progression, especially in the setting of clinical ambiguity.

**Supplementary material**

Supplementary material is available online at Neuro-Oncology Advances online.

**Keywords**

lower-grade glioma | magnetic resonance spectroscopic imaging | progression.

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**Authorship**

Experimental design (TG, LNA, TLL, PD, JEVM)  
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Analysis of data (LNA, JEVM, PD, JP)  
Interpretation of the data (LNA, TLL, JEVM, PD, NAOB, JWT, SMC, JP)

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