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Characterization of Active and Infiltrative Tumorous Subregions from Normal Tissue in Brain Gliomas Using Multi-Parametric MRI

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ABSTRACT—

Background—Targeted localized biopsies and treatments for diffuse gliomas rely on accurate identification of tissue subregions, for which current MRI techniques lack specificity.

Purpose—To explore the complementary and competitive roles of a variety of conventional and quantitative MRI methods for distinguishing subregions of brain gliomas.

Study Type—Prospective

Population—51 tissue specimens were collected using image-guided localized biopsy surgery from 10 patients with newly diagnosed gliomas.

Field Strength/Sequence—Conventional and quantitative MR images consisting of pre and post-contrast T1w, T2w, T2-FLAIR, T2-relaxometry, DWI, DTI, IVIM, and DSC-MRI were acquired pre-operatively at 3T .

Assessment—Biopsy specimens were histopathologically attributed to glioma tissue subregion categories of active tumor (AT), infiltrative edema (IE), and normal tissue (NT) subregions. For each tissue sample, a feature vector comprising of 15 MRI-based parameters was derived from pre-operative images and assessed by a machine learning algorithm to determine the best multi-parametric feature combination for characterizing the tissue subregions.

Statistical Tests— For discrimination of AT, IE, and NT subregions, one-way ANOVA test and for pairwise tissue subregion differentiation, Tukey-HSD and Games-Howell tests were applied ($P<0.05$). Cross-validated feature selection and classification methods were implemented for identification of accurate multi-parametric MRI parameter combination.

Results— After exclusion of 17 tissue specimens, 34 samples (AT=6, IE=20, and NT=8) were considered for analysis. Highest accuracies and statistically significant differences for discrimination of IE from NT and AT from NT were observed for diffusion-based parameters (AUCs>90%), and perfusion-derived

parameter as the most accurate feature in distinguishing IE from AT. A combination of “*CBV*, *MD*, *T2_ISO*, *FLAIR*” parameters showed high diagnostic performance for identification of the three subregions (AUC~90%).

Data Conclusion—Integration of a few quantitative along with conventional MRI parameters may provide a potential multi-parametric imaging biomarker for predicting the histopathologically-approved glioma tissue subregions.

KEYWORDS— Glioma, Imaging Biomarker, Intra-tumor Heterogeneity, Multi-parametric MRI, Machine Learning

INTRODUCTION

Diffuse gliomas manifest extensive diffuse infiltration of tumor cells in adjacent brain parenchyma (1). Even when low grade, diffuse gliomas are generally fatal after several years (1,2). It is widely known that characteristic spatial intra-tumor variability within microenvironment of gliomas account for grim prognosis of these patients (3,4). Within heterogeneous tumors like gliomas, multiple subregions with different phenotypic characteristics coexist, which often represent heterogeneous genetic and microenvironmental profiles, likely to respond variably to treatment (5,6). Lack of sensitive and specific quantitative imaging biomarkers for realizing spatial variations of gliomas and localizing the most aggressive portion of the tumor leads to inaccurate biopsy sampling, which hinders target-specific diagnosis and therapies (5).

Targeted biopsy procedures and surgical/treatment planning for gliomas most often rely on conventional contrast-enhanced T1-weighted (CE-T1w) and T2-weighted (T2w) and T2-FLAIR images (7-9). Nonetheless, as diffuse gliomas tend to invade the brain tissue in small cell groups, tumor expansion and progression may precede changes in contrast enhancement (1,8). Furthermore, CE-T1w cannot sufficiently localize the most aggressive or active tumor sub-compartment to ensure reliable biopsy outcome (10,11). On the other hand, T2w hyperintense regions are incapable of characterizing the infiltrative glioma sub-region (infiltrative edema, also referred to as non-enhancing tumor (NET)), and diffuse infiltrating cells can be found beyond the extent of T2w hyperintensity borders (12,13). Hence, based on conventional MRI, stratification of the most aggressive/active part of the tumor and infiltrative glioma and detection of the extent of invasion from normal tissue remains challenging.

Various advanced MRI contrasts have been explored for their potential in localizing and grading glioma brain tumors. Diffusion weighted imaging (DWI) and diffusion tensor imaging (DTI) reflect the water proton mobility related to changes in cellular density, permeability of the cell membrane, and tissue microstructure(14,15). Perfusion-weighted imaging (PWI) visualizes microvascular changes and cellular proliferation; particularly, regional cerebral blood volume (rCBV) derived from dynamic susceptibility

contrast enhanced (DSC)-MRI, correlates with the degree of neo-vascularization and histopathological grade surpassing the performance of CE-T1w MRI (16-18).

There have been sporadic attempts to study the role of quantitative MRI methods, including DSC-MRI, DWI/DTI, IVIM, and T2-relaxometry for characterizing infiltrating glioma regions through biopsy validation (18-20). Hence, we sought to address: (1) the roles of individual MRI-derived parameters in differentiation of the three subregions, i.e. AT, IE, and NT, from each other; (2) the relationship among MRI-derived parameters with histopathological cellular density; and (3) the best multi-parametric combination of MRI-derived features for discrimination of AT, IE, and NT through a quantitative methodology.

MATERIALS AND METHODS

Patients

Between July 2015 and February 2016, 10 adult patients (6 men and 4 women; mean age 40.4 years; age range, 20-76 years) with newly diagnosed gliomas were prospectively recruited in this study. The patients underwent pre-surgical MRI and CT examinations, followed by image-guided neurosurgery within 1-3 days from image acquisition. Institutional review board (IRB) approval was obtained and the patients provided their informed written consent to be included in this study, designed in compliance with Health Insurance Portability and Accountability Act (HIPAA) law. Inclusion criteria for this study consisted of patients: (1) with suspected glioma brain tumors based on their initial MR scan; (2) with no prior treatment, including surgery or radiotherapy; (3) having no contra-indications for surgery; (4) in whom the tumor was not in eloquent area; and (5) for whom there were no technical challenges for performing surgery.

Patients or biopsy specimens were excluded if (1) they did not undergo surgery; (2) imaging was performed inadequately or some image sequences were missing; and (3) if histopathological assessment of gross tumor or specimens were impossible or not provided.

Image Acquisition

All patients underwent pre-operative MRI acquisition within two days prior to their surgery. Structural and physiological MRI acquisitions were performed on a 3T MRI scanner (Siemens MAGNETOM Tim Trio, Erlangen, Germany). Sagittal T2w turbo spin echo (TE/TR=80/6000, Slice Thickness = 5 mm, Flip Angle = 140°, FOV = 230×230 mm², Image Matrix = 320×320, Pixel size = 0.72×0.72 mm²). Axial T2w turbo spin echo (TE/TR=106/5400, Slice Thickness = 5 mm, Flip Angle = 120°, FOV = 200×200 mm², Image Matrix = 232×256, Pixel size = 0.78×0.78 mm²). Axial fluid attenuated inversion recovery (FLAIR) (TE/TR/TI = 115/8400/2240, Slice Thickness = 5 mm, Flip Angle = 125°, FOV = 181×200 mm², Image Matrix = 232×256, Pixel size= 0.78×0.78 mm²). Axial 3D MP-RAGE T2w spin-echo as a part of image-guided surgery protocol with TE/TR = 200/2500, Slice Thickness = 1 mm, Flip Angle = 120°, FOV = 208×256 mm², Image Matrix = 420×512, Pixel size= 0.50×0.50 mm².

Axial DWI using echo-planar imaging (EPI) sequence with TE/TR = 100/4000, Slice Thickness = 5 mm, Flip Angle = 90°, FOV = 200×200 mm², Image Matrix = 136×136, Pixel size = 1.47×1.47 mm, b-value = 50, 1000 s/mm²). Axial multi b-value DWI using EPI method with TE/TR = 100/4000, Slice Thickness = 5 mm, Flip Angle = 90°, FOV = 200×200 mm², Image Matrix = 136×136, Pixel size = 1.47×1.47 mm², b-values = 0, 50, 200, 400, 600, 800, 1000 s/mm². Diffusion tensor imaging (DTI) with 64 directions and TE/TR = 90/9000, Slice Thickness = 5 mm, FOV = 256×256 mm², Flip Angle = 90°, NEX = 1, Image Matrix = 128×128×72, Pixel size = 1.72×1.72 mm², b-value = 50, 1000 s/mm².

Axial multi-echo T2w spin-echo (T2-relaxometry) was carried out with TR=4000 ms, TE=12, 24, 36, 48, 60, 72, 84, 96, 108, 120, 132, 144, 156, 168, 180, 192 ms, Slice Thickness = 5 mm, Flip Angle = 180°, FOV = 200×200 mm², Image Matrix = 232×256, Pixel size = 0.78×0.78 mm².

After injection of 0.2 $mmol/kg$ of Gadovist (Bayer Schering Pharma, Berlin, Germany) at a rate of 5 ml/s and followed by a 20 mL saline flush, dynamic susceptibility contrast enhanced (DSC-) MRI was acquired using GE-EPI sequence with TE/TR = 30/1600, Slice Thickness = 5 mm , FOV = 220×220 mm^2 , Flip Angle = 70°, NEX = 1, Image Matrix = 128×128×20, Pixel size = 1.72×1.72 mm^2 , Number of Dynamic Scans = 64, Temporal Resolution = 1.5 s. To minimize the effects of contrast agent extravasation a pre-load of contrast agent was administered with about 25% of the total contrast dose about 4 minutes of incubation time before the second injection for DSC-MR imaging.

Pre- and post-contrast axial 3D MP-RAGE T1w spin-echo were also acquired as a part of the protocol required for navigation system TE/TR = 5/17, Slice Thickness = 1 mm , Flip Angle = 25°, FOV = 208×256 mm^2 , Image Matrix = 208×256, Pixel size = 1×1 mm^2 .

MR imaging pulse sequences and their parameter adjustments are summarized in **Table 1**.

Computed tomography (CT) images were acquired for all patients prior to surgery to ensure accurate registration of the images with the position of the patient during image-guided neurosurgery. 3D images were acquired on a 64-slice CT scanner (GE Healthcare Technologies, WI, USA) with no gantry tilt, slice thickness of 0.625 mm, from maxilla to the top of the head.

Quantification of MR Images

Diffusion Tensor Imaging (DTI)

Quantification of DTI was executed in ExploreDTI software (v4.8.6)(21) with proper parameter adjustments, and after EPI and patient motion correction. Maps of mean diffusivity (MD), fractional anisotropy (FA), and $L1$, $L2$, and $L3$ eigenvalues were exported from the software. Pure isotropic diffusion and pure anisotropic diffusion maps denoted by P and Q , respectively, were calculated based on this formula (22):

$$P = \sqrt{3}MD = \frac{\lambda_1 + \lambda_2 + \lambda_3}{\sqrt{3}}$$

$$Q = \sqrt{(\lambda_1 - MD)^2 + (\lambda_2 - MD)^2 + (\lambda_3 - MD)^2}$$

Dynamic Susceptibility Contrast-Enhanced (DSC-) MRI

DSC-MRI analysis was carried out based on an algorithm developed in-house, which inputs signal intensity-time curves of T_2^* -based dynamic images in each pixel, converts them to ΔR_2^* curves, selects the arterial input function (AIF) based on a modified automated algorithm, performs decorrelation using block-circulant SVD, and generates regional CBV ($rCBV$) map.

T2/Proton Density Mapping

In T_2 -relaxometry, images are acquired at multiple echo times to obtain an estimate of T_2 and proton density by fitting the following equation to the signal:

$$S = S_0 e^{-\frac{TE}{T_2}}$$

where S_0 is the proton density and T_2 is the spin-spin relaxation. Quantitative T_2 and proton density (PD) maps were generated by pixelwise fitting of the mentioned equation to the corresponding multi-echo signal.

Quantification of Intra-Voxel Incoherent Motion

Diffusion-weighted images are highly affected by flow of blood and cerebrospinal fluids at very low b -values (i.e. less than 100-200 s/mm^2) (23,24). Here, to separate contributions of diffusion and perfusion, the bi-exponential intra-voxel incoherent motion (IVIM) model is which is denoted by:

$$S = S_0 (f e^{-D^*b} + (1-f) e^{-Db})$$

where S_0 is the signal at b -value of zero, f is the fraction of signal affected by flow, D^* describes the decay of signal caused by flow, D is the measured diffusion coefficient which is a function of diffusion time. To nullify the effect of flow on diffusion measurements, diffusion coefficients (D) for each voxel were calculated using conservative minimum b -values of 200 s/mm² (26). As diffusion parameters are highly affected by non-Gaussian behavior of diffusion at higher b -values, for calculation of D , this effect was conservatively cancelled by using maximum value of 600 s/mm².

To account for non-Gaussianity of the signal in higher b -values, the modified IVIM model including diffusion kurtosis imaging (IVIM-DKI) was considered (25):

$$S = S_0 \left(f e^{-bD^*} + (1 - f) e^{-bD + \frac{b^2 D^2 K}{6}} \right)$$

where K is the kurtosis factor that measures deviation from the Gaussian behavior. D for each voxel was estimated by fitting the equation to the segment of the signal at b -values of 200, 400, and 600 s/mm² and K was estimated on b -values of 600, 800, and 1000 s/mm² using nonlinear least-squares method.

Biopsy Site Selection

The generated maps from different modalities, including $rCBV$ -map from DSC-MRI, ADC -map from DWI, FA , MD , P , Q from DTI, D , D^* , f , and K -maps from multi b -value diffusion imaging, T_2 and PD maps from T2-relaxometry technique, and conventional images comprising of $FLAIR$, T_2_ISO (MP-RAGE T2w), and MP-RAGE pre-contrast T1w, were co-registered with post-contrast MP-RAGE T1w (CE-T1w) image using rigid registration method with normalized mutual information (NMI) as the similarity measure in SPM software (<http://www.fil.ion.ucl.ac.uk/spm/>). The difference between post- and pre-contrast T1w images was calculated to form $T1_SUB$ (subtracted T1w) image, which shows the enhanced regions (Fig. 1).

A senior radiologist (K. F. with 12 years of experience in neuroradiology) identified the biopsy targets, i.e. active tumor, infiltration, and normal regions, based on presence of hyperintensity on CE-T1w, hypointensity on T2w and FLAIR images, and according to quantitative *ADC* and *rCBV* maps. Rectangular regions of interest (ROIs) with sizes of at most 8x8 pixels were marked on CE-T1w images and imported into the navigation workstation to plan for image-guided biopsy intervention. The regions were ideally identified on areas with best accessibility for the neurosurgeon for specimen collection and distant from vessels, ventricles, and critical brain regions.

Image-Guided Tissue Biopsy Procedure

The specimens were sampled from the patients by a senior neurosurgeon (M. Z. with 10 years of experience in neurosurgery) using disposable biopsy needle (Stryker Inc., Switzerland). The coordinates of the ROIs were recorded by the navigation system (OpticVision™, Parseh Intelligent Surgical Systems Co. (PARSISS), Iran), which were used to overlay on the images and MRI-derived quantitative maps. Based on the guide provided by navigation, needle biopsy was performed on all target points. For each patient, 4 to 6 specimens with at least 1-cm of distance were collected for histopathological assessments.

These technical considerations were addressed during the biopsy to minimize intra-operative brain shift and error:

- 1- The target points were pre-surgically marked in the order of normal tissue, edema, tumor, and necrosis;
- 2- Serum mannitol, corticosteroids or any other diuretics pre-/intraoperative were not given to patients prior to biopsy procedure;
- 3- Hypo/hyperventilation during surgery was avoided;
- 4- Minimal dural opening was performed to prevent CSF egress resulting in brain shift;

- 5- The target points in proximity of the ventricles were the last points for biopsy, to avoid unwanted entrance into the ventricles resulting in CSF drainage and therefore changing navigation accuracy due to brain shift;
- 6- For the target points in proximity of cystic portions within the tumor, similar strategy was adopted.

After needle biopsy, the dura was opened and based on pre-operative images, complete resection of the tumor was performed and the resected gross tumor volume was transferred for histopathological grading.

Histopathological Assessment of Tissue Samples

After being removed from the brain, the specimens were fixed in 10% buffered formalin and routinely processed for histopathological assessment. Five micrometer sections were stained using hematoxylin and eosin (H&E) method and reviewed by a pathologist (F. A. with 20 years of experience in neuropathology). For each specimen, the pathologist determined the presence of tumor cells and scored the samples as “normal tissue” where no tumor cells were identified, “positive tumor cells” when infiltrating tumor cells were present, and “tumor core” when tumor constituted the majority of tissue. Tumor was identified, classified and graded based on morphological features, such as density and distribution of cells, nuclear atypia, mitotic activity, vascular proliferation and necrosis. Throughout this manuscript, we regard the pathological term “tumor core” as “active tumor (AT)”, “positive tumor cells” as “infiltrating edema (IE)”, and “normal tissue” as “normal tissue (NT)”.

For each specimen, the area with the highest cellularity was selected and one image was captured at x40 magnification. Quantitative assessment of histology images was performed automatically by a cell segmentation method which applies decorrelation and stretching of the colorspace in the preprocessing step for improving the performance of cell segmentation (**Fig. 2**) (27). Cellular count (CC) within the specified area was calculated from the segmented images and used as a representative parameter of cellular density. The relationship between the calculated CC on all regions and interpretation of the pathologist, as the gold standard, was examined.

Statistical Analysis

The statistical analysis comprised of the following steps: (1) evaluating the association of MRI-derived parameters with each other and with statistically significant quantitative histopathological feature(s); (2) assessment of each of the mean values of the extracted features among the three tissue subtypes, i.e. infiltrative edema (IE), normal tissue (NT), and active tumor (AT); (3) investigating the diagnostic performances of individual MRI-derived features for identifying the histopathologically-identified regions and their combinations based on a machine learning technique, comprising automatic feature selection and classification.

The computed MRI-based features ($n=15$) include morphological parameters, i.e. $T1_SUB$, $T2_ISO$, $FLAIR$, diffusion-related parameters, i.e. ADC , MD , FA , P , Q , D , D^* , K , perfusion-related parameters, i.e. CBV , and f , and T2-relaxometry parameters, i.e. T_2 and PD . Histopathological CC metric was also evaluated for its potential in discriminating the sub-regions. All statistical analysis and machine learning procedures were performed in R Statistical Software (R3.0.2, Vienna, Austria).

Comparison of Mean Values—One-way ANOVA method was applied to evaluate the discrimination of normal tissue (NT), infiltrative edema (IE), and active tumor (AT) tissue subtypes based on the MRI-derived parameters and cellular count. P -value <0.05 was considered significant for differentiation of the three tissue regions simultaneously. For each parameter, pairwise differentiation of the regions, namely NT from IE, NT from AT, and IE from AT, was assessed by post-hoc tests, namely Tukey's honest significant difference (Tukey-HSD) or Games-Howell methods, to avoid Type I errors by adjusting the calculated p-values for multiple comparisons. The HSD test is based on assumptions of normality and equality of variance, which the former was evaluated using normality tests and the latter by Bartlett method. If equality of variance assumption passed, Tukey-HSD was performed; otherwise, Games-Howell nonparametric test was applied.

Association of Parameters—Relationships between MRI-derived parameters with each other and also between them and cellular count were obtained using Spearman's rank correlation. To correct for multiple comparisons, Holm's method for multiple testing was applied (28) and after correction, a level of 0.05 was considered statistically significant.

Assessment of Diagnostic Performances of Individual MRI-derived Parameters—Fischer's linear discriminant analysis (LDA) method was used to assess performances of MRI-derived and CC parameters in discrimination of NT from IE, NT from AT, and IE from AT. LDA was implemented with leave-one-out cross validation (LOOCV), where in each iteration (number of iterations equals the number of samples), the dataset is partitioned into the training set composed of whole data excluding one sample, which is separated as the test object. Assessment metrics, namely sensitivity, specificity, accuracy, and area under the receiver operating characteristic (ROC) curve (AUC) were calculated on test samples and averaged over loops of cross-validation.

Combination of MRI-derived Parameters through Classification—To identify the complementary/competitive values of MRI-derived features, feature selection was carried out using Akaike Information Criterion (AIC) (29) and Schwarz Bayesian Information Criterion (BIC) (30) in forward selection, backward elimination and stepwise selection strategies (constructing 6 feature selection methods) on a feature space consisting of the 15 MRI-derived parameters.

The feature selection algorithm was adjusted to avoid choosing similar parameters in a set, e.g. the algorithm should select only one feature among diffusion-related features: *ADC*, *D*, *MD*, *P*; anisotropy-related features: *FA*, *Q*; perfusion-related features: *f*, *CBV*; T2-related features: *T₂*, *T₂_ISO*.

Feature selection was performed in loops of LOOCV and the most frequently selected feature subsets by each of the feature selection methods were the representative feature subset (31). This was followed by a classification step based on Fischer's linear discriminant analysis (LDA), quadratic discriminant

analysis (QDA), and support vector machine (SVM) methods (32) for discrimination of tissue subtypes. To avoid bias, classification was also performed using LOOCV.

RESULTS

Among the 10 recruited patients, 6 were diagnosed with low grade glioma (WHO Grade II: low-grade astrocytoma (n=3), diffuse oligoastrocytoma (n=2), oligodendroglioma (n=1)), 1 with anaplastic oligoastrocytoma (WHO Grade III), and 3 with glioblastoma multiforme (WHO Grade IV). A total of 51 tissue specimens were collected from ten patients. Three patients (2 GBM and 1 Grade II Oligodendroglioma) were excluded due to absence or distorted multi b-value, multi-echo, DTI or DSC-MRI. Finally, seven patients with 34 samples were included in the study. Based on qualitative histopathological assessment, 6 samples were diagnosed as the “tumoral core”, 20 samples as “positive tumor cells”, and 8 samples as “normal tissue”.

Relationship of MRI Parameters

The summary results for correlations among the MRI parameters with each other and with CC are provided in **Table 2**. The statistically significant correlations are stated in boldface type and are shaded in gray.

Diffusion-based parameters—ADC parameter significantly correlated with MD, P, D, T₂_ISO and T₂ and negatively with FA and K. This parameter showed strongest correlations with D (R=0.96), MD (R=0.91), P (R=0.88), and T₂ (R=0.80), showed no significant correlation with CC.

D is directly correlated with ADC, FA, MD, P, T₂_ISO, T₂, and inversely correlated with K. The strongest correlation of D was with ADC (R=0.96), MD (R=0.90), and P (R=0.88). It showed significant correlation with CC (R=0.40).

D^* was correlated poorly and inversely with T_2 . K correlated inversely with ADC , MD , P , and T_2 (in all, $R \leq 0.62$). Besides the correlations mentioned earlier that included MD , this parameter was strongly correlated with P ($R=0.98$) and T_2 ($R=0.84$), and showed moderate correlations with $T2_ISO$. P was highly related to T_2 ($R=0.81$), and moderately correlated with $T2_ISO$. Both MD and P were correlated with CC ($R=0.36$).

FA was strongly correlated with Q ($R=0.86$) and indicated modest inverse association with ADC , D , MD , P , and T_2 . Furthermore, FA showed an inverse correlation with CC ($R=-0.37$). Q was directly correlated with FA , and showed no relationships with CC .

Perfusion-based parameters— CBV only showed statistically significant correlation with f parameter derived from IVIM which was not strong ($R=0.35$). Additionally, it indicated a significant correlation with CC ($R=0.46$). Parameter f correlated with none of the parameters except CBV .

cMRI parameters— $FLAIR$ only correlated with $T2_ISO$ with correlation coefficient of 0.67. $T2_ISO$ was significantly associated with ADC , D , MD , FA , $FLAIR$, and P (for all, $R < 0.70$). $T1_SUB$ showed no significant correlations with any of the parameters.

T_2 -mapping parameters—As mentioned in previous sections, T_2 demonstrated correlations with ADC , D , K , MD , P , and $T2_ISO$. PD -map correlated with none of the parameters.

Association of Tissue Subtypes with Quantitative MRI-derived Parameters

Comparison of the mean values of individual parameters based on ANOVA test for differentiation of the three tissue subtypes from each other is stated in **Table 3**. The results suggest that among MRI-derived parameters, $T2_ISO$, $FLAIR$, ADC , CBV , MD , FA , P , D , and T_2 show statistically significant differences among the three subregions, i.e. NT, IE, and AT. Histopathologically-derived parameter, CC , showed statistically significant differences among the three tissue subregions ($P = 8.9E-06$).

The details of classification performances of individual parameters in differentiation of each of the two subregions from each other and based on cross-validated LDA method are given in **Table 3**. For differentiation of NT from IE (requiring high sensitivity), highest performances were for $T2_ISO$, $FLAIR$, ADC , MD , P , D , and T_2 . T2-based features, i.e. $T2_ISO$, $FLAIR$, and T_2 resulted in AUCs >84% for this discrimination. The four diffusion-related parameters, i.e. ADC , MD , P , and D resulted in AUCs of over 90% (and sensitivities >90%) for discrimination of NT from IE. In this context, $T1_SUB$, CBV , and f were highly sensitive (100%) but lack specificity.

All MRI-derived parameters except for D^* , f , and PD were accurate for separation of NT from AT, implying high sensitivity of these metrics.

The only statistically significant MRI-derived feature for identification of IE from AT was DSC-MRI-derived parameter, i.e. CBV , with high specificity of 95%.

CC showed high AUCs for differentiation of the three two-by-two subregions (>84%), although its specificity for separation of NT from IE was rather low.

Combination of Quantitative MRI-derived Parameters

For classification of the three tissue subtypes from each other, i.e. NT, IE, and AT, four feature combinations were selected by the feature selection methods: (1) CBV , MD , $FLAIR$, $T2_ISO$ (selected by Backward AIC); (2) CBV , MD (selected by Backward BIC); (3) CBV , D , $T2_ISO$ (Forward/Stepwise AIC); and (4) CBV , D (selected by Backward BIC) (**Table 4**). Three classification methods, namely, LDA, QDA, and SVM, were applied on the selected combinations. As **Table 4** summarizes, using the classifiers, the most accurate feature set was “ CBV , MD , $FLAIR$, $T2_ISO$ ” with AUC of ~90% for all three classifiers. Additionally, the feature combination “ CBV , D , $T2_ISO$ ” resulted in AUC of ~92% using LDA classification scheme.

Table 5 should assess the performances of classification schemes (each classifier on the selected feature subsets) for two-by-two discrimination of the three subregions from each other. Combination of “*CBV, MD, FLAIR, T2_ISO*” using either QDA or SVM classifiers produces accurate results for all the three discrimination purposes, i.e. NT from IE, NT from AT, and IE from AT. Using QDA, this combination indicates AUC of 92.3% for characterization of NT from IE, 100% for NT from AT, and 89.2% for IE from AT. SVM classification based on this combination results in AUC of 98.8% for identification of NT from IE, 100% for NT from AT, and 100% for IE from AT.

Additionally, “*CBV, D, T2_ISO*” using QDA classifier can generate AUC of 85.6% for differentiation of NT from IE, 100% for NT from AT, and 91.6% for IE from AT.

DISCUSSION

Conventional MR imaging lacks accuracy in assessing physiological variations and regional heterogeneity within tumors. During the past years, several quantitative methods, such as DWI/DTI and DSC-MRI, have been investigated for their potential as adjuncts to cMRI. Yet, the comparative and complementary roles of these methods and emerging techniques, such as T2-relaxometry and IVIM, are not explored especially through histopathologically-approved specimens and objective classification techniques.

In this preliminary study, we attempted to tackle this problem by evaluation of the aforementioned techniques on localized biopsy specimens, which were histopathologically assessed and attributed to active tumor (AT), infiltrative edema (IE), and normal tissue (NT) subregions. Among individual MRI-derived parameters, diffusion-related *ADC, MD, P, D* parameters, as markers of cellular proliferation, indicated high classification performance for identification of NT from IE, and NT from AT, but were not suitable for IE from AT separation. This finding along with significant correlation of *D, MD, and P* with histopathological *CC* parameter, suggests that changes in cellular distribution captured by diffusion-based parameters can be early markers of glioma infiltration (NT from IE discrimination). But, these

changes in low-grade glioma are not significantly different among IE and AT subregions; therefore, these parameters may not be diagnostically valuable for localizing the most active part of the tumor and other features must be explored. Among *FA* and *Q* as indicators of diffusion anisotropy, *FA* was statistically significant for discrimination of the three subregions and showed significant inverse correlation with *CC*, and accurately discriminated NT from AT.. However, *FA* did not result in statistically significant differences for isolation of NT from IE or IE from AT in our sample population.

In animal studies, DTI has been shown to identify microinfiltration of tumor cells in the surrounding brain tissue (33,34). Early investigations on human brain glioma indicate that DWI/DTI can provide helpful information for assessment of peritumoral edema and to define margins of tumor invasion (3,35,36). Specifically, apparent diffusion coefficient (ADC) (20,37) or mean diffusivity (MD) and fractional anisotropy (FA) (19,38,39) have shown correlations with NET based on histopathologically-approved biopsy specimens.

Unlike other MRI-derived parameters, *CBV* was the only statistically significant feature for discrimination of IE from AT . Hence, early vascularization is a discriminative factor for identifying the most active component of gliomas. This parameter was not statistically significant for identification of NT from IE, indicating that in early stages of infiltration and before transforming into the most active subpopulations, neo-angiogenesis may not have a role in gliomas. *CBV* was evidently statistically significant for characterization of NT from AT. Nonetheless, the classification performance was lower than that of diffusion-based parameters . Suboptimal performance of *CBV* in contrast to diffusion-based parameters in this context can be attributed to our sample population of low-grade gliomas, in which neo-vascularization has not become largely prominent in the infiltrative or active tumor regions.

The best combination of these parameters was sought using cross-validated feature selection and classification methods. Three classifiers with increasing complexity, namely LDA, QDA, and SVM, were implemented to get insights about capability of the selected feature combinations in tissue identification

irrespective of classifier formulation. In all four selected feature sets, *CBV* and a diffusion-based parameter like *D* or *MD* were present, signifying the importance of perfusion and diffusion parameters in subregion characterization. Incorporation of *T2_ISO* and *FLAIR* to perfusion and diffusion parameters in the form of “*CBV, MD, T2_ISO, and FLAIR*” combination resulted in high classification performance for characterization of the three tissue subregions from each other, using the classifiers (AUC~90%). This feature set revealed high classification performance in pairwise subregion classification, as well. *T2_ISO* and *FLAIR* are important factors for illustrating the pathogenic changes within glioma borders. Similar integration of parameters has been intuitively employed in brain tumor segmentation elsewhere (40-42).

A combination of fewer parameters including “*CBV, D, T2_ISO*” based on QDA classifier generated a relatively high classification performance for pairwise differentiation of tissue subregions. In occasions with high risk of patient discomfort and motion, IVIM acquisition combined with DSC-MRI perfusion and anatomical high-resolution T2w, required for image-guided neurosurgery, could be performed without addition of DTI acquisition, which is highly susceptible to motion.

Nonetheless, our study and the results achieved are limited by the included patient population and small sample size. Furthermore, despite using cross-validation for reducing bias in classification in our work, an independent validation dataset is required for generalizing the findings. Finally, in this work, our research imaging protocol had to be adapted to the clinical set-up constraints of our academic educational institution and therefore, some of the imaging parameters may not be optimal; for example, in quantitative imaging sequences, the voxel sizes are non-isotropic. Future works may benefit from optimization of parameter specifications for quantitative imaging sequences.

In conclusion, through investigation of a variety of conventional and quantitative MRI techniques applied on histopathologically-approved tissue specimens, we have demonstrated that incorporation of a few imaging techniques (comprising of conventional MRI (T2w and FLAIR) and DSC-MRI with DTI or IVIM)

can form a multi-parametric surrogate marker, predictive of tissue subregions prior to image-guided biopsy procedures.

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Table 1—MR Imaging Specifications

| Sequence | TE/TR | Slice Thickness (mm) | Flip Angle | FOV (mm ²) | Image Matrix | Other Specifications |
|-------------------------------|---|----------------------|------------|------------------------|--------------|--|
| Pre- and Post-contrast 3D T1w | 5/17 | 1 | 25° | 208×256 | 208×256 | N/A |
| 3D T2w | 200/2500 | 1 | 120° | 208×256 | 420×512 | N/A |
| T2-FLAIR | 115/8400 | 5 | 125° | 181×200 | 232×256 | TI=2240 |
| T2-Relaxometry | (12, 24, 36, 48, 60, 72, 84, 96, 108, 120, 132, 144, 156, 168, 180, 192)/4000 | 5 | 180° | 200×200 | 232×256 | N/A |
| DWI | 100/4000 | 5 | 90° | 200×200 | 136×136 | b-values=50, 1000 <i>s/mm²</i> |
| Multi b-value DWI | 100/4000 | 5 | 90° | 200×200 | 136×136 | b-values=0, 50, 200, 400, 600, 800, 1000 <i>s/mm²</i> |
| DTI | 90/9000 | 5 | 90° | 256×256 | 128×128 | b-values=50, 1000 <i>s/mm²</i> |
| DSC-MRI | 30/1600 | 5 | 70° | 220×220 | 128×128 | No. Dynamic Scans = 64, Temporal Resolution = 1.5 s |

DWI—Diffusion Weighted Imaging; DTI—Diffusion Tensor Imaging; DSC-MRI—Dynamic Susceptibility Contrast Enhanced MRI;

Table 2—Correlations (in terms of *R*) among the MRI-derived parameters with each other and between MRI-derived parameters and cell count (from histopathology) †.

| | <i>ADC</i> | <i>CBV</i> | <i>D</i> | <i>D*</i> | <i>FA</i> | <i>FLAIR</i> | <i>f</i> | <i>K</i> | <i>MD</i> | <i>P</i> | <i>PD</i> | <i>Q</i> | <i>T1_SUB</i> | <i>T2_ISO</i> | <i>T2</i> | <i>Count</i> |
|---------------|------------|------------|-------------|-----------|--------------|--------------|-------------|--------------|--------------|--------------|-----------|-------------|---------------|---------------|--------------|--------------|
| <i>ADC</i> | | -0.078 | 0.96 | -0.40 | -0.60 | 0.45 | -0.11 | -0.71 | 0.91 | 0.88 | 0.28 | -0.28 | 0.13 | 0.66 | 0.80 | 0.30 |
| <i>CBV</i> | | | 0.01 | 0.14 | -0.25 | 0.03 | 0.35 | 0.17 | -0.09 | -0.06 | 0.18 | -0.24 | 0.28 | -0.22 | -0.19 | 0.46 |
| <i>D</i> | | | | -0.31 | -0.64 | 0.46 | -0.09 | -0.68 | 0.90 | 0.88 | 0.26 | -0.31 | 0.15 | 0.64 | 0.78 | 0.40 |
| <i>D*</i> | | | | | 0.30 | -0.25 | 0.15 | 0.43 | -0.38 | -0.40 | -0.25 | 0.30 | -0.09 | -0.30 | -0.49 | -0.02 |
| <i>FA</i> | | | | | | -0.33 | 0.10 | 0.58 | -0.56 | -0.58 | -0.41 | 0.86 | -0.20 | -0.39 | -0.54 | -0.37 |
| <i>FLAIR</i> | | | | | | | 0.14 | -0.43 | 0.41 | 0.44 | 0.21 | -0.14 | 0.37 | 0.69 | 0.34 | 0.05 |
| <i>f</i> | | | | | | | | 0.36 | 0.01 | -0.01 | 0.07 | 0.08 | 0.27 | 0.10 | -0.07 | 0.32 |
| <i>K</i> | | | | | | | | | -0.60 | -0.59 | -0.18 | 0.41 | -0.19 | -0.51 | -0.61 | -0.09 |
| <i>MD</i> | | | | | | | | | | 0.98 | 0.46 | -0.28 | 0.19 | 0.62 | 0.84 | 0.36 |
| <i>P</i> | | | | | | | | | | | 0.47 | -0.31 | 0.26 | 0.59 | 0.81 | 0.36 |
| <i>PD</i> | | | | | | | | | | | | -0.47 | 0.27 | 0.02 | 0.39 | 0.00 |
| <i>Q</i> | | | | | | | | | | | | | -0.23 | -0.11 | -0.34 | -0.17 |
| <i>T1_SUB</i> | | | | | | | | | | | | | | 0.03 | 0.03 | 0.23 |
| <i>T2_ISO</i> | | | | | | | | | | | | | | | 0.69 | 0.15 |
| <i>T2</i> | | | | | | | | | | | | | | | | 0.18 |

† The cells with bold-face type and with gray-shaded color are those with significant correlation as indicated by Spearman's rank correlation.

ADC—Apparent Diffusion Coefficient (DWI); *CBV*—Cerebral Blood Volume (DSC-MRI); *D*—True diffusion coefficient (IVIM); *D**—Pseudo-diffusion coefficient (IVIM); *f*—perfusion fraction (IVIM); *K*—Diffusion Kurtosis; *MD*—Mean Diffusivity (DTI); *FA*—Fractional Anisotropy (FA); *P*—Pure isotropic diffusion coefficient (DTI); *Q*—Pure anisotropic diffusion coefficient (DTI); *PD*—Proton Density (T2-relaxometry); *T2*— T2 relaxation time (T2-relaxometry); *T1_SUB*—Subtracted high-resolution T1w from the corresponding high-resolution contrast-enhanced T1w image; *T2_ISO*—High-resolution T2w image; *FLAIR*—Fluid-attenuated inversion recovery image;

Table 3— Comparison of the Mean Values Based on ANOVA Test (Second Column), for Differentiation of Normal Tissue (NT) vs. Infiltrative Edema (IE) vs. Active Tumor (AT) Simultaneously, and Evaluating the Diagnostic Performances of Individual Parameters in Differentiating Pairwise Regions, i.e. “NT” from “IE”, “NT” from “AT”, and “IE” from “AT” Using Cross-validated LDA Method †.

| Feature | ANOVA Test (P-values) | NT from IE | | | | | NT from AT | | | | | IE from AT | | | | |
|----------------|-----------------------|--------------|-------|-------|------|---------------------------|----------------|-------|-------|------|--------------------------|----------------|-------|-------|------|--------------------------|
| | | P-values | Sens. | Spec. | Acc. | AUC (95% CI) | P-values | Sens. | Spec. | Acc. | AUC (95% CI) | P-values | Sens. | Spec. | Acc. | AUC (95% CI) |
| T1_SUB | 0.203 | 0.563 | 100 | 0 | 71.4 | 68.8 (45.0 – 92.5) | 0.176 | 50 | 100 | 78.6 | 81.2 (54.1 – 100) | 0.462 | 1.9 | 95.1 | 73.7 | 67.5 (36.4 – 98.1) |
| T2_ISO | 0.001 | 0.018 | 87.0 | 75.9 | 83.9 | 89.4 (77.0 – 100) | 0.000 | 77.9 | 87.5 | 83.5 | 95.8 (86.0 – 100) | 0.913 | 0.0 | 100 | 76.9 | 59.1 (35.5 – 82.8) |
| FLAIR | 0.001 | 0.016 | 89.8 | 64.7 | 82.7 | 90.6 (79.0 – 100) | 0.000 | 67.9 | 87.5 | 79.1 | 93.7 (81.2 – 100) | 0.958 | 0.0 | 100 | 76.9 | 49.2 (21.8 – 76.5) |
| ADC | 0.005 | 0.002 | 89.6 | 87.5 | 89.0 | 90.0 (76.9 – 100) | 0.001 | 100 | 87.5 | 92.9 | 100 (100 – 100) | 0.473 | 0.64 | 100 | 77.1 | 65.0 (41.9 – 88.1) |
| CBV | 0.004 | 0.963 | 100 | 0 | 71.4 | 50.7 (24.1 – 77.3) | 0.003 | 66.7 | 87.5 | 78.6 | 75.0 (41.1 – 100) | 0.018 | 51.3 | 95.4 | 85.2 | 80.0 (50.7 – 100) |
| MD | 0.000 | 0.001 | 93.9 | 63.8 | 85.3 | 93.8 (85.0 – 100) | 0.001 | 100 | 100 | 100 | 100 (100 – 100) | 0.601 | 1.3 | 100 | 77.2 | 68.3 (46.3 – 90.3) |
| FA | 0.010 | 0.082 | 94.6 | 12.9 | 71.3 | 70.6 (50.2 – 91.0) | 0.007 | 100 | 85.5 | 91.8 | 100 (100 – 100) | 0.209 | 0.0 | 100 | 76.9 | 86.7 (71.0 – 100) |
| Q | 0.116 | 0.215 | 100 | 0 | 71.4 | 51.9 (27.5 – 76.4) | 0.104 | 77.6 | 80.4 | 79.7 | 83.3 (58.6 – 100) | 0.990 | 5.8 | 98.6 | 77.2 | 82.5 (65.4 – 99.3) |
| P | 0.000 | 0.001 | 89.7 | 69.2 | 84.0 | 91.9 (81.0 – 100) | 0.002 | 100 | 99.1 | 99.5 | 100 (100 – 100) | 0.737 | 0.0 | 100 | 76.9 | 60.0 (35.3 – 84.7) |
| D | 0.000 | 0.002 | 90.0 | 71.4 | 84.9 | 90.6 (78.6 – 100) | 0.000 | 100 | 94.0 | 96.7 | 100 (100 – 100) | 0.251 | 9.9 | 99.0 | 78.5 | 68.3 (46.0 – 90.6) |
| D* | 0.748 | 0.918 | 98.9 | 0.9 | 70.9 | 60.6 (31.8 – 89.5) | 0.734 | 41.2 | 74.1 | 61.5 | 60.4 (24.4 – 95.9) | 0.855 | 0.0 | 100 | 76.9 | 52.2 (23.5 – 80.8) |
| f | 0.505 | 0.941 | 100 | 0 | 71.4 | 52.0 (27.1 – 76.8) | 0.504 | 32.6 | 86.1 | 63.7 | 59.5 (27.8 – 90.7) | 0.574 | 0.6 | 97.6 | 75.2 | 64.2 (39.9 – 88.4) |
| K | 0.1136 | 0.906 | 93.5 | 14.9 | 71.2 | 73.7 (52.9 – 94.6) | 0.165 | 83.3 | 86.5 | 85.2 | 87.5 (65.8 – 100) | 0.145 | 0 | 100 | 76.9 | 55.8 (32.4 – 79.3) |
| T ₂ | 0.018 | 0.024 | 93.3 | 66.8 | 86.0 | 84.4 (65.4 – 100) | 0.041 | 83.3 | 87.5 | 85.7 | 87.5 (67.8 – 100) | 0.890 | 0.0 | 100 | 76.9 | 50.1 (24.4 – 75.8) |
| PD | 0.282 | 0.461 | 100 | 0.51 | 71.6 | 66.9 (43.0 – 90.7) | 0.272 | 42.9 | 86.5 | 68.1 | 72.9 (42.7 – 100) | 0.738 | 2.1 | 100 | 77.4 | 55.1 (23.4 – 86.9) |
| Count | 8.9E-06 | 0.221 | 92.5 | 25.9 | 73.5 | 83.8 (68.5 – 98.9) | 1.0E-05 | 66.7 | 100 | 85.7 | 100 (100 – 100) | 6.1E-05 | 66.7 | 99.8 | 92.1 | 91.7 (79.2 – 100) |

† The values with bold-face type under the *P*-values column are those with significant *P*-values based on ANOVA test for simultaneous differentiation of the three regions from each other, and also pairwise differentiation of the regions based on Tukey-HSD or Games-Howell tests (whichever was appropriate for each of the parameters). The bold-face type values under other columns indicate high accuracy.

Table 4— Evaluating accuracy and diagnostic performances (AUC) of LDA, QDA, and SVM classifiers on the selected features †.

| Feature Selection Method | Selected Features | Accuracy (%) | AUC (95% CI) (%) |
|---------------------------------|-------------------------------|---------------------|--------------------------|
| Cross-Validated LDA | | | |
| Backward AIC | CBV, FLAIR, MD, T2_ISO | 82.8 | 89.3 (75.9 – 100) |
| Backward BIC | CBV, MD | 75.4 | 74.7 (55.0 – 94.4) |
| Forward/Stepwise AIC | CBV, D, T2_ISO | 82.1 | 91.9 (83.1 – 100) |
| Backward BIC | CBV, D | 74.8 | 73.2 (54.4 – 91.6) |
| Cross-Validated QDA | | | |
| Backward AIC | CBV, FLAIR, MD, T2_ISO | 85.1 | 92.4 (84.1 – 100) |
| Backward BIC | CBV, MD | 79.9 | 80.0 (61.8 – 98.0) |
| Forward/Stepwise AIC | CBV, D, T2_ISO | 83.4 | 85.3 (71.2 – 99.1) |
| Backward BIC | CBV, D | 81.2 | 84.8 (69.1 – 99.4) |
| Cross-Validated SVM | | | |
| Backward AIC | CBV, FLAIR, MD, T2_ISO | 91.3 | 91.9 (79.6 – 100) |
| Backward BIC | CBV, MD | 80.2 | 79.8 (61.4 – 98.1) |
| Forward/Stepwise AIC | CBV, D, T2_ISO | 79.3 | 83.0 (65.9 – 99.7) |
| Backward BIC | CBV, D | 79.3 | 81.4 (63.6 – 98.8) |

† The gray-shaded cells with bold-type face numbers or letters indicate highest diagnostic performances.

Table 5— Evaluating classification performances of LDA, QDA, and SVM classifiers on the selected features for differentiation of each of two tissue subregions from each other, i.e. NT from IE, NT from AT, and IE from AT †.

| Classifiers | Feature Combinations | NT from IE | | | | NT from AT | | | | IE from AT | | | |
|-------------|-------------------------------|-------------|-------------|-------------|---------------------------|------------|------------|------------|------------------------|-------------|-------------|-------------|--------------------------|
| | | Sens. | Spec. | Acc. | AUC (95% CI) | Sens. | Spec. | Acc. | AUC (95% CI) | Sens. | Spec. | Acc. | AUC (95% CI) |
| LDA | CBV, FLAIR, MD, T2_ISO | 90.0 | 86.8 | 89.1 | 94.8 (86.4 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 57.9 | 99.8 | 90.1 | 92.0 (80.0 – 100) |
| | CBV, MD | 91.1 | 65.9 | 84.0 | 92.6 (83.1 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 62.3 | 99.6 | 91.1 | 91.5 (79.0 – 100) |
| | CBV, D, T2_ISO | 86.1 | 94.1 | 88.5 | 91.7 (80.5 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 50.6 | 100 | 88.6 | 91.4 (78.6 – 100) |
| | CBV, D | 89.6 | 64.2 | 82.5 | 90.0 (77.7 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 50.6 | 100 | 88.6 | 91.3 (78.3 – 100) |
| QDA | CBV, FLAIR, MD, T2_ISO | 85.2 | 99.5 | 89.3 | 92.3 (83.9 – 99.9) | 100 | 100 | 100 | 100 (100 – 100) | 84.1 | 94.4 | 92.0 | 89.2 (72.5 – 100) |
| | CBV, MD | 86.6 | 73.4 | 82.9 | 80.0 (61.6 – 98.0) | 100 | 100 | 100 | 100 (100 – 100) | 66.7 | 99.6 | 92.0 | 83.1 (83.1 – 100) |
| | CBV, D, T2_ISO | 81.4 | 89.8 | 93.9 | 85.6 (71.7 – 99.0) | 100 | 100 | 100 | 100 (100 – 100) | 83.3 | 99.8 | 96.0 | 91.6 (91.6 – 100) |
| | CBV, D | 85.5 | 83.4 | 85.1 | 84.5 (68.6 – 99.3) | 100 | 100 | 100 | 100 (100 – 100) | 61.1 | 100 | 91.1 | 80.6 (59.1 – 98.9) |
| SVM | CBV, FLAIR, MD, T2_ISO | 94.6 | 89.8 | 93.3 | 98.8 (95.9 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 85.4 | 100 | 96.6 | 100 (100 – 100) |
| | CBV, MD | 93.7 | 65.2 | 85.6 | 94.6 (86.8 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 69.5 | 100 | 92.9 | 93.5 (80.2 – 100) |
| | CBV, D, T2_ISO | 88.3 | 86.9 | 88.0 | 92.9 (82.9 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 50.6 | 100 | 88.6 | 91.1 (77.8 – 100) |
| | CBV, D | 91.1 | 61.4 | 82.7 | 91.4 (79.8 – 100) | 100 | 100 | 100 | 100 (100 – 100) | 50.6 | 100 | 88.6 | 91.6 (78.7 – 100) |

† The gray-shaded cells with bold-type face numbers or letters indicate highest diagnostic performances of classifiers for all three discrimination

FIGURE CAPTIONS

Fig 1. MR images and quantitative maps of a 20-years old female with histopathologically-confirmed grade II Oligodendroglioma (images are coregistered with CE-T1w image)

Fig 2. Automated segmentation of cells in microscopic images (x40 magnification): (A) a sample image from tumor core of the same patient indicated in Fig 1; (B) automated cell segmentation result.