Accurate Patient-Specific Machine Learning Models of Glioblastoma Invasion Using Transfer Learning

L.S. Hu, H. Yoon, J.M. Eschbacher, L.C. Baxter, A.C. Dueck, A. Nespodzany, K.A. Smith, P. Nakaji, Y. Xu, L. Wang, J.P. Karis, A.J. Hawkins-Daarud, K.W. Singleton, P.R. Jackson, B.J. Anderies, B.R. Bendok, R.S. Zimmerman, C. Quarles, A.B. Porter-Umphrey, M.M. Mrugala, A. Sharma, J.M. Hoxworth, M.G. Sattur, N. Sanai, P.E. Koulberis, C. Krishna, J.R. Mitchell, T. Wu, N.L. Tran, K.R. Swanson, and J. Li

ABSTRACT

BACKGROUND AND PURPOSE: MR imaging–based modeling of tumor cell density can substantially improve targeted treatment of glioblastoma. Unfortunately, interpatient variability limits the predictive ability of many modeling approaches. We present a transfer learning method that generates individualized patient models, grounded in the wealth of population data, while also detecting and adjusting for interpatient variabilities based on each patient’s own histologic data.

MATERIALS AND METHODS: We recruited patients with primary glioblastoma undergoing image-guided biopsies and preoperative imaging, including contrast-enhanced MR imaging, dynamic susceptibility contrast MR imaging, and diffusion tensor imaging. We calculated relative cerebral blood volume from DSC-MR imaging and mean diffusivity and fractional anisotropy from DTI. Following image coregistration, we assessed tumor cell density for each biopsy and identified corresponding localized MR imaging measurements. We then explored a range of univariate and multivariate predictive models of tumor cell density based on MR imaging measurements in a generalized one-model-fits-all approach. We then implemented both univariate and multivariate individualized transfer learning predictive models, which harness the available population-level data but allow individual variability in their predictions. Finally, we compared Pearson correlation coefficients and mean absolute error between the individualized transfer learning and generalized one-model-fits-all models.

RESULTS: Tumor cell density significantly correlated with relative CBV \((r = 0.33, P < .001)\), and T1-weighted postcontrast \((r = 0.36, P < .001)\) on univariate analysis after correcting for multiple comparisons. With single-variable modeling (using relative CBV), transfer learning increased predictive performance \((r = 0.53, \text{mean absolute error } 15.19\%\) compared with one-model-fits-all \((r = 0.27, \text{mean absolute error } 17.79\%\)). With multivariate modeling, transfer learning further improved performance \((r = 0.88, \text{mean absolute error } 5.66\%)\) compared with one-model-fits-all \((r = 0.39, \text{mean absolute error } 16.55\%)\).

CONCLUSIONS: Transfer learning significantly improves predictive modeling performance for quantifying tumor cell density in glioblastoma.

ABBREVIATIONS: FA = fractional anisotropy; GBM = glioblastoma; LOOCV = leave-one-out cross-validation; MD = mean diffusivity; OMFA = one-model-fits-all; rCBV = relative CBV; T1 + C = T1-weighted postcontrast; TCD = tumor cell density; TL = transfer learning; EPI + C = post-contrast T2*WI

Surgical debulking and radiation therapy represent first-line treatments for glioblastoma (GBM), which rely heavily on image guidance to delineate tumor from adjacent nonneural brain. Contrast-enhanced MR imaging currently serves as the clinical standard for image guidance, but its diagnostic accuracy remains limited. Specifically, contrast-enhanced MR imaging localizes contrast-enhancing tumor for surgical resection and/or biopsy but poorly identifies nonenhancing invasive tumor in the sur-

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Please address correspondence to Leland S. Hu, MD, Department of Radiology, Mayo Clinic, 5777 E. Mayo Blvd., Phoenix, AZ 85054; e-mail: Hu.Leland@Mayo.edu

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This invasive tumor segment can represent a substantial proportion of overall burden for many GBM tumors and contributes to recurrent disease and poor survival if left unresected.\textsuperscript{3,4} In addition to surgical guidance, contrast-enhanced MR imaging also fails to localize nonenhancing tumor during dosimetric radiation treatment planning. To compensate, most radiation oncologists must apply submaximal doses across the entire nonenhancing T2/FLAIR volume, which risks toxicity to normal brain and undertreatment of nondetected bulk tumor.\textsuperscript{7,8} These issues underscore the need to improve the image-based detection and targeted treatment of the nonenhancing tumor segment in GBM.

Advanced MR imaging techniques can help characterize nonenhancing tumor by measuring an array of biophysical features that complement contrast-enhanced MR imaging. These include tissue cell density (TCD) on diffusion-weighted imaging,\textsuperscript{2,9} white matter infiltration on diffusion tensor imaging,\textsuperscript{2,10,11} and microvessel morphology on dynamic susceptibility contrast perfusion MR imaging.\textsuperscript{12–14} Multiple studies have used image-guided biopsies to compare these advanced MR imaging features with TCD in a spatially accurate manner.\textsuperscript{3,5,9,11,15–22} These studies have revealed promising trends between MR imaging signal and tumor abundance but also surprising discrepancies in correlations among studies. For instance, some groups have reported that higher TCD correlates negatively with lower fractional anisotropy (FA) on DTI, presumably from greater white matter tract disruption.\textsuperscript{2,9} Yet, opposite (positive) correlations between FA and tumor content have also been reported.\textsuperscript{21} Similarly, published studies have differed on whether TCD correlates negatively or positively with mean diffusivity (MD) measures on DWI.\textsuperscript{9,11,16,19} These discrepancies present obvious challenges for developing generalized MR imaging-based models to prospectively quantify TCD and extent of invasion.

We hypothesized that the relationship between MR imaging signal (for any given contrast) and regional TCD demonstrates patient-to-patient variability within a given cohort. This may underlie the aforementioned discrepancies among published reports in the literature. In this study, we set out to quantify interpatient variability and determine whether it can strengthen the predictive accuracy of MR imaging–based models for quantifying TCD and invasion. Specifically, we have developed a transfer learning (TL) approach that quantifies concordance and variability of MR imaging–histologic relationships across patients. TL builds 1 model for each patient to account for potential interpatient variabilities in MR imaging–histologic relationships, while coupling the estimation processes of each patient-specific model to allow knowledge transfer between models. As proof of concept, we trained and cross-validated this TL approach in a cohort of patients with primary GBM using multiparametric MR imaging and spatially matched image-guided biopsies. Our overarching goal was to optimize predictive models that guide targeted treatment for the problematic nonenhancing tumor segment of GBM.

**MATERIALS AND METHODS**

**Acquisition and Processing of Clinical MR Imaging and Histologic Data**

**Patient Recruitment.** We recruited patients with clinically suspected primary GBM undergoing preoperative stereotactic MR imaging for first-line surgical resection before any treatment, as per our institutional review board protocol at Barrow Neurological Institute. All patients provided written and informed consent before enrollment. The patient cohort presented here has also been described in previous studies.\textsuperscript{3,6}

**Preoperative MR Imaging Acquisition Protocol.** We acquired preoperative 3T MR imaging (Sigma HDx; GE Healthcare, Milwaukee, Wisconsin) within 1 day of stereotactic surgery, including T2-weighted, T1-weighted precontrast, and T1-weighted postcontrast (T1 + C) sequences.\textsuperscript{3,6,14} T1 + C images were acquired after completion of dynamic susceptibility contrast perfusion MR imaging following a total Gd-DTPA dosage of 0.15 mmol/kg.\textsuperscript{3,6,14} In brief, for the DSC protocol, we administered an IV preload dose (0.1 mmol/kg of Gd-DTPA) to minimize T1-weighted leakage errors, after which we administered a second IV bolus injection (0.05 mmol/kg of Gd-DTPA) during the 3-minute DSC acquisition (gradient-echo echo-planar imaging: TR/TE/flip angle = 1500/20 ms/60°, matrix = 128 × 128, thickness = 5 mm).\textsuperscript{3,6,14} We derived postcontrast T2WI (EPI + C) from the initial-source DSC volume.\textsuperscript{3,13} DTI acquisition consisted of spin-echo EPI (TR/TE = 10,000/85.2 ms, matrix = 256 × 256, FOV = 30 cm, thickness = 3 mm, 30 directions, array spatial sensitivity encoding technique, b = 0, 1000).\textsuperscript{3,6}

**Surgical Biopsy.** Neurosurgeons used T1 + C and T2-weighted imaging to guide stereotactic biopsies following the smallest possible diameter craniotomies to minimize brain shift, as previously described.\textsuperscript{3,6} On average, we collected 5–6 tissue specimens from each tumor, selecting targets ≥ 1 cm apart from both T1 + C enhancing regions and nonenhancing T2-weighted hyperintense regions (so-called brain-around-tumor) in pseudorandom fashion from different poles of the enhancing lesion periphery while avoiding centrally necrotic regions on the basis of the clinical feasibility as per clinical protocol. Enhancing and nonenhancing regions were distinguished on the basis of visual assessment by the neurosurgeon at the time of surgical biopsy and/or resection, as per clinical protocol. The target volume of each biopsy sample was approximately 125 mg. The neurosurgeons recorded biopsy locations via screen capture to allow subsequent coregistration with multiparametric MR imaging datasets. The neurosurgeon visually validated stereotactic imaging locations with corresponding intracranial anatomic landmarks, such as vascular structures and ventricle margins, before recording specimen locations.

**Histologic Analysis and TCD Measurements from Image-Local- ized Biopsies.** Tissue specimens (volume = 125 mg) were flash frozen in liquid nitrogen in the operating suite and stored in a −80°C freezer until subsequent retrieval for embedding in an optimal cutting temperature compound and sectioning (thickness = 4 mm) in a −20°C cryostat (Microm-HM-550; Richard-Allan Scientific Company, Canton, Michigan) using a Microtome Blade (Thermo Scientific, Waltham, Massachusetts).\textsuperscript{3,6} Hematoxylin-eosin–stained slides were reviewed blinded to diagnosis by our neuropathologist (J.M.E.) to assess tumor content. Taking into account all visible cells (neurons, inflammatory cells, reactive glia, tumor cells, and so forth), we estimated the percentage of tumor nuclei (ie, tumor cells relative to all visible cells) (0%–100%), rounded to the nearest fifth percentile, and recorded it for
each tissue sample as a ±5% range of TCD (eg, 30%–40%, 75%–85%, and so forth). This method served as a compromise between the resolution of TCD measurements and the precision of the neuropathologist’s estimates.

**Image Signal Normalization, DSC and DTI Analysis, Image Coregistration, and ROI analysis.** We normalized the signal for T1 + C, T2-weighted, and EPI+C image datasets using the Simple Insight Segmentation and Registration Toolkit (SimpleITK, Version 1.0.1; http://www.simpleitk.org/)

24 in Python (Version 3.6.2; https://www.python.org/). The CurvatureFlow ITK algorithm was applied to remove image noise, and the N4ITK algorithm to correct for image-intensity nonuniformity bias that could be due to factors such as local magnetic field heterogeneity.

Following these corrections, the CSF of the lateral ventricles was used as a reference tissue to normalize the intensity distributions of each dataset using a previously described linear scaling process. For DTI, we generated mean diffusivity and fractional anisotropy maps. For DSC, we generated relative cerebral blood volume (rCBV) maps using IB Neuro (Imaging Biometrics, Elm Grove, Wisconsin), using leakage correction, and white matter normalization.

We coregistered all images using ITK tools (https://itk.org/) and IB Suite (Imaging Biometrics), using the DTI B0 anatomic volume as the coregistration target. A board-certified neuroradiologist (L.S.H.) placed ROIs (8 × 8 voxels) for all coregistered multiparametric MR imaging datasets at the stereotactic biopsy locations. We selected this ROI size (0.93 × 0.93 cm) to encompass a broader area relative to the expected biopsy dimensions (125 mg). This selection helped to offset any potential minor shifts in biopsy localization (at the time of tissue sampling) that might otherwise result in misregistration. We measured mean signal values within ROIs for all MR imaging contrasts to correlate with spatially matched estimates of TCD from corresponding biopsies.

**Statistical Analysis and Predictive Modeling**

Univariate Statistical Analysis and Predictive Modeling. To broadly survey potential associations between MR imaging signal and TCD, we performed univariate analysis using linear regression and Pearson correlations for each contrast (eg, T1 + C, rCBV, and so forth) against spatially matched biopsies across the entire patient cohort. We used the false discovery rate to adjust for multiple testing.

To quantify intersubject variability, we also plotted MR imaging signal-versus-TCD in each patient separately.

Multivariate Statistical Analysis and Predictive Modeling Using the Conventional One-Model-Fits-All Approach. To evaluate the potential complementary function of multiple combined MR imaging contrasts to characterize TCD, we performed multivariable linear regression to correlate MR imaging contrasts with histologic quantification of TCD across all biopsies and patients with GBM.

This entailed the conventional approach in which a single static multivariable model was applied uniformly to all biopsy samples within the cohort (ie, one-model-fits-all [OMFA]). We evaluated a range of combined MR imaging contrasts to create various linear regression OMFA models. For instance, a model using signal intensities on rCBV, MD, and FA would be defined by the following: $F(x) = ax_1 + bx_2 + cx_3 + d$, where the predicted TCD, $F(x)$, was a linear function of signal intensity on rCBV, MD, and FA maps ($x_1$, $x_2$, and $x_3$, respectively).

**Transfer Learning Predictive Modeling.** Interpatient variability is a known limiting factor for many types of predictive models. For instance, a single model applied uniformly across all patients within a cohort (ie, one-model-fits-all) is unable to adjust for likely interpatient variabilities in MR imaging–histologic relationships. In contrast, individual models developed for each patient using only that patient’s data (ie, one-model-each-patient) are unable to benefit from general population patterns and thus inherently have small sample sizes.

TL represents a compromise between the 2 aforementioned approaches. TL is a subfield of Machine Learning (ML), with various algorithms having been developed to allow knowledge transfer in jointly building different-but-inherently related models.

We have previously published a TL algorithm under the Bayesian framework. With our TL algorithm, 1 model is built for each patient, but a Bayesian framework is used to bias the model parameter-estimation process toward the population pattern in the case in which the individual data are not sufficient to precisely determine the individual’s variation. In essence, the population pattern represents a generalized model, and TL uses each patient’s own MR imaging and biopsy data to modify that generalized model to more appropriately fit that particular individual patient. Thus, TL models can account for potential variabilities in MR imaging–histologic relationships across different patients, while also coupling the estimation processes of each patient-specific model to allow knowledge transfer between models. We detailed TL theory and methodology in the On-line Appendix and in our previous publication.

**Leave One-Out Cross-Validation.** To reduce overfitting, we used leave-one-out cross-validation (LOOCV) for all OMFA and TL model training. Briefly, all samples except 1 (randomly selected in each patient) were used to train the predictive model, while the excluded sample served as the test case to generate predicted TCD. We repeated this iteratively so that all 82 samples served as the test case.

We then plotted model-predicted TCD against actual TCD for all biopsies to determine cross-validated Pearson correlation coefficients and mean absolute error rates (the difference between predicted versus actual TCD) for each model.

**RESULTS**

**Patients and Biopsy Samples**

We collected 82 image-recorded biopsy samples (median = 4/patient, range = 2–14) from 18 patients with primary GBM (9.9 = female/male, median age = 60 years, range = 18–81 years). We collected ≥3 biopsies from 14/18 patients. In total, 33 biopsy samples originated from regions of nonenhancement (ie, brain-around-tumor), with 14 of 18 patients contributing at least 1 biopsy from nonenhancing regions. The remaining 49 biopsy samples originated from enhancing tumor regions.

**OMFA Univariate and Multivariate Correlations between TCD and MR Imaging**

We performed univariate analysis (Table 1) across all 82 samples comparing TCD with the 6 MR imaging features individually. We
found significant correlations for T1 + C (r = 0.36, P < .001),
rCBV (r = 0.33, P < .001), and FA (r = -0.24, P < .003), though
only T1 + C and rCBV remained significant after correcting for
multiple comparisons (P = .01). Our observed T1 + C correla-
tion with TCD supports the long-held assumption that regions of
MR imaging enhancement (and higher T1 + C signal) generally
correspond with higher tumor content compared with peripheral-
ally nonenhancing regions (with lower T1 + C signal).35 This
assumption underlies the clinical rationale for guiding surgical
cytoreduction based on MR imaging enhancement. However, be-
cause nonenhancing regions, by definition, lack MR imaging en-
hancement, then T1 + C would presumably have much lower
correlation with TCD within nonenhancing regions containing
invasive tumor content. In fact, separate subgroup analysis
(On-line Table) showed much lower correlation values between T1
+ C and TCD when restricted to only nonenhancing biopsy sam-
ple. Meanwhile, the same subgroup analysis showed more con-
sistent rCBV correlations across enhancing and nonenhancing
biopsy subgroups, suggesting the potential utility of rCBV as a
biomarker for both tumor segments. We also address any poten-
tial risks of overfitting by performing LOOCV to estimate the
predictive performance of rCBV for quantifying TCD. Using the
generalized one-model-fits-all approach, we generated a single
variable linear regression model, with rCBV as the sole predictor
input, and plotted predicted-versus-actual TCD (n = 82 sam-
ple). This generalized OMFA approach (based on rCBV alone)
demonstrated poor performance, with a low correlation coeffi-
cient (r = 0.27) and high error (mean absolute error = 17.79%)
(Table 2). We then evaluated other MR imaging features (com-
bined with rCBV) using multivariable linear regression analysis.
Regardless of the combination of MR imaging features (with
rCBV), the generalized multivariate OMFA models failed to sig-
nificantly improve performance (Table 2).

Individualized Patient Plots and Intertepatient Variability
To investigate interpatient variability, we separately plotted MR
imaging signal versus TCD for each of the 14 (of 18) patients who
contributed ≥3 separate biopsies (which allowed for LOOCV, as
described in the Materials and Methods section above). The other
4 contributed <3 biopsies, which was insufficient for LOOCV
analysis. Patient-by-patient plots are also shown in On-line Figs
1–6. Of the MR imaging correlations with TCD, rCBV demon-
strated the greatest consistency across patients, with 13/14 pa-
ients (92.5%) having a positive correlation (r > 0.00), though
these varied in strength from patient to patient (range of r = 0.07
to 0.95) (On-line Fig 1). Conversely, FA correlations showed
greater variability (r = -0.75 to +0.78), with only 57.1% (8/14)
of patients having negative correlations with TCD (versus 42.9%
with positive correlations) (On-line Fig 2). MD correlations were
also highly variable (range, r = -0.96 to +0.78), with 50% of
patients split between negative and positive correlations (On-line
Fig 3). These data suggest that while certain MR imaging features
may be highly correlated with tumor content in a particular pa-
tient (or subset of patients), other patients may exhibit weaker or
even opposite correlations that mask the overall effect in group
analysis. We recognize that some of these individual plots may
have small sample sizes that limit the statistical confidence of the
correlation coefficients. Therefore, the individual plot coefficient
values themselves should be viewed with this limitation in mind.
At the same time, the directionality of the individual correlation
plots (ie, positive-versus-negative) appears to show distinct pop-
ulation trends, and we intend to use the results here to illustrate
some of the interpatient variabilities that may exist between TCD
and imaging measurements. These interpatient variabilities mo-
tivate the use of the transfer learning approach detailed in the next
section.

Transfer Learning
On the basis of the results from univariate analysis and individual
scatterplots, we prioritized rCBV for training of TL and con-
strained knowledge transfer from patients with an arbitrary
threshold for correlation between rCBV and TCD (r ≥ 0.10). By
means of LOOCV, the TL model with rCBV as the sole model
predictor improved the correlation between predicted-versus-ac-
tual TCD (r = 0.53, P < .001, n = 82), compared with the gener-
alized OMFA model, which also used rCBV as the sole predictor
(r = 0.27 using LOOCV) (Table 2). We then quantified incremen-
tal gains from adding other MR imaging contrasts to the rCBV-
based TL model. As shown in Table 2, individualized transfer

| Table 1: Univariate correlations for spatially matched MRI features vs TCD (n = 82 biopsies, 18 patients) |
|----------------------------------------------------------|
| MRI Feature | Pearson Correlation Coefficient (r) | P Value | FDR-Corrected P Value |
| T1 + C | +0.36 | <.001 | .01 |
| T2WI | +0.13 | .25 | .38 |
| rCBV | +0.33 | <.001 | .01 |
| EPI + C | -0.02 | .85 | .85 |
| FA | -0.24 | .03 | .06 |
| MD | +0.03 | .78 | .85 |

Note:—FDR indicates false discovery rate; EPI = post-contrast T2*WI.

| Table 2: OMFA vs transfer learning for various MRI model predictorsa |
|----------------------------------------------------------|
| Predictors for Model | OMFA | TL, LOOCV |
| rCBV | 0.27 | 17.79 |
| rCBV, EPI + C | 0.25 | 18.03 |
| rCBV, FA | 0.34 | 17.24 |
| rCBV, MD | 0.28 | 17.74 |
| rCBV, T1 + C | 0.33 | 16.69 |
| rCBV, T2WI | 0.26 | 17.96 |
| rCBV, FA, MD | 0.32 | 17.47 |
| rCBV, T1 + C, T2WI | 0.35 | 16.61 |
| rCBV, T1 + C, FA | 0.39 | 16.55 |
| rCBV, T1 + C, MD | 0.35 | 16.77 |
| rCBV, T2WI, FA | 0.32 | 17.47 |
| rCBV, T2WI, MD | 0.26 | 18.02 |
| rCBV, T1 + C, FA, MD | 0.37 | 16.85 |
| rCBV, T2WI, FA, MD | 0.30 | 17.68 |
| rCBV, T1 + C, T2WI, FA | 0.37 | 16.79 |
| rCBV, T1 + C, T2WI, MD | 0.34 | 16.88 |
| rCBV, T1 + C, T2WI, FA, MD | 0.35 | 17.05 |
| rCBV, T1 + C, T2WI, FA, MD, EPI, C | 0.34 | 17.17 |

Note:—MAE indicates mean absolute error.

a Statistical significance after correcting for multiple comparisons.
Our study showed that rCBV was the only advanced MR imaging feature to significantly correlate with TCD after correcting for multiple comparisons (P = .01) (besides the conventional MR imaging feature T1 + C). This low-moderate positive correlation (r = 0.33, P < .001) remains concordant with other published studies on rCBV, which have all consistently reported positive rCBV correlations in the literature. Our results strongly suggest that TL should prioritize rCBV, given its high interpatient consistency and statistically significant correlations in univariate analysis. This assertion is supported further by the consistency of reported rCBV correlations in the literature. Learning models consistently improved correlation coefficients and mean error rates compared with corresponding generalized OMFA models. Figure 1 shows the scatterplots for actual-versus-model-predicted TCD using the highest performing TL model, which incorporated rCBV, T1 + C, FA, and MD as model predictors. After LOOCV, this model achieved a Pearson correlation of r = 0.88 (P < .001) across all samples (n = 82), which further increased among nonenhancing T2/FLAIR samples alone (r = 0.94, P < .001, n = 33). By comparison, the generalized OMFA approach, using the same model predictors, achieved much lower correlation coefficients for all samples (r = 0.39, n = 82) and nonenhancing samples alone (r = 0.09, n = 33). As shown in Figure 2, the TL model can be used to generate color overlay maps of predicted TCD that correspond with actual TCD from spatially matched biopsies throughout different nonenhancing regions in the same GBM tumor.

**DISCUSSION**

In this study, we correlated a panel of MR imaging features with regional tumor cell density using image-guided biopsies and spatially matched MR imaging measurements. In particular, the advanced MR imaging features (ie, rCBV, MD, FA) have been widely published for their potential to characterize TCD within the nonenhancing tumor segment of GBM. On univariate analysis, we found that rCBV was the only advanced MR imaging feature to significantly correlate with TCD after correcting for multiple comparisons (P = .01) (besides the conventional MR imaging feature T1 + C). This low-moderate positive correlation (r = 0.33, P < .001) remains concordant with other published studies on rCBV, which have all consistently reported positive correlations with TCD. In contrast, our analysis showed no significant correlation for MD. The correlation for FA (r = −0.24) showed a P value of .06 after correcting for multiple comparisons, which trends toward significance. These results are not entirely discordant with the literature because past studies have lacked general consensus on how MD and FA correlate with TCD. For instance, Stadlbauer et al reported negative correlations between MD and TCD, while Beppu et al reported positive correlations. Similarly, Ellingson et al and Chang et al reported negative correlations between MD and TCD, while Stadlbauer et al reported positive correlations.

We hypothesized that the correlations observed in our study (as well as in the literature) may be impacted by interpatient variability in how MR imaging signal relates to TCD. To gain insight to this variability, we analyzed individualized plots of MR imaging signal versus tumor content within our cohort. In keeping with trends in the literature, rCBV correlations were highly consistent with regard to direction, with 92.5% of patients showing positive correlations. We did, however, observe a wide range of correlation strengths (range, r = +0.07 to +0.95), which explains the low-moderate group correlation. In contrast to rCBV, we observed highly variable correlations for both FA (range, r = −0.75 to +0.78) and MD (range, r = −0.96 to +0.78), with conflicting directions across patients. These discrepancies echo the discordance among studies in the literature and underscore the challenge of using a one-model-fits-all approach to generalize heterogeneity across patients.

**FIG 1.** Transfer learning improves predictive performance for quantifying TCD. Scatterplots show the correlations between actual (y-axis) versus-model-predicted (x-axis) tumor cell densities using multivariable modeling (rCBV, T1 + C, MD, and FA) and the generalized OMFA versus the individualized TL approaches. Blue and red dots denote biopsy samples from enhancing (n = 49) and nonenhancing (n = 33) regions, respectively. Regression lines and Pearson correlations (r) are shown for all 82 samples (black lines and text) versus only the 33 nonenhancing samples (red lines and text). Correlations using TL are significantly higher compared with the generalized OMFA approach.
The application of TL to an rCBV-based model predictor significantly improved correlation (r = 0.53) compared with the one-model-fits-all approach (r = 0.27). TL further improved performance when combining rCBV with other MR imaging features in a multivariate fashion (r = 0.88), particularly for the nonenhancing T2/FLAIR samples (r = 0.94). This represents a substantial increase in performance compared with the conventional one-model-fits-all approach in our study (r = 0.30) (Table 2) and those previously published.16,22

Of note, the TL method presented here requires MR imaging and histologic input from at least 2 image-localized biopsies (per patient) to make predictive inferences for the remaining “unknown” (ie, unbiopsied) regions throughout each patient’s tumor. Our analysis shows that these patient-specific inputs dramatically increase predictive performance for quantifying regional TCD on an individualized basis. While retrospective in nature, these results offer proof of concept that TL can help prospectively guide surgical biopsy and/or resection in a patient-specific manner. As part of our intraoperative workflow, we are currently integrating real-time neuropathologists’ estimates of TCD (adding ~2–3 minutes of analysis per sample) from frozen surgical specimens to support prospective TL-based neurosurgical guidance. These real-time estimates allow active updating of the TL model intraoperatively, accomplished in <5 minutes given standard computing hardware, to guide surgical targeting for the remaining unresected tumor regions. Future prospective studies in a larger cohort of patients will also allow us to evaluate how the tissue-sampling methodology (eg, number and distribution of biopsy locations) in individual patients might be optimized to improve predictive accuracy and model uncertainty. Additionally, individualized TL-based models can also guide dosimetric radiation planning in the postoperative setting. By delineating regional populations of residual nonenhancing tumor in the otherwise nonspecific T2/FLAIR segment, individualized TL-based maps will facilitate more nuanced radiation-planning

FIG 2. TL maps and multiple biopsies in a 71-year-old patient with primary GBM. TL-based color map overlay on a T2-weighted image [A] shows predicted regional TCD ranging from 0% to 100% (blue to red) throughout the segmented tumor region (based on the T2-weighted signal abnormality). Histologic analysis of the top biopsy (purple boxes B and C) yielded 90% TCD, corresponding to red regions of high TL-predicted TCD (purple box, A). TL-based color map (D) from a more caudal T2-weighted slice shows blue regions of low TL-predicted TCD (blue/green, purple box), corresponding to 25% TCD on histologic analysis of the bottom biopsy (purple boxes, E and F).
strategies to prescribe higher doses with increasing TCD while sparing the dose to normal nontumoral brain.

We recognize several limitations to this study. First, like all other previously published reports, our study lacks a dedicated validation set. In fact, the conventional approach in the literature has been to report correlation coefficients from training data alone, which are prone to overfitting.3,5,9,11,15–22 By comparison, our study is the first to use more rigorous cross-validation testing (through LOOCV) to offset potential risks of overfitting, whereby each biopsy sample in our cohort is treated as an “unseen” or unknown case to test predictive performance.3 While cross-validation strengthens our confidence in these initial findings, prospective validation in future studies will ultimately be needed. Second, we have used neuropathologists’ estimates of TCD as the benchmark measure of tumor content. We believe that this represents a more clinically relevant metric compared with total cell density, which can comprise both tumor and nontumor components (eg, astrocytes, microglia).9,16 Nonetheless, we are currently pursuing automated quantification of cellular density in our histologic samples.16 This type of standardization will help facilitate TL implementation across different institutions in the future. Finally, we recognize that image distortions and brain shift following craniotomy could lead to misregistration errors. To compensate, neurosurgeons used small craniotomy sizes to minimize brain shift and also visually validated stereotactic image location with intracranial neuroanatomic landmarks to help correct for random brain shifts. Rigid-body coregistration of multiparametric imaging also helped reduce possible geometric distortions.2,3,12,18 Overall, our experience suggests that combined misregistration is approximately 1–2 mm from both brain shift and registration techniques, which is similar to that from previous studies using stereotactic needle biopsy.11 To help minimize the effects of these potential misregistration errors, we selected larger ROI sizes relative to the expected biopsy tissue volumes. Thus, any minor shifts in the recorded biopsy locations (at the time of tissue sampling) would, in all likelihood, remain colocalized to the specified ROI.

CONCLUSIONS

We present an MR imaging–based transfer learning approach that optimizes individualized models of TCD and extent for patients with GBM. These models show particularly high predictive performance for the nonenhancing infiltrative tumor segment that is problematic to the diagnosis and treatment of GBM. Particularly, relevant clinical applications include surgical guidance for the extent of resection and dosimetric radiation targeting of nonenhancing residual tumor during postoperative adjuvant care.

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