Edge Contrast of the FLAIR Hyperintense Region Predicts Survival in Patients with High-Grade Gliomas following Treatment with Bevacizumab


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ABSTRACT

BACKGROUND AND PURPOSE: Treatment with bevacizumab is standard of care for recurrent high-grade gliomas; however, monitoring response to treatment following bevacizumab remains a challenge. The purpose of this study was to determine whether quantifying the sharpness of the fluid-attenuated inversion recovery hyperintense border using a measure derived from texture analysis—edge contrast—improves the evaluation of response to bevacizumab in patients with high-grade gliomas.

MATERIALS AND METHODS: MRIs were evaluated in 33 patients with high-grade gliomas before and after the initiation of bevacizumab. Volumes of interest within the FLAIR hyperintense region were segmented. Edge contrast magnitude for each VOI was extracted using gradients of the 3D FLAIR images. Cox proportional hazards models were generated to determine the relationship between edge contrast and progression-free survival/overall survival using age and the extent of surgical resection as covariates.

RESULTS: After bevacizumab, lower edge contrast of the FLAIR hyperintense region was associated with poorer progression-free survival (P = .009) and overall survival (P = .022) among patients with high-grade gliomas. Kaplan-Meier curves revealed that edge contrast cutoff significantly stratified patients for both progression-free survival (log-rank χ² = 8.3, P = .003) and overall survival (log-rank χ² = 5.5, P = .019).

CONCLUSIONS: Texture analysis using edge contrast of the FLAIR hyperintense region may be an important predictive indicator in patients with high-grade gliomas following treatment with bevacizumab. Specifically, low FLAIR edge contrast may partially reflect areas of early tumor infiltration. This study adds to a growing body of literature proposing that quantifying features may be important for determining outcomes in patients with high-grade gliomas.

ABBREVIATIONS: EC = edge contrast; GBM = glioblastoma; GTR = gross total resection; HGG = high-grade glioma; MSRS = Maximally Selected Rank Statistics; OS = overall survival; PFS = progression-free survival; RANO = Response Assessment in Neuro-Oncology; STR = subtotal resection; VOL = volume

High-grade glioma (HGG) is among the most common primary brain tumors in adults and is associated with a poor prognosis. Moreover, monitoring the efficacy of different therapeutic agents in patients with HGG remains a challenge. The Response Assessment in Neuro-Oncology (RANO) and the Macdonald criteria are the current standards for response assessment to therapy in patients with HGG and have been shown to correlate with overall survival (OS). However, determining tumor response becomes particularly challenging following treatment with antiangiogenic agents such as bevacizumab, due to the tendency of these agents to decrease contrast enhancement and T2 fluid-attenuated inversion recovery hyperintensity in the absence of a true tumor response. Recent data have demonstrated that the RANO criteria may lead to earlier detection of tumor progression following treatment with bevacizumab because they take into account nonenhancing tumor progression. In a recent study, the RANO criteria were shown to be more sensitive to tumor progression than the Macdonald criteria with the detection of at least 35% of patients in an HGG cohort who had nonenhancing tumor progression.2,5-11

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Indicates article with supplemental on-line table.
Indicates article with supplemental on-line photos.
progression in T2 FLAIR regions. Thus, the evaluation of T2 FLAIR changes may be critical to the detection of early tumor progression following treatment with antiangiogenic agents.

Despite a growing appreciation of the importance of abnormal findings on T2 FLAIR for evaluating tumor response and progression, these features are typically evaluated qualitatively. This type of assessment relies on considerable reader expertise because T2 FLAIR changes can also reflect tumor growth or other processes that result in T2 prolongation, including radiation effects, edema, ischemic injury, infection, seizures, and postoperative gliosis. Given that subjectively differentiating such entities may be difficult or vary across experts, a quantitative assessment of T2 FLAIR abnormalities could aid in distinguishing early nonenhancing tumor progression from treatment-related changes.

Recent studies have shown that specific features of the FLAIR signal, including increased nodularity, blurring of tumor borders, and mass effect may improve the detection of tumor progression. These studies and others have demonstrated that specific FLAIR phenotypes alone or in combination with other imaging features (eg, contrast enhancement, perfusion) are predictive of survival following treatment with bevacizumab.

Nowosielski et al in 2014 found that “T2-circumscribed” tumors, characterized by a bulky appearance and sharp FLAIR borders, are associated with poorer survival, whereas Norden et al in 2008 suggested that a more infiltrative T2 FLAIR pattern may represent a more aggressive tumor. These studies suggest that sharpness of T2 FLAIR borders could have prognostic value following treatment with bevacizumab and provide a valuable biomarker of tumor progression. However, there appears to be discrepancy in the literature as to whether a “well-defined” versus “vague, ill-defined” FLAIR border indicates a poorer prognosis. Given the variability in terminology used in these qualitative studies and the ambiguous criteria for defining FLAIR borders, quantitative imaging metrics that are standardized across studies and independent of interrater and intrarater bias are needed to systematically address this question.

In this study, we used texture analysis to examine whether quantifying the sharpness of the FLAIR hyperintense border (ie, edge contrast [EC]) improves the evaluation of response to bevacizumab in patients with HGG. EC is a type of texture analysis that has been shown in previous studies to distinguish well-defined versus vague borders in grade II and III gliomas and has been validated against neuroradiologic reads.

On the basis of our previous work demonstrating that “ill-defined” FLAIR borders are associated with poorer survival in patients with grade II and III gliomas, we hypothesized that patients with low EC (ie, vague borders) of the FLAIR hyperintense region will have poorer progression-free survival (PFS) and OS compared with patients with high EC (ie, sharp borders). We also hypothesized that EC will be a stronger predictor of PFS and OS than the volume of the FLAIR or contrast-enhancement region following treatment with bevacizumab.

MATERIALS AND METHODS

Patients

This retrospective study was approved by the institutional review board. From 2011 to 2015, sixty patients with HGGs were identified at our institution who had both pre- and postbevacizumab MRIs, including T1 pre- and postcontrast and 3D-FLAIR. Patients were excluded if they received bevacizumab for any other reason than recurrent tumor (eg, radiation necrosis), the baseline scan was >60 days before initiating bevacizumab therapy or the follow-up scan was >60 days after initiation of treatment, there were significant artifacts on imaging, or the imaging sequence was not available at 1 of the 2 time points (pre- and postbevacizumab). Thirty-three patients with HGGs met the inclusion criteria to form the final study cohort (On-line Table). Twenty-nine patients had glioblastomas (GBMs), 2 patients had anaplastic astrocytomas, and 2 patients had anaplastic oligodendrogliomas. The final cohort included 23 men and 10 women; the average age was 55 years with a range of 31–74 years. All patients had at least 1 operation before treatment with bevacizumab, with the most recent operation before the initiation of bevacizumab being a subtotal resection (STR, n = 20) or a gross total resection (GTR, n = 13). All patients received chemoradiation with temozolomide (alone or in combination with other agents) as the initial treatment. On recurrence, 15 patients received bevacizumab monotherapy, whereas 18 received bevacizumab in combination with other agents (carboplatin, irinotecan, lomustine [CCNU], temozolomide). No MRIs obtained within the first 60 days following the operation were included, to avoid immediate postsurgical effects such as hemorrhage. The patients included in this study represented a subset (33 of 40) of the patients reported in a previous publication that examined the utility of advanced diffusion imaging (ie, restriction spectrum imaging) for the evaluation of the response to bevacizumab.

MR Imaging and Preprocessing

All the MR imaging scans were acquired on a 3T Signa Excite HDx scanner (GE Healthcare, Milwaukee, Wisconsin) using an 8-channel head coil. The imaging protocol included pre- and postgadolinium 3D volumetric T1-weighted inversion recovery echo-spoiled gradient-echo imaging with TE/TR = 2.8/6.5 ms, TI = 450 ms, flip angle = 8°, FOV = 24 cm, voxel size = 0.93 × 0.93 × 1.2 mm; and a 3D T2-weighted FLAIR sequence with TE/TR = 126/6000 ms, TI = 863, FOV = 24 cm, voxel size = 0.93 × 0.93 × 1.2 mm.

Before analysis, raw data were corrected for bias field and distortion. Then, correction for patient motion was performed using in-house software. The pre- and postcontrast 3D inversion recovery echo-spoiled gradient-echo and FLAIR images were registered to each other using rigid body registration at each of the 2 time points.

Volumes of Interest

Contrast-enhanced volumes (CE_VOL) and FLAIR hyperintense volumes (FLAIR_VOL) were segmented semiautomatically (Amira software package; Visage Imaging, San Diego, California) on the coregistered FLAIR and postcontrast 3D inversion recovery echo-spoiled gradient-echo images acquired both pre- and posttreatment, while regions of necrosis and the resection cavity were excluded. The final FLAIR_VOL also excluded the CE_VOL to obtain an estimate of the nonenhancing lesion. All volumes of interest were drawn by 2 trained image analysts.
(R.D. and N.F.) and approved by a board-certified neuroradiologist with expertise in neuro-oncology.  

**Edge Contrast Calculation**

The following postprocessing and image-enhancement steps were applied to the FLAIR\textsubscript{VOL} to extract the lesion surfaces and calculate the EC (On-line Figs 1 and 2, which present the flowchart for the EC extraction). We applied 3D analysis to the lesions to enhance the local precision and decrease the partial volume effect.\cite{27-30}

Step 1) Morphologic operations of erosion and dilation were applied to the FLAIR\textsubscript{VOL} binary mask using the spherical 3D mask \((r = 3)\) to remove the holes and small islands (On-line Fig 1A, -B).

Step 2) The contour of the FLAIR\textsubscript{VOL} binary mask was extracted in 3D, indexing the surface of the FLAIR\textsubscript{VOL} lesion.

Step 3) The gradients of the FLAIR image were calculated in 3D using the first derivative of the FLAIR image (On-line Fig 1C).

Step 4) The gradients of the FLAIR\textsubscript{VOL} from step 3 were overlaid on the surface of the 3D binary mask from step 2 to create the hyperintense surface with the initial EC (On-line Fig 1D). EC is defined as the gradient magnitude of the lesion edges.

Step 5) The initial EC was modified by removing the pixels with the highest 10% intensity to minimize the effect of edge magnitude arising from the CSF and skull (On-line Fig 1E, -F).

Step 6) Four EC parameters were calculated for each edge magnitude layer; \(EC_{100\%}\) = average magnitude of all points on the edge, \(EC_{75\%}\) = average magnitude of the lowest 75% of points, \(EC_{50\%}\) = average magnitude of the lowest half of points, and \(EC_{25\%}\) = average magnitude of the lowest 25% of points. This approach was selected to determine whether EC of the whole FLAIR border versus a subset of the FLAIR border with lower EC (ie, areas that could represent local tumor infiltration) predicts PFS/OS postbevacizumab. These 4 parameters were calculated on each scan (pre- and postbevacizumab) for each patient. The change in EC was defined as the change in EC parameters between the pre- and postbevacizumab scan.

**Progression-Free Survival and Overall Survival**

PFS and OS were calculated for all patients from the date of initiation of bevacizumab to tumor progression or death, respectively. Tumor progression was verified by direct pathologic confirmation when available or evidence of progression based on MR imaging and neurologic status jointly determined by a board-certified neuroradiologist (N.F.) and board-certified neuro-oncologist (D.P.) using the RANO criteria.

**Statistics**

Paired \(t\) tests were conducted to compare EC, FLAIR\textsubscript{VOL}, and CE\textsubscript{VOL} parameters pre- and postbevacizumab. Multivariate Cox proportional hazards models that included age and the extent of the resection (ie, subtotal resection versus gross total resection) as covariates were used to determine the relationship between each EC measure, FLAIR\textsubscript{VOL} and CE\textsubscript{VOL} postbevacizumab and the pre- and postchange in each of these measures with PFS/OS. Maximally Selected Rank Statistics (MSRS) were used to identify optimal cutoff points for EC that stratified patients according to PFS/OS. Kaplan-Meier survival analyses were conducted on the basis of the subgroups obtained from the MSRS split and compared using a log-rank test. In the case of no progression or death, the event time was censored at the date of last follow-up. In all cases, 2-tailed \(P < .05\) was considered statistically significant.

**RESULTS**

**Pre- to Postbevacizumab Changes in EC, FLAIR\textsubscript{VOL} and CE\textsubscript{VOL}**

At the group level, there were no significant differences between pre- and postbevacizumab EC parameters. However, inspection of individual subject data revealed that 37% of patients had a \(\geq 10\%\) decrease in \(EC_{100\%}\) postbevacizumab, whereas 46% of patients had a \(\geq 10\%\) decrease in \(EC_{75\%}\), \(EC_{50\%}\), and \(EC_{25\%}\). There was a significant decrease in both CE\textsubscript{VOL} ([\(t(32) = 6.34, P < .001\)] and FLAIR\textsubscript{VOL} ([\(t(32) = 4.28, P < .001\)] at the group level (Table 1).

**Table 1: Summary of the imaging parameters pre- and postbevacizumab**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Prebevacizumab Mean (SD) (\times 10^3)</th>
<th>Postbevacizumab Mean (SD) (\times 10^3)</th>
<th>(P) Value ((T) Value)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(EC_{100%})</td>
<td>2.52 (0.57)</td>
<td>2.43 (0.57)</td>
<td>(\leq .256) (1.15)</td>
</tr>
<tr>
<td>(EC_{75%})</td>
<td>1.91 (0.53)</td>
<td>1.77 (0.47)</td>
<td>(\leq .281) (1.08)</td>
</tr>
<tr>
<td>(EC_{50%})</td>
<td>1.51 (0.48)</td>
<td>1.38 (0.40)</td>
<td>(\leq .277) (1.11)</td>
</tr>
<tr>
<td>(EC_{25%})</td>
<td>1.06 (0.38)</td>
<td>0.98 (0.30)</td>
<td>(\leq .384) (1.35)</td>
</tr>
<tr>
<td>FLAIR\textsubscript{VOL}</td>
<td>121.88 (70.68)</td>
<td>77.31 (38.55)</td>
<td>(\leq .002) (4.28)</td>
</tr>
<tr>
<td>CE\textsubscript{VOL}</td>
<td>24.90 (15.26)</td>
<td>9.16 (9.17)</td>
<td>(\leq .001) (6.34)</td>
</tr>
</tbody>
</table>

\(a\) \(P < .05\).

**Table 2: Summary of the multivariate CPH models with age and surgical extent (subtotal, gross total resection) as covariates**

<table>
<thead>
<tr>
<th>Edge</th>
<th>Time</th>
<th>PFS (P) Value</th>
<th>HR (95%) CI</th>
<th>OS (P) Value</th>
<th>HR (95%) CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>(EC_{100%})</td>
<td>Postbevacizumab</td>
<td>(\leq .009)</td>
<td>0.37</td>
<td>0.18-0.78</td>
<td>(\leq .046)</td>
</tr>
<tr>
<td>(EC_{75%})</td>
<td>Postbevacizumab</td>
<td>(\leq .031)</td>
<td>0.33</td>
<td>0.13-0.79</td>
<td>(\leq .041)</td>
</tr>
<tr>
<td>(EC_{50%})</td>
<td>Postbevacizumab</td>
<td>(\leq .015)</td>
<td>0.29</td>
<td>0.10-0.78</td>
<td>(\leq .026)</td>
</tr>
<tr>
<td>(EC_{25%})</td>
<td>Postbevacizumab</td>
<td>(\leq .018)</td>
<td>0.21</td>
<td>0.05-0.77</td>
<td>(\leq .022)</td>
</tr>
<tr>
<td>FLAIR\textsubscript{VOL}</td>
<td>Postbevacizumab</td>
<td>(\leq .087)</td>
<td>1.00</td>
<td>1.00-1.00</td>
<td>(\leq .204)</td>
</tr>
<tr>
<td>CE\textsubscript{VOL}</td>
<td>Postbevacizumab</td>
<td>(\leq .075)</td>
<td>1.00</td>
<td>1.00-1.00</td>
<td>(\leq .258)</td>
</tr>
<tr>
<td>(\Delta EC_{100%})</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .018)</td>
<td>2.81</td>
<td>1.92-6.65</td>
<td>(\leq .033)</td>
</tr>
<tr>
<td>(\Delta EC_{75%})</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .039)</td>
<td>2.75</td>
<td>1.05-7.19</td>
<td>(\leq .050)</td>
</tr>
<tr>
<td>(\Delta EC_{50%})</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .058)</td>
<td>2.91</td>
<td>0.96-8.08</td>
<td>(\leq .054)</td>
</tr>
<tr>
<td>(\Delta EC_{25%})</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .067)</td>
<td>3.74</td>
<td>0.090-15.42</td>
<td>(\leq .056)</td>
</tr>
<tr>
<td>(\Delta FLAIR\textsubscript{VOL} )</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .036)</td>
<td>1.00</td>
<td>1.00-1.00</td>
<td>(\leq .487)</td>
</tr>
<tr>
<td>(\Delta CE\textsubscript{VOL} )</td>
<td>Pre- and postbevacizumab</td>
<td>(\leq .095)</td>
<td>1.00</td>
<td>1.00-1.00</td>
<td>(\leq .926)</td>
</tr>
</tbody>
</table>

**Note:** --CPH indicates Cox proportional hazards; HR, hazard ratio.

\(a\) \(P < .01\).

\(b\) \(P < .05\).
Seventy-six percent of patients showed a ≥50% decrease in CEVOL, whereas an additional 6% of patients showed a decrease of 25%–50%. Thirty percent of the patients showed a corresponding decrease of ≥50% in FLAIRVOL surrounding the contrast-enhancement region, whereas an additional 18% showed a decrease of 25%–50% following bevacizumab. There were no significant associations between change in any of the EC parameters and change in FLAIRVOL or CEVOL (all P values > .05). However, post hoc analysis revealed that 53% of patients with high postbevacizumab EC100% (EC100% > medianEC100% = 2117.62) had large decreases in CEVOL (ΔCEVOL > 71.2%) and 59% had large decreases in FLAIRVOL (ΔFLAIRVOL > 32.3%). In addition, 38% and 50% of patients with low postbevacizumab EC100% had large decreases in CEVOL and FLAIRVOL, respectively.

**Survival Analysis**

Median PFS following initiation of bevacizumab was 3.94 months (with 31 patients meeting the PFS end point), whereas the median OS following initiation of bevacizumab was 6.80 months (with 25 patients meeting the OS end point). Multivariate Cox proportional hazards models indicated that all EC thresholds postbevacizumab and ΔEC100% and ΔEC75% pre- to postbevacizumab were associated with poorer PFS, whereas all EC thresholds were associated with OS (Table 2). Post-EC100% showed the highest association with PFS, whereas ΔEC75% showed the highest association with OS. However, the overlap in confidence intervals among the EC parameters suggests that they did not significantly differ from one another in their associations with PFS or OS. In both cases, lower EC values were associated with poorer survival. Pre- and post-bevacizumab thresholds showed similar associations with PFS/OS in which ΔEC100% showed the highest association with PFS and ΔEC75% showed the highest association with OS. There were no associations between FLAIRVOL and CEVOL with PFS/OS postbevacizumab or pre- and postbevacizumab. Figure 1 shows an example of a patient with low EC and poor PFS/OS, whereas Fig 2 shows an example of a patient with high EC and good PFS/OS.

MSRS was used to identify optimal cutoff points for post-EC values that stratified the patients according to PFS/OS. A post-EC100% value of 2750.9 was determined to best stratify patients for PFS and OS. Kaplan-Meier survival curves revealed that the post-EC100% cutoff significantly separated the groups for PFS [log-rank \( \chi^2 (1) = 8.3, P = .003 \) (Fig 3A)]. Similarly, the post-EC100% cutoff significantly stratified the groups for OS [log-rank \( \chi^2 (1) = 5.5, P = .019 \)]. Both analyses categorized 8 patients with better PFS/OS and 25 patients with poorer PFS/OS (Fig 3B). Given the small number of patients with better OS and PFS, a median split was also used to divide patients into more balanced groups. Similarly, the post-EC100% cutoff significantly stratified groups for PFS and OS.

**FIG 1.** A 58-year-old man’s MR imaging post-subtotal resection with low edge contrast. The patient had a poor survival estimation, with a PFS of 1.8 months and OS of 7.3 months. A, FLAIR prebevacizumab. B, T1 postcontrast prebevacizumab. C, FLAIR postbevacizumab. D, T1 postcontrast postbevacizumab. E, Overlay of the EC contour over the postbevacizumab FLAIR image. F, 3D presentation of the surface of the hyperintense region. Darker areas on the surface indicate lower EC/more indistinct border, whereas the lighter areas toward red show higher EC/more distinct border.
DISCUSSION

In this study, we introduce a new, quantitative imaging technique for characterizing the FLAIR border in patients with HGG and highlight a clinical scenario in which it may have prognostic value.

We demonstrate that patients with vague, ill-defined FLAIR borders (low EC) have poorer PFS and OS compared with patients with sharper FLAIR borders (high EC). We also demonstrate that most patients with sharper EC post-bevacizumab had a greater decrease in the volume of the contrast enhancement and FLAIR regions despite the lack of a direct association among these measures. However, EC was the only imaging measure associated with PFS/OS, and this association was robust to the EC threshold used. These findings are in line with previous research suggesting that neither FLAIR nor contrast-enhancement volumes are significant predictors of PFS/OS following bevacizumab. Rather, quantitative estimates of FLAIR border patterns may have the potential to serve as a reliable biomarker for non-enhancing tumor progression in patients with HGG following treatment with bevacizumab.

In the current study, EC is a measure of the sharpness of the FLAIR hyperintense border and provides a quantitative measure that is not biased by inter- and intrarater variability. Although it is not clear what various EC levels of the FLAIR sharpness represent biologically, tumor infiltration following treatment with bevacizumab is one possible mechanism that reduces EC. Previous studies have shown that antiangiogenic therapy leads to the use of pre-existing cerebral blood vessels by tumor cells—a process known as vascular cooption—and consequently results in increased infiltrative growth. As a result, nonenhancing regions of FLAIR hyperintensity in patients with HGGs undergo a shift to an infiltrative pattern defined, in part, by a more ill-defined or vague FLAIR border. The association between an “invasive” FLAIR border and infiltrative tumor growth has been validated in histologic specimens of patients treated with bevacizumab. However, the sharpness of the FLAIR border may also be partially unmasked by decreases in edema, despite the lack of an association between changes in EC and FLAIR volumes. Thus, it is possible that changes in both edema and tumor infiltration influenced our postbevacizumab EC measures. Nevertheless, we provide initial evidence that this FLAIR border pattern can be quantified on 3D imaging.

FIG 2. A 60-year-old woman’s MR imaging post subtotal resection with high edge contrast. The patient had a high survival estimation with PFS of 11.5 months and OS of 13.6 months. A, FLAIR prebevacizumab. B, T1 postcontrast prebevacizumab. C, FLAIR postbevacizumab. D, T1 postcontrast postbevacizumab. E, Overlay of the EC contour over the postbevacizumab FLAIR image. F, 3D presentation of the surface of the hyperintense region. Darker areas on the surface indicate lower EC and more indistinct border, whereas the lighter areas toward red show higher EC and more distinct border.
and may serve as a valuable biomarker of early tumor progression, which may, in part, reflect infiltration in patients with HGGs.

In this study, we also demonstrate that the prognostic value of EC appears highly robust to the EC threshold used. Although different thresholds for EC showed slightly different associations with PFS and OS, the overlap in confidence intervals indicates that the differences among the thresholds are small and may not be clinically meaningful. EC robustness indicates that a global measure of FLAIR border sharpness may be just as sensitive as one that focuses on the “vaguest” regions of the border following treatment with bevacizumab. This is in line with previous studies that have shown that patterns of nonenhancing tumor progression are diffuse rather than focal following treatment with bevacizumab. EC consistency pattern may not be present in other clinical scenarios or following treatment with other chemotherapeutic agents in which more focal regions of tumor growth have been identified and different EC levels may have different prognostic values.

This study is the first to use a 3D texture analysis technique to quantify the sharpness of the FLAIR hyperintense border and demonstrate an association with PFS and OS in patients with HGG treated with bevacizumab. This method of 3D edge extraction may increase the level of precision and the reliability of FLAIR border pattern assessment relative to visual ratings used in previous studies. Such a measure may be particularly useful in large-scale clinical trials of antiangiogenic treatments in which standardized, high-throughput image analysis methods are critical and expert readings may introduce systematic bias and be too labor-intensive. This measure may eventually serve as an adjunct to standard qualitative MR imaging interpretation, especially in settings in which neuro-oncology volume and expertise in nuanced neuroradiology interpretation may be lower.

There are some limitations to this study. First, our sample size was relatively small, and all patients were recruited from the same institution. Second, we included a heterogeneous group of patients with HGG (ie, grade III and GBM). Given recent evidence that molecular information (eg, IDH status) may be as important or more important for predicting OS than histopathologic diagnosis, we included both groups. Unfortunately, we did not have molecular information on many patients (On-line Table). Although IDH status was available for 17 of our patients, only 3 patients had IDH-mutant status, precluding any meaningful analysis of these data. As a result, we were not able to determine how EC patterns relate to different molecular subtypes or how our imaging variables could be used in combination with molecular status to predict survival in patients with HGGs. These latter investigations are currently underway at our institution. Finally, the current resolution of MR imaging may be too low to adequately capture microscopic tumor infiltration in the brain. Although we propose that tumor infiltration is a likely reason for decreases in EC postbevacizumab, higher resolution MR imaging will help to increase the value of quantitative MR imaging metrics such as ours.

**CONCLUSIONS**

Our study introduces a quantitative measurement of FLAIR border patterns that may serve as a biomarker for detecting early nonenhancing tumor progression following treatment with bev-
acizumab in HGG. However, whether this measure has clinical utility following treatment with bevacizumab or other antiangiogenic agents will need to be established in further investigation with larger patient cohorts that are stratified according to molecular subtype and other important biologic and treatment-related variables.

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