Spatiotemporal heterogeneity of tumor vasculature during tumor growth and antiangiogenic treatment: MRI assessment using permeability and blood volume parameters

Cherry Kim1 | Ji-Yeon Suh2 | Changhoe Heo1 | Chang Kyung Lee2 | Woo Hyun Shim1,2 | Bum Woo Park2 | Gyunggoo Cho3 | Do-Wan Lee2 | Dong-Cheol Woo2 | Sang-Yeob Kim2,4 | Yun Jae Kim2 | Dong-Jun Bae5 | Jeong Kon Kim1,2

Abstract
Tumor heterogeneity is an important concept when assessing intratumoral variety in vascular phenotypes and responses to antiangiogenic treatment. This study explored spatiotemporal heterogeneity of vascular alterations in C6 glioma mice during tumor growth and antiangiogenic treatment on serial MR examinations (days 0, 4, and 7 from initiation of vehicle or multireceptor tyrosine kinase inhibitor administration). Transvascular permeability (TP) was quantified on dynamic-contrast-enhanced MRI (DCE-MRI) using extravascular extracellular agent (Gd-DOTA); blood volume (BV) was estimated using intravascular T2 agent (SPION). With regard to region-dependent variability in vascular phenotypes, the control group demonstrated higher TP in the tumor center than in the periphery, and greater BV in the tumor periphery than in the center. This distribution pattern became more apparent with tumor growth. Antiangiogenic treatment effect was regionally heterogeneous: in the tumor center, treatment significantly suppressed the increase in TP and decrease in BV (ie, typical temporal change in the control group); in the tumor periphery, treatment-induced vascular alterations were insignificant and BV remained high. On histopathological examination, the control group showed greater CD31, VEGFR2, Ki67, and NG2 expression in the tumor periphery than in the center. After treatment, CD31 and Ki67 expression was significantly suppressed only in the tumor center, whereas VEGFR2 and α-caspase 3 expression was decreased and NG2 expression was increased in the entire tumor. These results demonstrate that MRI can reliably depict spatial heterogeneity in tumor vascular phenotypes and antiangiogenic treatment effects. Preserved angiogenic activity (high BV on MRI and high CD31) and proliferation (high Ki67) in the tumor periphery after treatment may provide insights into the mechanism of tumor resistance to antiangiogenic treatment.

Keywords
magnetic resonance imaging, spatial heterogeneity, tumor vessels
1 | INTRODUCTION

Since neoangiogenesis was identified as a fundamental factor in tumor growth, many antiangiogenic agents have been developed.\(^1^,\)\(^5^\) Despite an early expectation that these therapeutic agents would successfully increase survival time in patients with solid tumors, validation of meaningful survival benefits has failed in a considerable number of clinical trials. To explain such unsatisfactory treatment results, a variety of molecular and histologic theories have been suggested. Among these, regional and biophysical heterogeneity of tumor vessels has been recognized as an important mechanism of tumor resistance to treatment. Non-invasive spatiotemporal analysis of tumor vascular phenotypes is, therefore, required to trace angiogenic alteration during tumor growth or treatment. For this assessment, magnetic resonance imaging (MRI) is useful for providing quantitative information on the spatiotemporal heterogeneity of tumor vasculature over a series of repeated examinations.

Transvascular permeability (TP) and blood volume (BV) are relevant vascular parameters used to characterize tumor vessel phenotype. TP is a concept incorporating plasma-to-tissue flux rate, interstitial volume, and perfusion. This parameter can be quantified from a leakage profile of an extracellular fluid contrast agent (CA) (eg, small-molecular gadolinium [Gd] chelates) on dynamic contrast-enhanced (DCE) MRI. BV is an intuitive indicator to quantify new vessel formation and entire tumor area. Finally, the intratumoral heterogeneity in antiangiogenic treatment effect was interpreted based on time- and region-oriented changes in TP and BV.

2 | MATERIALS AND METHODS

This study was approved by the Institutional Animal Care and Use Committee of the Asan Institute for Life Sciences (South Korea).

2.1 | Brain tumor model and study protocol

In 20 male mice (Balb/c, 8-week old, weight 24-26 g), C6 glioma cells (1.2 × 10^5 cells, Korean Cell Line Bank, Seoul, Korea) were injected into the right cerebral hemisphere using a stereotaxic injection kit (Hamilton Co, NV, USA). The holes were generated 1-mm inferior and 2.5-mm lateral to the bregma, leaving the dura intact, and the cell-containing needle tip was advanced 2.2 mm below the dura. Thereafter, cells were injected slowly over a period of 4 minutes.

After tumor cell injection, T2-weighted images (T2WIs) were obtained every 3 days to monitor tumor volume change. Day 0 was assigned when the tumor volume reached approximately 2 mm^3, and MRI examinations were performed on days 0, 4 and 7. The median time interval from cell injection to day 0 was 12 days. On day 0, the animals were randomly divided into control (n = 10) and treatment (n = 10) groups. Mice were treated daily with vehicle (Cremophor EL/95% ethanol [50:50], Sigma-Aldrich, St Louis, MO, USA) in the control group and sorafenib (orally, 30 mg/kg/d) in the treatment group. Sorafenib is a multireceptor tyrosine kinase inhibitor (TKI), and it has been applied in clinical trials for treating glioblastoma.\(^8^\)

2.2 | MRI examination

For MRI examinations, the mice were positioned on a custom-made mouse cradle and anesthetized using isoflurane (1.0%-1.5%) in 70% N2O and 30% O2. During MRI examination, respiration rate was monitored, and body temperature was maintained to be at 37°C using warm airflow.

All experiments were performed using a 9.4-T magnet (Agilent 160AS horizontal imaging system; Agilent Technologies, Santa Clara, CA, USA). Prior to intravenous CA injection, T2WIs were obtained using the fast-spin echo sequence (TR/TE, 4000/51 milliseconds; flip angle, 90°; echo train length, 8; slice thickness, 1 mm; number of slice, 12 or 13; field of view [FOV], 20 × 20 mm; matrix, 256 × 256). DCE-MRI was then performed before and after bolus injection of Gd-DOTA through the tail vein (Dotarem\(^\circledast\), Guerbet, France [molecular weight, 500 D]; injection dose, 0.1 mL/kg of body weight; injection approximately 30 seconds after scanning start) using the following parameters: TR, 33 milliseconds; TE, 2.8 milliseconds; flip angle, 30°; number of repeated acquisitions, 100; time for each dynamic set, 3.15 seconds; slice thickness = 1; number of slice, 12 or 13; FOV, 20 × 20 mm; matrix, 96 × 96.

Because Gd-DOTA has a short plasma half-life (approximately ~30 minutes) and a significantly low R2 relaxivity (4.17 mmol/L/s) compared with our SPION (40 mmol/L/s),\(^9^\) residual gadolinium after DCE-MRI examination would not affect SPION-induced ΔR2* and ΔR2 measurements.\(^10^\) Thus, SPION-induced ΔR2* and ΔR2 were measured after DCE MRI examinations. For measuring BV parameters (ie, ΔR2* and ΔR2), SPION (30 mg Fe/kg bodyweight) was used as an intravascular susceptibility CA. The pharmacokinetic profile of SPION was detailed in a recent article.\(^11^\) Before and after SPION injection, the R2 value was measured using...
the multiecho spin-echo sequence (TR, 3000 milliseconds; 15 TEs [10-150 milliseconds; ΔTE, 10 milliseconds]; flip angle, 90°; slice thickness, 1 mm; number of slice, 12 or 13; FOV, 20 × 20 mm; matrix, 128 × 128), and the $R_2^*$ was measured using the multiecho gradient-echo sequence (TR, 1500 milliseconds; 12 TEs [5-60 milliseconds; ΔTE, 5 milliseconds]; flip angle, 30°; slice thickness, 1 mm; number of slice, 12 or 13; FOV, 20 × 20 mm; matrix, 128 × 128), respectively.

2.3 | Tumor center vs periphery

An experienced radiologist drew three regions-of-interest (ROIs) on T2WI including the entire tumor, center, and periphery; areas with T2 hyperintensity and Gd enhancement were considered tumor tissue. By referring to previous studies that described the active proliferation and angiogenesis in tumor border, the outer one-third of the tumor area was assigned as periphery and the other area as the center. After drawing the ROI covering the entire tumor areas, the radiologist drew another ROI that passed through two-thirds of the tumor radius. The inner area of this ROI was considered as the tumor center, and the outer area as the tumor periphery.

2.4 | Vascular parameters

MRI vascular parameters were measured using Analysis of Functional NeuroImages (AFNI, http://afni.nimh.nih.gov/afni) software. DCE-MRIs were interpolated to have the same number of matrix and slice number as T2WIs. As mentioned above, the FOV, matrix size, slice thickness, and number of slice were the same among T2WIs, multiecho spin-echo, and multiecho gradient-echo images. Consequently, all MR images had the same geometry. To extract various MR signals from the same anatomic location, MR images were then co-registered to T2WIs.

Transvascular permeability parameters were quantified using non-PK-model-based analysis of DCE-MRI. To minimize machine- or sequence-induced variations, the time-intensity curve was normalized by dividing the signal intensity at each time point with the mean pre-contrast signal intensity. Thereafter, the maximum enhancement rate (MaxEnh) (maximum signal intensity/mean pre-contrast signal intensity) and initial area under the curve (IAUC) over 60 seconds were measured.

For measuring BV parameters, pre- and post-SPION $R_2^*$ and $R_2$ values were calculated according the following equation:

$$SI = M_0 \cdot e^{-R_2^* \cdot t}$$

where SI is the MR signal intensity, $M_0$ is the proton density, $t$ is the echo time, and $R_2^*$ is the transverse relaxation rate (ie, $R_2$ or $R_2^*$). The relative BV weighted for macrovascular and microvascular blood vessels was determined as SPION-induced $ΔR_2^*$ and $ΔR_2$, respectively.

To evaluate the regional variation of tumor vascular phenotypes, the center-to-periphery ratio of each vascular parameter was calculated (eg, MaxEnh center/periphery = MaxEnh center/MaxEnh periphery). The time-dependent changes in tumor volume and vascular parameters were estimated with the ratio between examination days (eg, MaxEnh_day4/day0 = MaxEnh_day4/MaxEnh_day0).

2.5 | Histopathologic examination

Immediately after the final MRI examination, the brains of the mice were removed and embedded in paraffin blocks. Cell density was measured using hematoxylin-eosin staining. Angiogenic activity was estimated using immunohistochemical staining for CD31 and vascular endothelial growth factor receptor 2 (VEGFR2), as described in a literature. The degree of cell proliferation and apoptosis was measured using Ki67 and α-caspase 3 staining methods, respectively. In addition, the amount of pericyte was assessed by NG2 staining. In the mid-coronal section of formalin-fixed glioma, brain sections were stained with CD31 (Abcam, 1:50), VEGFR2 (Cell Signaling, 1:50), Ki67 (Thermo Fisher, 1:100), and α-caspase 3 (Cell Signalling, 1:4000) using an automated slide preparation system Benchmark XT (Ventana Medical Systems Inc., Tucson, AZ, USA). The positive signals were amplified using ultraView Universal DAB detection kit (Ventana Medical Systems Inc.), whereas the sections were counterstained with hematoxylin reagent. NG2 staining (Abcam, 1:500) was performed using immunofluorescence multiplex system accomplished with PerkinElmer Opal kit (Perkin-Elmer, Waltham, MA). For NG2 staining, positive signals were visualized using Opal 520 TSA plus (1:3000) and scanned using the Vectra 3.0 Automated Quantitative Pathology Imaging System (Perkin-Elmer, Waltham, MA).

Similar to the MRI analysis, each tumor section was divided into the tumor center and periphery. Thereafter, the number of positive signals in the tumor ROI was analyzed on digitized TIF images using ImageJ software (National Institutes of Health, Bethesda, MD, USA). The cells on hematoxylin-eosin staining and positive signals from CD31, and VEGFR2 staining were automatically segmented according to thresholds of red-green-blue color values. For NG2 and α-caspase 3 staining, the positive signal was segmented by thresholds of hue-saturation-brightness values. Then, the fraction of positive signal area (ie, %area) was measured using the particle analyzer function in ImageJ.
2.6 Statistical analysis

All statistical analysis was performed using commercially available software (PRISM version 5.01, GraphPad Inc, La Jolla, CA USA). In all parameters, both inter- and intra-group data were compared using non-parametric methods, including the Wilcoxon rank sum test for two paired observations, the Mann-Whitney test for two unpaired observations, and the Friedman test for three paired observations. When there was a significant difference in the Friedman test, a post-hoc analysis was performed using the Dunn test. Differences with $P < .05$ were considered to be statistically significant.

3 RESULTS

Study results are summarized in Figure 1, and representative control and treatment mice are presented in Figures 2 and 3.

3.1 Tumor volume

Time-dependent changes in tumor volume are summarized in Figure 4. Tumor volume on day 0 was similar between the control and treatment groups ($P > .05$). The control group exhibited statistically significant tumor growth between day 0 and day 7 ($\text{volume}_{\text{day 7/day 0}} = 3.9 \pm 1.8$, $P = .01$). The
### A

| Day 0 | Day 4 | Day 7 |
|-------|-------|-------|
| ![T2WI](image1) | ![MaxEnh](image2) | ![IAUC](image3) |
| ![AR2](image4) | ![AR2](image5) | ![AR2](image6) |

| T2WI | MaxEnh | IAUC | AR2* | AR2 |
|------|--------|------|------|------|

### B

**Tumor centr**

![Graph](image7)

**Tumor Periphery**

![Graph](image8)

### C

- **H&E**
- **CD31**
- **VEGFR**
- **Ki67**
- **Caspase**
- **NG2**

*1000 µm*
treatment group showed no significant volume change during the follow-up period \( (P > .05) \), thereby validating successful suppression of tumor growth by antiangiogenic treatment. Accordingly, the control group showed significantly greater tumor volume than the treatment group on day 7 \( (P < .01) \).

3.2 | Spatial heterogeneity of tumor vasculature on initial MRI

In all mice, the initial MRI on day 0 demonstrated significant spatial heterogeneity in the distribution of TP and BV while all vascular parameters were similar between the control and treatment groups \( (P > .05 \) in all intergroup comparisons). On this MRI, TP parameters (MaxEnh and IAUC) were significantly greater in the tumor center than in the periphery (center-to-periphery ratio for TP parameters, 1.70-1.83; \( P < .05 \) for both parameters) (Figure 5A,B). In contrast, BV parameters (\( \Delta R_2^* \) and \( R_2 \)) were apparently greater in the tumor periphery than in center (center-to-periphery ratio for BV parameters, 0.71-1.86; \( P < .05 \) for both parameters) (Figure 5C,D).

3.3 | Heterogeneous vascular changes during tumor growth

As shown in Figure 5, in the control group, spatial distribution pattern of TP and BV (ie, prominent MaxEnh and IAUC in the tumor center, and high \( \Delta R_2^* \) and \( R_2 \) in tumor periphery) became more apparent as the tumor grew. This tendency can be clearly described by increasing center/periphery ratio of TP parameters (MaxEnh\_center/periphery: 1.71 on day 0 to 4.09 on day 7; IAUC\_center/periphery: 1.81 on day 0 to 5.59 on day 7) and decreasing center/periphery ratio of BV parameters (\( \Delta R_2^* \_center/periphery: 0.71 \) on day 0 to 0.34 on day 7; \( R_2 \_center/periphery: 0.78 \) on day 0 to 0.50 on day 7).

MaxEnh was greater on days 4 and 7 than on day 0 in the tumor center (MaxEnh\_day4/0 = 1.28 \pm 0.17; MaxEnh\_day7/0 = 1.90 \pm 0.64; \( P < .05 \) for both comparisons), but were similar in the periphery (\( P > .05 \) for all intra-group comparisons) (Figure 5A). IAUC in the tumor center was significantly greater on day 7 than on day 0 (IAUC\_day7/0 = 2.40 \pm 1.11; \( P = .04 \)), whereas IAUC in the tumor periphery exhibited no significant change during the follow-up period \( (P > .05) \) (Figure 5B). In accordance with such different temporal changes between tumor regions, the MaxEnh\_center/periphery was greater on day 7 than on day 0 \( (P = .03) \), and IAUC\_center/periphery gradually increased in every time window \( (P < .05 \) for all intra-group comparisons) (Figure 5A,B).

Blood volume parameters in the control mice demonstrated a tendency toward decrease in the tumor center according to tumor growth, whereas tumor periphery maintained high BV on three MRIs (Figures 2 and 5C,D). \( \Delta R_2^* \) in tumor center significantly decreased from day 0 to days 4 and 7 \( (\Delta R_2^* \_day4/0 = 0.68 \pm 0.20; \Delta R_2^* \_day7/0 = 0.50 \pm 0.21; \ P < .05 \) for both comparisons). \( R_2 \) in the tumor center also reduced from day 0 to day 7 \( (\Delta R_2 \_day7/0 = 0.59 \pm 0.23; \ P = .02) \). However, both \( \Delta R_2^* \) and \( R_2 \) in tumor periphery exhibited no significant change during the follow-up period \( (P > .05) \). Consequently, the control mice demonstrated a strong tendency toward decreasing \( \Delta R_2^* \_center/periphery \) in every time window \( (P < .05 \) for all intra-group comparisons). \( \Delta R_2 \_center/periphery \) was significantly decreased from day 0 to day 7 \( (P = .04) \).

3.4 | Spatial heterogeneity in antiangiogenic treatment effect

The effect of TKI treatment on MRI was spatially heterogeneous within each tumor as significant treatment-induced angiogenic alterations were identified predominantly in the tumor center (Figure 3). In the tumor center, TKI treatment significantly suppressed increases in TP parameters, and a decrease in BV parameters (ie, typical change pattern in control mice), thereby revealing similar values during the follow-up period (Figure 4). Accordingly, on days 4 and 7, the treatment group exhibited significantly lower MaxEnh (0.22 vs 0.35 on day 4; 0.24 vs 0.49 on day 7), lower IAUC (128 vs 230 on day 4; 135 vs 398 on day 7), and higher \( \Delta R_2^* \) (154 vs 84 on day 4; 151 vs 62 on day 7) in the tumor center than the control group \( (P < .01 \) for all intergroup comparisons). \( \Delta R_2 \) also exhibited no temporal change in tumor center in the
In contrast to the tumor center, the tumor periphery demonstrated no obvious treatment effect on TP and BV; no statistical significance was noted via inter- and intra-group comparisons (P > .05). In particular, high ∆R²* and ∆R² values were maintained in the tumor periphery during the follow-up period (Figure 3).

Due to different temporal changes of vascular parameters in the tumor center between the control and treatment groups, TP_{center/periphery} and BV_{center/periphery} were also clearly different between the two groups (Figure 5). The treatment group demonstrated significantly lower MaxEnh_{center/periphery} and IAUC_{center/periphery} compared with the control group, on days 4 and 7 (P < .01). ∆R²*_{center/periphery} was higher in the treatment group than in the control group on both days 4 and 7 (P < .01). ∆R² center/periphery was also higher in the treatment group on day 7 (P = .02).

3.5 Analysis of the entire tumor area

In contrast to region-oriented analysis, inter- and intra-group comparisons of the entire tumor area revealed no statistically significant differences (P > .05 in all intra- and intergroup comparisons).

3.6 Histopathological results

Representative histopathological findings are presented in Figures 2 and 3; the region-oriented comparison between the treatment group (P > .05) although such effect did not yield an intergroup difference.

In contrast to the tumor center, the tumor periphery demonstrated no obvious treatment effect on TP and BV; no statistical significance was noted via inter- and intra-group comparisons (P > .05). In particular, high ∆R²* and ∆R² values were maintained in the tumor periphery during the follow-up period (Figure 3).

Due to different temporal changes of vascular parameters in the tumor center between the control and treatment groups, TP_{center/periphery} and BV_{center/periphery} were also clearly different between the two groups (Figure 5). The treatment group demonstrated significantly lower MaxEnh_{center/periphery} and IAUC_{center/periphery} compared with the control group, on days 4 and 7 (P < .01). ∆R²*_{center/periphery} was higher in the treatment group than in the control group on both days 4 and 7 (P < .01). ∆R² center/periphery was also higher in the treatment group on day 7 (P = .02).

3.5 Analysis of the entire tumor area

In contrast to region-oriented analysis, inter- and intra-group comparisons of the entire tumor area revealed no statistically significant differences (P > .05 in all intra- and intergroup comparisons).

3.6 Histopathological results

Representative histopathological findings are presented in Figures 2 and 3; the region-oriented comparison between the control and treatment groups is summarized in Figure 6. The control group demonstrated higher expression of CD31 (3.4 vs 1.8), VEGFR2 (2.5 vs 1.2), Ki67 (13.4 vs 10.5), and NG2 (49 vs 41) in the tumor periphery than in the center, whereas cell density (ie, % area of cells) was not spatially different on hematoxylin and eosin staining. TKI treatment yielded significant suppression of CD31 and Ki67, predominantly in the tumor center, as their % area was lower in the treatment group than in the control group (P < .01). However, in the tumor periphery, the TKI treatment effect on these histopathological parameters was insignificant as they were similar between the two groups (P > .05). VEGFR2 and α-caspase 3 expression demonstrated apparent treatment-induced decrease in both the tumor center and periphery (P < .05), and NG2 expression exhibited significant treatment-driven increase in both the tumor center and periphery (P < .01).

4 DISCUSSION

Heterogeneity of the tumor microenvironment has emerged as a critical concept for understanding the mechanisms of disease progression and evasion from treatment. Accordingly, the intratumoral diversity of vascular phenotypes is an important characteristic now recognized to affect antiangiogenic treatment, and has been intensively investigated in preclinical and clinical studies. As an extension of these efforts, this study assessed region- and time-dependent changes in TP and BV in MRI with dual injection of intravascular and extracellular fluid CAs.

The most important finding of this study was the spatiotemporal heterogeneity in both vascular phenotypes and the treatment responses of glioma. In the control group, TP-dependent Gd-DOTA accumulation (ie, MaxEnh and IAUC) was greater in the tumor center than in the periphery, and BV-dependent contrast effect of SPION (ie, ∆R²* and ∆R²) was dominant in the tumor periphery. This region-dependent distribution of vascular parameters became more apparent with tumor growth, as clearly reflected by increasing TP_{center/periphery} and decreasing BV_{center/periphery} during the follow-up period. These MRI results can be biologically interpreted as upregulated, less-leaky vessel formation in the tumor periphery, and elevated plasma-to-interstitium flux in the tumor center.

The high BV and low TP in the tumor periphery during tumor growth may be explained by neoangiogenesis and vascular stabilization. It has been established that the tumor periphery exhibits highly active VEGF-associated angiogenesis, which increases the number and size of vessels by accelerating endothelial cell differentiation/proliferation, primitive vessel generation, vasodilation, and vessel lengthening. In our histopathological examinations, these processes manifested as high VEGFR2 and CD31 expression...
in the tumor periphery. In addition, the tumor periphery exhibits enhanced vascular stabilization under mediation of various growth factors such as PDGF, angiopoietins-1, and tumor growth factor-β. Newly generated primitive vessels are thus matured and covered by supporting matrix and cells, thereby suppressing uncontrolled vascular leakiness. Such vascular stabilization was also presented as increased NG2 activity (ie, pericyte amount) in the tumor periphery in our

**FIGURE 5** Spatiotemporal analysis of tumor vascular phenotypes. Statistically significant differences are indicated by solid lines (intra-group comparisons) and dashed lines (intergroup comparisons). A and B, Temporal changes and center-to-periphery ratios of maximum enhancement rate (MaxEnh) (A) and initial area under the curve (IAUC) (B). The initial magnetic resonance image day 0 revealed higher MaxEnh and IAUC in the tumor center than in the periphery. With tumor growth in the control group, both parameters increased in the tumor center and similarly maintained in the tumor periphery, thereby demonstrating tendencies toward increasing center-to-periphery ratios. Antiangiogenic treatment suppressed such temporal changes only in the tumor center but not in the periphery. Consequently, the treatment group exhibits lower MaxEnh and IAUC in the tumor center than in the control group. C and D, Temporal changes and center-to-periphery ratios of ΔR₂* (C) and ΔR₂ (D). On day 0, both groups demonstrated higher ΔR₂* and ΔR₂ in the tumor periphery than in the center. With tumor growth in the control group, both parameters demonstrated a tendency toward decrease in the tumor center from day 0 to day 7, whereas being similar in the tumor periphery. These changes demonstrated a strong tendency toward decreasing ΔR₂*/center:periphery with tumor growth. Antiangiogenic treatment suppressed such temporal changes in control mice. Consequently, the treatment group demonstrated higher center-to-periphery ratios of ΔR₂*(day 4 and 7) and ΔR₂ (day 7) than in the control group.
KIM et al. study. Therefore, our results of high BV and low TP in the tumor periphery are evidential MRI reflections of new vessel generation and maturation that constitute the invasive margin of the tumor.

Compared with the tumor periphery, the tumor center demonstrated higher TP and lower BV. These findings may be associated with vascular degeneration and destabilization. These vascular alterations—known as “vascular regression”—are characterized by progressive disengagement of endothelial cells from surrounding supportive structures in the absence of angiogenic stimulation.29-31 This phenomenon is primarily led by Ang-2, which induces endothelial cell apoptosis and detachment of endothelial cells from surrounding pericytes.24,25,30,31 Our histologic study also showed relatively low expression of CD31, VEGFR2, and NG2 in the tumor center in comparison with tumor periphery. In this sense, decreased BV and elevated TP in the tumor center are relevant MRI presentations of vascular regression.

As expected from the vascular heterogeneity in the control group, the effect of antiangiogenic treatment also led to apparent regional variations within tumors. The effect of TKI treatment on MRI was predominantly in the tumor center, whereas the tumor periphery maintained its initial features (ie, high BV and low TP) during treatment. Such regional differences in the effect of antiangiogenic treatment have been introduced in several studies.32-34 One explanation for these treatment effects in the tumor center may be “vascular normalization,” which suppresses “vascular regression” by restoring the balance between pro- and antiangiogenic factors, thereby converting abnormal tumor vessels to a more normal state.32,35 More specifically, vascular normalization processes “prune” immature vessels and improve the solidity of the remaining vasculature by enhancing the coverage of perivascular cell and basement membrane.35 This effect was demonstrated as increased NG2 expression (ie, increased pericyte coverage) and decreased α-caspase 3 expression...
(ie, suppressed apoptosis) in our histologic examination. As a supportive evidence, Dominietto et al.\textsuperscript{37} recently demonstrated that anti-VEGF treatment led to a reduced number of small vessels and apparently less chaotic organization of the vascular network. In addition to a vascular normalization effect, treatment-induced vessel disintegration and tumor apoptosis have also been proposed as factors that decrease TP in the tumor center.\textsuperscript{33} In fact, Obad et al.\textsuperscript{34} suggested that normalized vascular morphology may not improve vascular function and eventually results in poor tumoral blood flow and increased hypoxia.

Interestingly, compared with the tumor center, the tumor periphery exhibited no statistically meaningful treatment-induced change on MRI. Although our observations may depend on intrinsic characteristics of C6 glioma and sorafenib, spatially heterogeneous treatment response was also noted in several previous studies. For example, the tumor periphery exhibited maintenance of high BV and vascular integrity against monoclonal antibody to VEGF/VEGFR or vessel-targeting treatment.\textsuperscript{38-40} Therefore, preservation of inherent vascular phenotypes against antiangiogenic treatment in the tumor periphery can be regarded as a process in the tumor that potentiates drug resistance. The resistance to vessel-targeting agents has been intensively investigated in studies with vascular disrupting agents. In those studies, despite extensive necrosis in the tumor center, a viable rim of tumor cells survived in the tumor periphery, which subsequently led to regrowth.\textsuperscript{38,41,42} As an important mechanism of remaining viable rim, it has been discovered that vascular stabilization by high pericyte coverage in the tumor periphery provides inherent resistance to vascular disrupting agents. As mentioned, vessel generation and stabilization in the tumor periphery is reflected by high BV and low TP on MRI. Therefore, our MRI findings may be used as biomarkers to predict and monitor the tumor response to antiangiogenic treatment in preclinical studies.

In addition to the inherent potential against vessel-targeting agents, including high pericyte coverage of vessels, acquired drug resistance, such as over-expression of alternative signaling pathways, is recognized as an important mechanism of tumor evasion from angiogenic suppression.\textsuperscript{43,44} In this study, whereas the expression of VEGFR2, a major target of sorafenib, diminished in both the tumor center and periphery, the activities of CD31 (vessel formation) and Ki67 (cell proliferation) were preserved in the tumor periphery. This different response to treatment between VEGFR2 and CD31/Ki67 may be closely associated with acquired resistance to TKI treatment. We suggest that biologic processes other than VEGF-mediated angiogenesis may elicit the maintenance of angiogenesis to supply tumor growth in the tumor periphery.
This study applied dual injection of extravascular fluid (ie, Gd-DOTA) and intravascular (SPION) CAs to separately measure BV and TP, instead of simultaneously calculating TP and BV from a PK model-based analysis of Gd-DOTA using DCE-MRI. The main rationale of our approach was to avoid unreliability and inaccuracy of the PK-model-based method. It has been established that wide variability in estimating arterial input function leads to substantially deviations in calculating TP and BV parameters. Moreover, PK modeling is significantly influenced by $B_1$ field inhomogeneity. According to a recent simulation, pharmacokinetic parameters (ie, $K_{\text{trans}}$ and $v_p$) fluctuate up to eight times wider than non-model-based parameters (such as MaxEnh and IAUC) under the same $B_1$ inhomogeneity. Therefore, based on validated reliability in our previous study, we separately assessed the TP parameter from a non-model-based approach of DCE-MRI and BV parameter from SPION-induced $\Delta R_2^*$.

This study analyzed vascular phenotypes of orthotopic C6 glioma, which typically exhibited high TP in the tumor center and high BV in the tumor periphery. However, MRI features of tumor vasculature may vary according to numerous factors, including tumor cell properties, timing of evaluation, and method of tumor generation. For example, contrary to our results, brain metastasis models of Mel57-VEGF-A cells demonstrated only slight SPION-induced $R_2^*$ decrease but no Gd leakage at an early stage. Therefore, the angiogenic alterations observed in this study cannot be generally extended to all solid tumors. Nevertheless, this study provides a rationale for spatiotemporal vascular evaluation using both intravascular and extracellular fluid CAs. Regardless of the variety of angiogenic presentations, TP and BV are the main parameters to describe tumor angiogenesis, and their spatiotemporal manifestation can adequately reflect treatment-induced alterations of tumor vascular phenotypes.

As another potential limitation, the region-oriented analysis might be affected by rapid tumor growth, particularly between days 4 and 7. Indeed, paired comparison between time points may be incorrect because the former tumor periphery became tumor center on the follow-up MR images. Nevertheless, the temporal changes of center-to-periphery ratio of vascular parameters clearly present the time-dependent alteration in distribution of vascular characteristics.

In summary, this study quantitatively demonstrated intratumoral heterogeneity in the distribution of TP and BV parameters. This regional heterogeneity may be closely related to spatiotemporal variations in antiangiogenic treatment effects. Tumor centers exhibited apparent treatment-induced vascular normalization; however, the tumor periphery maintained its inherent vascular features against TKI treatment. Given the necessities to understand the mechanism of tumor resistance to antiangiogenic treatment, quantitative assessment of spatiotemporal heterogeneity in tumor vascular phenotypes may provide important information regarding tumor responses to therapeutic agents.

ACKNOWLEDGMENT

This work was supported by Basic Science Research Program through the National Research Foundation of Korea (NRF) funded by the Ministry of Education, Science and Technology (2017R1A2B3007567), by a grant of the Korea Health Technology R&D Project through the Korea Health Industry Development Institute (KHIDI), funded by the Ministry of Health & Welfare, Republic of Korea (HI14C1090), and by a grant [2016-602] from Asan Institute for Life Sciences, Asan Medical Center. We thank the optical imaging core facility at the ConveRgence mEDicine research cenTer (CREDIT), Asan Medical Center for support and instrumentation.

CONFLICT OF INTEREST

None declared.

ORCID

Jeong Kon Kim http://orcid.org/0000-0002-8101-8135

REFERENCES

1. Gerlinger M, Rowan AJ, Horswell S, et al. Intratumor heterogeneity and branched evolution revealed by multiregion sequencing. N Engl J Med. 2012;366:883-892.
2. Nakasone ES, Askautrud HA, Kees T, et al. Imaging tumor-stroma interactions during chemotherapy reveals contributions of the microenvironment to resistance. Cancer Cell. 2012;21:488-503.
3. Marusyk A, Almendro V, Polyak K. Intra-tumor heterogeneity: a looking glass for cancer? Nat Rev Cancer. 2012;12:323-334.
4. Nagy JA, Dvorak HF. Heterogeneity of the tumor vasculature: the need for new tumor blood vessel type-specific targets. Clin Exp Metastasis. 2012;29:657-662.
5. Nagy JA, Chang SH, Shih SC, Dvorak AM, Dvorak HF. Heterogeneity of the tumor vasculature. Semin Thromb Hemost. 2010;36:321-331.
6. Drett R, Foo J, Leder K, Mayberry J, Michor F. Intratumor heterogeneity in evolutionary models of tumor progression. Genetics. 2011;188:461-477.
7. Boxerman JL, Hamberg LM, Rosen BR, Weisskoff RM. MR contrast due to intravascular magnetic susceptibility perturbations. Magn Reson Med. 1995;34:555-566.
8. Wilhelm SM, Adnane L, Newell P, Villanueva A, Llovet JM, Lynch M. Preclinical overview of sorafenib, a multikinase inhibitor that targets both Raf and VEGF and PDGF receptor tyrosine kinase signaling. Mol Cancer Ther. 2008;7:3129-3140.
9. Guilfoyle DN, Dyakin VV, O’Shea J, Pell GS, Helpner JA. Quantitative measurements of proton spin-lattice (T1) and spin-spin (T2) relaxation times in the mouse brain at 7.0 T. Magn Reson Med. 2003;49:576-580.
10. Noh GT, Kim MH, Suh JY, et al. Sunitinib–CLIO conjugate: a VEGFR/PDGFR-targeting active MR probe. Mol Imaging Biol. 2014;16:340-349.

11. Kwon HJ, Shim WH, Cho G, et al. Simultaneous evaluation of vascular morphology, blood volume and transvascular permeability using SPION-based, dual-contrast MRE: imaging optimization and feasibility test. NMR Biomed. 2015;28:624-632.

12. Bru A, Albertos S, Luis Subiza J, Garcia-Asenjo JL, Bru I. The universal dynamics of tumor growth. Biophys J. 2003;85:2948-2961.

13. Kather JN, Zollner FG, Schad LR, et al. Identification of a characteristic vascular belt zone in human colorectal cancer. PLoS ONE. 2017;12:e0171378.

14. Farrar CT, Kamoun WS, Ley CD, et al. In vivo validation of MRI vessel caliber index measurement methods with intravital optical microscopy in a U87 mouse brain tumor model. Neuro Oncol. 2010;12:341-350.

15. Tropes I, Grimaud S, Vaeth A, et al. Vessel size imaging. Magn Reson Med. 2008;40:1130-1134.

16. Pathak AP, Ward BD, Schmainda KM. A novel technique for modeling susceptibility-based contrast mechanisms for arbitrary microvascular geometries: the finite perturber method. NeuroImage. 2008;40:1130-1143.

17. Kim JH, Suh JY, Woo DC, et al. Difference in the intratumoral distributions of extracellular-fluid and intravascular MR contrast agents in glioblastoma growth. NMR Biomed. 2016;29:1688-1699.

18. Mani NL, Schalper KA, Hatzis C, et al. Quantitative assessment of the spatial heterogeneity of tumor-infiltrating lymphocytes in breast cancer. Breast Cancer Res. 2016;18:78.

19. Graf JF, Zavodsky MI. Characterizing the heterogeneity of tumor tissues from spatially resolved molecular measures. PLoS ONE. 2017;12:e0188878.

20. Heindl A, Nawaz S, Yuan Y. Mapping spatial heterogeneity in the tumor microenvironment: a new era for digital pathology. Lab Invest. 2015;95:377-384.

21. Martinive P, De Wever J, Bouzin C, et al. Reversal of temporal and spatial heterogeneities in tumor perfusion identifies the tumor vascular tone as a tunable variable to improve drug delivery. Mol Cancer Ther. 2006;5:1620-1627.

22. Rich LJ, Sexton S, Curtin L, Seshadri M. Spatiotemporal optoacoustic mapping of tumor hemodynamics in a clinically relevant orthotopic rabbit model of head and neck Cancer. Transl Oncol. 2017;10:839-845.

23. Spagnolo DM, Al-Kofahi Y, Zhu P, et al. Platform for quantitative evaluation of spatial intratumoral heterogeneity in multiplexed fluorescence images. Can Res. 2017;77:e71-e74.

24. Holash J, Maisonneuve PC, Compton D, et al. Vessel cooption, regression, and growth in tumors mediated by angiopoietins and VEGF. Science. 1999;284:1994-1998.

25. Holash J, Wiegand SJ, Yancopoulos GD. New model of tumor angiogenesis: dynamic balance between vessel regression and growth mediated by angiopoietins and VEGF. Oncogene. 1999;18:5356-5362.

26. Bartha K, Rieger H. Vascular network remodeling via vessel cooption, regression and growth in tumors. J Theor Biol. 2006;241:903-918.

27. Carmeliet P, Jain RK. Angiogenesis in cancer and other diseases. Nature. 2000;407:249-257.

28. Carmeliet P, Jain RK. Molecular mechanisms and clinical applications of angiogenesis. Nature. 2011;473:298-307.

29. Simonavicius N, Ashenden M, van Weverwijk A, et al. Pericytes promote selective vessel regression to regulate vascular patterning. Blood. 2012;120:1516-1527.

30. Hardee ME, Zagzag D. Mechanisms of glioma-associated neo-vascularization. Am J Pathol. 2012;181:1126-1141.

31. Maisonneuve PC, Suri C, Jones PF, et al. Angiopoietin-2, a natural antagonist for Tie2 that disrupts in vivo angiogenesis. Science. 1997;277:55-60.

32. Gaustad JV, Simonsen TG, Smistad R, Wegner CS, Andersen LM, Rofstad EK. Early effects of low dose bevacizumab treatment assessed by magnetic resonance imaging. BMC Cancer. 2015;15:900.

33. Nguyen L, Fifis T, Christophi C. Vascular disruptive agent OXi4503 and anti-angiogenic agent Sunitinib combination treatment prolong survival of mice with CRC liver metastasis. BMC Cancer. 2016;16:533.

34. Obad N, Espedal H, Jirik R, et al. Lack of functional normalisation of tumour vessels following anti-angiogenic therapy in glioblastoma. J Cereb Blood Flow Metab. 2017. doi: 10.1177/0271678X17114656 [Epub ahead of print].

35. Sorensen AG, Batchelor TT, Zhang WT, et al. “A vascular normalization index” as potential mechanistic biomarker to predict survival after a single dose of cediranib in recurrent glioblastoma patients. Can Res. 2009;69:5296-5300.

36. Tong RT, Boucher Y, Kozin SV, Winkler F, Hicklin DJ, Jain RK. Vascular normalization by vascular endothelial growth factor receptor 2 blockade induces a pressure gradient across the vasculature and improves drug penetration in tumors. Can Res. 2004;64:3731-3736.

37. Dominietto M, Dobosz M, Burgi S, et al. Quantification of antiangiogenic treatment effects on tissue heterogeneity in glioma tumour xenograft model using a combination of DCE-MRI and 3D-ultramicroscopy. Eur Radiol. 2017;27:2894-2902.

38. Nguyen L, Fifis T, Malcontenti-Wilson C, et al. Spatial morphological and molecular differences within solid tumors may contribute to the failure of vascular disruptive agent treatments. BMC Cancer. 2012;12:522.

39. Zhu W, Kato Y, Artemov D. Heterogeneity of tumor vasculature and antiangiogenic intervention: insights from MR angiography and DCE-MRI. PLoS ONE. 2014;9:e86583.

40. Casanova O, Hicklin DJ, Bergers G, Hanahan D. Drug resistance by evasion of antiangiogenic targeting of VEGF signaling in late-stage pancreatic islet tumors. Cancer Cell. 2005;8:299-309.

41. Tozer GM, Kanthou C, Baguley BC. Disrupting tumour blood flow. Nat Rev Cancer. 2005;5:423-435.

42. Siemann DW, Horsman MR. Vascular targeted therapies in oncology. Cell Tissue Res. 2009;335:241-248.

43. Hida K, Kawamoto T, Ohga N, Akiyama K, Hida Y, Shindoh M. Altered angiogenesis in the tumor microenvironment. Pathol Int. 2011;61:630-637.

44. Hida K, Akiyama K, Ohga N, Maishi N, Hida Y. Tumour endothelial cells acquire drug resistance in a tumour microenvironment. J Biochem. 2013;153:243-249.

45. Beaumont M, Lemasson B, Farion R, Segebarth C, Remy C, Barbier EL. Characterization of tumor angiogenesis in rat brain using iron-based vessel size index MRI in combination with gadolinium-based dynamic contrast-enhanced MRI. J Cereb Blood Flow Metab. 2009;29:1714-1726.
46. Cheng HL. Investigation and optimization of parameter accuracy in dynamic contrast-enhanced MRI. *J Magn Reson Imaging*. 2008;28:736-743.

47. Park B, Choi BS, Sung YS, et al. Influence of B1-inhomogeneity on pharmacokinetic modeling of dynamic contrast-enhanced MRI: a simulation study. *Korean J Radiol*. 2017;18:585-596.

48. Leenders W, Kusters B, Pikkemaat J, et al. Vascular endothelial growth factor-A determines detectability of experimental melanoma brain metastasis in GD-DTPA-enhanced MRI. *Int J Cancer*. 2003;105:437-443.

How to cite this article: Kim C, Suh J-Y, Heo C, et al. Spatiotemporal heterogeneity of tumor vasculature during tumor growth and antiangiogenic treatment: MRI assessment using permeability and blood volume parameters. *Cancer Med*. 2018;7:3921–3934. [https://doi.org/10.1002/cam4.1624](https://doi.org/10.1002/cam4.1624)