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Blood Volume of Gliomas Determined by Double-Echo Dynamic Perfusion-Weighted MR Imaging: A Preliminary Study

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BACKGROUND AND PURPOSE: After bolus injection, gadopentetate dimeglumine causes a T2* rate change in permeable tissue that is contaminated by the T1 shortening effect due to the leakage of contrast agent. Therefore, tumor vascularity as reported in previous single-echo perfusion-weighted MR imaging studies has been underestimated. Our aim was to quantitatively and qualitatively evaluate the degree of blood volume of glioblastoma multiformes (GBMs) underestimated by this T1 shortening effect.

METHODS: We used double-echo dynamic MR imaging after a bolus injection of gadopentetate dimeglumine (double-echo perfusion-weighted MR imaging) to simultaneously determine tumor blood volume without (V_{TIU}) and with (V_{TIC}) T1 shortening correction. MR imaging was performed in five consecutive patients with GBMs. The ratios of V_{TIU} and V_{TIC} were calculated and compared by means of quantitative analysis. The degree of tumor blood volume as determined by V_{TIU} and V_{TIC} maps were qualitatively compared using a three-point scale.

RESULTS: All GBMs showed contrast enhancement on postcontrast T1-weighted images. In all subjects, the values of V_{TIU} were significantly lower than those of V_{TIC} (mean $\pm$ SD, 2.05 ± 1.01 vs. 3.62 ± 1.40 , respectively [$P < .05$]), indicating that tumor blood volume obtained by double-echo perfusion-weighted MR imaging was significantly higher than that by single-echo imaging. In the qualitative analysis, tumor blood volume on the V_{TIU} map was less conspicuous than that on the V_{TIC} map.

CONCLUSION: Careful attention should be paid to the underestimation of tumor blood volume resulting from T1 shortening effects when using single-echo perfusion-weighted MR imaging. Double-echo imaging may be more suitable for the analysis of blood volume in GBMs.

Studies of human neoplasms have shown that increased malignancy is associated with increased vascularity (1). The degree of vascular proliferation is an important parameter in determining the biological aggressiveness and histopathologic grading

of an astrocytoma (2). Leon et al (3) reported that the density of microvessels is a prognostic indicator for patients with astroglial brain tumors. The degree of vascular proliferation is one of the most important parameters in determining the histopathologic grade of astrocytomas, along with the degree of cellular or nuclear pleomorphism and the presence or absence of necrosis (4). Thus, it is recognized that the more malignant the astrocytoma, the greater the degree of vascular hyperplasia.

Contrast agents, such as gadopentetate dimeglumine, in the vascular space cause a signal intensity decrease on perfusion-weighted MR images resulting from the decreased homogeneity of local magnetic fields (T2* shortening effect). This characteristic, which is used to semiquantitatively evaluate tumor vascularity (5, 6), cannot be provided by conventional MR imaging. The T2* shortening effect offers a functional parameter for assessing glioma grade and regions of focal activity (7–9), information that may be important in determining treatment strategies (10–14).

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Although an examination of vascularity is useful for characterizing brain tumors, tumor vessels are different from normal blood vessels of the brain in that they do not usually express the properties of an intact blood-brain barrier. Therefore, after bolus injection, a T2* rate change caused by intravascular gadopentetate dimeglumine in a brain tumor is usually contaminated by the T1 shortening effect due to leakage of the contrast agent. Consequently, tumor vascularity as reported in previous studies with single-echo perfusion-weighted MR imaging may be underestimated.

The usefulness of double-echo perfusion-weighted MR imaging in the evaluation of the blood volume of brain tumors, such as meningiomas and neurinomas, has been reported (15, 16). It has been found that blood volume as determined by double-echo perfusion-weighted MR imaging is higher than that determined by single-echo imaging. Although glioblastomas may differ in barrier function of tumor vessels from that of extraaxial tumors, such as meningiomas and neurinomas, double-echo perfusion-weighted MR imaging may show a significant difference in the blood volume of glioblastomas as compared with that seen by single-echo imaging.

Our aim was to quantitatively and qualitatively evaluate how much vascularity is underestimated in glioblastomas by the T1 shortening effect due to the leakage of contrast material.

Methods

Theory

In previous studies with single-echo perfusion-weighted MR imaging (5–13), uncorrected tumor blood volume (V_{T1U}) contaminated by the T1 shortening effect due to the leakage of contrast agent can be estimated by the T2* shortening effect of the intravascular contrast agent ($\Delta R2^*_{T1U}$) by using the following equation, fitted with a gamma function to eliminate the second-pass effect (17):

$$V_{T1U} = \int_0^{\infty} -\ln(S/S_0)/TE(t) dt$$

where S represents the signal intensity at time t , S_0 the signal intensity before the arrival of the contrast agent, and TE the echo time. This T1 shortening effect can be corrected by the application of the double-echo technique (15, 16). The corrected tumor blood volume (V_{T1C}) can be estimated by means of the T2* shortening effect of the intravascular contrast agent, which is corrected by application of the double-echo technique ($\Delta R2^*_{T1C}$) using the following equation (double-echo perfusion-weighted MR imaging):

$$V_{T1C} = \int_0^{\infty} [\ln(S_1/S_2)/(TE_2 - TE_1)] - R2^*_{pre}(t) dt$$

where S_1 and S_2 represent the signal intensity of the first echo time (TE_1) and the second echo time (TE_2), respectively, and $R2^*_{pre}$ is the T2* rate before the arrival of the contrast agent.

Subjects

Human studies were performed under the guidelines of the Fukui Medical University Hospital Committee on clinical in-

vestigations. Written informed consent was obtained from all patients. Five consecutive patients (four men and one woman, 37–83 years old) with glioblastoma multiformes (GBMs) were examined. Two of the five patients with GBMs were examined twice on different days. In all cases, the histologic diagnosis was verified by means of surgical resection. One patient had previously undergone surgical resection followed by irradiation and chemotherapy for a GBM of the right frontal lobe. The maximum diameters of the GBMs ranged from 2.8 to 6.0 cm (mean, 4.16 cm).

MR Imaging Protocol

Axial T1-weighted spin-echo images (333/10/3 [TR/TE/excitations]) and T2-weighted fast spin-echo images (3500/88/2) were obtained before the dynamic study on a 1.5-T MR system using a rectangular 22 × 16-cm field of view, a 256 × 224 matrix, and a 5-mm slice thickness.

For the dynamic studies, we used two echoes with TEs of 7 and 23 ms of a spoiled gradient-recalled acquisition (SPGR) sequence (33.3/7, 23/0.75) with a 10° flip angle, a 256 × 128 matrix, a 7-mm slice thickness, and a rectangular 24 × 16-cm field of view. The slice displaying the largest solid portion was selected by the neuroradiologist from the information obtained from these spin-echo MR images. The apparent hemorrhagic portion was not selected. After five images were acquired, 0.15 mmol/kg body weight gadopentetate dimeglumine was rapidly injected intravenously at a rate of 4 mL/s with an MR-compatible power injector (MRS-50; Nemoto, Tokyo, Japan), followed by a 20-mL saline flush. After bolus administration of the contrast agent, a dynamic series of 50 sets of double-echo images were obtained at 2.4-second intervals. For this sequence, the total acquisition time was approximately 2 minutes. After the dynamic studies, we also obtained contrast-enhanced T1-weighted spin-echo images.

Data Processing

MR images were transferred to a Sun workstation (Ultra1 Creator 3D; Sun Microsystems, Mountain View, CA) for post-processing. For qualitative analysis, parametric maps of V_{T1U} and V_{T1C} were generated for each subject by using in-house software. On a pixel-by-pixel basis, the signal intensity was converted to changes in $\Delta R2^*_{T1C}$ and $\Delta R2^*_{T1U}$. We used a simple numerical integration of time- $\Delta R2^*_{T1C}$ and $\Delta R2^*_{T1U}$ curves instead of gamma fitting (17), because gamma fitting on a pixel-by-pixel basis is difficult owing to the relatively low signal-to-noise ratio of the dynamic images. One neuro-radiologist visually determined when the $\Delta R2^*_{T1U}$ curve reached zero, and this was considered the end point of the curve for a V_{T1U} map. The end point for the integration of the time- $\Delta R2^*_{T1C}$ curve was chosen according to regions-of-interest analysis.

Two experienced neuroradiologists qualitatively compared the degree of tumor blood volume by using only V_{T1U} and V_{T1C} maps. A consensus was reached if there was disagreement. The degree of tumor blood volume was rated on a three-point scale as follows: 1 = tumor blood volume on the V_{T1C} map was definitely more prominent than that on the V_{T1U} map, 2 = tumor blood volume on the V_{T1C} map was equivalent to that on the V_{T1U} map, and 3 = tumor blood volume on the V_{T1U} map was definitely more prominent than that on the V_{T1C} map.

For quantitative analysis, regions of interest were placed in the solid portion of the tumors and in the contralateral white matter on the SPGR images. The time series of signal intensity of SPGR images were converted to time- $\Delta R2^*_{T1U}$ and $\Delta R2^*_{T1C}$ curves. Gamma fitting (17) of time- $\Delta R2^*_{T1U}$ and $\Delta R2^*_{T1C}$ curves was performed to generate the values of V_{T1U} and V_{T1C} . However, because the time- $\Delta R2^*_{T1U}$ curve showed concentration values below zero during and after the passage of contrast agent, owing to its leakage into an interstitial space, we used

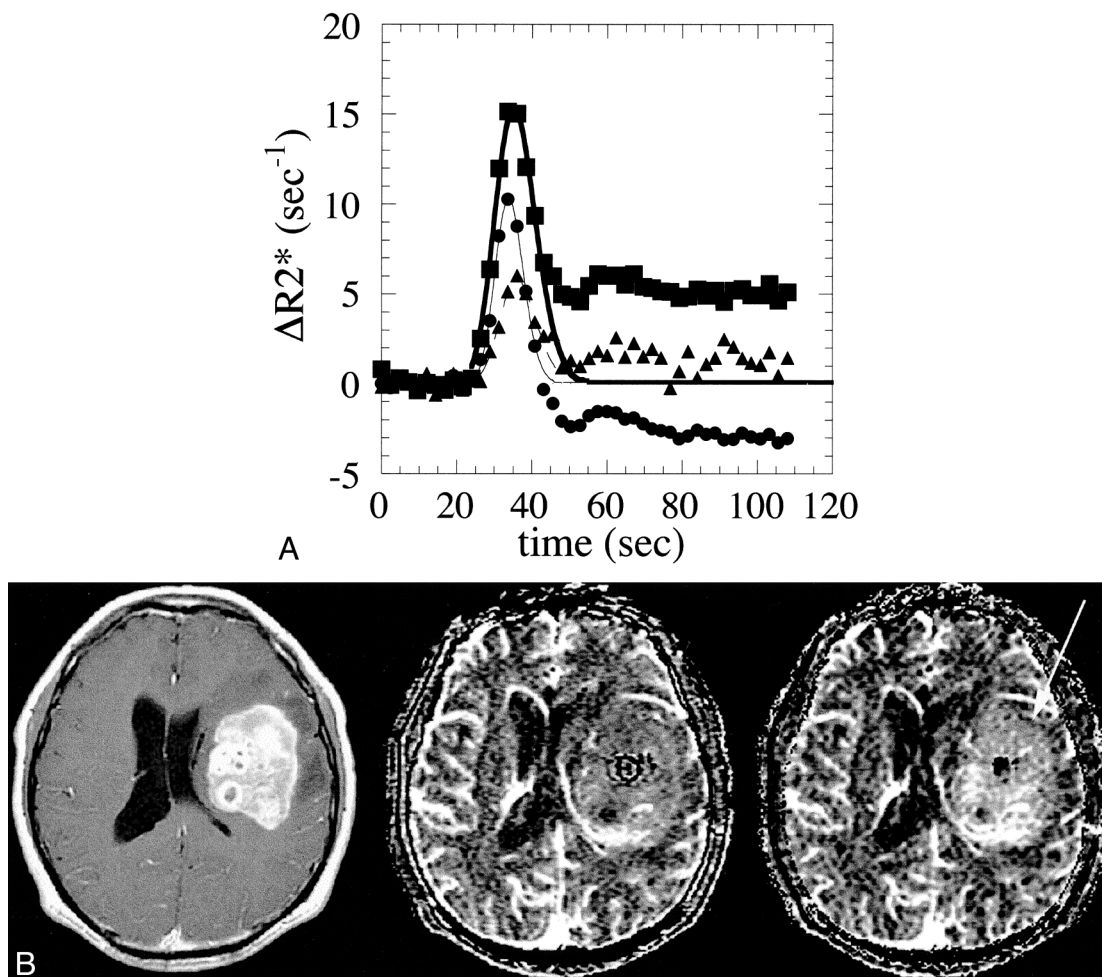


FIG 1. Case 2: 54-year-old man with a GBM.

A, Time course of the $\Delta R2^*$ values of the GBM and normal white matter. The $\Delta R2^*_{T1C}$ values of the GBM are represented by *solid squares* and its gamma-fitted curve by a *thick line*; $\Delta R2^*_{T1U}$ is represented by *solid circles* and its gamma-fitted curve a *thin line*. The $\Delta R2^*_{T1C}$ values of normal white matter are represented by *solid triangles* and its gamma-fitted curve by a *dashed line*.

B, A stereotactic biopsy was performed before the MR study. A transaxial section of a contrast-enhanced T1-weighted (333/10/3) spin-echo image is shown on the *left*, the corresponding V_{T1U} map is in the *middle*, and the V_{T1C} map is on the *right*. The GBM shows nonhomogeneously dense contrast enhancement in the left frontotemporal lobe. Note that blood volume of the tumor is much more prominent on the V_{T1C} map (*arrow*) than on the V_{T1U} map. The qualitative analysis was rated as 1 in this case. The V_{T1U}/V_{T1C} ratio is 0.41.

only sections of the curve with concentration levels above zero to perform the gamma fitting. The end point for integration was determined by gamma fitting. V_{T1U} and V_{T1C} were normalized by blood volumes of the contralateral white matter.

A statistical review of V_{T1U} and V_{T1C} was done with the Wilcoxon signed rank test, in which a P value of less than .05 was considered to be statistically significant.

Results

In this study, all GBMs showed contrast enhancement on postcontrast T1-weighted MR images (Figs 1 and 2). As shown in Figure 1A, a GBM was characterized by a high $\Delta R2^*_{T1C}$ curve and a relatively lower $\Delta R2^*_{T1U}$ curve, resulting in a lower V_{T1U} compared with V_{T1C} . The time- $\Delta R2^*_{T1C}$ and $\Delta R2^*_{T1U}$ curves of normal white matter were almost identical in all subjects. For quantitative analysis, the mean V_{T1U} of the GBMs was significantly lower than the mean V_{T1C} (mean

$\pm$ SD, 2.05 ± 1.01 for V_{T1U} vs. 3.62 ± 1.40 for V_{T1C} ; $P < .05$).

As shown in Figures 1B and 2, tumor blood volume on the V_{T1C} map was definitely more prominent than that on the V_{T1U} map in four patients. The results of the qualitative and quantitative analysis are summarized in the Table.

Discussion

Recent MR techniques have enabled the assessment of blood volume in brain tumors (5–16). Estimations of vascularity have relied on tissue characterization, preoperative assessment of tumor grade, radiation treatment planning, and subsequent follow-up care of patients with malignant gliomas. Aronen et al (7), Sugahara et al (8), and Knopp et al (9) reported that tumor blood volume was significantly different between low- and high-grade

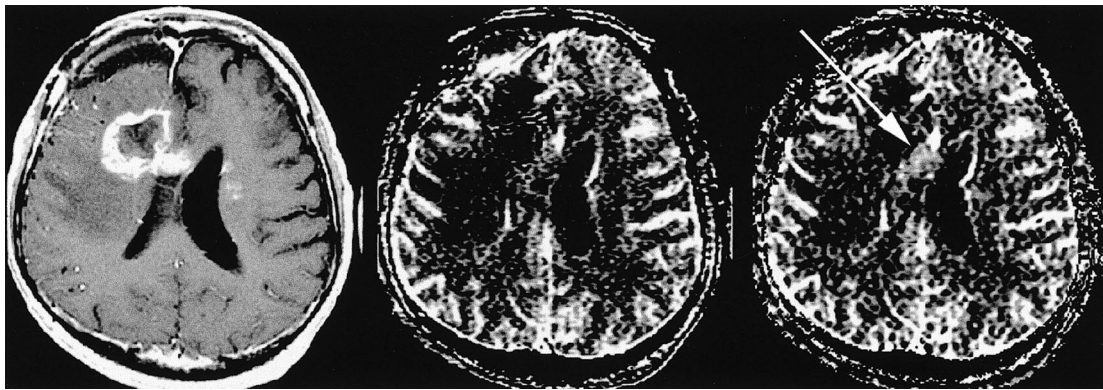


FIG 2. Case 5: 37-year-old man with a GBM who had previously undergone surgical resection, irradiation, and chemotherapy for a GBM of the right frontal lobe. Recently, a recurrent tumor was found on follow-up CT studies. A transaxial section of a contrast-enhanced T1-weighted spin-echo MR image (333/10/3) is shown on the *left*, a corresponding V_{T1U} map is in the *middle*, and the V_{T1C} map is on the *right*. The recurrent GBM shows dense contrast enhancement in the right frontal lobe, extending to the corpus callosum. Small, enhanced foci are also depicted in the left frontal lobe. Note that the lesion in the corpus callosum is depicted as an area of hypervascular blood volume on the V_{T1C} map (arrow), and the blood volume of this lesion is similar to that of gray matter on the V_{T1U} map, indicating the tumor blood volume is underestimated on the V_{T1U} map. The qualitative analysis was rated as 1 in this case. The V_{T1U}/V_{T1C} ratio is 0.64.

Results of quantitative and qualitative analysis of blood volume in GBMs

Patient No.	Age (y)/Sex	Qualitative Evaluation of Parametric Maps	Quantitative Evaluation of Regions of Interest		
			V_{T1U}	V_{T1C}	Ratio of V_{T1U}/V_{T1C}
1	83/M	1	3.80	5.89	0.64
2	54/M	1	1.50	3.62	0.41
		1	1.32	3.22	0.41
3	51/M	2	2.40	3.75	0.64
		2	1.22	2.19	0.56
4	77/F	1	2.92	4.81	0.61
5	37/M	1	1.21	1.88	0.64

Note.—Patients 2 and 3 were examined twice on different days; V_{T1U} and V_{T1C} are expressed as normalized blood volumes of the contralateral white matter.

gliomas as measured on single-echo perfusion-weighted MR images. Although examination of the vascularity of a brain tumor is useful for characterizing grades of gliomas, tumor vessels are different from normal blood vessels of the brain in that they do not usually express intact blood-brain barrier functions. Therefore, after bolus injection, gadopentetate dimeglumine causes a T1 shortening effect due to the leakage of contrast material into the tumor tissue as well as a T2* rate change in the tumor vessels on single-echo perfusion-weighted MR images.

Our findings showed that V_{T1U} was significantly lower than V_{T1C} in GBMs, suggesting that tumor blood volume is underestimated by single-echo perfusion-weighted MR imaging. Regional tumor blood volume information has been used in targeting areas for stereotactic biopsy and in debulking operations in patients with malignant gliomas (7). Complete removal of a malignant tumor is usually not possible without resecting functioning tissue.

Therefore, an underestimation of tumor blood volume may potentially cause diagnostic errors in the selection of biopsy targeting sites, leading to an erroneous assessment of tumor grade.

Because gray matter distal to a GBM might be involved in tumor extension, we used contralateral deep white matter as the reference tissue. The time- $\Delta R2^*_{T1C}$ and $\Delta R2^*_{T1U}$ curves of normal white matter were almost identical in this study, such that the question of normal brain tissue with an intact blood-brain barrier was minimized.

Concerning the relationship between the degree of contrast enhancement on postcontrast images and that of vascularity on perfusion-weighted images in brain tumors, meningiomas show high vascularity as well as prominent enhancement effect (6, 15). In a previous report by Sugahara et al (8), all GBMs showed high vascularity as well as prominent contrast enhancement. These authors also reported that the vascularity of anaplastic glioma with enhancement was significantly higher than that of the anaplastic gliomas without enhancement (8). In the case of low-grade gliomas with minimal blood-brain barrier breakdown, these lesions showed no contrast enhancement as well as low vascularity (8, 10). Thus, there may be a concordance between the degree of contrast enhancement and that of vascularity. On the other hand, Knopp et al (9) reported that in four (19%) of their 21 patients with GBMs, the lesions showed only minimal contrast enhancement but high vascularity. Cerebral lymphomas usually have lower vascularity, although these tumors produce a strong enhancement effect (18). In juvenile pilocytic astrocytoma, the tumor enhances well on postcontrast images; however, perfusion-weighted MR images show hypovascularity (5). Therefore, in some brain tumors, a discordance between the degree of contrast enhancement and that of vascularity is possi-

ble. When we apply perfusion MR techniques to brain tumors, a concordance or discordance between the degree of contrast enhancement on postcontrast images and that of vascularity on perfusion-weighted images should be kept in mind.

According to Roberts et al (19), tumor blood volume and tumor microvascular permeability can be quantified with dynamic contrast-enhanced 3D MR imaging. In 22 cases of glioma, signal intensity changes in blood and tissue were kinetically analyzed using a bidirectional two-compartment model, yielding estimates of blood volume and microvascular permeability (19). They found a strong correlation between microvascular permeability and tumor grade. Zhu et al (20) have also reported on quantification of microvascular permeability, leakage space, and blood volume in various brain tumors using combined T1- and T2*-weighted contrast-enhanced dynamic MR imaging. Three-dimensional mapping of the endothelial permeability surface area product and leakage space were generated on the basis of T1-weighted images. Tumor blood volume maps are obtained from T2*-weighted images (20). On the other hand, double-echo dynamic MR image acquisition after a bolus injection of gadopentetate dimeglumine can separate the T2* shortening effect due to intravascular localization of the contrast agent and the T1 shortening effect that occurs as a result of extravascular localization (15, 21). The combination of these two fractions can be used to measure the vascularity and the microvascular permeability of the tumors simultaneously. In the future, we may be able to characterize the gliomas by their vascularity and microvascular permeability.

Echo-planar imaging would have been desirable for this experiment; however, double-echo echo-planar imaging is not currently available in most institutions. Even if echo-planar imaging is used for the double-echo method, only two or three slices per 1.1 second can be imaged (22). We used the best technique possible with the equipment available.

Conclusion

Careful attention should be paid to the underestimation of brain tumor blood volume resulting from T1 shortening effects when using single-echo perfusion-weighted MR imaging. Double-echo perfusion-weighted MR imaging may provide a more accurate evaluation of the tumor blood volume in GBMs and may be more suitable for the purposes of analysis.

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