Diffusion Tensor MR Imaging of Cerebral Gliomas: Evaluating Fractional Anisotropy Characteristics

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Gliomas are the largest group of primary cerebral neoplasms. Low-grade gliomas by conventional MR imaging are usually homogeneous without necrosis. Edema, hemorrhage, and contrast enhancement are relatively uncommon. High-grade gliomas, by contrast, present as inhomogeneous masses with edema, hemorrhage, and irregular contrast enhancement. Signs of frank necrosis are highly suggestive of malignancy. However, hemorrhage, and irregular contrast enhancement. Signs of frank necrosis are highly suggestive of malignancy. However, hemorrhage, and irregular contrast enhancement. Signs of frank necrosis are highly suggestive of malignancy. However, hemorrhage, and irregular contrast enhancement. Signs of frank necrosis are highly suggestive of malignancy. However, hemorrhage, and irregular contrast enhancement. Signs of frank necrosis are highly suggestive of malignancy.

Materials and Methods
Thirty-four patients with gliomas (9 grade II, 8 grade III, and 17 grade IV) underwent diffusion tensor imaging at 3T. Mean FA, maximum FA, and minimum FA values were measured within the solid tissue components of the tumors. The variations of FA were evaluated by determining the range of FA values and the maximum SDs of FA. The variations of FA values among different tumor grades were compared statistically. We also correlated FA variations with minimum FA and maximum FA.

Results: The maximum FA, FA range, and maximum SD for grade II tumors were significantly lower than those for grade III and IV tumors (P < .0001 → P = .0164). A very good correlation of maximum FA to FA range (r = 0.931) and maximum SD (r = 0.889) was observed.

Conclusions: The FA range and maximum SD appear useful for differentiating low- and high-grade gliomas. This analysis added value to the findings on conventional MR imaging. In addition, focal maximum FA is a key factor contributing to the larger FA variation within high-grade gliomas.

Abbreviations: ANOVA = analysis of variance; ASSET = array spatial sensitivity encoding technique; FA = fractional anisotropy; FLAIR = fluid-attenuated inversion recovery; FSE = fast spin-echo; ROC = receiver operating characteristic analysis
six consecutive patients with pathologically proved gliomas underwent pretreatment 3T MR imaging, including diffusion tensor imaging. Thirty-four were included in this retrospective study. The remaining 2 patients had grade IV tumors that presented as extensive necrotic areas surrounded by thin rings of enhancement without obvious solid components, and the FA of a solid-tissue component could not be measured. The pathologic diagnosis was determined with specimens removed at surgical resection (n = 18) or biopsy (n = 16) by a board-certified neuropathologist (S.A.J.K.) (Table). The interval between the preoperative MR imaging studies and the pathologic diagnosis was 1–36 days (mean, 7 days). Institutional review board approval was obtained, and informed patient consent was not required for the retrospective review of the medical records or the MR images for this study.

**MR Imaging**

MR imaging was performed on a 3T MR imaging unit (Signa HDx; GE Healthcare, Milwaukee, Wisconsin) with an 8-channel head coil. The gradient system was used in zoom mode with a gradient strength of 50 mT/m and a slew rate of 150 mT/m/s. Axial diffusion tensor imaging was performed by using a single-shot spin-echo echo-planar imaging sequence (TR, 9000; TE, 80.8; bandwidth, 250 kHz; matrix, 128 × 128 zero-filled to 256 × 256; FOV, 220 × 220 mm; section thickness, 3 mm; section gap, 0 or 0.5 mm; NEX, 1; ASSET factor, 2; b = 1000; directions of gradient sampling, 25). Studies obtained were associated with preoperative brain mapping MR imaging protocols that included thin-section axial T2-weighted images (TR, 3500; TE, 98.4; flip angle, 90°; bandwidth, 41.7 kHz; matrix, 256 × 256 zero-filled to 512 × 512; FOV, 250 × 250 mm; section thickness, 2 mm; section gap, 0 mm; NEX, 1; echo-train length, 18; ASSET factor, 2), thin-section postgadolinium axial T1-weighted FLAIR images (TR, 2026; TE, 9.2; TI, 883; bandwidth, 31.2 kHz; matrix, 256 × 256 zero-filled to 512 × 512; FOV, 250 × 250 mm; section thickness, 2 mm; section gap, 0 mm; NEX, 1; echo-train length, 5; ASSET factor, 2), and/or volumetric postgadolinium T1-weighted sagittal images (TR, 7.3; TE, 2.1; flip angle, 20°; bandwidth, 31.2 kHz; matrix, 256 × 256 zero-filled to 512 × 512; FOV, 250 × 250 mm; section thickness, 1.6 mm; section gap, −0.8 mm; NEX, 1; ASSET factor, 2). These examinations were performed with a single dose of gadobenate dimeglumine (n = 19) (0.1 mmol/kg of body weight; MultiHance, Bracco, Milan, Italy) or gadopentate dimeglumine (n = 15) (0.1 mmol/kg of body weight; Magnevist, Schering, Berlin, Germany). Standard diagnostic MR imaging studies were available for comparison, which included pregadolinium sagittal T1-weighted, axial T1-weighted, axial T2-weighted FSE, axial T2-weighted FLAIR FSE, axial gradient-echo, and axial diffusion-weighted images. Postgadolinium axial and coronal T1-weighted images were also obtained.

**MR Imaging Data Analysis**

The MR images of these patients were analyzed in conference by a board-certified neuroradiologist (M.L.W.) and a radiologist (Y.Z.) with knowledge of the diagnosis of glioma but without knowledge of the histologic grade. The radiologists reached a consensus regarding the imaging findings and region-of-interest determination. Areas evaluated were designated as whole tumor, enhanced tumor component, and nonenhanced tumor component. The maximum diameters of the tumors were recorded. The sensitivity and specificity of conventional MR imaging for preoperative grading were calculated. The preoperative grading had been made clinically before this study. Tumors without enhancement, edema, and necrosis were diagnosed as grade II gliomas. Tumors with inhomogeneous intensity, enhancement, and edema were diagnosed as grade III gliomas. Tumors that showed strong inhomogeneous enhancement and/or necrosis were diagnosed as grade IV gliomas.

All diffusion-tensor data were processed on a workstation (ADW 4.3, GE Healthcare) with FuncTool software (Version 4.5.1, GE Healthcare) for determining the FA measurements.

The mean FA of a tumor was measured by placing regions of interest on the FA maps in FuncTool software on the basis of the b = 0, b = 1000 isodiffusion, mean diffusivity maps, and conventional MR images. The 3D MR imaging spatial coordinates of tumor components delineated from the gadolinium-enhanced sequences were used to direct region-of-interest placement when needed in FuncTool. When measuring the mean FA, we drew regions of interest as large as possible to cover a maximum of tumor tissue without contacting the tumor border. Careful inspection of the images and cross-referencing of the 3D coordinates of tumor cysts, necrosis, and hemorrhage allowed us to avoid these structures. Ambiguous nonenhancing areas of tumor versus edema were not included in the analysis. All imaging sections that depicted tumor were selected for measuring FA. The mean tumor FA value represents averaged FA values from all image sections that contained tumor. This was calculated for the whole tumor, the enhanced component, and the nonenhanced component.

To evaluate the variation of FA, we calculated an FA range by subtracting the minimum FA from the maximum FA value measured in a tumor. Maximum SD of FA, which is supposed to reflect maximum histologic heterogeneity, was also measured. When measuring minimum FA, maximum FA, and maximum SD of FA, we used uniform ovoid regions of interest of approximately 30 mm³ in obtaining measurements from the enhanced component, the nonenhanced component, and the whole tumor. An FA rainbow-colored map was used for analysis, which demonstrated the FA values in shades of blue, green, yellow, and red. The dark blue represented areas of low FA and the dark red represented areas of high FA. With visual inspection of the hue on FA maps, regions of interest were used to measure areas with high (red) or low (blue) FA values on each tumor section. From the values of these regions of interest, the highest or lowest one was chosen as the maximum or minimum FA of the corresponding areas of tumor, respectively. Areas with high contrast of hue (high FA variability) were measured to determine the SD of the FA. The largest SD measured was selected as the maximum SD of a tumor (Figs 1–4).
We positioned the region of interest multiple times (usually in the range of 10–30 times) in each area of interest to make sure we measured the maximum/minimum FA or maximum SD of FA. The need to use a small region of interest for calculating maximum SDs is critical so that small areas of presumably very heterogeneous pathologic change would not be overlooked. These areas of high heterogeneity may be a small component of an otherwise very homogeneous low-grade tumor. This technique roughly correlates to the neuropathologic approach in which the grade of the tumor is based on the highest grade of tumor found, even if it is a very small component of the whole tumor.

When the contrast enhancement was small, punctate, or a thin ring, the FA for the enhanced component was not measured. When the enhanced and nonenhanced components were mixed and could not be evaluated separately, the FAs were only recorded as being from the whole tumor.

**Statistical Analysis**

Data analysis was performed by a biostatistician (F.Y.) by using a commercial statistical software, SAS, Version 9.2 (SAS Institute, Cary, North Carolina). FA measurements were quantitatively compared among different tumor grades and tumor components. Normality assumption was first checked for the FA measurements of each tumor grade and component. The minimum FA, the maximum FA, and the range of FA failed to meet the normal assumption. Accordingly, the ANOVA with unequal variance was used for overall comparison of the mean value of maximum SD and mean FA, respectively. Alternatively, the Kruskal-Wallis test, a nonparametric version of ANOVA, was used to compare the minimum FA, maximum FA, and FA range, respectively, among all grades and components. To evaluate differences between each pair of groups, we used the Tukey multiple comparison method. \( P < .05 \) was considered significant.

When a significant difference was observed in a metric between

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**Fig 1.** A, T2-weighted \((b = 0)\) MR image shows a grade IV glioma in the right frontal lobe. B, FA map shows the different hues from blue to red in the tumor. C and D, FA maps at the same level show the method of region-of-interest selection. Region-of-interest 1 and region-of-interest 2 in C delineate red and blue areas that represent where the maximum FA and the minimum FA were measured. Region-of-interest 3 includes both blue and red portions and has a high contrast of color hue. The maximum SD is measured by region-of-interest 3 \((SD = 0.125)\). The areas with homogeneous color hue have a low SD of FA. Region-of-interest 4 is an example of an area with a low SD \((0.028)\).

**Fig 2.** A, T2-weighted \((b = 0)\) MR image shows a grade II glioma. B, Postgadolinium axial T1-weighted image shows whole-tumor contrast enhancement. C, FA map shows tumor FA with a homogeneous blue hue. The darker blue (cool color tone) represents low FA and the red and yellow (warm color tones) correspond to high FA. Region-of-interest selection for FA measurement is based on visually inspecting the hue. Maximum FA, minimum FA, FA range, and maximum SD are 0.145, 0.061, 0.094, and 0.0411, respectively. Arrows point to arteries.
low-grade (II) and high-grade (III and IV) tumors, an ROC was further applied to identify the cutoff value of the metric that yielded the best combination of sensitivity and specificity, to distinguish low- and high-grade tumors.

We also correlated FA variations (FA range and maximum SD) with minimum FA and maximum FA. In addition, the effect of tumor size on the FA variation was assessed. Because the data were not normally distributed, a Spearman rank correlation was calculated for the evaluation.

Results

Conventional MR Imaging Findings

The mean maximum diameters of the tumors were 4.5 cm (range, 3–6.3 cm) for grade II, 6.1 cm (range, 3–9.2 cm) for grade III, and 4.9 cm (range, 2.7–8.0 cm) for grade IV tumors. Only 1 grade IV tumor was <3 cm with a maximum diameter of 2.7 cm. Contrast enhancement was noted in only 1 grade II tumor that was pathologically proved as a pilomyxoid astrocytoma. The enhancement was total and homogeneous. There was no cystic or necrotic change in the grade II tumors. Four grade III tumors were nonenhancing, and 4 had punctate nodular-like areas of contrast enhancement. One grade III tumor contained a cystic area. There was no obvious cystic or necrotic change in the other 7 grade III tumors. Irregular contrast enhancement was noted in all 17 grade IV tumors. Four grade IV tumors were nonenhancing, and 4 had punctate nodular-like areas of contrast enhancement. One grade III tumor contained a cystic area. There was no obvious cystic or necrotic change in the other 7 grade III tumors. Irregular contrast enhancement was noted in all 17 grade IV tumors. Four grade IV tumors had nonenhanced tumor components. All grade IV tumors had various degrees of cyst formation and necrosis. Five of 8 grade III tumors were preoperatively diag-
nosed as low-grade gliomas, and the pilomyxoid astrocytoma was suspected to be a germinoma. The preoperative grading of all the other tumors (including grade II and IV) was consistent with histopathology. The sensitivity and specificity of conventional MR imaging in differentiating low-grade from high-grade (grade III and IV) tumors were 83.3% and 90%, respectively. However, the sensitivity and specificity of conventional MR imaging for differentiating grade II from grade III tumors were 61.5% and 90%, respectively.

Tumor FA Values

Only whole-tumor FA measurements were made in grade III tumors in which the contrast enhancement consisted of small nodules. Among grade IV tumors, 2 had mixed nonenhanced and enhanced components, and 2 had very small areas of enhancement. Therefore, 9 grade II tumors, 8 grade III tumors, 17 grade IV tumors, 4 nonenhanced components (grade IV), and 13 enhanced components (grade IV) were included for FA measurement. Only 4 nonenhanced components of grade IV tumors were identified for analysis because ambiguous nonenhancing areas of tumor versus edema were not included.

The ANOVA with unequal variance was used for comparison of the mean FA value among all grades and components. The mean FA value for grade II tumors (mean, 0.124 ± 0.035) was significantly lower than that for grade IV tumors (0.171 ± 0.054) (P = .0363) and nonenhanced components (0.203 ± 0.039) (P = .0275), but not significantly different from that for grade III tumors (0.148 ± 0.040) (P = .4027). The difference between grade II tumors and enhanced components (0.180 ± 0.055) approached significance (P = .0540). There were no significant differences among any pair of grade III, IV, or components (Fig 5A).

Maximum and Minimum FA Values

The mean values of the maximum FAs were 0.161 ± 0.054 for grade II, 0.237 ± 0.073 for grade III, 0.262 ± 0.090 for grade IV, 0.264 ± 0.053 for nonenhanced components, and 0.280 ± 0.093 for enhanced components. By using the Kruskal-Wallis test, statistically significant differences were found between grade II and grade III (P = .0164), grade IV (P = .0007), the nonenhanced components (P = .0187), and the enhanced components (P = .0007). No significant difference was found among any pair of grade III, IV, or components (Fig 5B).

With regard to the mean value of the minimum FAs, no significant difference was found among any pair of grade II, III, or IV tumors.

FA Range

The Kruskal-Wallis test was used to compare the FA range among all grades and components. The FA range for grade II tumors (0.084 ± 0.036) was significantly lower than that for grade III (0.168 ± 0.067) (P = .0034), grade IV (0.168 ± 0.070) (P = .0004), and the enhanced component (0.182 ± 0.068) (P = .0001). We also observed that the FA range for grade II tumors was lower than that for the nonenhanced component (0.137 ± 0.064), close to significance (P = .0692).
No significant difference was found among any pair of grade III, IV, or components (Fig 5C).

A very good correlation (correlation coefficient = 0.931) between FA range and maximum FA was observed. In contrast, only a fair correlation (correlation coefficient = 0.409) was noted between FA range and minimum FA.

**Maximum SD**

The ANOVA with unequal variance was used for comparison of the maximum SDs among all grades and components. The mean values of the maximum SDs were found between grade II and grade III (P = .0008), grade IV (P < .0001), nonenhanced components (P = .0048), or enhanced components (P = .0305). No significant difference was found for maximum SD among any pair of grade III, IV, or components (Fig 5D). A very good correlation (correlation coefficient = 0.889) between maximum SD and maximum FA was observed. In contrast, only a fair correlation (correlation coefficient = 0.467) was noted between maximum SD and minimum FA.

No correlation was found between tumor size and FA variation with a correlation coefficient < 0.1.

**Cutoff Value of FA from ROC**

The cutoff value of 0.17 for the maximum FA differentiated low- from high-grade tumors with a sensitivity of 92% and a specificity of 77.8% (Fig 6A). The cutoff value for the FA range was 0.0971, which yields a sensitivity of 96% and a specificity of 77.8% (Fig 6B). The cutoff value of 0.0492 for the maximum SD had both a sensitivity and specificity of 100% and showed complete agreement with histopathologic grading (Fig 6C). As to differentiating grade II from grade III tumors, the cutoff values were set to 0.17, 0.109, and 0.0492 for the maximum FA, the FA range, and the maximum SD, respectively. The sensitivity and specificity were 87.5% and 77.8% for the maximum FA, 87.5% and 88.9% for the FA range, and both were 100% for the maximum SD.

**Discussion**

Our study revealed that the mean FAs were significantly higher in grade IV than in grade II gliomas. However, all the grade IV gliomas had cyst formation, necrosis, and/or irregular contrast enhancement in various degrees. None were preoperatively diagnosed as a lower grade glioma. By contrast, 5 of the 8 grade III gliomas were preoperatively diagnosed by conventional MR imaging as grade II gliomas due to benign imaging features. However, mean FA did not significantly differ between grade III and grade II gliomas either. Therefore, there was no added value of mean FA evaluation over conventional MR imaging in grading gliomas despite a mean FA difference between grade IV gliomas with typical malignant imaging features and grade II gliomas. A study by Inoue et al showed that grade III gliomas had significantly higher FA than grade II gliomas. However, the FA values were measured within enhanced components when the tumor had contrast enhancement. In contrast, all the grade III gliomas in our study were nonenhanced or had miniscule contrast enhancement.

Tumor cells that infiltrate fiber tracts cause displacement, deviation, and destruction of these tracts and lead to reduction of FA. With more active tumor proliferation and consequently more obvious fiber tract disorganization, it might be hypothesized that high-grade gliomas are expected to reveal lower FA values than low-grade gliomas. However, our result, higher mean FA values in high-grade gliomas, is contrary to this expectation. This difference could be explained by the increases of cell density, cellularity, and vascularity in high-grade gliomas. It has been previously reported that tumor FA positively correlates to cell density, cellularity, and vascularity.

Sometimes high-grade gliomas, especially grade III, have focal hypercellularity and higher cell density within an otherwise low-grade-appearing mass. The malignant foci might not be large enough to induce mean FA changes in the whole tumor but could induce higher FA values in these foci. In our study, the mean values of maximum FA of both grade III and IV gliomas were significantly higher than those of grade II gliomas.

When comparing minimum FAs, we found no difference between low- and high-grade gliomas. This could be attributed to the fact that most high-grade gliomas arise from dedifferentiated lower grade ones. The low-grade glioma tissue, which has low FA values to start with, is likely due to less cellularity and density and may still exist in high-grade gliomas.

Like maximum FA, the FA range and maximum SD in grade II gliomas were significantly smaller than those in grade...
IV gliomas with typical malignant features as well as in the grade III gliomas with benign findings on conventional imaging. A strong correlation of maximum FA with FA range and maximum SD was noted statistically. Our study suggested that the larger FA ranges and maximum SDs in higher grade gliomas might be mainly caused by the increased maximum FA values. Because most grade III gliomas in this study were lacking in malignant features on conventional imaging, the difference of maximum FA, FA range, and maximum SD between grade II and III gliomas in our study population provided added value to conventional MR imaging in the differentiation of these tumors. By using the specified cutoff values of maximum FA, maximum SD, and FA range, ROC revealed that the sensitivities increased from 61.5% on conventional imaging to 87.5%–100% in differentiating grade II from grade III gliomas and from 83.3% to 92%–100% in differentiating grade II from grade III and IV gliomas, respectively. This finding is also clinically significant because of the dramatically different treatment and prognosis between the low- and high-grade gliomas.23,24 Further study with more cases is necessary to evaluate the ROC in a larger population. We were not able to evaluate FA characteristics in patients with highly enhancing grade III gliomas because the grade III gliomas in this study were nonenhancing or had only punctuate nodular-like areas of contrast enhancement.

Theoretically, the larger the tumor, the more likely it is to have more areas of histologic variability. A large low-grade glioma could potentially have greater FA variation than a small high-grade glioma. However, only 1 high-grade glioma was <3 cm in this study. Given this finding, we were not able to specifically compare the FA variation between large low-grade gliomas and small high-grade gliomas. When analyzing all our cases, we found no correlation between tumor size and FA variation.

To our knowledge, there has been very limited analysis of heterogeneity using diffusion imaging in gliomas. In 1 study, the analysis focused on apparent diffusion coefficient values but did point to a potential broader application of our tumor analysis approach.25 In another study, visual FA map analysis was used and grade II tumors were homogeneous but overall FA evaluation did not help grade tumors.26 Our analytic approach of evaluating focal maximum SD is rudimentary at this point, but the results are promising.

We speculate that the tumor components with maximum SD or high maximum FA might reflect regional heterogeneity of histopathology and/or the more malignant foci. The need to define focal changes in a glioma rather than average values of metrics can be very important because a small focal area of high-grade glioma in an otherwise low-grade glioma makes a critical diagnostic difference. Our approach of focal SD analysis lends itself to creating maps of the SD metrics in a mass by calculating the focal SD across a lesion on a pixel-by-pixel basis. This SD map could then provide guidance for further noninvasive analysis or even biopsy. Of course, the point of maximum FA in a glioma may also provide a target for biopsy and directed neuropathologic analysis. However, future studies correlating diffusion tensor imaging to tissue histopathology are needed to confirm this speculation.

Positive correlation between FA and vascularity has been reported previously.11 However, no significant difference of FA was found among nonenhanced and enhanced high-grade tumors or components in this study. This is likely because tumor enhancement results mainly from disruption of the blood-brain barrier, not tumor vascularity.27,28 Our FA evaluation of the nonenhanced tumor components of grade IV tumors was likely affected by the fact we did not include areas in the brain that had high T2-weighted signal intensity but were ambiguous for representing tumor or only edema. This limited our analysis of nonenhanced components of grade IV tumors to n = 4.

There were limitations to our study. The results were obtained on the basis of a small number of patients. Some patients only had biopsies that could be associated with the potential for tumor undergrading. On the other hand, the regions of interest for measuring maximum FA and SD were not assessed histologically. Therefore, we can only speculate that areas with maximum FA might be the most metabolically active portions of tumor, which have hypercellularity, higher cell density, and increased vascularity. Also, we can only speculate that maximum SD might represent regional heterogeneity of histopathology. In applying our technique more broadly, we found that obtaining an accurate FA measurement is also likely limited in small tumors or very infiltrative tumors because the FA values of these tumors would likely be markedly affected by infiltrating white matter fibers. Region-of-interest mismapping may have occurred when evaluating the FA characteristics of the whole tumors and the tumor components and when avoiding areas of cysts, necrosis, and hemorrhage. If mismapping occurred, then the FA values measured would not be representative of the soft-tissue components that we were aiming to measure. Because we carefully placed the regions of interest on the basis of multiple MR imaging sequences, we have minimized the potential for such errors.

Conclusions

The results of this study reveal that high-grade gliomas have a greater variation of FA values. In addition, focal maximum FA, the key factor for causing larger FA variation, itself appears useful in differentiating glioma grades. Further study that correlates maximum FA and maximum SD with histologic features is needed to make definite conclusions about the clinical application of these diffusion tensor metrics in grading gliomas.

References