Utility of Diffusion Tensor Imaging in Evaluation of the Peritumoral Region in Patients with Primary and Metastatic Brain Tumors

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ABSTRACT

SUMMARY: In the brain, diffusion tensor imaging is a useful tool for defining white matter anatomy, planning a surgical approach to space-occupying lesions, and characterizing tumors, including distinguishing primary tumors from metastases. Recent studies have attempted, with varying success, to use DTI to define the extent of tumor microinfiltration beyond the apparent borders on T2-weighted imaging. In the present review, we discuss the current state of research on the utility of DTI for evaluating the peritumoral region of brain tumors.

ABBREVIATIONS: FA = fractional anisotropy; MD = mean diffusivity; TII = tumor infiltration index

Malignant brain tumors such as glioblastoma multiforme have an infiltrative pattern of growth, often invading the surrounding tissue.1–3 Actual tumor margins can extend microscopically for several centimeters past the radiographically detected margin of disease.3 Furthermore, infiltrative brain tumors are surrounded by extensive areas of edema, as detected on T2-weighted imaging. The transition between the tumor edge and peritumoral edema is ambiguous. In metastatic tumors, this peritumoral edema is thought to consist of pure water, which can obscure tumor borders. However, the peritumoral edema of infiltrative tumors such as glioblastoma multiforme has often been shown to contain tumor cells that have spread into the edematous tissue.4 Conventional imaging techniques such as MR imaging and CT are not able to detect this minute cellular infiltration, and studies show that tumor cells have been demonstrated beyond tumor borders as defined by these modalities.5–8 This limitation of conventional CT and MR imaging presents a serious challenge for treatment planning. In successful surgical resection or biopsy with adjuvant therapy, the tumor must be excised to the greatest extent possible while minimizing injury to nearby healthy tissue.

Standard imaging techniques tend to underestimate the extent of tumor involvement, a problem that can lead to suboptimal treatment and a worse prognosis.9

Studies of animal models provide insight into the mechanisms of change in DTI metrics and imply that DTI may have a role in more precisely defining the extent of tumor cell infiltration. Among other applications, DTI has been successfully used in animals for mapping white matter tracts,10 for characterizing the directionality of diffusion within tumors,11 and as a marker of pathologic states such as demyelination.12 Several animal studies have also used DTI to detect tumor microinfiltration into surrounding tissue, by using histologic analysis as the criterion standard.13,14 For example, Lope-Piedrafita et al (2008)15 implanted glioma cells (C6) into the brains of 6 adult rats and, by using DTI metrics, showed that fractional anisotropy (FA) values in the tissue contralateral to the tumor were relatively high and did not change significantly with time, while the FA values adjacent to the tumor, though they were initially similar to those of the contralateral side, tended to decrease as the tumor grew. This reduction in fractional anisotropy became significant by day 6 of growth, suggesting that as the tumor grew, cancer cells infiltrated the tissue surrounding the tumor. The introduction of these more randomly distributed cells among white matter axons caused a reduction in the uniformity of water diffusion within tissue, leading to lower FA values of the peritumoral white matter on DTI.

In a similar study, Kim et al (2008)16 used DTI to monitor the growth of rapid-growing F98 and the slower growing 9L tumor models in rats. In evaluating the peritumoral regions, they found that due to tissue compression from rapid growth, the coefficient of planar anisotropy was higher surrounding
Summary table of DTI studies of the peritumoral region

| Article                        | No.   | Design            | Measures | Tumor types                                      | Criterion Standard | Conclusion |
|-------------------------------|-------|-------------------|----------|-------------------------------------------------|--------------------|------------|
| Deng et al, 2010              | 20    | Prospective cohort study | FA, ADC  | Primary Metastasis Extra-axial Tissue | Yes                |            |
| Wang et al, 2009              | 28    | Prospective cohort study | FA, p, q, L | Primary Metastasis Extra-axial None | Yes                |            |
| Price et al, 2006             | 24    | Prospective cohort study | FA, p, q  | Primary Metastasis Extra-axial Tissue | Yes                |            |
| Sinha et al, 2002             | 9     | Retrospective analysis | FA, MD   | Primary Metastasis Extra-axial None | Yes                |            |
| Provenzale et al, 2004        | 17    | Retrospective analysis | FA, ADC  | Primary Metastasis Extra-axial None | Yes                |            |
| Lu et al, 2005                | 20    | Prospective cohort study | RAI      | Primary Metastasis Extra-axial Follow-up | Yes                |            |
| Sundgren et al, 2006          | 28    | Prospective cohort study | FA, ADC  | Primary Metastasis Extra-axial None | Yes                |            |
| van Westen et al, 2006        | 30    | Prospective cohort study | FA, ADC  | Primary Metastasis Extra-axial None | No                 |            |
| Kinoshita et al, 2010         | 14    | Retrospective analysis | FA, ADC, TII | Primary Metastasis Extra-axial None | No                 |            |
| Tropine et al, 2004           | 12    | Prospective cohort study | FA, MD   | Primary Metastasis Extra-axial None | Inconclusive       |            |

**Note:** p indicates pure isotropic vector component; q, pure anisotropic vector component; L, total magnitude of the diffusion tensor; RAI, relative anisotropy index.

*Whether the study data suggest that DTI has utility in determining the extent of brain tumor infiltration.*

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FIG 1. The peritumoral region and corresponding imaging modalities. Tumor cells can extend beyond the bulk tumor into vaguely concentric subdivisions of the peritumoral region, including the transitional zone (1), the region of tumor infiltration (2), the peritumoral edema (3), and even the normal-appearing white matter (4).

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be needed. The purpose of this review is to highlight the available literature as a basis for further research.

**Literature Search Approach and Results**

Using variations of the search term “diffusion tensor imaging brain tumor,” we acquired our raw list of articles from the PubMed database. For completeness, we also looked for relevant articles cited by these publications. Our search yielded 252 articles that discussed DTI. Of those, there were 5 animal studies, 35 case reports, 2 letters, 1 meta-analysis, 137 randomized controlled trials, 21 retrospective studies, and 51 reviews. In terms of content, 33 articles concerned the peritumoral region; 108, white matter tractography and/or surgical approach; 46, bulk tumor characterization; and the rest covered miscellaneous topics. Of the peritumoral articles, 12 directly addressed the question of whether DTI has utility in evaluating tumor infiltration in the peritumoral region (Table). These studies typically used ROIs and an expert rater to evaluate DTI results. A notable limitation in discussing these studies is that they are not consistent in precisely which subdivision of the peritumoral region (Fig 1) they address. FA values vary depending on location, such as whether they are measured in compact or noncompact regions of tissue. This must be taken into account in any DTI analysis so as not to compromise the specificity of study results.

Finally, the tractography results reviewed here are based on deterministic algorithms. Other approaches, such as probabilistic tractography and multishell dMRI, may overcome obstacles to tractography such as crossing-fiber zones and abnormal tissue. While these techniques may improve the delineation of tract-based ROIs in the peritumoral region, this review specifically focuses on assessments of scalar metrics of anisotropy because these comprise the methods that have been reported to date.
Distinguishing Primary Brain Tumors from Metastases

Brain tumors, regardless of whether primary or metastatic, are surrounded by some degree of vasogenic edema, detectable as hyperintensity on T2-weighted images. Diffusion metrics have been shown to be altered within this surrounding edema, compared with internal controls, in both high-grade gliomas and metastatic tumors, reflecting the presence of, at minimum, increased extracellular water. It has been shown further, by using DWI, that the diffusivity of the peritumoral edema varies depending on the characteristics of the tumor it encompasses. High-grade gliomas are associated with edema of high trace value (mean diffusivity [MD]), computed by averaging the diagonal elements of the matrix \[17\]), whereas the low-grade gliomas or nonglial tumors are associated with edema of low trace value. Morita et al (2005)\[18\] found a similar range of anisotropy in both types of edema, yet the diffusivity was much greater in the edema surrounding high-grade gliomas. They concluded that the "higher diffusivity of water molecules within the area of edema associated with high-grade gliomas is likely to reflect destruction of the extracellular matrix ultrastructure by malignant cell infiltration."\[18\] This conclusion was supported by a subsequent study by Kimura et al\[19\] by using changes in glutamate levels (as a marker of the integrity of the extracellular matrix) as the central measure. Figure 2 illustrates the factors behind comparable changes in peritumoral FA.

Current research demonstrates that surrounding both primary brain tumors and metastatic tumors, there is an increase in mean diffusivity and a decrease in FA compared with normal-appearing white matter, measurements best explained by increased extracellular water. The greater the amount of free water in tissue, the greater the magnitude of diffusion will be (increased MD) and the more disorganized the diffusion becomes (decreased FA).\[19\] Lu et al (2003)\[20\] found that the peritumoral MD of metastatic lesions was greater than that of gliomas, while the peritumoral FA between the 2 tumor types demonstrated no significant difference. This finding implies that the reduced FA surrounding gliomas can be attributed not only to increased extracellular water but also to axonal disorganization caused by infiltration of the tumor beyond its detectable margin.

FIG 2. Schematic illustrating the factors behind the comparable change in peritumoral FA. A, Water content contributing to decreased FA in metastatic lesions. B, Water content and axonal disorganization contributing to decreased FA in high-grade gliomas. Reprinted from Lu et al.\[20\]

Distinguishing Tumor Extension from Edema of the Peritumoral Region

External support for this notion that reduced FA in peritumoral edema is a marker of tumor infiltration comes from the Sundgren et al (2006)\[21\] study of recurrent and nonrecurrent brain tumors. Tumors that have great infiltrative potential are more likely to recur than tumors lacking such potential.\[22\] Sundgren et al measured the FA values in the normal-appearing white matter bordering the peritumoral edema of recurrent and nonrecurrent tumors. These investigators reported significantly lower FA values in the normal tissue surrounding recurrent tumors compared with nonrecurrent tumors,\[23\] suggesting the presence of microinfiltration by the recurrent tumors into the peritumoral white matter. This hypothesis could be neatly reconciled with the known correlation between tumors of high infiltrative capacity and the associated recurrence risk. Earlier work has similarly shown that changes in DTI metrics can identify subtle white matter disruption due to occult infiltration by aggressive cancers.\[23\]

In a second study in 2004, Lu et al\[24\] looked retrospectively at DTI data from 40 patients and attempted to determine whether the MD and FA could be used to distinguish intra-axial from extra-axial lesions, metastatic lesions from gliomas, and high-grade (World Health Organization grades III-IV) from low-grade (World Health Organization grade II) gliomas. However, to better define the extent of tissue infiltration, these investigators used a third metric that they termed the "tumor infiltration index (TII)," which was calculated from DTI data by the following formula:

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TII = \frac{FA_{exp} - FA_{obs}}{3} \times 10^3
\]

where \(FA_{exp}\) is the expected FA for the corresponding MD, assuming no tumor infiltration (as determined by a linear regression analysis of such data) and \(FA_{obs}\) is the currently measured FA. Thus, in order for the results to be consistent with those of the above study, the TII of noninfiltrative tumors should be approximately zero, while the TII of infiltrative tumors should be considerably higher. That is indeed the outcome achieved. Gliomas had a mean TII of 64, while metastases and meningiomas had a mean TII of 0 (Fig 3).
A number of other studies have emerged that lend support to the notion that DTI metrics might be a useful marker of tumor infiltration into the surrounding tissue. For example, Tropine et al (2004)\textsuperscript{25} found a reduction of FA in the peritumoral, T2 signal-enhancing region of glioblastomas compared with meningiomas and concluded that this may be an indicator of tumor cell infiltration, though they questioned the reliability of such a test given the evidence available. DTI results from a patient with glioblastoma are shown in Fig 4.

A similar study by Provenzale et al (2004)\textsuperscript{26} further demonstrated the efficacy of DTI in differentiating normal and tumor-infiltrated peritumoral tissue. The investigators found that mean FA values in peritumoral hyperintense regions were 43% of the normal white matter value for gliomas and 65% of the normal white matter value for meningiomas (\(P = .05\)), while the mean FA values for normal-appearing white matter neighboring the tumors were 83% of the normal value for patients with gliomas and 100% of the normal value for those with meningiomas (\(P = .01\)).\textsuperscript{26} In other words, there was a noticeable but statistically insignificant difference in FA in the hyperintense regions of gliomas and meningiomas. However, the magnitude of the decrease in FA in the normal-appearing, neighboring white matter was significant: The decrement was greater in patients with glioma (\(P = .01\)).

The fact that DTI analysis reveals a disparity in the FA of peritumoral normal-appearing tissue and that this disparity is greater in the tissue neighboring the gliomas than in neighboring meningiomas perhaps implies that DTI has the potential to detect tumor infiltration that cannot be discerned by standard MR imaging techniques.

Recent evidence further supports the notion that DTI, specifically the FA value, is a useful tool in precisely delineating tumor margins. Deng et al (2010)\textsuperscript{27} correlated fractional anisotropy with the distance from visible tumor margins in gliomas and found that the value increases as the distance from the tumor increases. In addition, by obtaining tissue samples from the regions of interest as determined on MR imaging, these investigators observed a negative association between FA values from the peritumoral edema of gliomas and the degree of tumor infiltration (categorized as mild, moderate, severe, or no infiltration).\textsuperscript{27} These results are consistent with past findings that a reduction in FA in the peritumoral region could be used as a quantitative index of tumor cell infiltration.

In addition to FA and MD, several eigenvalues derived from DTI have been successfully used for studying the extent of brain tumor infiltration. These include the pure isotropic component (\(p\)), the pure anisotropic component (\(q\)), and the total magnitude of the diffusion tensor (\(L\)), where \(L = \sqrt{p^2 + q^2}\).\textsuperscript{28} It has been found, for example, that by using the pure diffusion components, one can distinguish gross tumor (reduction of \(q > 12\%\) from contralateral region) from tumor infiltration (increase \(p > 10\%\) from contralateral region).\textsuperscript{29} These metrics may perhaps strengthen the role of DTI in the analysis of the peritumoral region.

**Limitations of DTI in Delineating Tumor Margins**

In contrast to those studies discussed previously, several studies did not find conclusive evidence of the utility of DTI for detecting tumor infiltration. In a study of 9 patients by Sinha et al (2002),\textsuperscript{13} the FA values were significantly different in only 7 of the 9 patients studied and DTI was not found to have any advantage over the more conventional diffusion-weighted MR imaging (the FA differences paralleled the diffusivity differences and did not add any additional insight). With such a small study population, however, this result may not be representative.

A larger study conducted by van Westen et al (2006)\textsuperscript{14} measured the tumor-to-brain ratios of FA and ADC in T2 hyperintense regions surrounding gliomas, meningiomas, and metastases and the adjacent normal-appearing white matter of 30 patients. They found no difference among the 3 tumor types in terms of ADC magnitude, the lesion-to-brain ratios of ADC, the FA magnitude, the lesion-to-brain ratio, or FA within the peritumoral areas with T2 signal changes. Furthermore, they found no difference in the ADC or tumor-to-brain ratios of FA in the adjacent normal-appearing white matter. van Westen et al, therefore, con-
cluded that the FA was not helpful in identifying tumor-infiltrated edema. In addition, Kinoshita et al (2010) re-evaluated the utility of the previously discussed tumor infiltration index in 14 patients, comparing DTI with $^{11}$C-methionine PET, which has been shown to have utility in identifying tumor infiltration. In glioma cases ($n = 8$), the mean TII ranged from $–127$ to $96$, with an overall mean of $5.4 \pm 65$. On the other hand, the TII for meningiomas ($n = 5$) and metastatic brain tumors ($n = 1$) ranged from $–24$ to $44$, with an overall mean of $6.8 \pm 29$. However, these differences were not statistically significant—a finding that questions TII (a measure derived from the FA) as a tool for distinguishing tumor-infiltrated edema from vasogenic edema. However, voxel-by-voxel comparison of TII and $^{11}$C-methionine PET in the areas of gliomas demonstrating T2-signal hyperintensity showed a positive correlation between the two modalities. This suggests that though TII may perhaps be unable to delineate tumor infiltration within vasogenic edema, it is still useful for identifying tumor cell infiltration into nonedematous regions. It has been further suggested that $^{11}$C-methionine PET could be used as an adjunctive study with DTI to help distinguish tumor-infiltrated edema from typical vasogenic edema.

The concern may arise that the integration of results from multiple diffusion MRI studies creates the potential for systematic bias due to differences in data acquisition and analysis methods among studies. This concern is germane when quantitative measures are compared across studies. However, when study parameters (such as hardware, software, data acquisition, and analysis methods) are standardized across all subjects within a study, as in the articles reviewed herein, this concern is minimized, provided that the temporal stability of the diffusion MRI measurement is maintained during the period of data acquisition for the study sample. Because this review was based on the assessment of the findings of other studies but does not compare their quantitative measurements, this concern should be minimal in the context of this review.

**CONCLUSIONS**

From the evidence discussed above (Table), we contend that DTI is a sophisticated and useful tool to aid in the delineation of tumor margins and the detection of brain tumor infiltration. The evidence suggests that decreased fractional anisotropy in peritumoral tissue, compared with the corresponding contralateral brain region, is an indicator of tumor invasion past the tumor borders as detected on T2-weighted images. Although the architecture of the tissue can be highly variable and therefore create inconsistent or nonspiculate results, the use of adjunctive DTI metrics can improve the quality of findings and facilitate accurate and specific conclusions.

We believe that the technique is promising but that the literature is currently insufficient to establish a definitive clinical role for DTI in the characterization of the peritumoral region. The primary reason for this is that though there have been many studies of the utility of DTI for this novel purpose (Table), few have used a criterion standard such as tissue histopathology to confirm their results. Future studies by using a criterion standard could lay the groundwork for an eventual meta-analysis that could conclusively quantify the overall effectiveness of the technique. In addition, automated translation of DTI metrics to visual anatomic maps may make the process of evaluating the peritumoral region by using DTI more uniform and therefore more clinically relevant. We expect that continued research in this area will further advance the quality and utility of DTI for delineating tumor extent.

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