Diffusion Tensor MR Imaging and Fiber Tractography: Theoretic Underpinnings

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AJNR Am J Neuroradiol 2008, 29 (4) 632-641
doi: https://doi.org/10.3174/ajnr.A1051
http://www.ajnr.org/content/29/4/632
**Diffusion Tensor MR Imaging and Fiber Tractography: Theoretic Underpinnings**

**SUMMARY:** In this article, the underlying theory of clinical diffusion MR imaging, including diffusion tensor imaging (DTI) and fiber tractography, is reviewed. First, a brief explanation of the basic physics of diffusion-weighted imaging (DWI) and apparent diffusion coefficient (ADC) mapping is provided. This is followed by an overview of the additional information that can be derived from the diffusion tensor, including diffusion anisotropy, color-encoded fiber orientation maps, and 3D fiber tractography. This article provides the requisite background for the second article in this 2-part review to appear next month, which covers the major technical factors that affect image quality in diffusion MR imaging, including the acquisition sequence, magnet field strength, gradient amplitude and slew rate, and multichannel radio-frequency coils and parallel imaging. The emphasis is on optimizing these factors for state-of-the-art DWI and DTI based on the best available evidence in the literature.

Diffusion MR imaging of the brain was first adopted for use in clinical neuroradiology during the early 1990s and was found to have immediate utility for the evaluation of suspected acute ischemic stroke. Since that time, enormous strides forward in the technology of diffusion imaging have greatly improved image quality and enabled many new clinical applications. These include the diagnosis of intracranial pyogenic infections, masses, trauma, and vasogenic-versus-cytotoxic edema. Furthermore, the advent of diffusion tensor imaging (DTI) and fiber tractography has opened an entirely new noninvasive window on the white matter connectivity of the human brain. DTI and fiber tractography have already advanced the scientific understanding of many neurologic and psychiatric disorders and have been applied clinically for the presurgical mapping of eloquent white matter tracts before intracranial mass resections.

This 2-part review explores the current state of the art for the acquisition and analysis of clinical diffusion imaging, including DTI and fiber tractography. This first article begins with an explanation of the basic physics and underlying theory of diffusion-weighted imaging (DWI) and apparent diffusion coefficient (ADC) mapping. This is followed by an overview of the additional information that can be derived from the diffusion tensor, including diffusion anisotropy, color-encoded fiber-orientation maps, and 3D fiber tractography. This article provides the necessary background for the second article, which focuses on the major technical factors that affect image quality in diffusion imaging, with an emphasis on optimizing these factors for state-of-the-art DWI and DTI based on the best available evidence in the literature.

**Theoretic Underpinnings of Diffusion Imaging**

**Brownian Motion and Tissue Ultrastructure**

Diffusion, also known as “Brownian motion,” refers to constant random microscopic molecular motion due to heat. At a fixed temperature, the rate of diffusion can be described by the Einstein equation:

\[ \langle r^2 \rangle = 6Dt, \]

where \( \langle r^2 \rangle \) refers to the mean squared displacement of the molecules, \( t \) is the diffusion time, and \( D \) is the diffusion constant, a constant of proportionality for the particular substance being measured. In clinical diffusion imaging, the type of diffusion being investigated is water self-diffusion, meaning the thermal motion of water molecules in a medium that itself consists mostly of water. The diffusion constant, typically expressed in units of square millimeters per second, relates the average displacement of a molecule over an area to the observation time, with higher values of this constant indicating more mobile water molecules. The ADC is the diffusion constant measured in the clinical setting, reflecting the limitation that in vivo diffusion cannot be separated from other sources of water mobility, such as active transport, flow along pressure gradients, and changes in membrane permeability.

Imaging molecular water diffusion confers the ability to probe the microstructural properties of biologic tissues. The typical diffusion times used for clinical DWI are 10–50 ms, corresponding to average molecular displacements on the order of 10 µm. This microscopic spatial scale is in the same range as that of cellular dimensions. This sensitivity to cellular processes has been exploited clinically for improving the detection of acute cerebral ischemia, distinguishing vasogetic from cytotoxic edema, identifying foci of axonal shearing injury after acute head trauma, characterizing cellularity in brain tumors, discriminating between metastases and gliomas and between tumor recurrence and postsurgical injury, differentiating pyogenic abscesses from tumors, and for the noninvasive mapping of white matter connectivity by using the diffusion tensor model among other applications.

**Stejskal-Tanner Diffusion Encoding**

A diffusion-weighted pulse sequence is constructed by the addition of a pair of diffusion-sensitizing gradients, also known as motion-probing gradients, to a T2-weighted spin-echo sequence. The diffusion gradients are applied along the same directional axis both before and after the 180° refocusing pulse (Fig 1). This is known as Stejskal-Tanner diffusion encoding. Molecular motion thus results in loss of signal intensity due to incomplete rephasing of water proton spins, which change...
position between and during the applications of the 2 diffusion-sensitizing gradients. This diffusion-weighted contrast can be fit to an exponential model:

$$S_i = S_0 \cdot e^{-b \cdot \text{ADC}}.$$  

In the above expression, $S_i$ is the diffusion-weighted signal intensity observed at a given voxel with the diffusion-sensitizing gradients applied along direction $i$, $S_0$ is the signal intensity at the same voxel measured without diffusion-sensitizing gradients, and ADC$_i$ is the ADC in the $i$ direction. Within the exponential term, the $b$ factor is a measure of diffusion-weighting that is a function of the strength, duration, and temporal spacing of the diffusion-sensitizing gradients, assuming rectangular gradient pulses:

$$b = \gamma^2 G^2 \delta^2 (\Delta - \delta/3),$$

where $\gamma$ is a physical constant known as the gyromagnetic ratio, $G$ is the amplitude of the diffusion gradient typically measured in milliteslas per meter, $\delta$ is the duration of each diffusion gradient in milliseconds, and $\Delta$ is the interval between the onset of the diffusion gradient before the refocusing pulse and that after the refocusing pulse, measured in milliseconds (Fig 1). The units of $b$ are seconds per square millimeter, reciprocal to the units of ADC. It can be seen from equation 2 that raising $b$ increases the degree of diffusion-weighting (ie, the loss of signal intensity $S_i$ along direction $i$ produced by application of the diffusion gradients). From equation 3, it is apparent that boosting the gradient amplitude $G$ and the gradient duration $\delta$ are the most effective ways of increasing the diffusion-weighting, due to the quadratic dependence of $b$ on these 2 parameters. Typical values of $b$ used in clinical applications range from 600 to 1500 seconds per square millimeter. The lowest values of $b$ are used for imaging of fetuses in utero or premature neonates, due to the very high ADC values of these structurally immature brains.38,39 The highest $b$ factors have been advocated for hyperacute stroke imaging in adults at 1.5T, to maximize sensitivity to the often subtle reductions in ADC in this very early stage of cerebral ischemia, 40,41 though a recent 3T study42 found no benefit of $b = 1500$ s/mm$^2$ DWI for stroke detection compared with the traditional adult b-value of 1000 s/mm$^2$.

**Diffusion-Weighted Image Contrast and ADC Calculation**

In equation 2, ADC$_i$ is the ADC of water along the direction of the diffusion-sensitizing gradient. Higher ADC values, reflecting greater rates of water diffusion in the tissue under investigation, result in lower signal intensity $S_i$ in the diffusion-weighted image (Fig 2). Conversely, reduced ADC values result in higher observed signal intensity on DWI. The direct calculation of ADC values has assumed considerable clinical importance because of the T2-weighting inherent in the long TE of a DWI pulse sequence:

$$S_i = k \cdot e^{-\frac{TE}{T2} \cdot e^{-b \cdot \text{ADC}_i}},$$

where $k$ is a proportionality constant and $T2$ is the transverse relaxation time of the interrogated tissue. Hence, T2 prolongation within pathologic tissues may cause DWI signal hyperintensity, even in the absence of reduced ADC values. This T2 shinethrough effect renders DWI hyperintensity less specific for clinical diagnosis than decreases of ADC that are confirmed on an ADC map (Fig 3). Conversely, decreases in T2 may mask a true reduction in ADC values; this phenomenon has been referred to as “T2 blackout”. 43 T2 blackout may be at least in part responsible for the relatively low sensitivity of DWI signal intensity compared with ADC maps for the detection of acute hypoxic-ischemic injury in neonatal white matter because such lesions typically reduce the very long T2 relaxation times of white matter in neonates. However, T2 effects can also be beneficial: The “lightbulb” brightness of DWI signal intensity in acute infarcts of the adult brain is caused by the multiplicative effect of T2 prolongation and reduced ADC (equation 4). Therefore, DWI has better sensitivity than ADC maps for small acute ischemic lesions in children beyond infancy and in adults.

ADC values may be calculated from DWI by solving for ADC in equation 2:

$$\text{ADC}_i = -\ln(S_i/S_0)/b,$$

where $\ln$ is the natural logarithm. The signal intensity $S_0$ in the absence of a diffusion-sensitizing gradient can be derived from an image acquired at $b = 0$ s/mm$^2$. In general, ADC maps can be derived from a set of diffusion-weighted images acquired at a high $b$ factor and a second set of images acquired with a low $b$ factor, which can be zero. This only applies in the range of $b$-values used clinically, because for $b$-values greatly exceeding 1000 s/mm$^2$, DWI signal intensity $S_i$ no longer conforms to the monoexponential relationship with $b$ described in equation 2. 44,45

Another popular way of examining diffusion images without the contaminating influence of T2 effects is with an attenuation coefficient (AC) map, also known as the exponential diffusion-weighted image, 46 the exponential image, or the attenuation factor map (Fig 2). The AC map can be computed by dividing the diffusion-weighted image $S_i$ by the T2-weighted $b = 0$ s/mm$^2$ image $S_0$. This yields

$$\text{AC}_i = S_i/S_0 = e^{-b \cdot \text{ADC}_i},$$

Note from the last expression, compared with equation 4, that only the diffusion-related exponential term remains, eliminating the T2-weighting effects. Like DWIs, areas of reduced diffusion appear bright on AC maps and areas of elevated diffusion appear dark. This is the opposite of the contrast polarity of ADC images. However, like ADC maps, AC maps remain less sensitive than DWI signal hyperintensity for
detecting small acute infarcts because they lack the multiplicative effect of T2 prolongation combined with reduced diffusion that produces the “lightbulb” effect on DWI.

Isotropy, Anisotropy, and Rotational Invariance
In isotropic diffusion, molecular motion is equal in all directions. Pure water at body temperature demonstrates isotropic diffusion, with an ADC of approximately 3.0 × 10⁻³ mm²/s. In the human brain, isotropic diffusion may be found in the CSF spaces, except in areas of bulk flow such as the jets emanating from the foramina of Monro or through the cerebral aqueduct. Due to its complex microarchitecture, the gray matter of the cerebral cortex in adults is also thought to exhibit isotropic diffusion at the clinical range of b-values, though some studies differ. When diffusion is isotropic, the choice of direction for the diffusion-sensitizing gradient is not important because ADCᵢ is identical for all directions i.

In anisotropic diffusion, molecular mobility is not equal for all directions. White matter tracts with tightly packed coherently oriented fiber bundles hinder water displacement perpendicular to the direction of the fibers, resulting in larger ADCᵢ values parallel to the tracts rather than orthogonal to them. Hence, more than 1 diffusion-encoding direction is required to characterize regions of anisotropic diffusion. If only a single diffusion direction was probed, interpretation of the DWIs would be complicated by variable signal intensity in white matter tracts, depending on their orientation relative to the direction of the diffusion gradient, which could also be affected by changes in the orientation of the patient’s head. Although this anisotropy can be exploited in DTI, to be discussed in detail below, it is not desirable for routine clinical DWI. To avoid this problem, one can compute parameters with the property of rotational invariance from the original DWI data because these measures mathematically eliminate any directional dependence that might confuse clinical interpretation.

Computation of commonly used rotationally invariant measures, such as the geometric mean DWI and the trace ADC, requires at least 4 separate image acquisitions: 1 without diffusion-sensitizing gradients (b = 0 s/mm²), labeled S₀, and 3 probing motion along 3 mutually orthogonal directions, labeled S₁, S₂, and S₃ respectively. The DWIs submitted to the radiologist for interpretation are not typically the sets of unidirectional DWIs S₁, S₂, and S₃ but rather the rotationally invariant geometric mean computed from these 3 image sets, also known as the “isotropic DWI” or the “combined DWI”:

\[
S_{\text{DWI}} = \left( S₁ \cdot S₂ \cdot S₃ \right)^{1/3} = S₀ \cdot e^{-b ∙ \frac{(ADC₁ + ADC₂ + ADC₃)}{3}} = S₀ \cdot e^{-b ∙ \frac{ADC}{3}}
\]

where ADC₁, ADC₂, and ADC₃ are the ADCs along the directions of the 3 diffusion-sensitizing gradients and, in the final expression, ADC = (ADC₁ + ADC₂ + ADC₃)/3. This last variable, ADC, is the average of the ADC values along the 3 orthogonal directions and is also rotationally invariant. It is commonly known as the mean diffusivity, the trace ADC, Dₑ, or, simply, the ADC. Henceforth in this discussion, as is typical in the clinical setting when the acronym ADC is used, it will refer to this averaged value.
Isotropic DWI and trace ADC maps can be constructed from more than 3 diffusion-encoding directions. However, it is preferable that the diffusion-encoding directions not be collinear (ie, pointing in opposite directions along the same axis) and that they be as widely distributed in 3D space as possible. This latter condition is satisfied by mutual orthogonality in the case of 3 diffusion-encoding directions. Diffusion along 1 direction cannot be distinguished from diffusion in the polar opposite direction by Stejskal-Tanner diffusion encoding, a property known as “antipodal symmetry”. Therefore, 2 collinear diffusion-sensitizing gradients do not contribute independent information; hence, collinearity should be avoided in DWI acquisitions.

**ADC Mapping of Healthy Human Brain and of Cerebral Pathology**

In the healthy adult brain at the clinical range of b-values (<1500 s/mm²), the ADC of gray and white matter is very similar, such that DWI has little intrinsic gray-white matter contrast. Any gray-white contrast observed is primarily due to underlying differences in T2 (ie, the contrast of the b = 0 s/mm² component of the image). Measured mean diffusivities in adult brain in vivo are 0.67–0.83 × 10⁻³ mm²/s for gray matter and 0.64–0.71 × 10⁻³ mm²/s for white matter. This differs from neonates and children, in whom ADC is initially very high at term gestational age, approximately 50% greater in rapidly developing gray matter structures such as thalamus and basal ganglia, and more than twice as high in slowly maturing lobar white matter regions. Normal ADC values drop steeply during the first 2 years of postnatal life, during which time the initially higher ADC in most white matter becomes more equal to that in gray matter. ADC values continue to decrease more gradually throughout childhood, adolescence, and young adulthood, with variable rates of decline that reflect heterogeneity in the speed of maturation in different parts of the brain. Conversely, in the healthy aging brain, mild increases in average brain ADC values may be seen with advancing age beyond 40 years, especially in white matter.

Most types of pathology in the human brain cause increases of ADC values. Causes of abnormally reduced ADC in the human brain are less numerous, often allowing a more specific diagnosis. The most common ones are acute cerebral ischemia and other causes of cytotoxic edema, such as status epilepticus and hypoglycemia. The conventional explanation for reduced ADC in acute ischemia is that energetic failure leads to membrane permeability changes, resulting in cellular swelling and decreased volume and increased tortuosity of the extracellular space. However, this remains controversial because biophysical evidence has been presented that extracellular ADC and intracellular ADC are both reduced in acute cerebral ischemia, with the major contribution arising from the intracellular space.

The viscosity of purulent material also reduces diffusion below that of the surrounding brain or CSF spaces. In vitro measurements of ADC values in pus indicate that viable cell attenuation is the most important determinant of hindered diffusion in pus. Hence, infectious etiologies for intracranially reduced diffusion include pyogenic abscess, subdural or epidural empyema, and intraventricular empyema. Similarly, low ADC values due to hindered water motion also produce very bright signal intensity on DWI within extra-axial epidermoid masses and within acute clotted hematomas, except in T2 dark hematomas due to T2 blackout. Highly cellular tumors, especially those with high nuclear-to-cytoplasmic ratios, can show reduced diffusion. Examples include the entire class of “small round blue cell tumors” such as lymphoma, neuroblastoma, and primitive neuroectodermal tumors such as medulloblastoma. There have been case reports of brain metastases with restricted diffusion and basal ganglia, and more than twice as high in slowly maturing lobar white matter regions.

Fig 3. T2 shinethrough artifact in DWI. A, T2-weighted fluid-attenuated inversion recovery image shows numerous nonspecific periventricular and subcortical hyperintense foci as well as a larger wedge-shaped region of hyperintensity in the right parietal lobe suggestive of infarct. B, Combined DWI image at the same axial level reveals that the right parietal abnormality is hyperintense, suggesting acute ischemia. C, Corresponding ADC map demonstrates elevated mean diffusivity within the abnormal region, consistent with chronic infarction. D, Low signal intensity within the lesion on the attenuation coefficient image confirms that the DWI signal-intensity hyperintensity is due to T2 shinethrough and does not reflect reduced diffusion as would be seen in an acute infarct.
sion, even those without intralosomal hemorrhage. Reduced diffusion has been described in the acute phase of primary demyelinating diseases, metabolic disorders, and toxic exposures. Rare causes of reduced ADC include Creutzfeldt-Jakob disease.

The Diffusion Tensor

As discussed previously, variations in diffusion-weighted signal intensity due to the anisotropy of white matter tracts can be troublesome for the interpretation of clinical DWI, unless rotationally invariant geometric mean DWI and trace ADC maps are computed. However, the anisotropic diffusion of coherently oriented axonal fibers can also be exploited for quantitative characterization and anatomic mapping of white matter tracts. If at least 6 diffusion-encoded image sets are acquired along noncollinear directions, in addition to at least one \( b = 0 \, \text{s/mm}^2 \) (or low \( b \)) image set, the diffusion tensor can be calculated.

The diffusion tensor, a 3 \( \times \) 3 matrix of vectors, is a mathematical model of the 3D pattern of diffusion anisotropy of white matter tracts. The tensor, \( D \), can be related to the diffusion-weighted signal intensity, \( S \), and the \( b = 0 \, \text{s/mm}^2 \) signal intensity, \( S_0 \), as follows:

\[
\frac{S}{S_0} = e^{-\left( \sum_{i=x,y,z} \sum_{j=x,y,z} b_{ij} \rho_{ij} \right)}, \quad \text{where} \quad b_{ij} = \gamma^2 G_i G_j \left[ \delta^2 \left( \Delta - \delta \right) \right].
\]

In this formalism, the diffusion-weighting factor, \( b \), incorporates the direction and magnitude of the applied diffusion gradient vector \((G_x, G_y, G_z)\). Six or more measurements of \( S \) by using noncollinear diffusion gradients are necessary to estimate \( D \) by using linear regression methods.

The most intuitive way to conceptualize the information provided by the diffusion tensor is to view it geometrically (Fig 4). The tensor effectively fits the angular variation of the ADC values to the shape of a 3D ellipsoid. The diffusion ellipsoid is a shape defined by 6 variables that describes the ADC of water molecules in each direction at a particular time. For isotropic diffusion, the diffusion ellipsoid is a sphere, because the ADC in every direction is equal. Anisotropic diffusion is modeled with an elongated ellipsoid, indicating a greater mean diffusion distance along the longest axis of the ellipsoid. The elements of the tensor above the diagonal are always equal to those below the diagonal (Fig 4, bottom row), a characteristic termed “conjugate symmetry”, which reflects the antipodal symmetry of Brownian motion. Thus, there are only 6 independent elements of the tensor, necessitating a minimum of 6 noncollinear diffusion-encoding measurements to solve for the tensor elements. The diagonal terms of the tensor indicate the magnitude of diffusivity in each of 3 orthogonal directions. In the case of anisotropic diffusion, the off-diagonal terms of the diffusion tensor indicate the magnitude of diffusion along 1 direction arising from a concentration gradient in an orthogonal direction.

**Diffusion Tensor Parameters**

A number of diffusion tensor metrics are used to characterize the dimensions and shape of the diffusion ellipsoid associated with the microstructure of a particular voxel. The 3 principal axes of the diffusion tensor, termed the “eigenvectors,” can be calculated by diagonalizing the diffusion tensor:

\[
\Lambda = \begin{bmatrix}
\lambda_1 & 0 & 0 \\
0 & \lambda_2 & 0 \\
0 & 0 & \lambda_3
\end{bmatrix} = R \cdot D \cdot R^\top.
\]

The diffusion tensor is rotated by matrix \( R \) to produce the diagonal matrix, \( \Lambda \). The columns of \( R \) are composed of the eigenvectors of the system, and \( R^\top \) is the matrix transpose of \( R \). The 3 eigenvectors and rotationally invariant eigenvalues \( \lambda_1, \lambda_2, \lambda_3 \) describe the directions and lengths of the 3 diffusion ellipsoid axes, respectively, in descending order of magnitude (Fig 5, top row). The largest eigenvector, termed the “primary eigenvector,” and its associated eigenvalue \( \lambda_1 \), indicate, respectively, the direction and magnitude of greatest water diffusion. The primary eigenvector is important for fiber tractography algorithms because this vector indicates the orientation of axonal fiber bundles. Therefore, \( \lambda_1 \) is also termed “longitudinal diffusivity,” because it specifies the rate of diffusion along the orientation of the fibers. The second and third eigenvectors are orthogonal to the primary eigenvector, and their associated eigenvalues \( \lambda_2 \) and \( \lambda_3 \) give the magnitude of diffusion in the plane transverse to axonal bundles. Hence, the mean of \( \lambda_2 \) and \( \lambda_3 \) is also known as “radial diffusivity”.

A number of additional rotationally invariant diffusion metrics, derived from the 3 eigenvalues, are commonly used to describe the microstructure in a particular voxel. The mean
Diffusivity \( (D_{av}) \) is the mean of the 3 eigenvalues and describes the directionally averaged diffusivity of water within a voxel:

\[
D_{av} = \frac{\lambda_1 + \lambda_2 + \lambda_3}{3} = \text{trace} (D)/3. \tag{10}
\]

The trace of \( D \) is the sum of the 3 eigenvalues and is another measure of orientationally averaged diffusion. The fractional anisotropy (FA) and relative anisotropy (RA) indices measure the degree of directionality of intravoxel diffusion:

\[
FA = \frac{\sqrt{(\lambda_1 - \lambda_2)^2 + (\lambda_2 - \lambda_3)^2 + (\lambda_3 - \lambda_1)^2}}{\sqrt{2} \sqrt{\lambda_1^2 + \lambda_2^2 + \lambda_3^2}} \tag{11}
\]

\[
RA = \frac{\sqrt{(\lambda_1 - \lambda_2)^2 + (\lambda_2 - \lambda_3)^2 + (\lambda_3 - \lambda_1)^2}}{\lambda_1 + \lambda_2 + \lambda_3} \tag{12}
\]

FA and RA can both also be defined by equivalent formulas where the numerators are expressed in terms of the difference of each eigenvalue from the mean diffusivity:

\[
FA = \frac{\sqrt{3} \sqrt{(\lambda_1 - D_{av})^2 + (\lambda_2 - D_{av})^2 + (\lambda_3 - D_{av})^2}}{\sqrt{\lambda_1^2 + \lambda_2^2 + \lambda_3^2}} \tag{13}
\]

\[
RA = \frac{\sqrt{3D_{av}}}{\sqrt{3D_{av}}} \frac{\sqrt{(\lambda_1 - D_{av})^2 + (\lambda_2 - D_{av})^2 + (\lambda_3 - D_{av})^2}}{\sqrt{3D_{av}}} \tag{14}
\]

When the primary eigenvalue is much larger than the second and third eigenvalues, anisotropy measures such as FA and RA will be high, indicating a preferred direction of diffusion. This corresponds to a prolate (cigar) shape of the diffusion ellipsoid (Fig 4, right column), with the preferred direction of diffusion indicated by the long axis of the ellipsoid or, equivalently, by the primary eigenvector of the tensor. In the brain, prolate diffusion within a voxel usually indicates a single coherently organized white matter fiber bundle, with the axonal orientation coincident with the primary eigenvector. FA has become by far the most widely used measure of anisotropy in the DTI literature, and its value varies from zero, in the case of isotropic diffusion, up to a maximum of 1, indicating perfectly linear diffusion occurring only along the primary eigenvector. The fiber orientation information inherent in the primary eigenvector can be visualized on 2D images by assigning a color to each of 3 mutually orthogonal axes, typically red to left-right, green to anteroposterior, and blue to up-down (Fig 5, bottom right).

**DTI Fiber Tracking: Methodology**

The objective of DTI fiber tracking is to determine intervoxel connectivity on the basis of the anisotropic diffusion of water.\(^ {34, 56, 83, 86} \) The clinical and scientific utility of DTI fiber tracking is found in both the localization and the quantitative assessment of specific neuronal pathways, as applied to basic neuroscience,\(^ {87} \) cognitive neuroscience,\(^ {88, 89} \) and diagnostic neuroradiology.\(^ {90-92} \) In each brain voxel, the dominant direction of axonal tracts can be assumed to be parallel to the primary eigenvector of the diffusion tensor. Fiber tracking uses the diffusion tensor of each voxel to follow an axonal tract in 3D from voxel to voxel through the human brain. Because DTI provides only microstructural information at relatively low spatial resolution, DTI fiber tracking is often combined with functional and/or higher resolution anatomic information to delineate specific pathways.\(^ {34, 91, 93} \) In this way, 3D DTI tractog-
raphy has opened up a whole new dimension to the ability to depict human neuroanatomy noninvasively. This new anatomic connectivity information has been summarized in review articles, as well as presented in a book-length DTI atlas.

DTI fiber tracking algorithms can be divided into deterministic and probabilistic methods. Fiber assignment by continuous tracking (FACT) is a deterministic method, which initiates fiber trajectories from user-defined voxels (Fig 6). Fiber trajectories, also known as “streamlines”, follow the primary eigenvector from voxel to voxel in 3 dimensions. When the fiber trajectory reaches the edge of the voxel, the direction of the trajectory is changed to match the primary eigenvector of the next voxel. Constraints on the maximum turning angle of the streamline between voxels and on the minimum FA within a voxel for propagation of the streamline can be applied to contain the fiber tracks to regions of the brain where the diffusion tensor model realistically represents the white matter pathways. Figure 7 shows how user-defined regions of interest, based on prior anatomic knowledge, can be used to restrict fiber tracks to the corticospinal tract. This multiple-region-of-interest technique for isolating anatomically specific fiber pathways by using DTI tractography has been called “virtual dissection.”

Noise, patient movement, and distortion from imaging artifacts produce uncertainty in the orientation of the diffusion ellipsoid and are detrimental to deterministic streamline fiber tracking. Probabilistic fiber tracking methods incorporate the expected uncertainty into the tracking algorithm and can be used to produce a connectivity metric for each voxel. The probability density function of the orientation of a neuronal fiber can be estimated with an empiric function based on the FA, or a Bayesian model, or bootstrap statistics. Probabilistic DTI fiber tracking techniques tend to disperse trajectories more than deterministic methods and have the potential to delineate a greater portion of a white matter tract. However, the accuracy of these probabilistic methods is still limited by the information contained in the diffusion tensor and the method of constructing the probability density function. Another fundamental limitation of diffusion tractography is that it cannot distinguish antegrade from retrograde along a fiber pathway, due to the antipodal symmetry inherent to the diffusion process.

**DTI Fiber Tracking: Clinical Applications**

The visualization of an entire white matter tract in 3D has clinical and scientific value for detecting gross changes to the anatomic course and the microstructural integrity of specific pathways. DTI has recently been used to identify and characterize novel defects in axonal growth and guidance in developmental disorders, such as the absence of normally decussating pontocerebellar fibers in horizontal gaze palsy with progressive scoliosis due to mutation of the ROBO3 gene and the ectopic location of pontocerebellar fibers in pontine tegmental cap dysplasia. An aberrant fiber pathway known as the “asymmetric sigmoid bundle” was first identified with DTI fiber tracking in individuals with dysgenesis of the corpus callosum. Brain tumors and other intracranial mass lesions are capable of dramatically altering the position of the corticospinal tract; hence, knowledge of the location of this important tract within deep white matter is critical for resecting mass lesions without postoperative motor deficit (Fig 8). DTI fiber tracks can be used in conjunction with stereotactic navigation to identify a safety margin of approximately 1 cm around the motor tract, which can be used for surgical planning.

In addition to basic 3D visualization, many studies have used fiber tracking to delineate specific white matter tracts for quantitative analysis. Quantitative DTI tractography studies have examined the microstructure of white matter tracts in pediatric subjects, in elderly subjects, and in patients with schizophrenia, brain tumors, Alzheimer disease, and many other disorders. DTI fiber tracking can objectively create 3D regions of interest specific to an entire white matter tract. However, most white matter tracts have heterogeneous structural characteristics, and fiber tracking can be used to segment a pathway into several regions for quantitative analysis. Figure 9 shows the optic radiations in a premature infant color-coded according to the underlying FA values in the voxels contained within the tracts. The anterior portion of the optic radiation adjacent to the thalamus was observed to have significantly higher FA than the posterior portions of the tract. The spatial and temporal heterogeneity of DTI metrics has also been observed with fiber tracking of sensorimotor tracts in premature infants, indicating the maturation of the delineated tract and the emergence of other tracts crossing the sensorimotor pathway in the centrum semiovale.

**Conclusion**

This review article has explored the theoretic background needed to understand clinical DWI and DTI, including fiber tractography, and their application to neuroradiology. These diffusion MR imaging techniques provide microstructural information about biologic tissues that is not available from other imaging techniques. In the central nervous system, this has yielded important new tools for diagnosis in ischemia, infection, tumors, and demyelinating disease, among other pathologies, as well as for presurgical mapping of white matter pathways to avoid postoperative injury.
However, even as sophisticated a mathematic construct as the diffusion tensor is an oversimplification of the properties of water diffusion in the brain. The limitations of the diffusion tensor in areas of complex white matter architecture, where fiber tracts intersect, branch, or are otherwise partial volume averaged within a voxel, affect the ability of DTI fiber tractography to fully delineate an axonal pathway and may also lead to the generation of spurious tracks.\textsuperscript{95} Measurements of quantitative DTI parameters such as FA are also difficult or impossible to interpret in regions of complex white matter. These problems have led to the introduction of more advanced methods such as probabilistic tractography\textsuperscript{126-128} and whole-brain connectivity networks.\textsuperscript{129} Hence, these newer techniques that transcend the diffusion tensor are likely to be adopted in clinical neuroradiology during the years to come.

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