SHORT REPORT

Diffusion tensor imaging can detect and quantify corticospinal tract degeneration after stroke

David J Werring, Ahmed T Toosy, Christopher A Clark, Geoffrey JM Parker, Gareth J Barker, David H Miller, Alan J Thompson

Abstract

Diffusion tensor imaging (DTI) fully characterises water molecule mobility in vivo, allowing an exploration of fibre tract integrity and orientation in the human brain. Using DTI this study demonstrates reduced fibre coherence (anisotropy) associated with cerebral infarction and in the corticospinal tract remote from the lesion, in five patients 2 to 6 months after ischaemic stroke. The study highlights the potential of DTI to detect and monitor the structural degeneration of fibre pathways, which may provide a better understanding of the pattern of clinical evolution after stroke.

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Keywords: magnetic resonance imaging; diffusion tensor; stroke; pathophysiology

Wallerian degeneration—the anterograde degeneration of axons and their myelin sheaths after proximal axonal or cell body injury—commonly follows CNS pathology. Some studies using MRI suggest that the presence or absence of wallerian degeneration may influence the clinical outcome after stroke.1 However, conventional MRI is only abnormal in a proportion of cases,2 and cannot easily quantify the extent or severity of fibre damage. Newer MRI methods that are sensitive to water diffusion can be used to study aspects of tissue structure and orientation not accessible to conventional MRI. Magnetic resonance diffusion imaging has recently generated considerable interest due to an ability to disclose early pathophysiological changes in acute stroke,3 but less attention has been paid to structural changes in fibre tracts beyond the area of infarction.

It is generally accepted that a full description of water molecule diffusion requires the determination of the diffusion tensor, a nine element mathematical matrix quantity from which indices describing the magnitude (diffusivity) and directional bias (anisotropy) of water molecule motion may be derived.4 It is possible to estimate the diffusion tensor by measuring changes in the magnetic resonance signal collected with diffusion sensitising gradients applied along at least six non-collinear directions.4 In brain regions where the diffusion of water varies significantly with direction, the tissue exhibits the property of anisotropy; by contrast, regions where diffusion is similar in all directions have low anisotropy (and are said to be isotropic). In an axon cylinder water diffusion is faster along the axon than across it, probably due to the presence of directionally subcellular structures including the axonal membrane and the neurofilamentary cytoskeleton which behave as barriers to diffusion.5 As intact fibre tracts are composed of collections of similarly oriented axons, they generally exhibit high anisotropy.6 Anisotropy measurement therefore shows promise in detecting the degree of fibre damage in diseases affecting white matter.7 Furthermore, diffusion within each volume of tissue (voxel) may be most fully and naturally described by three mutually perpendicular eigenvectors, the magnitudes of which are given by three corresponding eigenvalues; the eigenvalues are the three principal diffusion coefficients measured along the three (intrinsic) eigenvector directions which define the local “fibre” frame of reference for that voxel.8 The direction of fibres is thus given by the eigenvector of the largest eigenvalue of the diffusion tensor, allowing in vivo maps of white matter fibre trajectories to be constructed.9 We used DTI to investigate structural changes (including in diffusivity, anisotropy, and fibre direction) within the infarction and associated descending corticospinal tracts in five patients after stroke.

Methods

We used DTI (echo planar imaging (EPI) TE=78 ms, 96×96 matrix, four diffusion b values increasing from 0 to~700 smm⁻² applied in seven non-collinear directions) and conventional T2 weighted MRI (TR=2000, TE=120, matrix 256×256) to study five patients (three men, two women; mean age 58 years) with resolving hemiplegia 2 to 6 months after middle cerebral artery territory infarction (four right sided; one left sided), and five normal controls (three men, two women; mean age 37 years). All patients gave written informed consent to participate in the study, which had been approved by the joint ethics committee of the Institute of Neurology and the National Hospital of Neurology and Neurosurgery.
Diffusion tensor imaging of a 45 year old man 2 months after right middle cerebral artery territory stroke. (A), (B), and (C) show different axial levels through the brain (left side of figure). The coronal images (right side of figure) provide direct visualisation of the descending corticospinal tracts; the horizontal yellow lines show the locations of the axial slices. The grey scale images are maps of fractional anisotropy (FA), one of the anisotropy indices derived from DTI. Overlaid on these are the principal eigenvectors (red lines), which depict the local fibre direction at each voxel in the image. For greater clarity, the eigenvectors have been given a threshold so that only those in regions of anisotropy >0.5 are displayed. The large open arrow in (A) indicates the infarct, which has reduced anisotropy (dark pixels). The small closed white arrows in (A), (B), and (C) indicate the location of the damaged corticospinal tract in which anisotropy is lower than on the opposite side. In the coronal images (right side of figure) the intact corticospinal tract on the left side of the brain has high anisotropy (bright pixels) and coherent fibre orientation (red lines). This is in contrast with the loss of tract anisotropy and direction on the right side of the brain ipsilateral to the infarct. In all images the right side of the brain is to the left of the panel.
Mean values (SD) of fractional anisotropy (FA), a measure of the directionality of water diffusion (dimensionless units); and mean diffusivity (MD), a measure of the amount of water diffusion (×10−6 mm2/s), in the infarct region in patients, and at different corticospinal tract levels in patients and controls.

<table>
<thead>
<tr>
<th>Patient group (n=5)</th>
<th>Control group (n=5)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Right FA</td>
</tr>
<tr>
<td>Infarct</td>
<td>0.29 (0.08)*</td>
</tr>
<tr>
<td>Internal capsule</td>
<td>0.65 (0.09)*</td>
</tr>
<tr>
<td>Cerebral peduncles</td>
<td>0.66 (0.08)*</td>
</tr>
<tr>
<td>Pons</td>
<td>0.61 (0.12)*</td>
</tr>
<tr>
<td>All corticospinal tract regions</td>
<td>0.61 (0.13)*</td>
</tr>
</tbody>
</table>

*Right and left sides show a statistical difference at p<0.01, Mann-Whitney U test.

**Discussion**

Our results have shown reduced FA with preservation of MD in the corticospinal tract distal to cerebral infarction. This pattern of diffusion changes is similar to that found in experimental animal models of wallerian degeneration in the peripheral nervous system. Wallerian degeneration involves breakdown of the myelin sheath and disintegration of axonal microfilaments. Loss of integrity of these directional structures is consistent with our finding of reduced anisotropy in the corticospinal tract ipsilateral to cerebral infarction. Although disruption of myelin and axons might be expected to increase the MD of water molecules, an accumulation of cellular debris from the breakdown of axons may hinder water molecule motion. Furthermore, wallerian degeneration is characterised histologically by glial proliferation, which would also be expected to restrict the mobility of water molecules. Both of these factors may contribute to the relative preservation of MD in the affected pathways. Replacement of the intact anisotropic microstructure by disorganised glial proliferation may also underlie the marked reduction in FA we have demonstrated in affected corticospinal tracts.

The primary infarct showed a different pattern of DTI abnormalities from those found in the associated corticospinal tract. Large MD increases were seen in all patients compared with matched contralateral brain regions (group mean 1.75 v 0.91, p<0.001), consistent with previous reports of increased water diffusion after the acute phase (>10 days) of cerebral infarction. Moreover, anisotropy was markedly reduced in the infarct by comparison with contralateral regions (FA group mean 0.29 v 0.48, p<0.001), a finding to our knowledge not previously reported in human subjects but in keeping with experimental data in animals after transient cerebral hypoxia-ischaemia showing reduced white matter anisotropy. In a longstanding cerebral infarct it is thought that cell lysis and loss of normal tissue architecture expand the extracellular space, allowing water molecules to diffuse more freely. These changes would account for the increased diffusivity and reduced anisotropy that we have demonstrated in regions of infarction.

The difference in diffusion properties between the primary lesion and the degenerated tract (reduced FA with increased MD in the infarct; reduced FA with preserved MD in the corticospinal tract) may allow DTI to distin-
guish between the primary lesion and associated wallerian degeneration (not possible on conventional MRI). A second advantage of DTI is the ability to generate eigenvector maps, which provide a striking visual comparison between the trajectories of the affected and the intact corticospinal tracts (figure). Finally, the finding of minor T2 weighted signal change in only one patient, by contrast with DTI abnormalities in all five cases indicates the higher sensitivity for DTI to detect wallerian degeneration.

This preliminary study indicates that DTI can visualise and quantify changes in the integrity and orientation of white matter tracts transected by a focal ischaemic lesion that are consistent with wallerian degeneration. Our findings highlight the potential of DTI to detect and monitor the structural changes in cerebral infarction and associated degeneration of fibre pathways; longitudinal DTI studies may provide a better understanding of the pattern of clinical evolution after stroke.

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