Diffusion kurtosis imaging of microstructural changes in brain tissue affected by acute ischemic stroke in different locations

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Abstract
The location of an acute ischemic stroke is associated with its prognosis. The widely used Gaussian model-based parameter, apparent diffusion coefficient (ADC), cannot reveal microstructural changes in different locations or the degree of infarction. This prospective observational study was reviewed and approved by the Institutional Review Board of Xiamen Second Hospital, China (approval No. 2014002). Diffusion kurtosis imaging (DKI) was used to detect 199 lesions in 156 patients with acute ischemic stroke (61 males and 95 females), mean age 63.15 ± 12.34 years. A total of 199 lesions were located in the periventricular white matter (n = 52), corpus callosum (n = 14), cerebellum (n = 29), basal ganglia and thalamus (n = 21), brainstem (n = 21) and gray-white matter junctions (n = 62). Percentage changes of apparent diffusion coefficient (ΔADC) and DKI-derived indices (fractional anisotropy [ΔFA], mean diffusivity [ΔMD], axial diffusivity [ΔDa], radial diffusivity ΔDr, mean kurtosis [ΔMK], axial kurtosis [ΔKa], and radial kurtosis [ΔKr]) of each lesion were computed relative to the normal contralateral region. The results showed that (1) there was no significant difference in ΔADC, ΔMD, ΔDa or ΔDr among almost all locations. (2) There was significant difference in ΔMK among almost all locations (except basal ganglia and thalamus vs. brainstem; basal ganglia and thalamus vs. gray-white matter junctions; and brainstem vs. gray-white matter junctions. (3) The degree of change in diffusional kurtosis in descending order was as follows: corpus callosum > periventricular white matter > brainstem > gray-white matter junctions > basal ganglia and thalamus > cerebellum. In conclusion, DKI could reveal the differences in microstructure changes among various locations affected by acute ischemic stroke, and performed better than diffusivity among all groups.

Key Words: nerve regeneration; apparent diffusion coefficient; diffusion weighted imaging; diffusion kurtosis imaging; acute ischemic stroke; mean kurtosis; microstructure changes; white matter; 1.5 Tesla magnetic resonance system; neural regeneration

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Introduction
Stroke, which has become the second leading cause of death among the global population aged > 60 years (Johnston et al., 2009; Thrift et al., 2014), has large impacts on patients and their families (Chen, 2008; Zhao et al., 2010; Liu et al., 2011). The incidence of ischemic stroke, a common form accounting for 43–79% of all strokes (Jia et al., 2010), is rising. Acute ischemic stroke, which is often fatal, is caused by thrombotic or embolic occlusion of a cerebral artery. Ischemia ensues, resulting in loss of neurological function (Writing Group Members et al., 2016; Khatri et al., 2018). Thrombolytic therapy is the treatment of choice for acute ischemic stroke if caught early enough (Yang et al., 2014; Liu and Zhang, 2017; Li et al., 2018b). The therapy is always based on the onset time reported by the patient or onlookers; however, few recognized radiological techniques can reveal whether there is recoverable tissue. (Domingues-Montanari et al., 2008). If the therapy timing is inappropriate late, complications such as cerebral hemorrhage and reperfusion injury may exacerbate tissue damage (Khatri et al., 2012; Kanazawa et al., 2017).

Studies have indicated that magnetic resonance imaging (MRI), especially diffusion imaging, has great potential in this field (Weber et al., 2015; Khalil et al., 2016; Piliszek et al., 2016; Li et al., 2018a). Diffusion weighted imaging (DWI) and its metric, the apparent diffusion coefficient (ADC), have played an important role in diagnosis and assessment of stroke since the 1990s (Sevick et al., 1990; Warach et al., 1992). Diffusion tensor imaging (DTI), a special kind of DWI based on the two-order 3D tensor model, is widely used to evaluate white matter changes in the brain (Basser et al., 1994; Auriel et al., 2014; Li et al., 2016). Both DWI and DTI are based on the fact that water molecules diffuse in a free and unrestricted environment (Basser and Jones, 2002). In biological tissues, an extremely complicated environment and structure affects water molecule diffusion, such that the diffusion displacement distribution occurs in a non-Gaussian form (Tuch et al., 2003). Several technologies, such as Q-space imaging (Assaf et al., 2002), diffusion spectrum imaging (Wedeen et al., 2005), and diffusion kurtosis imaging (DKI) (Jensen et al., 2005), have been proposed to characterize non-Gaussian diffusion patterns. Compared with similar approaches, DKI has the feature of being the least inconvenient in terms of scanning time and hardware demands.

DKI, proposed by Jensen et al. (2005), is a straightforward extension of diffusion tensor imaging. It uses a 2nd-order diffusivity tensor together with a 4th-order kurtosis tensor to provide an enhanced depiction of non-Gaussian water diffusion. Fieremans et al. (2011) and Hui et al. (2008) reported that DKI can detect subtle changes occurring in tissues affected by tumor or inflammation, and believed that DKI had great potential in the assessment of stroke lesions. Cheung et al. (2012) evaluated ischemic brain lesions using an animal model, and the results showed that DKI has great advantages in characterizing ischemic stroke. Whereas Rudrapatna et al. (2014) found that DKI could detect structural tissue changes and sensitively reflected angiogenesis and functional reconstruction around ischemic areas after the stroke occurred. The aim of our study was to evaluate the performance of DKI in detecting differences in microstructural changes at different locations of the brain after acute ischemic stroke.

Subjects and Methods
Subjects and design
This prospective observational study was reviewed and approved by the Institutional Review Board of Xiamen Second Hospital, China (approval No. 2014002). Written informed consent was obtained from all subjects before MRI. Inclusion criteria for subjects were as follows:

- Clinical symptoms, onset duration and neurological deficit symptoms of each subject were highly consistent with the diagnosis of acute ischemic stroke (Chung et al., 2014), and subjects did not undergo clinical treatment.
- This was the first onset for each subject.
- Cerebral hemorrhage or brain neoplasm were absent by brain computerized tomography (CT).
- There were no contraindications to MRI.

The exclusion criteria were:

- Patients could not finish the scanning.
- High signal intensity on DWI mapping and corresponding low signal on ADC mapping were not found.
- Maximum measurable lesion area was < 50 mm².
- There were artifacts that influenced the measurement.

One hundred sixty-eight patients (67 females and 101 males) aged 66.62 ± 14.88 years, with acute ischemic stroke, underwent routine MRI and DKI scans (b = 0, 1000, 2000 s/mm², 15 diffusion directions), were enrolled (Figure 1). Four patients with motion artifacts, six patients with maximum measurable lesion areas < 50 mm², and two patients with negative DWI and ADC were excluded. As a result, a total of 199 lesions were detected in the remaining 156 patients (61 females and 95 males) aged 63.15 ± 12.34 years. These lesions were divided into six groups according to commonly affected locations, including periventricular white matter (52 lesions), corpus callosum (14 lesions), cerebellum (29 lesions), basal ganglia and thalamus group (21 lesions), brainstem (21 lesions), and gray-white matter junctions (62 lesions) groups.

Data acquisition
Data acquisition was performed with a 1.5 T clinical MRI scanner (Signa HDe®, GE Medical System, Milwaukee, WI, USA) equipped with an eight-channel head and neck coil. Tight, but comfortable, foam padding was used to fix the head in place and minimize motion. Routine anatomical MRI sequences, including T1-weighted images (T1WI), T2-weighted images (T2WI), fluid attenuation inversion recovery T2WI, and DWI (b = 0, 1000 s/mm²), were performed. DKI images were acquired using a spin-echo single-shot diffusion-weighted echo planar imaging sequence along 15 encoding diffusion directions at three b values (0, 1000, and 2000 s/mm²). Scanning parameters were as follows: repetition time/echo time: 6000/162 ms; field of view: 24 cm × 24 cm; thickness/spacing: 5 mm/1.5 mm; number
of excitations: 2; acquisition matrix: 96 × 130; total acquisition time was 6 minutes and 18 seconds. All metrics were obtained using DKI software on the Functool® platform of GE ADW4.3 workstation (GE Medical System).

For reliable results, we acquired these metrics in the following ways:

(i) Two experienced radiologists (FNW and QHC with 6 and 9 years of experience in stroke MR imaging, respectively) assessed the DKI-derived maps independently,

(ii) Each lesion was outlined on the diffusion-weighted image at b = 1000,

(iii) For each lesion, a mirrored contralateral normal region of interest (ROI) was placed;

(iv) The mean changes in percentages of all metrics in lesions with reference to the normal contralateral ROIs were computed using the measurements by the above two radiologists.

Mean values of ADC and DKI-derived metrics were measured. Percentage changes of all metrics (including apparent diffusion coefficient (ΔADC), fractional anisotropy (ΔFA), mean diffusivity (ΔMD), axial diffusivity (ΔD⊥), radial diffusivity (ΔD∥), mean kurtosis (ΔMK), axial kurtosis (ΔK⊥), and radial kurtosis (ΔK∥)) were computed relative to normal contralateral ROI using the formula:

\[
\text{ΔMETRIC} = 100\% \times \frac{\text{METRIC}_{\text{affected}} - \text{METRIC}_{\text{normal}}}{\text{METRIC}_{\text{normal}}}.
\]

(1)

Theoretical model of DWI

ADC is calculated according to Eq. 2, (Le Bihan et al., 1988; Gray and MacFall, 1998), where S(b) and S(0) represent the signal intensity at b and minimum-diffusion-weighted b₀. Typically, b₀ is set to zero and Eq. 1 can be rewritten as Eq. 3.

\[
S(b) = S(b₀) e^{b₀ b - b \text{ADC}}.
\]

(2)

\[
ADC = \frac{\ln(S(b)/S(0))}{b}.
\]

(3)

Commonly, the value of b is set to 1000 s/mm² for the brain.

Theoretical model of DKI

DKI describes non-Gaussian water diffusion using a 2nd-order diffusion tensor together with a 4th-order kurtosis tensor. The diffusion-weighted signal S(b) decays with increasing b value as the following equation (Le Bihan et al., 1986; Jensen et al., 2005; Jensen and Helpern, 2010):

\[
\ln \frac{S(b)}{S(0)} = \frac{-b D_{\text{app}}}{6} + \frac{b^2 K_{\text{app}}}{12} K_{\text{app}}
\]

(4)

where \( D_{\text{app}} \) is the apparent diffusion coefficient and \( K_{\text{app}} \) is the apparent diffusional kurtosis along the special direction. \( D_{\text{app}} \) and \( K_{\text{app}} \) along each direction are obtained by fitting diffusion-weighted signal intensities into Equation 4 with the least squares method. Diffusion tensor values and kurtosis tensor values can then be estimated using \( D_{\text{app}} \) and \( K_{\text{app}} \) along all directions. FA, MD, \( D_{\text{app}} \), and \( K_{\text{app}} \) are derived from the diffusion tensor, while the diffusional kurtosis tensor is used to calculate MK, \( K_{\|} \), and \( K_{\perp} \).

The value of MD is determined by \( D_{\text{app}} \) in all directions, while MK is determined by \( K_{\text{app}} \) in all directions. For example, the standard DKI sequence typically uses 15 directions and three b values (b = 0, 1000, 2000 s/cm²) to collect data, and the calculation sketch map of MD and MK can be expressed as shown in Figure 2. Figure 3 shows the DWI and DKI maps from a normal subject.

Statistical analysis

All statistical analyses were performed using commercial software (Statistical analysis software, SPSS® Version 19.0.
for Windows; IBM SPSS, Chicago IL, USA). Mean values of all metrics were compared among different lesions. Meanwhile, Kolmogorov-Smirnov testing of ADCs and DKI-derived indices among metrics and groups were performed. Values of \( P < 0.05 \) were considered statistically significant.

## Results

### Comparison of DKI-derived metrics among six groups

Figures 4–9 show the DKI-derived indices within lesions in periventricular white matter, corpus callosum, cerebellum, basal ganglia and thalamus, brainstem, and gray-white matter junctions, respectively. As shown in Figures 4, 5, and 8, values of MD, \( D_r \), and \( D_s \) decreased substantially, while MK, \( K_r \), and \( K_s \) increased markedly compared with the normal contralateral side. MK, \( K_r \), and \( K_s \) of lesions in the cerebellum (Figure 6) and gray-white matter junctions (Figure 9) demonstrated different patterns. Notably, some lesions showed increased metrics as compared with the normal contralateral side. However, all diffusional kurtosis metrics were slightly increased in lesions of the basal ganglia and thalamus (Figure 7) compared with the normal contralateral side.

### Comparison between ADC and DKI matrices among groups

As exhibited in Figure 10, as ADC and diffusivity metrics (\( FA, MD, D_r \), and \( D_s \)) decreased, and kurtosis metrics (MK, \( K_r \), and \( K_s \)) increased in all groups. In all groups, kurtosis metrics exhibited larger changes than ADC and diffusivity metrics, which indicated their potential in distinguishing differences among groups. Absolute percentage changes along the axial direction were significantly larger than along the radial direction in all groups: \( \Delta D_r (\sim 51.01 \pm 10.77\%) \) vs. \( \Delta D_s (\sim -37.55 \pm 43.14\%) (P = 0.013) \); \( \Delta K_r (113.04 \pm 57.14\%) \) vs. \( \Delta K_s (58.63 \pm 85.36\%) (P = 0.002) \).

Table 1 illustrates that there was no significant difference in \( \Delta A D C \) between almost all group pairings (except periventricular white matter vs. cerebellum; gray-white matter junction vs. cerebellum; \( P > 0.05 \)); whereas there was a significant difference in \( \Delta M K \) between almost all group pairings (except basal ganglia and thalamus vs. brain stem; basal ganglia and thalamus vs. lobes mixes with gray and white matter; and brain stem vs. lobes mixed with gray and white matter) (\( P < 0.05 \)). Notably, \( \Delta K_s \) performed almost the same as \( \Delta M K \), and there were no significant differences in \( \Delta M D \) or \( \Delta D_r \) between groups (except periventricular white matter vs. cerebellum; periventricular white matter vs. basal ganglia and thalamus) (\( P > 0.05 \)). \( \Delta D_s \) demonstrated no statistically significant changes among all groups (\( P > 0.05 \)).

### Discussion

Several previous studies found that the prognosis of stroke is dependent upon the location (Galovic et al., 2013; Phan et al., 2013; Tang et al., 2013). Moreover, researchers have long been looking for appropriate radiological parameters to describe the degree of infarction in specified locations. Although ADC is widely used in the clinic for diagnosis of acute and hyperacute ischemic stroke (Bevers et al., 2018; Nakajo et al., 2018), it cannot reveal microstructural changes in tissue or the degree of infarction. DWI assumes that water molecules diffuse in a free and non-restricted environment (Pilszuk et al., 2016), while in tissues, water molecule diffusion is restricted by complicated microstructures and follows a non-Gaussian form of displacement distribution (Tuch et al., 2003).

In this study, we found that \( MD, D_r \), and \( D_s \) obviously decreased in all groups, while MK, \( K_r \), and \( K_s \) dramatically

### Table 1

|       | \( \Delta A D C \) | \( \Delta F A \) | \( \Delta M D \) | \( \Delta D_r \) | \( \Delta D_s \) | \( \Delta M K \) | \( \Delta K_r \) | \( \Delta K_s \) |
|-------|------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| PWM vs. CC | 0.948            | 0.353          | 0.549          | 0.479          | 0.734          | 0.033*         | 0.229          | 0.074          |
| PWM vs. CB | <0.001*         | 0.557          | 0.013*         | 0.007*         | 0.405          | <0.001*        | <0.001*        | <0.001*        |
| PWM vs. BG | 0.135            | 0.002*         | 0.002*         | 0.002*         | 0.051          | 0.029*         | 0.002*         | 0.94           |
| PWM vs. BS | 0.28             | 0.281          | 0.07           | 0.061          | 0.07           | 0.024*         | 0.043*         | 0.043*         |
| PWM vs. LGW | 0.852            | 0.012*         | 0.285          | 0.064          | 0.882          | 0.001*         | 0.001*         | 0.032*         |
| CC vs. CB | 0.06             | 0.639          | 0.075          | 0.093          | 0.708          | 0.001*         | 0.001*         | 0.021*         |
| CC vs. BG | 0.534            | 0.009*         | 0.063          | 0.074          | 0.286          | 0.028*         | 0.028*         | 0.174          |
| CC vs. BS | 0.270            | 0.138          | 0.354          | 0.084          | 0.536          | 0.005*         | 0.049*         | 0.084          |
| CC vs. LGW | 0.941            | 0.126          | 0.4             | 0.126          | 0.769          | 0.007*         | 0.025*         | 0.111          |
| CB vs. BG | 0.066            | 0.051          | 0.956          | 0.663          | 0.61           | 0.001*         | <0.001*        | 0.003*         |
| CB vs. BS | 0.515            | 0.24           | 0.24           | 0.954          | 0.112          | <0.001*        | 0.006*         | 0.017*         |
| CB vs. LGW | <0.001*         | 0.202          | 0.077          | 0.227          | 0.162          | 0.001*         | 0.002*         | 0.014*         |
| BG vs. BS | 0.668            | 0.207          | 0.376          | 0.163          | 0.749          | 0.68           | 0.61           | 0.207          |
| BG vs. LGW | 0.367            | 0.214          | 0.054          | 0.165          | 0.053          | 0.452          | 0.219          | 0.214          |
| BS vs. LGW | 0.38             | 0.183          | 0.159          | 0.159          | 0.056          | 0.202          | 0.917          | 0.782          |

Value with the superscript character (*) in the table indicates that the \( P \) value was < 0.05. PWM: Periventricular white matter; CC: corpus callosum; CB: cerebellum; BG: basal ganglia and thalamus; BS: brain stem; LGW: gray-white matter junctions; ADC: apparent diffusion coefficient; FA: fractional anisotropy; MD: mean diffusivity; \( D_r \): axial diffusivity; \( D_s \): radial diffusivity; MK: mean kurtosis; \( K_r \): axial kurtosis; \( K_s \): radial kurtosis; DKI: diffusion kurtosis imaging.

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Figure 2 Calculation sketch map of MD and MK (DKI sequence using 15 directions and three b values typically).
The value of MD was decided by the values of $D_{app}$ from all directions, $K_{app}$ while the MK was decided by the values of from all directions. MD: Mean diffusivity; MK: mean kurtosis; DKI: diffusion kurtosis imaging.

Figure 3 DWI and DKI metrics maps from a 46-year-old healthy female volunteer.
MD, $D_a$, and $D_r$ maps were almost the same as the ADC map. DWI: Diffusion-weighted imaging; ADC: apparent diffusion coefficient; FA: fractional anisotropy; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Increased in all groups. The behaviors of diffusion-derived parameters and kurtosis-derived parameters were quite different. In corpus callosum, periventricular white matter, and brainstem areas, diffusion-derived parameters decreased remarkably, while MK, $K_a$ and $K_r$ increased noticeably compared with the normal side. The kurtosis-derived parameters of lesions located in the basal ganglia and thalamus region increased slightly compared with the normal side. In lesions located at lobe areas mixed with gray and white matter, diffusion-derived parameters decreased slightly, with a corresponding slight increase in kurtosis-derived parameters. In lesions located in the cerebellum, diffusion-derived parameters decreased sharply, but the performances of MK, $K_a$ and $K_r$ mapping were quite different from those of other locations; only parts of lesions increased in signal change.

Figure 4 DKI-derived metric maps with lesion (pink arrow) located in periventricular white matter from a 73-year-old female patient suffering from an acute ischemic stroke (onset duration < 72 hours). Signal intensity of the lesion is decreased markedly on MD, $D_a$, and $D_r$ maps, while it is increased on MK, $K_a$, and $K_r$ maps. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Figure 5 DKI-derived metric maps with a lesion (pink arrow) located in the corpus callosum from a 38-year-old female acute ischemic stroke patient (onset duration < 72 hours). The signal intensity change of the lesion was the same as the lesion in the periventricular white matter area in Figure 4. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.
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Figure 6 DKI-derived metric maps with lesion (pink arrow) involving the left cerebellar hemisphere from an 85-year-old male patient suffering from acute ischemic stroke (onset duration < 72 hours).

Signal intensity of the whole lesion decreased markedly on MD, $D_a$, and $D_r$ maps, while part of the lesion’s signal intensity was higher than its mirror normal side on MK, $K_a$, and $K_r$ maps. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Figure 7 DKI-derived metric maps with focal lesion (pink arrow) located in the basal ganglia and thalamus from an 86-year-old female acute ischemic stroke patient (onset duration < 72 hours).

The signal intensity of the lesion was slightly higher than its mirror normal side on MK, $K_a$, and $K_r$ maps. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Figure 8 DKI-derived metric maps with lesion (pink arrow) involving the right half of the brainstem from a 69-year-old female patient suffering from acute ischemic stroke (onset duration < 72 hours).

The signal intensity change of the lesion was the same as the ones in the periventricular white matter (Figure 4) and corpus callosum (Figure 5) areas. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Figure 9 DKI-derived metric maps with lesion (pink arrow) located at parietooccipital lobe from a 60-year-old female acute ischemic stroke patient (onset duration < 72 hours).

Part of the lesion’s signal intensity decreased on MD, $D_a$, and $D_r$ maps, and increased on MK, $K_a$, and $K_r$ maps correspondingly. DWI: Diffusion weighted imaging; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; FA: fractional anisotropy; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.

Figure 10 Comparison of ADC and DKI matrices among groups.

The upper three lines represent kurtosis-derived parameters, while the lower five lines represent the diffusion parameters. The upper three lines underwent larger changes, which meant that they had greater potential in distinguishing the differences among groups. PWM: Periventricular white matter; CC: corpus callosum; CB: cerebellum; BG: basal ganglia and thalamus; BS: brain stem; LGW: gray-white matter junctions; ADC: apparent diffusion coefficient; FA: fractional anisotropy; MD: mean diffusivity; $D_a$: axial diffusivity; $D_r$: radial diffusivity; MK: mean kurtosis; $K_a$: axial kurtosis; $K_r$: radial kurtosis; DKI: diffusion kurtosis imaging.
compared with the normal side. In descending order, changes in percentage of kurtosis-derived parameters ($\Delta MK$ and $\Delta K$) in different locations was as follows: corpus callosum, periventricular white matter, brainstem, lobes mixed with gray and white matter, basal ganglia and thalamus, and cerebellum. This may indicate that when an acute ischemic stroke affects tissue mostly containing white matter, the microstructural changes of tissue are much more complex than those in other affected locations consisting of a higher proportion of gray matter.

As already known, the corpus callosum, brainstem, and periventricular white matter areas primarily contain white matter, while less white matter is found in lobes containing mixed gray and white matter. The basal ganglia and thalamus contain only gray matter. The cerebellum consists of a tightly folded layer of cortex with white matter underneath. These findings illustrate that in regions mostly containing white matter, the complexity of microstructural changes resulting from acute stroke is much higher than other affected regions, except the cerebellum. Although we have no distinct evidence to explain why affected tissue in the cerebellum performed quite differently from other regions, it should have some relationship with its microstructure.

In conclusion, when acute ischemic stroke affects tissues mostly containing white matter, the complexity of microstructural changes of the tissue is much higher than in other affected regions. DKI-derived kurtosis metrics, especially MK, have potential for characterizing different locations of brain tissue affected by acute ischemic stroke. Moreover, DKI technology can reveal differences in microstructural changes among different locations affected by acute ischemic stroke. For this purpose, kurtosis-derived parameters performed better than diffusivity-derived parameters.

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Institutional review board statement: This study was approved by the Institutional Review Board of Xiamen Second Hospital, China (approval No. 2014002). The study was performed in accordance with the relevant laws and regulations of the Declaration of Helsinki, and the hospital's relevant ethical principles.

Declaration of patient consent: The authors certify that they have obtained all appropriate patient consent forms. In the form, patients have given their consent for their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity.

Reporting statement: This study followed the Standard Protocol Items: STrengthening the Reporting of Observational studies in Epidemiology (STROBE) statement.

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