Clinical application value of 3.0T MR diffusion tensor imaging in grade diagnosis of gliomas

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Abstract. The clinical value of MR diffusion tensor imaging (DTI) in grade diagnosis of gliomas was investigated. A total of 31 patients with glioma were administered 3.0T MR convention and DTI examination, with quantitative measurement of anisotropy coefficient fractional anisotropy (FA) and apparent dispersion coefficient (ADC) value, and the comparison of quantitative parameters of glioma between low- and high-grade, which was detected by Mann-Whitney U test. The receiver operation characteristic (ROC) curve was drawn to take the value of ADC and FA in tumor ROI as a critical point, to calculate the area under the curve and to confirm the diagnosis threshold value and evaluate its diagnostic efficiency. The FA value of 14 low-grade glioma cases was 139.4±81.3, with an ADC value of (1.36±0.21) x10^{-3} mm^2/sec. The FA value of 17 high-grade glioma cases was 103.1±41.5, with ADC value of (1.09±0.28) x10^{-3} mm^2/sec; the difference between the two groups was statistically significant (P<0.05). The ADC value was taken as the critical point to judge tumor grade and draw the ROC curve; the area under the curve was 0.79. As the diagnosis threshold value, the ADC value of 1.11x10^{-3} mm^2/sec was used to distinguish between low- and high-grade tumor with a sensitivity of 58.8% and specificity of 92.9%. The FA value was taken as a critical point to judge tumor grade and draw the ROC curve; the area under the curve was 0.62. As the diagnosis threshold value, the FA value of 178.9 was applied to distinguish between low- and high-grade tumor sensitivity of 94.1% and specificity of 35.7%. Therefore, the FA value and ADC value in DTI has an important estimated value for the pathological grade of glioma.

Introduction

Glioma is one of the most common primary central nervous system malignant tumors. With relatively high mortality and disability rate, glioma can seriously influence the living quality of patients and cause them and their families heavy economic burden (1,2). Glioma patients account for ~50% of intracranial tumor patients. Its pathogenesis involves various factors, including genetics, ionizing radiation, biochemical circumstance and environmental pollution and infection. But its exact pathogenetic molecular mechanism remains unclear (3,4). Clinically, glioma is usually diagnosed at the middle and advanced stages with poor prognosis. Therefore, the exploration of imaging features of glioma is of important significance for indications of tumor occurrence, development and prediction of early prognosis.

Diffusion tensor imaging (DTI) mainly reflects the corrosion and damage situation of white matter fiber bundles caused by glioma, and its grade malignancy is associated with the proliferation of tumor cells, and tumor angiogenesis to damage the completeness and continuity of a fiber bundle. The present study quantitatively measured the DTI index of 31 cases with glioma and discuss the clinical application value of DTI technology in establishing glioma grade.

Materials and methods

Sample selection. The DTI data of 31 patients that have a confirmed diagnosis of glioma by operation and pathology from June, 2013 to January, 2015 was analyzed retrospectively. The cohort included 15 males and 16 females aged 4-70 years and a median age of 43 years. According to Classification and Grading Standard of WHO Central Nervous System Tumors (2007) (2), there are 14 cases of grade I and II glioma, namely low-grade glioma (LGG) and there are 17 cases of grade III and IV, which belong to high-grade glioma (HGG), among which there were 2 cases of pilocytic astrocytoma (grade I), 3 cases of protoplasmic astrocytoma (grade II), 4 cases of diffuse astrocytoma (grade II), 5 cases of oligoastrocytomas (grade II), 9 cases of anaplastic astrocytoma (grade III), 3 cases of anaplastic oligoastrocytomas (grade III), 3 cases of anaplastic ependymoma (grade III) and 2 cases of glioblastoma (grade IV). The main clinical symptoms include headache, dizziness, nausea, sickness, convulsion, acroagnosis and movement.
disturbance. The general clinical situations of the patient in the two groups is shown in Table I. Images of two cases with astrocytoma or spongioblastoma are shown in Figs. 1 and 2.

**Table I. General clinical situations of patients.**

| Grade of glioma | Total case no. | Detailed types of glioma | Case no. |
|-----------------|----------------|--------------------------|---------|
| Low             | 31             | Pilocytic astrocytoma(grade I) | 2 (2/31) |
|                 | 31             | Protoplasmic astrocytoma (grade II) | 3 (3/31) |
|                 | 31             | Diffuse astrocytoma (grade II) | 4 (4/31) |
|                 | 31             | Oligoastrocytomas (grade II) | 5 (5/31) |
| High            | 31             | Anaplastic astrocytoma (grade III) | 9 (9/31) |
|                 | 31             | Anaplastic oligoastrocytomas (grade III) | 3 (3/31) |
|                 | 31             | Anaplastic ependymoma (grade III) | 3 (3/31) |
|                 | 31             | Glioblastoma (grade IV) | 2 (2/31) |

Figure 1. Male, 59 years old, left frontal lobe astrocytoma (WHO grade II). (A) Enhancement scan of left frontal lobe focus, without clear reinforcement; (B) Directionally encoded color image showed low signal of focus and shifted alba under oppression; (C) pathological examination displayed that tumor consisted of neoplastic astrocyte with better differentiation, with loose cell arrangement and mild atypia; (D) CD34 immunohistochemistry stain showing micrangium dispersion, with regular tubular.

**MR device and scan scheme.** Skyra 3.0T superconductivity MR scanner was obtained from Siemens (Munich, Germany), with scanning sequence including conventional plain scan T1WI and T2WI, fluid-attenuated inversion-recovery sequence (FLAIR), DTI and conventional enhanced MRI scan.

The DTI sequence is as follows: Single-shot echo planner imaging was applied, imaging FOV taking 220x220 cm, with TR time of 5,400 msec, a total of 40 layers, with layer thickness of 3.0 mm, and interlamellar spacing of 0 mm. The phase encoding direction was taken from front to back and a collection technique GRAPPA was run to shorten the time of collecting signal, with 2 accelerated factors and resolution ratio of 128x128. Multi-direction diffusion imaging scan was applied to exert diffusion sensitive gradient in 30 directions, a b value taken as 0 and 1,000 sec/mm², respectively. The average time of collecting signal was 12 times for b value of 0 sec/mm² and the average time of collecting signal was twice for b value of 1,000 sec/mm². The DWI, apparent dispersion coefficient (ADC), index ADC and fractional anisotropy (FA)
were processed automatically after scanning and setting in the parameter tab control.

A magnevist solution injection was used as contrast-enhancing agent, with a dosage of 0.2 mmol/kg and injection rate of 2.0 ml/sec. After injection, normal saline of the same dosage was injected. The patients of the present study did not develop an allergic reaction to the contrast agent.

**Image processing.** The processing and analysis of the DTI image: The original data of DTI from scan was transmitted to Siemens Syngo MMWP workstation and processed by the software package Neuo 3D. Combining directionally encoded color and FA image, 4 ROIs of 30 mm² were drawn manually on the solid area of the tumor cross section's largest layer. The selection of solid area was referred to as T1, T2, FLAIR and DCE image of three directions, with combination of multiple weighted imaging to observe, judge and avoid edema area, measuring FA value and ADC, and tumor cysts, necrosis and hemorrhage area were avoided when measurement was conducted, the FA value of normal alba taken from the focus as a reference value, and the ratio of lesion and normal alba as the FA value. The ADC value was measured on the ADC image, and 4 ROIs sized 30 mm² were drawn manually on the solid area of tumor with the largest cross section. The location selection of ROI was brought into correspondence with that of the FA value measurement, taking the average value of the observed value (Fig. 3).

**Statistical analysis.** The statistical software SPSS 16.0 (SPSS, Inc., Chicago, IL, USA) was used for statistical analysis. The Kolmogorov-Smirnov method was applied to conduct a normality test for the value of FA and ADC, which met normal distribution. The quantitative data are shown as the mean ± standard deviation and comparison between groups utilized t-test or the \( \chi^2 \) test. The receiver operation characteristic (ROC) curve was used to analyze the FA and ADC value, as well as diagnose the specificity and sensitivity of the threshold value.

**Results**

**Analysis of FA and ADC value of different pathological patterns of glioma (mean ± standard deviation).** Statistical analysis of the FA and ADC value of 31 glioma cases of different pathological patterns demonstrated that there was no statistical difference among the patients with different pathological patterns (P<0.05; Table II).

**Analysis of FA and ADC value of different pathological patterns of glioma.** Furthermore, FA and ADC values of different grades of glioma were statistically analyzed, and the results showed that the average FA of LGG was higher than that of HGG group. Moreover, the ADC of LGG was higher than that of HGG group; results were statistically significant (Table III).
Results of the analysis of area under ROC curve. The ADC/FA value in tumor ROI was taken as the critical point to judge tumor grade and draw the ROC image (Fig. 4); the area under the curve was 0.79/0.62. As the diagnosis threshold value, the ADC/FA value of $1.11 \times 10^{-3}$ mm²/sec/178.9 was used to distinguish between low- and high-grade tumors. Using this threshold value, high-grade malignant tumors were diagnosed with sensitivity of 58.8/94.1% and specificity of 92.9/35.7%. Compared to FA, ADC has greater advantage in distinguishing tumor grade.
Analysis of sensitivity and specificity of conventional and DTI combination scan.

Table IV. Analysis of sensitivity and specificity of conventional and DTI combination scan.

| Groups                              | Pathological results |
|-------------------------------------|----------------------|
|                                     | High-grade glioma    | Low-grade glioma | Total |
| Conventional MRI                    |                      |                  |       |
| High-grade glioma                   | 13                   | 2                | 15    |
| Low-grade glioma                    | 4                    | 12               | 16    |
| Total                               | 17                   | 14               | 31    |
| Conventional MRI combined with DTI  |                      |                  |       |
| High-grade glioma                   | 17                   | 0                | 17    |
| Low-grade glioma                    | 0                    | 14               | 14    |
| Total                               | 17                   | 14               | 31    |

DTI, diffusion tensor imaging.

Table V. Comparison of accuracy, sensitivity, specificity, positive and negative predictive value of different detection methods (%).

| Variables                  | Sensitivity (%) | Specificity (%) | Positive predictive value (%) | Negative predictive value (%) | Accuracy (%) |
|----------------------------|-----------------|-----------------|-------------------------------|-------------------------------|--------------|
| Conventional MRI           | 76.1            | 86.2            | 81.1                          | 0.75                          | 81.3         |
| Conventional MRI combined DTI | 100             | 100             | 55.0                          | 45.0                          | 100          |
| T-value                    | 1.27            | 1.88            | 1.49                          | 1.57                          | 2.17         |
| P-value                    | <0.05           | <0.05           | <0.05                         | <0.05                         | <0.05        |

DTI, diffusion tensor imaging.

Discussion

The guiding significance of the MR DTI index is as follows: The subsidiarity that iconography can provide for tumor grading is correspondingly indispensable due to the importance of tumor grading for treatment and prognosis. DTI mainly reflects the integrity of the nerve fiber anatomical structure which can assess the microcosmic physiology and pathological state of white matter fiber tracts using non-invasive techniques. Moreover, DTI technology can also judge tumor cell density and evaluate the ability of tumor cells to invade white matter degree efficiently.

The change of ADC value in tumors is influenced by tumor cell structure to a great extent (5). With the increasing ascendance of grade malignancy and the density of tumor cell structure, the tumor form becomes more complicated. Brownian movement of molecules is limited even further, eventually leading to decline of the ADC value directly. In sensitive, specific, positive and negative predictive value of conventional MRI and DTI combination scan was conducted. Our analysis demonstrated that the sensitivity, specificity and accuracy of DTI combination scan were higher than that of conventional scans; differences were statistically significant (P<0.05; Tables IV and V).

Figure 4. Receiver operation characteristic curve that takes ADC/FA value as critical point to judge low- and high-grade tumor, with area under curve of 0.79 (ADC)/0.62 (FA). ADC, apparent dispersion coefficient, FA, fractional anisotropy.

Analysis of sensitivity and specificity of conventional and DTI combination scan. Analysis of HGG as the positive, sensitive, specific, positive and negative predictive value of conventional MRI and DTI combination scan was conducted. Our analysis demonstrated that the sensitivity, specificity and accuracy of DTI combination scan were higher than that of conventional scans; differences were statistically significant (P<0.05; Tables IV and V).
the present study, the ADC value in HGG is lower than that in LGG, which indicates that tumor cell structures will become more complicated with an increase of grade malignancy, albeit indirectly. In addition, HGG with high-grade malignancy can lead to abnormal tumor blood vessel hyperplasia and the micro-structure completeness of the tumor cell is broken, which results in free water infiltration nerve fiber axon gap and a decline in the FA value. Through the comparison of FA values, the present study found that when compared to LGGs, the FA value of HGG is reduced significantly, which reflects blood vessels in HGG mesenchyme erodes or damage to nerve fiber bundle. This leads to free water infiltration to nerve fiber axon gap and decline in the FA value, which suggests that it indirectly leads to the spatial distribution situation of tumor capillaries (5-7).

Previous research results indicate that the difference between FA and ADC in high- and low-grade glioma was statistically significant; our study corroborated the results that both of them can distinguish different grades of the tumor (4). However, in terms of the comparison of diagnostic efficiency, the FA value of different grades as the critical point to distinguish tumor grade had a curve area of 0.62, which was lower than that of ADC; this suggests that ADC has a better diagnostic efficiency. However, FA has a stronger sensitivity and a lower specificity and ADC has lower sensitivity and stronger specificity; the difference is attributed to the individual theory behind these two parameters (8). FA mainly reflects anisotropy of the internal structure of tumor, while ADC reflects a limited degree of molecular diffusion into the tumor, which is a comprehensively average effect, excluding anisotropy (9-11). Tumors are heterogeneous, and therefore, there is coexistence of edema and substance as well as coexistence of low- and high-grade. If we evaluated the FA and ADC from two different starting points, there would be a different analysis result (12-14).

The present study utilized the FA and ADC to evaluate high- and low-grade of tumors, revealing that the two indexes can distinguish between the grades of tumor. The combination of these is more helpful in distinguishing the grade of tumor and understanding the internal structure, which can also be applied for a study with a larger sample size. We analyzed the sensitivity, specificity, accuracy, positive predictive value and negative predictive value of conventional MRI and DTI combination and found that the sensitivity, specificity and accuracy of DTI combination scan was higher than that of conventional MRI scan; differences were statistically significant (P<0.05). We speculated that DTI method was superior in the evaluation of blood vessel corrosion and damage to the nerve fiber bundle, which results in free water infiltration to nerve fiber axon gap. Therefore, it has a favorable reminder for clinical diagnosis, which is helpful for a pathologist to diagnose the glioma grade.

A serious limitation of this study is that there was a limited number of clinical cases, and therefore, there was a possible existence of a bias in the evaluation of diagnostic efficiency. In conclusion, conventional and DTI combination scan can assess glioma with new vessel hyperplasia and the degree of nerve fiber bundle damage more comprehensively, which can provide a new method for the monitoring of tumor diagnosis, growth, transfer and postoperative recurrence.

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