Validity of Acute Stroke Lesion Volume Estimation by Diffusion-Weighted Imaging–Alberta Stroke Program Early Computed Tomographic Score Depends on Lesion Location in 496 Patients With Middle Cerebral Artery Stroke

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**Background and Purpose**—Alberta Stroke Program Early Computed Tomographic Score (ASPECTS) has been used to estimate diffusion-weighted imaging (DWI) lesion volume in acute stroke. We aimed to assess correlations of DWI-ASPECTS with lesion volume in different middle cerebral artery (MCA) subregions and reproduce existing ASPECTS thresholds of a malignant profile defined by lesion volume ≥100 mL.

**Methods**—We analyzed data of patients with MCA stroke from a prospective observational study of DWI and fluid-attenuated inversion recovery in acute stroke. DWI-ASPECTS and lesion volume were calculated. The population was divided into subgroups based on lesion localization (superficial MCA territory, deep MCA territory, or both). Correlation of ASPECTS and infarct volume was calculated, and receiver-operating characteristics curve analysis was performed to identify the optimal ASPECTS threshold for ≥100-mL lesion volume.

**Results**—A total of 496 patients were included. There was a significant negative correlation between ASPECTS and DWI lesion volume (r = −0.78; P < 0.0001). With regards to lesion localization, correlation was weaker in deep MCA region (r = −0.19; P = 0.038) when compared with superficial (r = −0.72; P < 0.001) or combined superficial and deep MCA lesions (r = −0.72; P < 0.001). Receiver-operating characteristics analysis revealed ASPECTS ≤ 5 as best cutoff to identify ≥100-mL DWI lesion volume; however, positive predictive value was low (0.35).

**Conclusions**—ASPECTS has limitations when lesion location is not considered. Identification of patients with malignant profile by DWI-ASPECTS may be unreliable. ASPECTS may be a useful tool for the evaluation of noncontrast computed tomography. However, if MRI is used, ASPECTS seems dispensable because lesion volume can easily be quantified on DWI maps. *(Stroke. 2014;45:3583-3588.)*

**Key Words:** brain ischemia ▪ diffusion magnetic resonance imaging ▪ magnetic resonance imaging ▪ middle cerebral artery ▪ neuroimaging ▪ severity of illness index ▪ stroke

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*A list of all STIR and VISTA Imaging study participants is given in the Appendix.

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stroke. Although originally designed for evaluation of CT ASPECTS may also be applied for standardized scoring of magnetic resonance diffusion-weighted imaging (DWI),2 DWI-ASPECTS was found to be superior in detecting ischemic changes when compared with noncontrast CT alone,2 reflecting the higher sensitivity of DWI for acute cerebral ischemia.

It has been reported that patients with large lesions on DWI-MRI have a high risk of developing symptomatic intracerebral hemorrhage (SICH) and a worse clinical outcome after intravenous thrombolysis. The Diffusion and Perfusion Imaging Evaluation for Understanding Stroke Evolution (DEFUSE) study identified a threshold of ≥100-mL lesion volume,4 characterizing a malignant course (ie, high risk of SICH and poor outcome). It has been suggested to use DWI-ASPECTS to identify patients with acute stroke and large DWI lesions, thus avoiding the need for measuring lesion volumes in the acute setting. DWI-ASPECTS was already found to predict SICH after thrombolysis.5 In addition, several studies have tried to define ASPECTS cutoff values identifying large DWI lesion volumes.6,7

However, there are limitations to ASPECTS. First, the score is limited to the anterior circulation. Moreover, the anatomic regions covered by ASPECTS are not equally distributed across the middle cerebral artery (MCA) territory. There are 3 score items allotted to subcortical regions, which are located in close proximity, whereas 7-score items represent cortical brain areas distributed across the entire MCA territory. It has already been argued that ASPECTS regions are weighed unequally and in favor of the striatocapsular region.8

The aim of the present study was to assess the performance of DWI-ASPECTS as surrogate for ischemic lesion volume with regard to the affected region of the MCA territory and to reproduce existing threshold values to identify patients with a malignant profile defined by extensive DWI lesion volume.

Methods

We analyzed patient data from the Predictive Value of Fluid-Attenuated Inversion Recovery (PRE-FLAIR) study database. The details of this study have been published previously.9 PRE-FLAIR was a multicenter observational study of patients with acute ischemic stroke who underwent multiparametric MRI within 12 hours of symptom onset.

Demographic data, time from symptom onset to MRI, severity of neurologic deficit on admission as assessed with the National Institutes of Health Stroke Scale (NIHSS), stroke cause according to Trial of Org 10172 in Acute Stroke Treatment (TOAST)10 definitions on admission were recorded. DWI-ASPECTS was rated by a neurologist experienced in stroke imaging. DWI lesion volumes were calculated by a semiautomated thresholding approach using an in-house developed software tool (AnToNiA), as described previously.9 For the present analysis, we excluded patients with anterior or posterior cerebral artery infarction and bifrontal ischemic lesions. Thus, we included only patients with MCA infarction. We further divided the population into 3 subgroups based on the localization of DWI lesions: involvement of superficial MCA territory alone, involvement of the deep MCA territory alone, and finally involvement of both superficial and deep MCA territories.

Correlations among DWI-ASPECTS, DWI lesion volume, and NIHSS were calculated using Pearson correlation coefficient. Group comparison was performed using the Kruskal–Wallis or Mann–Whitney U test. Receiver operating characteristic curves were generated to determine the optimal DWI-ASPECTS cutoff point to characterize a DWI lesion volume ≥100 mL.4 All statistical analysis was performed using SPSS 21.

Results

Of 543 patients included in the final analysis in the PRE-FLAIR study, we excluded 47 patients with DWI lesion outside the MCA territory or no visible DWI lesion. Thus, 496 patients with MCA territory stroke were included in the present analysis (47% women; mean age, 66 years; SD±15 years). Mean onset-to-MRI delay was 244±170 minutes (see the Table for patient characteristics). Figure 1 shows the distribution of DWI-ASPECTS values.

The subgroups defined by involvement of the different MCA regions showed significant differences in mean DWI lesion volume, NIHSS, and median ASPECTS (Table). Patients with DWI lesions in both superficial and deep MCA territories had larger DWI lesion volumes, higher NIHSS values, and lower ASPECTS scores than both other groups. In addition, patients

| Table. Baseline Clinical and Imaging Parameters in Different Subgroups and Overall Study Population |
|---|---|---|---|---|
| All Patients MCA | Superficial MCA (n=279) | Deep MCA (n=128) | Both (n=88) | Group Comparison |
| Age (mean, SD) | 66±15 | 67.1±14.7 | 64.3±15.8 | 65±14.8 | P=0.203 |
| Sex | | | | |
| Men | 263 (53%) | 163 (58.4%) | 55 (43%) | 45 (51.1%) | |
| Women | 233 (47%) | 116 (41.6%) | 73 (57%) | 43 (48.9%) | P=0.014 |
| NIHSS (mean, SD) | 10.3±7.1 | 8.7±6.5 | 9.4±6.4 | 16±6.6 | P=0.0001 |
| Cause (TOAST) | | | | |
| Large artery atherosclerosis | 139 (28%) | 71 (25.4%) | 41 (32%) | 26 (29.5%) | |
| Cardioembolism | 150 (30.2%) | 95 (34.1%) | 28 (21.9%) | 27 (30.7%) | |
| Small-vessel occlusion | 21 (4.2%) | 4 (1.4%) | 17 (13.3%) | 0 | |
| Stroke of other determined cause | 36 (7.3%) | 14 (5%) | 11 (8.6%) | 11 (12.5%) | |
| Stroke of undetermined cause | 64 (12.9%) | 43 (15.4%) | 14 (10.9%) | 7 (8%) | |
| ASPECTS (median, IQR) | 9, 7–9 | 9, 7–9 | 9, 9–9 | 6, 5–7 | P=0.0001 |
| DWI lesion volume (mean, SD), MI | 25±38.6 | 21.2±30.8 | 7.2±17.7 | 67.8±54.6 | P=0.0001 |

Group comparison was conducted using the Kruskal–Wallis Test. ASPECTS indicates Alberta Stroke Program Early Computed Tomographic Score; DWI, diffusion-weighted imaging; IQR, interquartile range; MCA, middle cerebral artery; NIHSS, National Institutes of Health Stroke Scale; and TOAST, Trial of Org 10172 in Acute Stroke Treatment.
with superficial MCA stroke lesions had higher DWI lesion volumes than patients with deep MCA stroke lesions (21.2 versus 7.2 mL), whereas median ASPECTS seemed comparable between groups (9 versus 9).

There was a significant negative correlation between DWI-ASPECTS and DWI lesion volume for all patients ($r = -0.78; P < 0.0001$). Looking at the different subgroups, strong negative correlations between DWI-ASPECTS and DWI lesion volume were detected for superficial stroke lesions ($r = -0.72; P = 0.0001$) and in the combined superficial and deep MCA region group ($r = -0.72; P < 0.0001$). In the deep MCA region group, correlation of DWI-ASPECTS with lesion volume was considerably weaker ($r = -0.19; P = 0.0383$). This is also reflected in the plots of DWI lesion volume by ASPECTS value (Figure 2).

Correlations of DWI-ASPECTS and initial NIHSS revealed a similar pattern. A significant negative correlation was found for the entire sample ($r = -0.49; P = 0.0001$) and for the groups with involvement of superficial MCA regions alone ($r = -0.41; P = 0.0001$) or superficial and deep MCA regions ($r = -0.39; P < 0.0001$), whereas correlation was much weaker in patients with involvement of deep MCA regions alone ($r = -0.18; P = 0.0455$).

Figure 3 shows the receiver operating characteristic curve for ASPECTS to identify a DWI lesion $\geq 100$ mL. Area under the curve was 0.938 with a 95% confidence interval of 0.891 to 0.985. ASPECTS $\leq 6$ was the best predictor of DWI lesion $\geq 100$ mL with high sensitivity 0.935 (95% confidence interval, 0.772–0.989), specificity 0.877 (0.841–0.905), and negative predictive value 0.995 (0.979–0.999), but low positive predictive value 0.349 (0.250–0.463).

The previously described ASPECTS threshold of $\leq 36.7$ to identify lesion volume $\geq 100$ mL applied to our sample resulted in the following predictive values: sensitivity 0.258 (0.125–0.449), specificity 0.990 (0.975–0.997), negative predictive value 0.950 (0.924–0.967), and positive predictive value 0.666 (0.354–0.887).

![Figure 1](image1.png)

**Figure 1.** Distribution of diffusion-weighted imaging (DWI) Alberta Stroke Program Early Computed Tomographic Score (ASPECTS) values across 496 patients.

![Figure 2](image2.png)

**Figure 2.** Scatterplot of diffusion-weighted imaging (DWI) lesion volume per Alberta Stroke Program Early Computed Tomographic Score (ASPECTS) value in all middle cerebral artery (MCA) strokes and different subgroups. A, For all analyzed MCA strokes; B, superficial MCA territory involved; C, deep MCA territory involved; and D, superficial and deep MCA territory involved.
Discussions

In our analysis of DWI-ASPECTS and DWI lesion volume in a large sample of patients with acute ischemic stroke, we found that the performance of ASPECTS as substitute for stroke lesion volume depends on lesion location. Although there was a strong correlation of ASPECTS with DWI lesion volume in the stroke lesions involving the superficial MCA region, a much weaker correlation was found for lesions confined to the deep MCA territory. Thus, our study demonstrates for the first time in a large sample of patients an unequal weighing of ASPECTS, treating lesions in the basal ganglia differently from those in the superficial MCA territory. This is in line with a previous theoretical demonstration of an unequal distribution of tissue volume per ASPECTS region of interest.

ASPECTS was originally designed to overcome limitations of noncontrast CT imaging in acute stroke, including high inter-rater variability and to standardize assessment of early ischemic changes. It is increasingly used to identify patients likely to benefit from reperfusion treatment both in clinical routine as in clinical stroke trials. ASPECTS predicts functional outcome and risk of SICH after thrombolysis, a threshold value of ≤7 on CT and ≤6 on DWI-MRI was proposed to isolate patients at high risk. Although originally designed to be used with CT imaging, it has also been used for standardized assessment and quantification of acute stroke lesions on DWI. In contrast to CT, DWI allows accurate delineation and volume measurement of acute ischemic lesions and thus enables comparison of ASPECTS with effective lesion volumes. In addition, DWI-ASPECTS was found to be superior to noncontrast CT ASPECTS in detecting ischemic changes, which again highlights the superiority of MRI in detecting acute ischemic brain lesions.

In the present data, we found a wide range of lesion volumes per ASPECTS value. Nevertheless, there was a strong correlation of lesion volume and ASPECTS when all patients with MCA infarction were included in the analysis. This is in line with previous studies, which also report a strong negative correlation between ASPECTS and DWI lesion volume.

However, already in 2006 Phan et al discussed an unequal weighing of different MCA regions in ASPECTS, favoring the basal ganglia. They speculated that this could lead to unjustified exclusion of patients from recombinant tissue-type plasminogen activator treatment based on ASPECTS thresholds that do not account for different lesion locations. Our study supports this concern by the finding of a striking disparity of ASPECTS-DWI lesion volume correlations between the superficial and the deep MCA regions. For stroke lesions in the deep MCA territory, ASPECTS showed only a poor correlation to DWI lesion volume. This means that estimation of lesion volume in striatocapsular stroke by ASPECTS is unreliable. Whether ASPECTS favors lesions in the basal ganglia, as suggested before or whether it should be considered as a disadvantage depends on the point of view. In any case, ASPECTS performs rather poorly as a surrogate of lesion volume in striatocapsular stroke, which raises concerns about the use of ASPECTS to exclude patients from treatment or stroke trials without consideration of lesion location. However, it has to be considered that virtually all patients with stroke lesions restricted to the deep MCA territory score ≥7 on DWI-ASPECTS, a value that usually allows inclusion in stroke trials using ASPECTS thresholds. Nevertheless, as soon as ASPECTS is used to compare lesion volumes between patients groups, differences in stroke location between those groups may introduce a significant bias.

Of note, a similar phenomenon was found for correlation of ASPECTS with NIHSS with strong correlations for lesions involving the superficial MCA territory but rather weak correlation for strokes confined to the deep MCA region. Thus, evaluation of striatocapsular stroke by ASPECTS seems to be inaccurate in multiple regards.

The volume of the ischemic lesion is a strong predictor of clinical outcome and is also correlated with the risk of SICH. In DEFUSE, DWI lesions ≥100 mL were associated with a poor outcome and a high risk of SICH. A tissue volume of 100 mL is also a widely accepted estimate for one third of the MCA territory, a cutoff that is frequently used to define large infarctions, which may not benefit from reperfusion treatment, but which is subject to notorious discussions and poor interobserver agreement. Previous studies have already aimed at determining an ASPECTS cutoff value for ≥100-mL lesion volume. In our study, a threshold value of ASPECTS ≤6 was the best predictor for ≥100-mL DWI lesion volume. This is close to the originally proposed cutoff value of ASPECTS ≤7 for identifying patients at high risk for ICH and poor clinical outcome irrespective of an explicit tissue volume. However, a low positive predictive value of only 0.35 limits the value of ASPECTS ≤6 as predictor of large DWI lesion volume in our study. Using this cutoff to exclude patients might thus lead to unjustified exclusion of patients from treatment or clinical trials if exclusion of patients with large (≥100 mL) DWI lesion is intended. However, negative predictive value is high (ie, one can assume that patients with ASPECTS >6 have a lesion volume <100 mL). Previous publications considering ASPECTS as a possible substitute for lesion volume measurement...
suggested a threshold value of \( \leq 3.6 \). Using this value in our data set resulted in only low sensitivity (0.26).

There are limitations to our study. We studied patients with acute stroke and no lower threshold for symptom severity or lesion volumes including patients where thrombolysis was not considered. This probably leads to a higher number of patients with relatively low lesion volume and the different distribution could explain at least part of the disparity in predictive values when compared with previous studies. Additional studies should assess the validity of DWI-ASPECTS in a balanced population of infant volumes.

Conclusions

If MRI or multimodal CT are not available, ASPECTS may be considered a simple and useful tool to help in the assessment of acute ischemic lesions on noncontrast CT in a semi-quantitative way and in the identification of patients with large acute stroke who may not benefit from reperfusion treatment. Nevertheless, comparison of ASPECTS with lesion volume measurements reveals relevant limitations, including a wide variation of lesion volumes for each score value and rather poor correlation with stroke lesion volume in the deep MCA territory. These limitations may impair decisions aiming at the exclusion of patients with large stroke lesions. Finally, the use of ASPECTS for evaluation of DWI seems somewhat anachronistic. Acute stroke lesions are visible with high contrast on DWI and tools for fast and easy volume measurement on MRI images are now widely available. Thus, if MRI is used for stroke imaging, there should be no need to use a substitute for stroke lesion volume measurement because lesion volume can easily be quantified on DWI maps.

Appendix


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References


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