Microstructural Integrity of Salvaged Penumbra after Mechanical Thrombectomy

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ABSTRACT

BACKGROUND AND PURPOSE: There are sparse data on the microstructural integrity of salvaged penumbral tissue after mechanical thrombectomy of large-vessel occlusions. The aim of the study was to analyze possible microstructural alteration in the penumbra and their association with clinical symptoms as well as angiographic reperfusion success in patients undergoing mechanical thrombectomy.

MATERIALS AND METHODS: All patients who underwent mechanical thrombectomy for large-vessel occlusions in the anterior circulation and who received an admission CT perfusion together with postinterventional DTIs were included (n = 65). Angiographic reperfusion success by means of modified Thrombolysis in Cerebral Infarction (mTICI) scale and clinical outcome were recorded. Microstructural integrity was assessed by DTI evaluating the mean diffusivity index within the salvaged gray matter of the former penumbra.

RESULTS: The mean diffusivity index was higher in completely recanalized patients (mTICI 3: 0.001 ± 0.034 versus mTICI <3: 0.030 ± 0.055, P = .03). There was a positive correlation between the mean diffusivity index and NIHSS score improvement (r = 0.49, P = .003) and the mean diffusivity index was associated with midterm functional outcome (r = –0.37, P = .04) after adjustment for confounders. In mediation analysis, the mean diffusivity index and infarction growth mediated the association between reperfusion success and clinical outcomes.

CONCLUSIONS: The macroscopic salvaged penumbra included areas of microstructural integrity changes, most likely related to the initial hypoperfusion. These abnormalities were found early after mechanical thrombectomy, were dependent on angiographic results, and correlated with the clinical outcome. When confirmed, these findings prompt the evaluation of therapies for protection of the penumbral tissue integrity.

ABBREVIATIONS: IQR = interquartile range; LVO = large-vessel occlusion; MD = mean diffusivity; MT = mechanical thrombectomy; SD = standard deviation; mTICI = modified Thrombolysis in Cerebral Infarction; mRS = modified Rankin Scale; NIHSS = National Institutes of Health Stroke Scale
MATERIALS AND METHODS

Sample and Patient Description
All patients included in a prospective stroke registry of a single comprehensive stroke center and treated with MT between April 2016 and December 2018 were reviewed. Four-hundred-thirty-nine patients with ischemic stroke presented with an LVO of the anterior circulation (MCA or carotid-T). Patients were included if they had postinterventional MR imaging, including DTI and structural 3D T1WI ($n = 192$). Patients with motion artifacts ($n = 20$) or space-occupying malignant infarction were excluded ($n = 7$). From the remaining 165 patients, 65 patients received quality-sufficient CT perfusion imaging on admission. Prospectively collected baseline demographic, clinical, and interventional data were extracted from the registry. Board-certified neurologists assessed NIHSS scores on admission and at the time of MR imaging acquisition. The functional outcome was evaluated by applying the modified Rankin Scale (mRS) 90 days after admission, either on a routinely scheduled clinical visit or by a structured telephone interview. This study was approved by the local ethics committee, and the need for patient consent was waived.

Angiographic Data
The modified Thrombolysis in Cerebral Infarction (mTICI) score was determined by 2 experienced neuro-interventionalists in consensus (C.M., B.F.). Complete reperfusion was defined as mTICI 3. The time of groin puncture, time of reperfusion, and corresponding procedure times were extracted from the data base. The time elapsed between groin puncture to reperfusion is referred to as reperfusion time. If no successful recanalization was achieved (mTICI < 2b), then the control series after the last maneuver was used as the time point of procedure termination.

Assessment of the Penumbra in Admission CT Imaging
Standard nonhelical cerebral CT was performed on a 64-row CT scanner equipped with a 40-mm detector (Brilliance 64, Philips Healthcare, Best, the Netherlands). The scanner was calibrated weekly by air calibration and regularly by using phantom scans (CT calibration phantom; Mindways Software, Austin, Texas). Besides noncontrast images and CT angiography, a perfusion image was acquired by using the following parameters: 120 kV, 400 mAs, with 5-mm section thickness. A dual-head power injector (Medrad, Indianola, Pennsylvania) with an 18-gauge IV access was used for contrast injection. A delay of 5 seconds was applied after injecting 40 mL Imeron (400 mg I/mL Imeron 350, iomeprol; Bracco, Milan, Italy) at a flow rate of 6 mL/s, followed by 90 mL NaCl. For postprocessing of the acquired CT perfusion images, RAPID (iSchemaView, Stanford, California) was used. This software is an operator-independent fully automated image processing and visualization tool, and allows on-line estimation of perfusion maps and mismatch masks. These maps were verified by a neuroradiologist (M.T.B.) with 3 years of experience. For assessing penumbral tissue, the individual hypoperfusion mask was extracted. With this, the hypoperfused tissue was estimated at threshold values of Tmax delay of >6 seconds and the admission infarct core was estimated at threshold values of relative CBF < 30%.

Assessment of Microstructural Changes Within the Penumbra
For analyses of microstructural tissue integrity within the above-identified penumbral tissue, an MR imaging examination was acquired in the acute poststroke phase, including DTI and structural T1WI. These data were postprocessed, coaligned with admission CT perfusion imaging, and analyzed with respect to microstructural tissue alterations within the salvaged penumbra by using the mean diffusivity (MD) index for gray matter. The salvaged penumbra was defined as formerly hypoperfused tissue in CT perfusion imaging, which, in the end, did not show visible infarction, which was semiautomatically segmented in MR imaging (Fig 1). A detailed methodologic description is found in the On-line Appendix.

Statistical Analysis
The mean value comparison of the MD index and infarction growth was performed by means of a 2-sample t-test for independent samples. Wilcoxon rank sum tests were used for comparison of the NIHSS scores and mRS values between the different reperfusion groups.

To test the association of the MD index and infarction growth with the NIHSS score (at the MR imaging date and the percentage improvement compared with admission) as well as with midterm functional outcome (mRS after 90 days), the following statistical procedures were applied. After performing linear regression to get residuals of the MD index/infarction growth (corrected for age, sex, time between recanalization and MR imaging scan, reperfusion time, admission infarct core), the Spearman rank bivariate correlation was performed between residuals of the MD index/infarction growth and the NIHSS score at the time of MR imaging as well as percentage improvement of the NIHSS score between admission and time of MR imaging and mRS at 90 days. In a second analysis, new residuals of the MD index that additionally control for infarction growth were calculated and used for Spearman rank bivariate correlation analyses with clinical outcome parameters.

In a mediation model, reperfusion success, measured by the mTICI score and dichotomized into complete and incomplete, was entered as the causal variable; percentage improvement of the NIHSS score between admission and the time of MR imaging as the outcome variable; and infarction growth and the MD index as mediator variables. The analysis was corrected for age, sex, and the time between recanalization and MR imaging by calculating residuals of infarction growth and the MD index by using linear regression. Path coefficients were estimated by using unstandardized regression coefficients of multiple regression analyses. Statistical significance of the indirect pathway, which reflects the impact of mediation, was evaluated by using a nonparametric bootstrap approach with 10,000 replication samples to obtain a 95% CI. All statistical analyses were performed by using SPSS Statistics, version 25 (IBM, Armonk, New York).
RESULTS

Patient Characteristics

Sixty-five patients were included (median age, 75 years; 52% female). Patients presented with severe symptoms (median [interquartile range (IQR)] NIHSS score, 12 [6–17]) and postinterventional MR imaging was scheduled at a median (IQR) of 3 days (3–4 days) after reperfusion. Most of the patients ($n = 39/65, 60\%$) had complete reperfusion (mTICI 3), whereas only 4 patients (6.2%) were unsuccessfully recanalized (mTICI < 2b). Further details can be found in the Table.

Microstructural Changes of the Salvaged Penumbra

For the whole study cohort, the MD index of the salvaged penumbra showed mainly negative values (mean ± standard deviation [SD], $-0.013 \pm 0.046$). The MD index was higher in patients with complete reperfusion (mean ± SD, $-0.001 \pm 0.034$ versus $-0.030 \pm 0.055$, $P = .03$ for mTICI 3 versus mTICI 0–2b). A higher MD index means a smaller decrease of MD values within the salvaged gray matter in comparison with the contralateral side.

Association of Micro- and Macrostructural Changes to the Clinical Outcome

The following analyses showed the association of studied microstructural changes in the salvaged penumbra as well as infarction growth with clinical parameters: a significant negative correlation was found between the MD index and the NIHSS score at the time of MR imaging ($r = -0.43, P = .01$) and NIHSS score between admission and the time of MR imaging ($r = -0.43, P = .01$) and mRS after 90 days ($r = -0.37, P = .04$). A significant positive correlation was found between the MD index and percentage improvement of the NIHSS score between admission and the time of MR imaging ($r = 0.49, P = .003$). A significant positive correlation was found between infarction growth and the NIHSS score at the time of MR imaging ($r = 0.34, P = .05$) and mRS after 90 days ($r = 0.43, P = .02$). A significant negative correlation was found between the infarction growth and the percentage improvement of NIHSS scores between admission and the time of MR imaging ($r = -0.409, P = .02$).

After controlling for infarction growth, the MD index was negatively correlated with the NIHSS score at the time of MR imaging ($r = -0.28, P = .11$) and mRS after 90 days ($r = -0.35, P = .05$) and positively correlated with percentage improvement of the NIHSS score between admission and the time of MR imaging ($r = 0.37, P = .03$). All analyses were corrected for the above-mentioned covariates under the precondition of complete
reperfusion. The association between the MD index and the NIHSS score as well as NIHSS score improvement are shown in Fig 2.

**Impact of Tissue Integrity on the Relationship between Complete Reperfusion and Clinical Outcome**

The percentage improvement of the NIHSS scores between admission and the time of MR imaging differed significantly between the subgroups of complete reperfusion (mTICI 3: median [IQR] 0.9 [0.7–1]) and incomplete reperfusion (mTICI 0–2: median [IQR] 0.47 [0.24–0.87], P = .02). In addition, an association between reperfusion success and imaging parameters, for example, the MD index, was found (P = .03) (see the section Results: Microstructural Changes of the Salvaged Penumbra).

The mean ± SD volume of infarction growth also showed significantly different values, depending on the reperfusion success (28.4 ± 48.9 mL for incomplete and 9.24 ± 17.0 mL for complete reperfusion, P = .03).

In a mediation analysis (Fig 3), the effect of reperfusion success (measured by mTICI) on the clinical presentation (measured by percentage improvement of the NIHSS score between admission and the time of MR imaging) (total effect c = 0.22, SE = 0.09, P = .02) was absent when controlling for imaging parameters such as infarction growth and the MD index (direct effect c’ = 0.03, SE = 0.08, P = .72); critically, the bootstrapped 95% CI for the indirect effect (ie, mediation: total − direct effect) was different from zero for the mediator variable infarction growth (95% CI, 0.04–0.22) as well as for the MD index (95% CI, 0.004–0.19) under consideration of the covariates, which indicated that both imaging parameters significantly mediated the relationship between reperfusion success and clinical improvement.

**DISCUSSION**

The present study shows that the macroscopic salvaged penumbra after MT includes areas of microstructural integrity changes related to the preceding hypoperfusion. The existence of
microstructural integrity alterations was evaluated by using tissue integrity measurements by assessing diffusion parameters, such as the MD in a sample of 65 patients early after MT of an LVO of the anterior circulation. Compared with the contralateral side, a relative decrease of gray matter MD values within the salvaged penumbra was found. There was an explicit dependency on reperfusion success, which suggests that incomplete reperfusion results in further tissue abnormalities beyond macrostructural ischemia. Microstructural changes in the salvaged penumbra were further associated with neurologic and functional deficits, underlining their potential clinical importance.

The salvaged penumbra was identified by applying a co-registration process between DTIs in the acute poststroke phase and admission CT perfusion images, which visualize the tissue at risk. This penumbral tissue was assessed via RAPID software, which has been used in numerous studies.\(^2\),\(^3\),\(^17\),\(^18\) Penumbral tissue, which did not undergo final infarction in poststroke DTIs, was defined as the salvaged penumbra. Previous analyses have suggested that microstructural alterations also occur in the salvaged penumbra,\(^8\)-\(^10\) which was affirmed in the present structural imaging study.

The MD alterations found in the acute poststroke phase with lowered values in the gray matter suggest cytotoxic-like processes with a decrease of the MD but on the microstructural, not the apparent level.\(^19\) As a result of this, assessing the cortical MD asymmetry compared with the contralateral nonaffected hemisphere has the potential to detect changes of the molecular diffusion rate, which can be affected by subtle integrity alterations.\(^20\) The asymmetry index was already used in several previous DTI studies, mostly in the context of FA (fractional anisotropy) calculations, and appeared as an accessible predictor variable.\(^21\)-\(^23\) MD loss within the salvaged penumbra that was temporarily undersupplied could either be a consequence of the primary ischemic processes when the tissue was at risk (eg, microembolization) or could arise out of ischemic reperfusion, which could exacerbate a primary ischemic injury. Results of studies with animal models found diffusion restriction early after stroke within the reversible penumbra similar to the findings in the present study in humans, and the studies discuss the phenomenon of reperfusion injury extensively.\(^24\)

The observed “secondary deterioration” of diffusivity could be explained by a secondary energy failure after reperfusion,\(^25\),\(^26\) but the pathophysiological mechanism and processes of reperfusion injury are not yet conclusively clarified in humans. Existing pathophysiologic concepts explain the loss of diffusivity by a decline in extracellular water induced by a water shift from extracellular to intracellular space. This shift comes from cell depolarization because of a failure of adenosine triphosphate (ATP)-dependent pumps, which ATP levels are deteriorated by the over-release of toxic intermediates in reperfusion injury.\(^24\),\(^27\)-\(^29\) Despite the underlying processes that cause such an observed integrity change in the salvaged penumbra, these alterations may be substantial with concern for technical, procedural, or clinical aspects.

A theoretic attempted explanation of microstructural changes of the salvaged penumbra would relate to clot fragmentation and distal microemboli, which occur within previously unaffected vascular territories as well as within the occlusion-dependent brain tissue during mechanical recanalization. Thus, the penumbral tissue would be less affected by lessening microembolic damage during recanalization. This explanation would fit the above-mentioned theory of primary ischemic processes as a reason for MD loss caused by microembolization into the penumbral tissue. It encourages the usage of proximal flow arrest during mechanical recanalization, which was recently recommended by reporting improved procedural, angiographic, and clinical outcomes.\(^30\)

The study results show a relationship between the MD values of the penumbral tissue and the clinical presentation at the time of MR imaging acquisition with a loss of MD correlated to a more significant concern measured by the NIHSS score. It is also associated with less percentage improvement of the NIHSS score between admission and the time of MR imaging and higher mRS after 90 days. Thus, MD loss has a negative association with the clinical presentation measured in the present study within the clinical workflow. Consequently, effort should be invested to minimize penumbral integrity changes within the therapeutic setting of an LVO.

A focus should be on gaining complete reperfusion because this is associated with a better clinical outcome as well as a reduced infarction growth and a better-preserved penumbral integrity as shown in the present study. By performing a mediation analysis on the relationship between reperfusion success and clinical outcome, imaging parameters of the acute poststroke phase, such as the infarction growth as well as the penumbral integrity, turned out to be significant mediators. By the disappearance of this known relationship, such imaging parameters gain importance because they can explain the impact of recanalization on clinical outcome, which again raises the meaningfulness of applying procedures with the aim of complete reperfusion and less macro- and microstructural tissue damage.

However, it is important to discuss that the observed microstructural integrity changes within the penumbra are measured at a single time within the acute poststroke phase. Reversibility of these abnormalities, which are well known even for macrostructural diffusion restrictions, was not proved. The results present microstructural diffusion restrictions within the penumbra, which is most probably based on the primary ischemic process and influenced by the therapeutic intervention (which was shown for reperfusion success). Follow-up imaging is required to examine reversibility, which was not the content of the present study. The first evidence of sustainable changes in terms of permanent damage could be derived from the existing association between microstructural penumbral changes and midterm functional outcome.

With regard to methodologic aspects, the analyses of tissue alterations were restricted to the gray matter and analyzing white matter integrity was not the content of the present study. By constructing the methodologies like this, the following aspects were considered: during ischemic processes, gray matter is more sensitive to infarction, determined by the ischemic time, than white matter, which is secondarily dependent on the related gray matter in respect to processes like early disintegration. Thus, measuring the processes within the gray matter may reflect the ischemic processes more accurately and have an impact on further progressions, such as degeneration, which is important for the...
clinical outcome. However, studying white matter integrity by using fiber-tracking should be considered in further studies, which would, additionally, gain importance by creating a longitudinal approach.

CONCLUSIONS

The macroscopic salvaged penumbra included areas of microstructural integrity changes early after MT of an LVO in the anterior circulation. Penumbral microstructural integrity seemed to be dependent on recanalization success and was associated with clinical outcome. If confirmed, it harbors another rationale for further development of procedural processes of MT, with the aim for complete reperfusion, less ischemic macro- and microstructural injury, and, as a result of this, a better clinical outcome. The method used in this work enables the early consideration of the outcome on the microstructural level beyond the classic clinical scores.

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