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Multiple Inflow Pulsed Arterial Spin-Labeling Reveals Delays in the Arterial Arrival Time in Minor Stroke and Transient Ischemic Attack

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TECHNICAL NOTE

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SUMMARY: Our purpose was to use multiple inflow pulsed ASL to investigate whether hemodynamic AAT information is sensitive to hemispheric asymmetry in acute ischemia. The cohorts included 15 patients with acute minor stroke or TIA and 15 age-matched controls. Patients were scanned by using a stroke MR imaging protocol at a median time of 74 hours. DWI lesion volumes were small and functional impairment was low; however, perfusion abnormalities were evident. Prolonged AAT values were more likely to reside in the affected hemisphere (significant when compared with controls, P < .048). An advantage of this ASL technique is the ability to use AAT information in addition to CBF to characterize ischemia.

ABBREVIATIONS: AAT = arterial arrival time; ASL = arterial spin-labeling; AUH = area under the histogram; CBF = cerebral blood flow; DSC = dynamic susceptibility contrast; DWI = diffusion-weighted imaging; FMRIB = Functional MRI of the Brain; GRASE = gradient and spin-echo; ICA = internal carotid artery; MCA = middle cerebral artery; NIHSS = National Institutes of Health Stroke Scale; PASL = pulsed arterial spin-labeling; TIA = transient ischemic attack

A SL is a noninvasive MR imaging technique capable of providing perfusion information without the use of gadolinium-based contrast agents. ASL relies on contrast by magnetically labeling blood water and detecting the signal intensity as a tracer bolus; it has shown promise in clinical studies involving acute stroke^{1,2} and ICA occlusion. The shortcomings of the ASL techniques include the following: 1) limited brain coverage, 2) low signal intensity—to-noise ratio, and 3) accounting for delays in AAT (the time duration for blood to move from tagging to imaging locations). The current study attempts to address each of these issues.

An acute cerebrovascular event may increase the AAT by the following mechanisms: Labeled blood must travel via either collateral pathways of the circle of Willis or secondary collateral pathways, such as those that are provided by leptomeningeal vessels. Labeled blood that remains in the large-vessel intravascular space at the time of imaging, due to an insufficient postlabel delay as in continuous ASL or an insufficient TI as in pulsed ASL, will affect the CBF image. A recent ICA occlusion study highlighted the relevance of AAT in clinical perfusion ASL. The goal in the current study is to demonstrate the clinical utility of a whole-brain 3D-GRASE-PASL implementation with multiple-inflow periods to derive maps of CBF and AAT in patients with minor stroke/TIA. Analo-

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gous to hemodynamic timing parameters used in DSC perfusion for acute stroke diagnosis, AAT maps, hypothetically, can be used to characterize the extent of perfusion abnormalities.

Technique

MR imaging data were collected on a 3T scanner (Tim Trio; Siemens, Erlangen, Germany) with a 12-channel head receive coil in a 2-cohort study: 1) 15 patients with acute minor stroke or TIA, and 2) 15 age-matched controls. Carotid Doppler sonography was performed on patients only. Relevant sequences in the protocol were the following: fluid-attenuated inversion recovery, DWI, and pulsed ASL with flow-sensitive alternating inversion recovery preparation and a 3D GRASE readout. D-GRASE-PASL data were collected at 9 inflow times (TI: start = 500 ms, end = 2500 ms, increments = 250 ms; imaging volume: $200 \times 200 \times 120 \text{ mm}^3$; $64 \times 64 \times 24 \text{ matrix}$; voxel dimensions: $3.1 \times 3.1 \times 5.0 \text{ mm}^3$; TR/TE = 3156/39.9 ms). ASL parameters for patients were 8 controls/tags, 9 TIs, and 7:30 scan duration. ASL parameters for the age-matched healthy cohort were 11 controls/tags, 11 TIs, and 11:30 scan duration.

ASL images were spatially smoothed (full width at half maximum = 6 mm). Difference images were averaged at each TI, producing an ASL kinetic curve to which a 2-parameter model⁸ was fit for each voxel by using least squares in Matlab (MathWorks, Natick, Massachusetts), and fit confidence intervals were analyzed for CBF and AAT. Units for AAT are output as seconds, while CBF milliliters per 100 grams of tissue per minute quantification requires additional steps, as described elsewhere.⁹

AAT patterns were characterized in the age-matched cohort to create an expected distribution. It has recently been shown that AAT varies regionally and is correlated with brain volume and sex in young adults. AAT patterns are defined by the vascular territories, and we surmise that AAT is age-dependent, thus the motivation for the age-matched cohort. AAT maps were analyzed in Talairach coordinate space. Absolute AAT depends on the thickness of the imaging volume, but relative differences between brain regions or hemispheres are independent of imaging parameters. Therefore, an asymmetry metric was devised to compare the distribution of AAT values in the affected and unaffected hemispheres. The AAT asymmetry index was calculated by using the AAT area AUH (AAT_{AUH} in the af-

Table 1: Patient demographics with diagnoses

				Stenosis						
Patient		Age	((%)	DWI	NIHSS	Barthel	Time to	Clinical	
No.	Sex	(yr)	Left	Right	(mL)	Score	Index ^b	Scan (hr)ª	Diagnosis	Report (Outcome)
1	F	48	96	0	49.10	1	100	104	Stroke	Left frontal embolic
2	M	66	15	0	6.10	3	60	17	Stroke	Left embolic MCA
3	M	85	0	100	6.25	3	50	103	Stroke	Right embolic
4	M	81	70	30	3.21	1	90	72	Stroke	Right embolic
5	M	92	50	30	1.56	0	90	74	Stroke	Left subcortical
6	M	70	30	0	0.89	0	100	194	TIA	Left, unknown etiology
7	M	71	50	10	0.00	0	100	45	TIA	DWI negative, no lesion
8	M	81	60	40	0.76	0	55	60	Stroke	Left cortical
9	F	70	65	35	0.73	0	100	44	TIA	Right frontal embolic
10	M	69	65	100	5.97	1	95	96	Stroke	Left frontal embolic
11	F	68	10	70	0.00	0	100	112	TIA	DWI negative, no lesion
12	M	79	60	90	0.46	0	100	44	Stroke	Right embolic MCA
13	M	82	70	30	0.48	0	100	68	Stroke	Left embolic MCA
14	M	62	40	70	0.86	1	100	143	Stroke	Left thalamic
15	M	84	45	50	4.21	6	55	131	Stroke	Right frontal embolic

a Time to scan is the time in hours from symptom onset to MR imaging

^b Barthel Index of Activities of Daily Living

Table 2: Patient statistics ^a											
Statistics	Age (yr)	DWI (mL)	NIHSS Score	Barthel Index ^b	Time to Scan (hr)						
Mean	74	5.4	1	86	87						
Minimum	48	0	0	50	17						
Maximum	92	49	6	100	194						
Median	71	0.9	0	100	74						
SD	11	12.3	2	20	46						

^a Time to scan is the time in hours from symptom onset to MR imaging ^b Barthel Index of Activities of Daily Living

fected hemisphere - AAT $_{\rm AUH}$ in the unaffected hemisphere) / (total AAT_{AUH}). A positive index indicates prolonged AAT in the affected hemisphere (asymmetry range: -1 to 1). A similar index was devised for controls by using (left AAT $_{\rm AUH}-{\rm right\,AAT}_{\rm AUH})$ / total AAT $_{\rm AUH}$. An unpaired 2-tailed t test was used to compare AAT hemispheric asymmetry between patients and controls.

Discussion

Tables 1 and 2 show demographic data for the 15 patients. DWI lesion volumes were small (median volume = 0.9 mL; range = 0-49 mL), consistent with the clinically minor strokes in this group. 12 Mean NIHSS score was 1 ± 2 . There was no significant difference in age between cohorts (patients = 74 ± 11 years, controls = 69 ± 5 years; P = .24).

Figure 1 shows the DWI, CBF (milliliters per 100 grams per minute), and AAT (seconds) maps for 3 patients scanned acutely, ordered with decreasing DWI lesion volumes. Patient 1 had a large DWI infarct despite an NIHSS score of 1. The corresponding CBF and AAT maps show regions of hypoperfusion and delayed AAT in the affected hemisphere. Patients 12 and 9 had small DWI infarcts. While Patient 12 has regions of pronounced AAT delay, findings on CBF and AAT maps for patient 9 are relatively normal.

Figure 2 shows AAT asymmetry data for patients and controls. The control cohort demonstrates a symmetric AAT hemispheric index that is centered near zero with low variance. Patient AAT hemispheric metrics favor the affected hemisphere, suggesting prolonged AAT, and are significantly different compared with those in controls (P < .048).

A whole-brain noninvasive ASL technique capable of map-

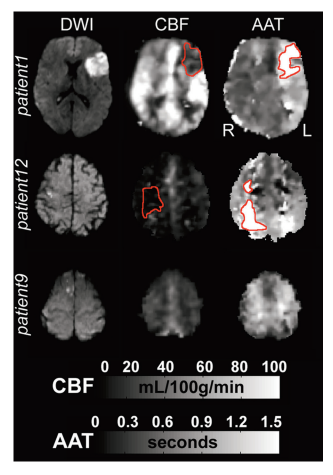


Fig 1. Acute images for 3 patients with decreasing infarct volumes. Patient 1 shows a delayed AAT region (hyperintensity) in the affected hemisphere, where CBF is reduced (hypointensity). Patient 12 shows reduced CBF in the affected hemisphere and delayed AAT, while maps for Patient 9 have normal findings. Abnormal CBF and AAT regions are outlined in red

ping CBF and AAT in patients with minor stroke/TIA is presented. CBF data were consistent with previous ASL studies.^{2,3} Our PASL acquisition, however, was designed as a bolustracking experiment and provided evidence that AAT can be used to characterize perfusion abnormalities.

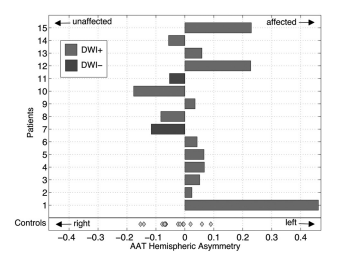


Fig 2. AAT hemispheric asymmetry index is displayed as a bar plot for each patient. Control cohort data are more symmetric about zero with less variance. Patient data are significantly different from those of controls (P < .048). Light gray indicates DWI-positive; dark gray, DWI-negative; diamond datum, control participant.

CBF and AAT maps revealed asymmetric perfusion patterns in some patients, defined by delayed AAT in patients in whom acute symptoms were minor or had resolved. This result is not unexpected because previous studies have shown perfusion deficits in 32% of patients with TIA. This technical report demonstrates the feasibility of multi-TI PASL in acute stroke and provides support for future efforts to characterize longitudinal changes.

There are 2 main advantages of this multi-TI 3D-GRASE-PASL: 1) whole-brain coverage allowing visualization of perfusion patterns in patients with only a small embolic infarct in superior regions (eg, patient 9), and 2) calculation of AAT, with a temporal resolution of 250 ms. Others have reported previously that AAT artifacts were found in 7 of 15 patients due to an insufficient postlabeling delay in some patients when using continuous ASL.² This issue is minimized in the current study because both CBF and AAT are estimated.

There are some important caveats: 1) Stroke may result in an AAT that is beyond the maximum TI in the current study (ie, $AAT_{max} = 2500 \text{ ms}$), 2) The presence of large-vessel atherosclerosis likely influenced our AAT estimates. In DSC per-

fusion MR imaging, timing parameters are often used to delineate perfusion deficits, though they tend to overestimate the ischemic region. It is conceivable that the AAT also shares this same issue of overestimating the perfusion deficit. The degree to which extracranial and intracranial stenoses influence AAT is underinvestigated, and further studies are required.

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