The Effects of Acetazolamide on the Evaluation of Cerebral Hemodynamics and Functional Connectivity Using Blood Oxygen Level–Dependent MR Imaging in Patients with Chronic Steno-Occlusive Disease of the Anterior Circulation

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The Effects of Acetazolamide on the Evaluation of Cerebral Hemodynamics and Functional Connectivity Using Blood Oxygen Level–Dependent MR Imaging in Patients with Chronic Steno-Occlusive Disease of the Anterior Circulation

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ABSTRACT

BACKGROUND AND PURPOSE: Measuring cerebrovascular reactivity with the use of vasodilatory stimuli, such as acetazolamide, is useful for chronic cerebrovascular steno-occlusive disease. The purpose of this study was to evaluate the effects of acetazolamide on the assessment of hemodynamic impairment and functional connectivity by using noninvasive resting-state blood oxygen level–dependent MR imaging.

MATERIALS AND METHODS: A 20-minute resting-state blood oxygen level–dependent MR imaging scan was acquired with infusion of acetazolamide starting at 5 minutes after scan initiation. A recently developed temporal-shift analysis technique was applied on blood oxygen level–dependent MR imaging data before and after acetazolamide infusion to identify regions with hemodynamic impairment, and the results were compared by using contrast agent–based DSC perfusion imaging as the reference standard. Functional connectivity was compared with and without correction on the signal by using information from temporal-shift analysis, before and after acetazolamide infusion.

RESULTS: Visually, temporal-shift analysis of blood oxygen level–dependent MR imaging data identified regions with compromised hemodynamics as defined by DSC, though performance deteriorated in patients with bilateral disease. The Dice similarity coefficient between temporal-shift and DSC maps was higher before (0.487 ± 0.150 by using the superior sagittal sinus signal as a reference for temporal-shift analysis) compared with after acetazolamide administration (0.384 ± 0.107) (P = .006, repeated-measures ANOVA). Functional connectivity analysis with temporal-shift correction identified brain network nodes that were otherwise missed. The accuracy of functional connectivity assessment decreased after acetazolamide administration (P = .015 for default mode network, repeated-measures ANOVA).


ABBREVIATIONS: ACZ = acetazolamide; BOLD = blood oxygen level–dependent MR imaging; DMN = default mode network; SMN = sensorimotor network; SSS = superior sagittal sinus; Tmax = time-to-maximum of the residue function; TS = temporal-shift

The measurement of cerebral perfusion can aid in the characterization of patients with cerebral ischemic diseases.1,2 Recent studies have demonstrated that the determination of diffusion-perfusion mismatch provides a valuable paradigm for selecting a subpopulation of patients with acute stroke most likely to benefit from reperfusion therapies.3–8 However, MR perfusion imaging is typically based on DSC with bolus injection of a gadolinium-based contrast agent.5,8–10 Although the risk of nephrogenic systemic fibrosis associated with the use of gadolinium-based contrast agents may be minimized through renal function screening, there are recent concerns about chronic deposition of gadolinium in the brain.11 The use of a contrast agent may furthermore preclude repeat perfusion scans in the same session,12 which are needed in clinical settings such as the evaluation of cerebrovascular reactivity.

The development of noninvasive approaches without the need for contrast agent administration can provide useful alternatives. Although arterial spin-labeling is a noninvasive method for measuring CBF, it is prone to errors in regions with a long arterial
transit time of blood,\textsuperscript{12-14} which is particularly problematic in patients with steno-occlusive disease. Recently, temporal-shift (TS) analysis of the resting-state blood oxygen level–dependent MR imaging (BOLD) signal, which is sensitive to local blood flow and oxygen metabolism,\textsuperscript{15} has been shown to depict regions with cerebrovascular impairment in acute stroke and chronic cerebral hypoperfusion.\textsuperscript{16-18} In addition, compared with the measurement of hemodynamic parameters, the assessment of the functional status of such hypoperfused brain is underinvestigated. A growing body of work supports resting-state BOLD signal possibly being used to evaluate functional brain networks.\textsuperscript{19-23} Leveraging different aspects of the same BOLD acquisition, simultaneous assessment of cerebral hemodynamics and functional connectivity therefore becomes an attractive application of resting-state BOLD.

Traditionally, cerebrovascular reactivity has been an important measure in patients with chronic steno-occlusive disease. The measurement of cerebrovascular reactivity is performed by quantifying cerebrovascular responses to vasodilatory stimuli, such as the administration of acetazolamide (ACZ) or inhalation of air with increased \(CO_2\) concentration (eg, 5\%).\textsuperscript{24-26} Examining the effects of vasodilatory stimuli on TS and functional connectivity analyses may shed light on their physiologic basis and allow development of an operationalized approach to their evaluation. In this study, we aimed to assess the effects of ACZ on the evaluations of hemodynamic impairment and functional brain connectivity by using resting-state BOLD in patients with chronic steno-occlusive disease of the anterior circulation. We hypothesized that TS analysis of BOLD data could identify regions with hemodynamic compromise in patients with chronic cerebrovascular disease, similar to those shown in acute stroke and Moyamoya disease. We further hypothesized that the use of ACZ would affect the results of TS and functional connectivity analyses due to its alteration in neurovascular coupling.

**MATERIALS AND METHODS**

**Participants**

Fourteen patients with chronic steno-occlusive disease of the anterior circulation (mean age, 48 years; range, 31–70 years; 3 men, 11 women) were included in this study. These included 4 patients with idiopathic Moyamoya disease (3 with bilateral disease), 7 patients with atherosclerotic occlusion of the MCA or ICA (2 with bilateral disease), and 3 patients with unilateral atherosclerotic high-grade stenosis of the ICA (see the On-line Table for details). The study was approved by our institutional review board; and all patients underwent specialized acetazolamide–challenge MR imaging. Ten had follow-up scans the next day without ACZ administration.

**Data Acquisition and Preprocessing**

MR imaging was performed at 3T (Tim Trio; Siemens, Erlangen, Germany). BOLD images were acquired by using a gradient-echo EPI sequence: TR/TE = 2000/30 ms, flip angle = 78°, FOV = 220 × 220 mm\(^2\), matrix = 64 × 64, section thickness = 4 mm, 30 sections. The entire BOLD scan spanned 20 minutes. At 5 minutes after the initiation of the scan, 1 g of ACZ dissolved in 10 mL of normal saline was slowly infused intravenously for 3–5 minutes without interruption of the scanning session.

At the conclusion of the BOLD acquisition, DSC imaging was performed by using a gradient-echo EPI sequence (TR = 1500 ms, TE = 40 ms, flip angle = 60°, FOV = 240 × 240 mm\(^2\), matrix = 128 × 128, section thickness = 5 mm, 19 sections) with the injection of 0.1 mmol/kg of gadobenate dimeglumine (MultiHance; Bracco Diagnostics, Princeton, New Jersey) delivered by a power injector at 4 mL/s through an antecubital intravenous access and followed by a normal saline flush at the same rate. Automated arterial input function and venous output function detection, followed by a delay-insensitive deconvolution with a regularization threshold of 15% of the maximum singular value,\textsuperscript{27} were implemented to generate perfusion maps, including CBF, CBV, MTT, and the time-to-maximum of the residue function (\(T_{\text{r}_{\text{max}}}\)). The perfusion maps were further spatially normalized to standard Montreal Neurological Institute space.

Additionally, T1-weighted MPRAGE imaging (TR = 1900 ms, TE = 3.52 ms, flip angle = 9°, FOV = 216 × 256 mm\(^2\), matrix = 216 × 256, section thickness = 1 mm, 176) was performed for anatomic localization.

Preprocessing of BOLD images was performed by using SPM8 software (http://www.fil.ion.ucl.ac.uk/spm/software/spm8). The first and last 5 minutes of data (150 volumes) were used for analysis, representing data acquired before (pre-ACZ) and after (post-ACZ) administration of ACZ, respectively. The last 5 minutes of images were used for post-ACZ assessment because we found that the ACZ effect plateaued approximately 10 minutes after the initiation of infusion. For both pre-ACZ and post-ACZ data, after removal of the first 10 volumes, BOLD images were corrected for the timing of the section acquisition, realigned to the mean image, normalized to the Montreal Neurological Institute space, resampled to 3-mm isotropic voxel size, and spatially smoothed with a Gaussian kernel of 6-mm full width at half maximum. After removal of the linear temporal trend from the processed images, we regressed out the effects of head motion by using estimates (6 parameters) from the above realignment step as confounding factors. Last, the data were bandpass-filtered to retain signal components with temporal frequency between 0.01 and 0.1 Hz.

**Temporal-Shift Maps**

TS maps were calculated from the BOLD data by determining, for each voxel, the temporal offset that maximizes the correlation coefficient between the time-shifted (−6 TR to +6 TR; ie, −12 seconds to +12 seconds) reference signal and the temporal signal of each voxel (On-line Fig 1). The temporal offset was then assigned as the value of the respective voxel in the TS map. We considered 2 options of reference signals: the global mean signal and the average time-series over an ROI within the superior sagittal sinus (SSS). The global mean signal was obtained by calculating the mean temporal signal across the entire brain, including contributions from GM, WM, and CSF. To obtain the SSS reference signal, we calculated a temporal SD map of the BOLD signal to reflect the magnitude of spontaneous signal fluctuation in each voxel. Because the SSS has 100% blood volume compared with a maximum of 5% blood volume in brain tissue, the BOLD signal of the SSS has a large variance compared with other regions. An ROI was therefore
FIG 1. DSC $T_{\text{max}}$ (A) and temporal-shift maps (B–E) derived from resting-state BOLD data by using the global mean signal (B and D) and the superior sagittal sinus (C and E) as references, as well as before (B and C) and after (D and E) acetazolamide administration in a patient with occlusion of the left ICA. Visually, TS maps exhibited a global resemblance to $T_{\text{max}}$ maps. This agreement decreased after ACZ administration. Compared with using the global signal as reference, TS maps generated by using the SSS signal as the reference signal showed a better agreement with $T_{\text{max}}$ maps. The images are in radiologic convention.

manually placed in a brain area with a high SD in the SSS without including adjacent brain tissues (4 voxels for each subject).

While the success of TS analysis depends on the optimal selection of a reference signal (ie, the identification of “normal” tissues), this information could, in turn, be provided by the results of TS analysis. We therefore proposed an iterative approach to optimize the computation of TS maps by improving estimation of the reference signal as shown below. The average time-series of all voxels with a zero time delay (offset = 0 second) from the previous iteration was calculated and used as the new reference signal for the next iteration. The mean TS value over the whole brain was also calculated. The iteration was repeated until convergence in that the absolute difference of mean values between the TS maps in the successive iterations was lower than 0.001 second.

**Functional Connectivity Analysis**

The default mode network (DMN) and sensorimotor network (SMN) were identified and evaluated by using a seed-based functional connectivity analysis method on the BOLD data.\(^19,21,28\) Using a functional ROI atlas,\(^29\) we defined the seed region for the DMN in the precuneus/posterior cingulate cortex: Montreal Neurological Institute coordinate = (0, −56, 28), radius = 10 mm. The seed ROI for the SMN was placed in the precentral gyrus of the unaffected or less affected hemisphere: Montreal Neurological Institute coordinate = (±44, −16, 46), radius = 6 mm. For each network, the average temporal signal in the seed ROI of the preprocessed images was calculated, and then the Pearson correlation coefficient $r$ with this seed temporal signal was calculated for all brain voxels to generate the functional connectivity maps. To evaluate the effect of correction by using temporal-shift information from the TS analysis on functional connectivity analysis, we shifted the temporal signal of each voxel according to the value in the TS map; the functional connectivity analysis was then repeated. This correction is termed “TS correction.”

**Image and Statistical Analysis**

A mixed-effects model was used to study the correlation between the TS and $T_{\text{max}}$ maps and the effect of ACZ administration (pre-ACZ versus post-ACZ) on this correlation. The mean values of BOLD TS and $T_{\text{max}}$ were calculated in regions with $T_{\text{max}}$ values binned in different ranges (0–1 second, 1–2 seconds, 2–3 seconds, 3–4 seconds, 4–5 seconds, 5–6 seconds, and >6 seconds). The linear relationship between the mean TS and $T_{\text{max}}$ was estimated by modeling subject-specific slope as a random effect.

To further assess the potential clinical value of the TS map, we evaluated the spatial overlap between regions with perfusion deficits as defined by the TS and DSC $T_{\text{max}}$ maps by using the Dice similarity coefficient.\(^18\) The Dice coefficient between regions A and B was defined as the following: Similarity = $2 |A \cap B|/(|A| + |B|)$, where $|A \cap B|$ represents the area of the overlapped region between A and B and $|A|$ and $|B|$ represent the areas of A and B, respectively. Because the threshold of $T_{\text{max}}$ for quantifying pathologic tissue volumes is yet to be established in patients with chronic cerebrovascular disease, we used 4 seconds as the threshold, which was an established threshold for estimating hypoperfused tissue volumes in patients with acute stroke.\(^27,30\) Additional analysis by using $T_{\text{max}} > 3$ seconds as a threshold was also performed to improve the sensitivity for identifying areas with hemodynamic compromise. The optimal threshold for a TS map that maximized its Dice similarity with its respective $T_{\text{max}}$ map was derived by varying the threshold from −12 to 12 seconds with increments of 2 seconds (ie, TR). The values of the Dice similarity were compared by using 2-way repeated-measures ANOVA to test the effect of ACZ administration (pre-ACZ versus post-ACZ) and the choice of reference signal (the global signal versus the SSS signal) on TS analysis.

The Dice similarity coefficient between patient functional networks and templates of brain networks derived from healthy-subject resting-state BOLD data\(^29\) was calculated to assess the effects of TS correlation and ACZ on functional connectivity assessment. Two-way repeated-measures ANOVA was performed to test whether ACZ administration (pre-ACZ versus post-ACZ) and TS correction (original versus TS-corrected analysis) influenced the assessment of functional connectivity.

**RESULTS**

The mixed-effects model showed significant correlation between the TS and $T_{\text{max}}$ maps when using either the global signal or the SSS as the reference signal for calculating TS maps ($P < .001$ for both tests). The slopes of change in TS with respect to $T_{\text{max}}$ were significantly lower after ACZ administration when using the
global signal ($P = .001$) and SSS ($P = .026$) as the reference signal, respectively (On-line Fig 2).

Figure 1 and On-line Fig 3 show the comparison of resting-state BOLD TS maps and DSC $T_{\text{max}}$ maps in representative patients. Brain regions with a long $T_{\text{max}}$ were associated with a positive TS value in the temporal-shift analysis (a delay in the time course with respect to the reference signal). The On-line Table shows the Dice similarity coefficient between the $T_{\text{max}}$ and TS maps in defining brain regions with compromised perfusion for each patient. Using $T_{\text{max}} > 4$ seconds as a threshold for compromised perfusion, 2-way repeated-measures ANOVA showed that after ACZ administration, the Dice similarity coefficient between TS and $T_{\text{max}}$ maps was significantly reduced compared with that before ACZ administration ($P = .006$) (Fig 2A; see the On-line Table for detailed statistics).

When we used $T_{\text{max}} > 3$ seconds as a threshold for compromised perfusion, significant effects were found related to the choice of different reference signals ($P = .008$), ACZ administration ($P = .003$), and the interaction between them ($P = .020$) on the similarity coefficient between TS and $T_{\text{max}}$ maps (Fig 2B and On-line Table). Post hoc Bonferroni-corrected paired $t$ tests showed the following: 1) before ACZ administration, TS maps derived by using SSS as the reference signal demonstrated higher similarity with $T_{\text{max}}$ maps, compared with those using the global signal as the reference signal ($P < .001$); and 2) lower similarity between $T_{\text{max}}$ and TS maps was found after ACZ administration compared with before ACZ administration when using either the global signal ($P = .011$) or SSS ($P = .002$) as the reference signal (Fig 2B and On-line Table). For the above analysis, the DSC perfusion after ACZ administration was used. The same analysis was repeated in a subset of 10 patients who underwent a second-day MR imaging examination without the administration of ACZ, and in these cases, the pre-ACZ second-day $T_{\text{max}}$ was used. Similar results were obtained (On-line Fig 4).

The default mode network and sensorimotor network of representative patients are shown in Fig 3 and On-line Fig 5. Although the results of functional connectivity changed after ACZ administration, both the DMN and SMN remained identifiable. TS correction uncovered new network nodes in areas with hemodynamic compromise in some patients. In general, these emerged nodes could be observed in the corresponding functional networks of healthy subjects. Temporal-shift information obtained from TS analysis by using the SSS signal as the reference was used to correct the BOLD signal for functional connectivity analysis because of its better performance in delineating regions with hemodynamic impairment.

Two-way repeated-measures ANOVA showed significant effects of TS correction on the functional connectivity assessment of the DMN ($P < .001$) and the SMN ($P = .024$) (Fig 4). Accuracy in the assessment of both functional networks (as measured by the overlap with template network masks) increased after correction of the signal by using time-shift estimates from the TS analysis. Assessment accuracy decreased after ACZ administration, with statistically significant effects for the DMN ($P = .015$) and a trend toward significance for the SMN ($P = .080$).

**DISCUSSION**

The present study demonstrated that temporal-shift maps derived from noninvasive resting-state BOLD scans could identify brain regions with abnormal per-
ever, differences between TS and Tmax maps were also observed.
The exact mechanism for potential discrepancies is not completely
understood, but the differences are not surprising because TS analy-
sis and DSC perfusion imaging likely reflect different aspects of the
same pathophysiologic process. Further development of the tech-
nique and validation within larger cohorts are necessary to further
establish the nature of this relation and to further assess TS maps as
an adjunct or potential alternative to DSC perfusion imaging in pa-
ients with neurovascular disease.

Two options of the reference signals were considered in the
temporal-shift analysis: the global mean signal and the signal
from the SSS. While both approaches produced TS maps compa-
rable with $T_{max}$ maps, a higher similarity was observed between
$T_{max}$ maps and TS maps obtained by using the SSS signal as the
reference. This suggests that the largely fluctuating signal in the
venous sinus and the iterative strategy ensured a better assessment
of hemodynamic compromise. Furthermore, while global signal
can be easily calculated even without operator input, the accuracy
of the reference may be contaminated by inclusion of neural ac-
tivity31 and the hypoperfused brain regions. The problem became
particularly apparent when hypoperfused regions were large, such
as in patients with bilateral steno-occlusive disease. We also found
that the time delay evaluated by using the global signal as a refer-
cence tended to be smaller compared with that calculated with the
SSS as reference. This finding could be because the global signal
showed a time lag from normal areas because of the contribution
from the hypoperfused areas, which is weighted by the volumes of
these areas. In contrast, although the SSS receives blood from
hypoperfused areas as well, the contribution from these areas is
weighted by their blood flow; and because of the reduced blood
flow in these areas, the contribution from them is smaller.

When we used the temporal delay information calculated
from resting-state BOLD data to correct for the functional con-
nectivity analysis of the DMN and the SMN, new areas of both
functional networks were found, particularly in regions with hemo-
dynamic compromise. Such areas generally emerged in locations
where brain networks of healthy participants typically reside. These
findings suggest caution when evaluating functional connectivity in
patients with cerebrovascular compromise. For example, an appar-
cient reduction in connectivity could be recovered by TS correction
and might not be caused by changes in the underlying neuronal ac-
tivity but rather by changes in neurovascular coupling.

It has been reported that neuronal function and underlying
vascular coupling could not be correctly evaluated by
BOLD MR imaging in the setting of neurovascular ischemia, pur-
purportedly due to the uncoupling of the positive relationship be-
tween CBF and BOLD responses under physiologic baseline con-
ditions.26,32,33 In this study, changes in the detected DMN and
SMN were observed after ACZ administration, though spontane-
ous neural activity should be unaltered without explicit external
neural stimuli. Our finding consequently provides further evi-
dence for the altered assessment of neuronal function of BOLD
MR imaging after a vasodilatory challenge. Also, TS analysis needs
to be applied to evaluate neuronal activity by using BOLD MR
imaging in patients with steno-occlusive disease, due to the pos-
sibility of compromised neurovascular coupling. Moreover, we
found that TS maps exhibited a decreased similarity with $T_{max}$
after ACZ administration. This is consistent with previous find-
ings showing that BOLD responses are reduced in task-based

![FIG 4. The accuracy of functional connectivity assessment, mea-
sure by the Dice similarity coefficient between functional networks
and network templates, for default mode (A) and sensorimotor (B)
networks. The assessment accuracy of both functional networks in-
creased after temporal-shift correction. The assessment accuracy de-
creased after acetazolamide administration for the DMN, while a
trend toward significance was found for the SMN. Error bars repre-
sent the SD of the mean. Asterisks indicate significant differences as
determined by 2-way repeated-measures ANOVA.](ajnr.aacrjournals.org/content/38/1/139/F1.large.jpg)
fMRI after ACZ challenge. 26,33 Although ACZ is useful for interrogating cerebrovascular reactivity, there should be caution in interpreting the results of functional connectivity and TS analyses in BOLD studies after ACZ administration.

Several issues remain to be addressed in future studies. First, while the SSS reference approach provided a good estimate of hemodynamic compromise, the selection of ROIs within the SSS could affect the results. An algorithm for automatic determination of the SSS ROI is required to derive optimal results. Second, a relatively low spatial (voxel size = 3.4 × 3.4 × 4 mm³) and temporal resolution (TR = 2 seconds) BOLD sequence was used in the current study. Higher spatial and temporal resolutions can be achieved via multiband acquisitions, thereby improving the accuracy of the assessment of hemodynamic compromise, similar to that found in resting-state fMRI. 34,35 Third, although there is variability in the TS correction in unveiling otherwise occult nodes in functional networks, whether this reflects the severity of functional compromise is not clear. Further studies are required to systematically investigate whether functional connectivity with temporal-shift correction is associated with functional/cognitive symptoms and whether this information can be used to better predict functional recovery among these patients.

CONCLUSIONS
Temporal-shift analysis of non-contrast agent–based resting-state BOLD data can delineate brain areas with hemodynamic compromise as measured by DSC T max maps in patients with chronic cerebrovascular disease, though the performance of temporal-shift analysis deteriorates in patients with bilateral disease. Temporal-shift correction recovers nodes of functional brain networks in some hypoperfused areas. The use of ACZ changes the effectiveness of temporal-shift analysis in evaluating cerebral hemodynamic compromise as well as functional connectivity analysis from resting-state BOLD data.

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