Acetazolamide-Loaded Dynamic 7T MR Quantitative Susceptibility Mapping in Major Cerebral Artery Steno-Occlusive Disease: Comparison with PET

K. Fujimoto, I. Uwano, M. Sasaki, S. Oshida, S. Tsutsui, W. Yanagihara, S. Fujiwara, M. Kobayashi, Y. Kubo, K. Yoshida, K. Terasaki, and K. Ogasawara

BACKGROUND AND PURPOSE: Dynamic changes in cerebrovascular reactivity after acetazolamide administration vary markedly among patients with major cerebral arterial steno-occlusive disease. MR quantitative susceptibility mapping can dynamically quantify the cerebral magnetic susceptibility. The purpose of this study was to determine whether dynamic changes in susceptibility after administration of acetazolamide on 7T quantitative susceptibility mapping are associated with pre-existing states of CBV and the cerebral metabolic rate of oxygen in major cerebral arterial steno-occlusive disease.

MATERIALS AND METHODS: Sixty-five patients underwent 7T MR imaging at baseline and at 5, 10, 15, and 20 minutes after acetazolamide administration. Differences between the susceptibility of venous structures and surrounding brain tissue were calculated in the quantitative susceptibility mapping images. Susceptibility differences at 5, 10, 15, and 20 minutes after acetazolamide administration relative to baseline were calculated in 97 cerebral hemispheres with major cerebral arterial steno-occlusive disease. CBV and the cerebral metabolic rate of oxygen were also calculated using 15O-gas PET in the resting state.

RESULTS: Dynamic changes of susceptibility after acetazolamide administration were classified into 3 patterns: abnormally increasing 5 or 10 minutes after acetazolamide administration; abnormally decreasing within 20 minutes after acetazolamide administration; and remaining unchanged after acetazolamide administration. CBV was significantly greater in the first pattern than in the latter 2. The cerebral metabolic rate of oxygen differed significantly in descending order from the first to middle to last pattern.

CONCLUSIONS: Dynamic changes of susceptibility after acetazolamide administration on 7T MR quantitative susceptibility mapping are associated with pre-existing states of CBV and the cerebral metabolic rate of oxygen in major cerebral arterial steno-occlusive disease.

ABBREVIATIONS: ACZ = acetazolamide; BOLD = blood oxygen level–dependent; CMRO2 = cerebral metabolic rate of oxygen; CVR = cerebrovascular reactivity; OEF = oxygen extraction fraction; QSM = quantitative susceptibility mapping; RS = relative susceptibility

When the chronic progression of major cerebral arterial steno-occlusive disease reduces cerebral perfusion pressure, cerebrovascular autoregulatory mechanisms attempt to preserve CBF through the dilation of precapillary resistance vessels; this is referred to as stage 1 ischemia. In stage 1 ischemia, cerebrovascular reactivity (CVR) to acetazolamide (ACZ), which reflects the degree of cerebrovascular autoregulatory vasodilation, starts to decrease. However, autoregulatory capacity cannot compensate for further reductions in cerebral perfusion pressure, which result in a decline in CBF; this is referred to as stage 2 ischemia, or misery perfusion. CVR to ACZ is exhausted in this stage. In the clinical setting, CVR to ACZ is measured using perfusion SPECT, dynamic susceptibility contrast perfusion MR imaging, and transcranial Doppler ultrasonography.

MR imaging is capable of quantifying the blood oxygen level–dependent (BOLD) changes in venous structures and/or brain parenchyma induced by differences in magnetic susceptibility between oxy- and deoxyhemoglobin. Wu et al continuously measured CVR to ACZ using BOLD imaging with 3T MR imaging in patients with steno-occlusive disease of major cerebral arteries and demonstrated that overall, BOLD signal began...
increasing immediately after ACZ administration, approaching a plateau at ~8.5 minutes after administration. However, in some patients, BOLD signal showed a more severely reduced relative reactivity at 6–7 minutes after the initiation of ACZ infusion and progressive partial recovery toward a pre-ACZ administration value at 10–11 minutes.16 These findings suggested that dynamic changes to susceptibility after ACZ administration vary markedly among such patients. CVR to ACZ itself may depend on the pre-existing state of vasodilatation, which can be assessed using CBV, and pre-existing dilated vessels show little or no response to ACZ.1–3 We hypothesized that a pre-existing state of cerebral metabolism is another condition impacting CVR to ACZ; ACZ acts via enzyme reaction, and a reduction in this enzyme reaction causes a slow, decreased CVR to ACZ. Cerebral metabolism, which can be assessed using the cerebral metabolic rate of oxygen (CMRO2), may be reduced in such a state.

A postprocessing technique known as quantitative susceptibility mapping (QSM) on MR imaging can quantitatively measure the magnetic susceptibility of venous structures and/or brain parenchyma from T2*-weighted magnitude/phase images, which are easily obtained using a commercial scanner.11 Several investigators have measured CVR to ACZ using QSM on 3T MR imaging and suggested that QSM is sensitive to dynamic modulation of the oxygen extraction fraction (OEF) during hemodynamic augmentation.12 A recent study using QSM with a 7T scanner reported that it offered profound susceptibility effects and optimized postprocessing techniques.13 The scan time for this mapping is relatively short (3 minutes 25 seconds), and it may also enable imaging of dynamic changes in blood oxygenation levels in venous structures and/or brain parenchyma.

Therefore, the purpose of the present study was to determine whether dynamic changes in susceptibility after ACZ administration on 7T QSM are associated with pre-existing CBV and CMRO2 on PET in patients with ICA or MCA steno-occlusive disease.

**MATERIALS AND METHODS**

All procedures performed in studies involving human participants were in accordance with the institutional ethics committee, and written, informed consent was obtained from all subjects or their next of kin before patient participation.

**Healthy Subjects**

To obtain healthy control values from brain 15O-gas PET studies, we enrolled 10 healthy adult men (mean age, 47 ± 8 years; age range, 35–60 years) who had no history of diabetes mellitus, dyslipidemia, or hypertension and had an absence of asymptomatic lacunar infarction and leukoaraisis on conventional brain MR imaging.

**Patient Inclusion Criteria**

We prospectively selected patients who provided written informed consent to participate and who met the following basic inclusion criteria: presence of clinical symptoms suggesting ischemic episodes in the MCA or ICA territory at 1–6 months before visiting our institution; useful residual function (0, 1, or 2 on the mRS); presence of uni- or bilateral MCA (M1 portion) or extra- or intracranial ICA stenosis (>70% or 50% for the ICA and MCA, respectively) or occlusion on angiography; and absence of infarcts in the entire cortical area supplied by the M4 branch of the MCA on T1WI and T2WI, and DWI.

**MR Imaging Protocol and Generation of QSM**

A 7T MR scanner (Discovery MR950; GE Healthcare) with quadrature transmission and 32-channel receive head coils was used. QSM source data were obtained using a 3D spoiled gradient recalled acquisition technique with flow compensation (TR, 30 ms; TE, 15 ms; flip angle, 20°; FOV, 256 mm; acquisition matrix size, 512 × 256; section thickness, 2 mm; number of slices, 160; reconstruction voxel size after zero-fill interpolation, 0.5 mm3; and scan time, 3 minutes 25 seconds). The scan time for this mapping is relatively short (3 minutes 25 seconds), and it may also enable imaging of dynamic changes in blood oxygenation levels in venous structures and/or brain parenchyma. The susceptibility difference was finally displayed with a surface rendering technique with flow compensation (TR, 30 ms; TE, 15 ms; flip angle, 20°; FOV, 256 mm; acquisition matrix size, 512 × 256; section thickness, 2 mm; number of slices, 160; reconstruction voxel size after zero-fill interpolation, 0.5 mm3; and scan time, 3 minutes 25 seconds).13 Magnitude and real/imaginary phase images were regenerated from this acquisition.

First, each patient underwent the above-mentioned MR imaging at baseline. Next, ACZ (1000 mg; range, 13–19 mg/kg body weight) was dissolved in physiologic saline (20 mL); this solution was then administered intravenously for 1 minute. The MR imaging was performed in the same fashion 4 times so that each midscan time was 5, 10, 15, and 20 minutes after the end of intravenous administration of ACZ.

Before the start of this ACZ challenge study, 30 other cerebral hemispheres with ICA or MCA steno-occlusive disease underwent 5 MR imaging scans with the same timing without ACZ administration to assess the minute-to-minute test-retest variability of susceptibility as measured using QSM.

QSM images were generated from the source images using an in-house program with a multiple dipole-inversion combination with k-space segmentation14 and regularization-enabled sophisticated harmonic artifact reduction for phase data methods,15 as described previously.13 A 2D Gaussian low-pass filter with a kernel size of 60% of the total image power in each section was applied to extract iron deposition in deep nuclei, hemosiderin deposition, dural sinuses, and large venous structures, and a 2D Gaussian high-pass filter of 2% was applied to extract small venous structures. Subsequently, small venous structures were determined by multiplying the Gaussian high-pass filter–processed binary images and the logical negations of Gaussian low-pass filter–processed binary images under the threshold for binarization of ±2 SDs.13,15

The difference between the average susceptibility of venous structures and surrounding brain tissue was calculated in each voxel of interest of 25 mm3 in the processed QSM images. The susceptibility difference was finally displayed with a smoothing procedure. To validate a correlation between baseline OEF obtained from QSM images and pre-existing OEF obtained on PET, we generated OEF maps from baseline QSM images before ACZ challenge in accordance with a previous study.13

**Brain 15O-Gas PET Study**

Patients who were scheduled to undergo MR imaging studies with the ACZ challenge underwent PET studies in a resting state without ACZ challenge. PET studies were performed using a SET-3000GCT/M scanner (PET/CT; Shimadzu) (full width at half maximum for in-plane and axial spatial resolutions of 3.5 and 4.2 mm, respectively) within 5 days before MR imaging. Before the PET scans, a 3-minute transmission scan with a
FIG 1. Diagrams showing the 318 ROIs for the 3D stereotaxic ROI template. White ROIs indicate territories perfused by bilateral MCAs.

cesium 137 ($^{137}$Cs) point source was conducted using a bismuth germanate transmission detector ring that was coaxially attached to the gadolinium silica oxide emission detector ring. Next, CBF was measured while the patient inhaled $^{15}$O$_2$ continuously through a mask. The CMRO$_2$ and OEF were measured during continuous inhalation of $^{15}$O$_2$; these data were collected for 5 minutes. CBV was measured during a single breath of $^{13}$O. Finally, CBF, OEF, and CMRO$_2$ were calculated using the steady-state method and then corrected on the basis of CBV.

Image Data Analyses
Image data analyses were performed according to previously reported methods. By means of SPM, Version 12 (http://www.fil.ion.ucl.ac.uk/spm/software/spm12), PET images that were coregistered to QSM source images, as well as QSM images after Gaussian smoothing ($\sigma = 10$ pixel), were warped to Montreal Neurological Institute coordinates. Next, 318 constant ROIs were placed in both the cerebral and cerebellar hemispheres automatically using a 3D stereotaxic ROI template with SPM. The ROIs were then grouped into a total of 10 segments—callosomarginal, pericallosal, precentral, central, parietal, angular, temporal, posterior, hippocampal, and cerebellar—in each hemisphere according to the arterial supply. Five of these 10 segments (precentral, central, parietal, angular, and temporal) perfused by the MCA were then combined and defined as an MCA ROI (Fig 1).

Mean susceptibility differences on QSM images and CBF, CBV, OEF, and CMRO$_2$ on PET images were measured using image-analysis software (ITK-SNAP; www.itksnap.org) in the MCA ROI in the cerebral hemisphere with ICA or MCA steno-occlusive disease. In each MCA ROI of each cerebral hemisphere with ACZ challenge, relative susceptibility (RS, %) was calculated as follows: 100 × susceptibility difference at each time (5, 10, 15, or 20 minutes) after ACZ administration on QSM images. In each MCA ROI of each cerebral hemisphere without ACZ challenge, RS was also calculated as follows: 100 × susceptibility difference on QSM images on the basis of the presence or absence of an abnormal increase or decrease in RS at any time point.

RESULTS
During the 30-month study period, a total of 89 patients successfully underwent 7T MR imaging without severe adverse effects. In 4 of these patients, data sufficient to generate QSM were not obtained because of motion artifacts; these 4 patients were excluded from the analysis. Therefore, 85 patients (65 and 20 with and without ACZ challenge, respectively) were finally enrolled in the study.
Among these 85 patients (46 men, 39 women; mean age, 55 ± 12 years; age range, 30–76 years), 60 had hypertension, 32 had diabetes mellitus, and 55 had dyslipidemia. Thirty-seven, 27, 3, and 17 patients had bilateral ICA (bilateral extracranial diseases in 27; bilateral intracranial diseases in 5; unilateral extracranial disease and contralateral intracranial disease in 5), unilateral ICA (extracranial disease in 22; intracranial disease in 5), the M1 portion diseases of bilateral MCA, and unilateral M1 portion steno-occlusive disease, respectively. The remaining patient had unilateral extracranial ICA and contralateral M1 portion steno-occlusive diseases. Therefore, 127 MCA ROIs of the 127 cerebral hemispheres with MCA or ICA steno-occlusive disease (97 and 30 with and without ACZ challenge, respectively) were eventually analyzed.

The 95% CI of RS values in the second, third, fourth, and fifth MR imaging in 30 MCA ROIs without ACZ challenge were 91.85%–108.55%, 90.81%–109.33%, 90.59%–109.49%, and 90.70%–109.22%, respectively. No difference was observed among each RS.

A significant correlation was observed between baseline QSM-OEF and pre-existing PET-OEF ($r = 0.865$; $P \leq .0001$) (Fig 2).

**Figure 2.** Correlations between baseline QSM-OEF before ACZ administration and pre-existing PET-OEF in 97 MCA ROIs.

**FIG 3.** Chronological changes in RS after ACZ administration in 97 MCA ROIs. Left, middle, and right graphs show the 47 MCA ROIs with abnormally increased RS at ≥1 time point, the 22 MCA ROIs with abnormally decreased RS at ≥1 time point, and the 28 MCA ROIs without abnormally increased or decreased RS at any time point, respectively. Red, blue, and black lines denote ROIs with abnormally elevated CBV, an abnormally decreased cerebral metabolic rate of oxygen, and neither, respectively. Upper and lower dotted horizontal lines denote the upper and lower limits of the 95% CI of controls without ACZ challenge, respectively.
MCA ROIs with abnormally increased RS than in those with abnormally decreased RS, unchanged RS, or controls. No significant differences in CBV were identified in other any comparisons. Of the 22 MCA ROIs with abnormally increased RS, 17 (77%) had abnormally elevated CBV. In contrast, none of the patients with abnormally decreased RS or unchanged RS had abnormally elevated CBV. No significant differences in CMRO₂ were apparent between controls and MCA ROIs with abnormally increased RS. Regarding other comparisons, the CMRO₂ differed significantly in descending order, as follows: controls or abnormally increased RS, abnormally decreased RS, and unchanged RS. Of the 28 MCA ROIs with unchanged RS, 8 (29%) showed abnormally reduced CMRO₂.

In contrast, of the 47 MCA ROIs with abnormally decreased RS, only 7 (14%) had abnormally reduced CMRO₂, and no MCA ROIs with abnormally increased RS had abnormally reduced CMRO₂. Furthermore, in the 47 MCA ROIs with abnormally decreased RS, whereas CBV did not differ significantly in the 11 MCA ROIs with unchanged RS₂ (3.35 ± 0.84 mL/100 g) and the 36 MCA ROIs with abnormally decreased RS₂ (3.40 ± 0.3862 mL/100 g), CMRO₂ was significantly lower in the former MCA ROIs (2.65 ± 0.52 mL/100 g/min) than in the latter (3.06 ± 0.38 mL/100 g/min; P = .0182).

Representative chronological images of susceptibility difference obtained from QSM after ACZ administration and PET images performed before MR imaging in a patient with abnormally increased CBV in the symptomatic cerebral hemisphere are shown in Fig 4.

**DISCUSSION**

The present study demonstrated that dynamic changes in susceptibility after ACZ administration on 7T QSM are associated with pre-existing CBV and CMRO₂ on PET in patients with ICA or MCA steno-occlusive disease.

In this study, ACZ was administered intravenously for 1 minute. This method has been applied for measuring CVR on brain perfusion SPECT in adult patients with ICA or MCA steno-occlusive disease. Considering the adverse effects of ACZ administration performed in the same fashion, 63% of patients reportedly developed only minor symptoms such as headache, nausea, dizziness, tinnitus, numbness of the extremities, and general malaise, and these symptoms resolved within 72 hours after ACZ administration. These patients experienced neither severe general symptoms nor neurologic deficits. Similarly, no patients in the present study experienced such severe adverse effects due to ACZ administration.

A study using ¹⁵O PET demonstrated that while CBF increased at 10 and 20 minutes after ACZ administration in healthy humans, CMRO₂ was unchanged compared with predadministration. Changes in susceptibility after ACZ administration on QSM theoretically reflect both changes in CBF and oxygen metabolism by ACZ. Although whether CMRO₂ after ACZ administration is constant in patients with cerebrovascular diseases remains unknown, dynamic changes in susceptibility after ACZ administration on QSM may more strongly reflect changes in CBF than changes in oxygen metabolism: Increases and decreases in the susceptibility may predominantly indicate decreases and increases in CBF, respectively. One study of CVR to ACZ using QSM on 3T MR imaging showed that OEF decreased from baseline to 16 minutes after ACZ administration in the cerebral hemisphere with anterior circulation steno-occlusive disease, as well as in the cerebral hemisphere without such disease, and the decrease was more substantial in the latter. These changes in susceptibility might be caused by changes of CBF.

In the present study, dynamic changes of susceptibility after ACZ administration were classified into 3 patterns: abnormally increasing, peaking at 5 or 10 minutes after ACZ administration and later returning toward baseline; abnormally decreasing within 20 minutes after ACZ administration and maintaining this decrease; and remaining unchanged after ACZ administration. A dynamic change comprising an abnormal increase in susceptibility early after ACZ administration and subsequent return was comparable with a finding obtained using continuous measurements of BOLD signal after ACZ administration. In MCA ROIs with such dynamic changes, pre-existing CBV was abnormally elevated and pre-existing CMRO₂ was normally maintained. Mean transit time (MMT) is a function of CBV/CBF, and an increase in CBV usually implies a long MMT in MCA or ICA steno-occlusive disease. A long MMT can result in low drug delivery, leading to little or no CVR immediately after ACZ administration in the region affected by such a condition, whereas CVR in surrounding or other brain tissues with a normal MMT reacts rapidly because of normal drug delivery. In addition, the small arterioles or intraparenchymal vessels are dilated in regions with an increase in CBV. These pre-existing dilated
vessels cannot respond immediately after ACZ administration, whereas normal vessels in surrounding or other brain tissue rapidly dilate. These differences in CVR to ACZ administration lead to a sudden reduction in cerebral perfusion pressure of normal vessels relative to pre-existing dilated vessels. This dynamically developed pressure gradient may result in the early steal of blood flow from pre-existing dilated vessels to normal vessels. Subsequently, pre-existing dilated vessels may gradually respond to ACZ, and a vasodilation effect of ACZ in normal vessels may gradually reduce with time. Such mechanisms could explain the abnormal increase in susceptibility early after ACZ administration (early steal phenomenon) and subsequent return.

A dynamic change of the abnormal decrease in susceptibility within 20 minutes after ACZ administration and maintenance of this decrease might imply an increase in CBF—that is, normal CVR to ACZ. Actually, in MCA ROIs showing such a dynamic change, pre-existing CBV was normal. Pre-existing CMRO2 was reduced compared with that in controls but remained within normal limits in 86% of MCA ROIs showing that susceptibility change. On the other hand, pre-existing CMRO2 was reduced more in MCA ROIs with an abnormal decrease in susceptibility ≥10 minutes after ACZ administration than in those with an abnormal decrease in susceptibility 5 minutes after ACZ administration. Because CBV did not differ significantly between these 2 groups, the difference in dynamic change after ACZ administration might be attributable to pre-existing CMRO2. ACZ acts via carbonic anhydrase, and a reduction in its activity slows CVR to ACZ. The activity of carbonic anhydrase may depend on cerebral metabolism. This hypothesis may also explain the maintenance of unchanged susceptibility after ACZ administration. In MCA ROIs with such a dynamic change in susceptibility, pre-existing CBV was normal, but pre-existing CMRO2 was quite markedly reduced; this finding suggests that the activity of carbonic anhydrase was severely reduced and cerebral vessels were unresponsive to ACZ.

Using continuous measurement of CVR to ACZ with BOLD imaging, Wu et al demonstrated that BOLD signal began increasing gradually following ACZ administration, approaching a plateau at ~8.5 minutes after administration in the cerebral hemisphere without ICA or MCA steno-occlusive disease. This dynamic change in susceptibility differed compared with that of MCA ROIs with normal CBV and normal CMRO2 in the present study; an abnormal decrease in susceptibility 5 minutes after ACZ administration and maintenance of this decrease. The difference in peak susceptibility may be due to the method of ACZ administration. ACZ (1000 mg) was administered intravenously for 3–5 minutes in the study by Wu et al and for 1 minute in the present study.

A preoperative decrease in CVR to ACZ or a preoperative increase in CBV in the cerebral hemisphere ipsilateral to carotid endarterectomy or carotid artery stent placement for cervical ICA stenosis or arterial bypass surgery for Moyamoya disease has been identified as a powerful and independent predictor of postoperative cerebral hyperperfusion. Postoperative cerebral hyperperfusion induces unilateral headache, face and eye pain, seizure, and focal symptoms that occur secondary to cerebral edema or intracerebral hemorrhage. This phenomenon, even when asymptomatic, induces slight-but-diffuse brain damage, resulting in postoperative cognitive decline. Identification of reduced CVR to ACZ or elevated CBV therefore enables improved risk stratification for such patients. However, the predictive accuracy of CVR to ACZ on brain perfusion SPECT is not always high, with a positive-predictive value of <50%. Our findings that dynamic changes in susceptibility, particularly susceptibility 5 minutes after ACZ administration, detected increased CBV with high positive (77%) and negative (100%) predictive values suggests that these dynamic changes may predict the risk of cerebrovascular adverse events more accurately than conventional measurements of CVR to ACZ. In addition, the examination time of the present method using 7T QSM was <12 minutes if the susceptibility change was measured until a midscan time of 5 minutes after ACZ administration.

QSM source data were obtained with a section thickness of 2 mm. Such a thick section may reduce the accuracy of susceptibility on QSM due to the partial volume effect for small veins. As another limitation, the scan time for 7T QSM is 3 minutes 25 seconds longer than that for BOLD or QSM imaging with 3T MR imaging. Shortening the scan time may help clarify dynamic changes in susceptibility after ACZ administration. Furthermore, considerable variation in cerebral hemodynamics might be anticipated within an MCA ROI, especially in the borderzone, and use of the MCA ROI might reduce such hemodynamic variation. Ranges of CBV in the cerebral gray matter are quite large. This high heterogeneity might also be reduced using MCA ROI analysis.

CONCLUSIONS

The present study demonstrated that the dynamic changes in susceptibility after ACZ administration on 7T QSM are associated with pre-existing CBV and CMRO2 on PET in patients with ICA or MCA steno-occlusive disease.

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