INTRODUCTION

Intracranial vessel wall MRI (VW-MRI) has recently attracted widespread interest and been increasingly adopted for visualization of extra- and intracranial VW pathology in clinical practice (1). VW-MRI offers advantages for the detection of symptomatic non-stenotic lesions and further characterization of stenotic lesions (2, 3). For intracranial VW-MRI, high signal-to-noise ratio (SNR), contrast-to-noise ratio (CNR), and spatial resolution are needed to visualize thin arterial VWs and characterize VW lesions (1, 4, 5), which inevitably prolong scan time; therefore, it is clinically important to perform VW-MRIs in a reasonable time while maintaining high resolution. To achieve this, scan coverage...
is often compromised with focus on target vessels either in the anterior or posterior circulation, rather than covering all intracranial arteries.

Recently introduced compressed sensing (CS) enables accelerated MRI acquisition using sparse k-space sampling by discarding redundancy in the data acquisition process (6-8). This helps avoid the undersampling artifacts by using incoherent sampling with nonlinear reconstruction (9-11). CS has been adopted for MRI for one of the following benefits: 1) reduced scan time, 2) increased spatial resolution with no increase in scan time, and 3) increased coverage while maintaining spatial resolution and scan time. Most previous studies of VW-MRIs adopted CS in order to reduce scan acquisition time. CS reduced the scan time without significantly sacrificing image quality in both intr- and extracranial VW-MRIs (12-14), while also providing robust and reproducible images (15). Most of the study subjects, however, were healthy volunteers and the clinical usefulness of CS in intracranial VW-MRIs must be validated in a large patient cohort. In addition, little research has been dedicated to whether CS is useful when increasing scan coverage while maintaining a similar scan time.

The assessment of VW enhancement is important for reporting VW-MRI findings. Enhancement pattern details are required to differentiate between intracranial atherosclerosis, arterial dissection, and other causes of luminal narrowing, such as reversible cerebral vasoconstriction syndrome and vasculitis (2, 16). In addition, the presence of enhancement helps identify culprit lesions, most importantly, unstable or ruptured aneurysms (17, 18). Thus, high quality post-contrast images are a prerequisite for VW-MRI interpretation.

In the present study, we applied CS to the post-contrast images from VW-MRI to increase scan coverage to cover both anterior and posterior circulations while maintaining a scan time similar to those without CS. We hypothesized that CS might provide comparable image quality, both quantitatively and qualitatively. The purpose of our study was to compare post-contrast images with and without CS from intracranial VW-MRIs in a large patient cohort.

MATERIALS AND METHODS

Study Population

This retrospective study was approved by the Institutional Review Board of our institution, Severance Hospital (4-2019-0534) and the requirement for informed consent was waived. We searched for all VW-MRIs performed between April 2017 and July 2018 in our institution and found 100 patients' VW-MRIs. The VW-MRIs were performed for various reasons, but the majority of them evaluated the cause of ischemic stroke, the etiology of intracranial stenosis, and the differentiation between atherosclerosis and arterial dissection. All patient MRIs included 3D fat-suppressed post-contrast T1-weighted volumetric isotropic turbo spin echo acquisition (VISTA) with CS (VISTA-CS). An exclusion criteria was established: 1) patients lacking post-contrast T1-weighted VISTA without CS (VISTA-nonCS, n = 13), 2) MRIs with severe motion artifacts (n = 10), and 3) MRIs with errors in the registration process due to significantly different scan ranges between VISTA with and without CS (n = 5). Finally, 72 patients with both VISTA-CS and VISTA-nonCS were included (Fig. 1).

MRI Acquisition

VW-MRI was performed using a 3T MRI scanner (Ingenia CX, Philips Medical Systems) with a 32-channel sensitivity-encoding (SENSE) head coil. 3D time-of-flight (TOF) MR angiography (MRA) and 3D fat-suppressed VISTA-CS and VISTA-nonCS were performed. 3D T1-weighted VISTA was obtained with improved motion-sensitized driven-equilibrium preparation (velocity encoding = 3 cm/s) for blood suppression. The imaging parameters of 3D TOF MRA were as follows: repetition time/echo time = 19/3.5 ms, flip angle = 18°, matrix = 640 x 330, slice thickness = 1.2 mm (interpolated to 0.6 mm), field of view = 210 mm, 9.6 cm

![](Fig. 1. Flow chart of patient enrollment. VISTA-CS = post-contrast T1-weighted volumetric isotropic turbo spin echo acquisition with compressed sensing, VISTA-nonCS = post-contrast T1-weighted volumetric isotropic turbo spin echo acquisition without compressed sensing, VW = vessel wall)
coverage on axial plane with maximum intensity projection (MIP) reconstruction, and acquisition time = 5 minutes 10 seconds. The imaging parameters of 3D VISTA-nonCS and VISTA-CS are presented in Table 1. Different anatomical coverages of VISTA-nonCS and VISTA-CS on a sagittal MIP image of contrast-enhanced MRA are demonstrated in Figure 2. Due to limited coverage of VISTA-nonCS, it was focused either on the anterior or posterior circulation according to the target vessel. Post-contrast 3D T1WI images were acquired 5 minutes after intravenous injection of a contrast agent. VISTA-CS was performed first and was followed by VISTA-nonCS (Dotarem [gadoterate meglumine]; Guerbet; 0.1 mmol/kg body weight).

**Image Analyses**

For VW-MRI focused on the anterior circulation, the distal internal carotid artery (ICA) from the supraclinoid ICA and middle cerebral artery (MCA) M1 segment was chosen for analysis. For VW-MRI focused on the posterior circulation, the vertebral artery (VA) V4 segment and basilar artery (BA) were chosen for analysis. Distal MCAs or posterior cerebral arteries were not included in the analysis since the diameter of these arteries is frequently small; it is not feasible to draw volume of interests (VOIs) in a considerable portion of patients. Extracranial arteries were also excluded from the analysis because extracranial regions, located off-center, provided relatively low SNR compared to intracranial regions.

**Table 1. Scan Parameters of Intracranial VW-MRI**

|                      | VISTA-NonCS | VISTA-CS               |
|----------------------|-------------|------------------------|
| Repetition time/echo time (ms) | 650/34.8    | 650/34.8               |
| Turbo spin echo factor         | 32          | 32                     |
| Sensitivity-encoding factor | 2           | 2                      |
| Extra reduction factor        | NA          | 1.5 (sampling pattern = hybrid, denoising level 10%) |
| Number of signal averages    | 1           | 1                      |
| Matrix                      | 304 x 304   | 304 x 304              |
| Slice thickness (mm)         | 0.6 (interpolated to 0.3) | 0.6 (interpolated to 0.3) |
| Field of view (mm²)          | 180 x 180   | 180 x 180              |
| Coverage on coronal plane    | 4           | 6                      |
| (antero-posterior direction) (cm) |             |                        |
| Acquisition time            | 8 minutes 21 seconds | 7 minutes 14 seconds |

VISTA-CS = post-contrast T1-weighted volumetric isotropic turbo spin echo acquisition with compressed sensing, VISTA-nonCS = post-contrast T1-weighted volumetric isotropic turbo spin echo acquisition without compressed sensing, VW = vessel wall.

**Fig. 2. Different anatomical coverages in antero-posterior direction of VISTA-CS and VISTA-nonCS.**

VISTA-CS (red box) and VISTA-nonCS (blue box) are demonstrated on sagittal maximum intensity projection image of contrast enhanced MR angiography (A), with coronal imaging slab being rotated slightly to cover both carotid and vertebo-basilar arterial systems. Axial images corresponding to horizontal line (light blue) (A) depict larger coverage in antero-posterior direction in VISTA-CS (B) than that in VISTA-nonCS (C).
Quantitative Analyses

VISTA-CS images were co-registered to VISTA-nonCS images using rigid body transformation with normalized mutual information as a cost function. Then, VOIs were placed by a neuroradiologist with 2 years of experience with VW-MRI using a 3D slicer (3D Slicer v4.10.2) (19). Lumen contours and outer wall boundaries were manually drawn. If the VW-MRI showed enhancing VW lesions, VOIs were drawn on both the normal and lesion sites; otherwise, VOIs were drawn on normal sites only. The lesions were defined as VW thickening with enhancement, first determined by the junior neuroradiologist, then confirmed by a senior neuroradiologist with 9 years of experience. Normal site VOIs were drawn, for VW-MRI focused on the anterior circulation, on two continuous axial slices in a unilateral distal ICA and two continuous sagittal slices in a unilateral MCA M1 segment and, for VW-MRI focused on the posterior circulation, on two continuous axial slices in a unilateral distal VA and two continuous axial slices in BA. Lesion site VOIs were drawn on three continuous slices where wall thickening and enhancement were most prominent. In cases with multiple lesions, the three continuous slices with the most prominent enhancing wall thickening were selected. Normal site VOIs were drawn contralateral to the lesion site or any non-lesioned sites in the case of bilateral lesions. The proximal portion of the V4 segment was excluded for analysis due to normal dural enhancement or vasa vasorum that could interfere with assessment. Lesion sites and types were recorded.

Due to the inhomogeneous noise distribution as a result of parallel imaging, we could not directly measure the noise simply in the air. Instead, we used the standard deviation (SD) of the white matter (WM) to calculate SNR (5, 13, 20). A 10-mm diameter VOI was placed in the normal WM at the anterior temporal lobe for VW-MRI focused on the anterior circulation and the middle cerebellar peduncle for VW-MRI focused on the posterior circulation. The SNR was calculated as follows: $\text{SNR} = 0.695 \times \frac{\text{signal intensity of VW or lumen}}{\text{noise}}$. The CNR was calculated: $\text{CNR} = \text{SNR}_{\text{wall}} - \text{SNR}_{\text{lumen}}$. Wall and lumen volume, SNR, and CNR were calculated for normal and lesion sites.

Qualitative Analyses

Two neuroradiologists independently evaluated VISTA-nonCS and VISTA-CS. Overall image quality and normal wall and lesion wall delineation were evaluated with a four-point visual scale (13, 21). Overall image quality was graded as follows: 1) artifacts significantly interrupt VW assessment, 2) artifacts cause some difficulty in VW assessment, 3) artifacts exist without affecting VW assessment, and 4) negligible artifacts and excellent image quality (Fig. 3). In VISTA-nonCS images, ripple-like artifacts were frequently noted that might

![Fig. 3. Representative figures demonstrating different grades of overall image quality.](image-url)

Grade 1: artifacts in posterior fossa significantly interrupt assessment of VW lesions in left distal VA (VISTA-nonCS) (A). Grade 2: blurring of vessel contour with SNR decrease causes some difficulty in assessment of VW lesions in right distal VA (VISTA-CS) (B). Grade 3: ripple like artifacts exist without affecting assessment of VW lesions in left MCA (VISTA-nonCS) (C). Grade 4: negligible artifacts and excellent image quality (VISTA-CS) (D). MCA = middle cerebral artery, SNR = signal-to-noise ratio, VA = vertebral artery
be associated with SENSE (Fig. 3C). Instead of ripple-like artifacts, the blurring of vessel contours and decreased SNR predominantly in the center of the images were frequently noted in the VISTA-CS (Fig. 3B). Normal VW delineation was graded as follows: 1) VW discernible in < 1/2 lengths of the analyzed areas, 2) VW discernible in ≥ 1/2 and < 3/4 lengths of the analyzed areas, 3) VW discernible in ≥ 3/4 lengths of the analyzed areas with a few indiscernible sites, and 4) VW discernible along the entire length. Lesion VW delineation was graded as follows: 1) < 180 degrees of the VW is clearly visible, 2) ≥ 180 and < 270 degrees of the VW is clearly visible, 3) ≥ 270 and < 360 degrees of the VW is clearly visible, and 4) ≥ 360 degrees of the VW is clearly visible. No patients received a grade of 1 in the lesion VW delineation. Representative figures of grades 2–4 are demonstrated in Figure 4. Significant blurring of the lesion site led to a one-score reduction. After calculating the average scores from reader 1 and reader 2, acceptable images were defined as those with scores ≥ 3 for overall image quality and normal and lesion wall delineation.

Statistical Analyses

Statistical analyses were performed using R statistical software version 3.4.2 (R foundation for Statistical Computing). Paired t test and McNemar’s test were used to compare the two groups. Weighted kappa was calculated for interobserver agreement. An interclass correlation coefficient (ICC) was calculated for VW and lumen volumes. Bland-Altman plots were obtained to delineate the reproducibility of wall and lumen volumes from VISTA-nonCS and VISTA-CS. Violin plots were used to visualize the distribution of SNR and CNR from VISTA-nonCS and VISTA-CS. P values < 0.05 indicated statistical significance.

RESULTS

Study Cohort

Baseline patient characteristics and VW-MRI findings are shown in Table 2. A total of 72 patients (39 men [mean age

| Characteristics                  | Values |
|----------------------------------|--------|
| Number of patients               | 72     |
| Sex (male:female)                | 39:33  |
| Age (years), mean ± SD           | 50.9 ± 14.9 |
| Anterior:posterior circulation   | 46:26  |
| Cases with lesions               | 45     |
| Lesion location, n (%)           |        |
| Right ICA                        | 6 (13.3) |
| Left ICA                         | 2 (4.4)  |
| Right MCA                        | 10 (22.2) |
| Left MCA                         | 9 (20.0)  |
| Right VA                         | 10 (22.2) |
| Left VA                          | 6 (13.3)  |
| Basilar artery                   | 2 (4.4)  |
| Lesion type, n (%)               |        |
| Atherosclerosis                  | 25 (55.6) |
| Dissection                       | 17 (37.8) |
| Moyamoya disease                 | 1 (2.2)  |
| Aneurysm                         | 1 (2.2)  |
| Dysplasia                        | 1 (2.2)  |

ICA = internal carotid artery, MCA = middle cerebral artery, SD = standard deviation, VA = vertebral artery

Fig. 4. Representative figures demonstrating different grades of VW lesion delineation.

Grade 1 is not shown as no patients received grade 1 in lesion wall delineation. Grade 2: ≥ 180 and < 270 degrees of lesion in right MCA is clearly visible (black arrowheads) (A). Grade 3: ≥ 270 and < 360 degrees of lesion in right distal VA is clearly visible (black arrowheads) (B). Grade 4: ≥ 360 degrees of lesion in left MCA is clearly visible (white arrowheads) (C).
± SD, 48 years ± 13) and 33 women (52 years ± 17) were enrolled. Forty-six patients underwent VW-MRI focused on the anterior circulation, while 26 patients underwent VW-MRI focused on the posterior circulation. Forty-five patients had identifiable enhancing VW lesions. The most common type of lesion was atherosclerosis (55.6%), followed by dissection (37.8%). Representative cases are shown in Figure 5.

Quantitative Analyses
The results of the quantitative analyses are summarized in Table 3. In both normal and lesion sites, wall and lumen volumes did not differ significantly between VISTA-nonCS and VISTA-CS. In normal sites, ICCs for wall and lumen volume were 0.968 and 0.964, respectively. In lesion sites, ICCs for wall and lumen volume were 0.997 and 0.996, respectively, suggesting excellent reproducibility between the two sequences. Bland-Altman plots of wall and lumen volumes from both normal and lesion sites are shown in Figure 6. In normal sites, VISTA-CS presented significantly higher wall SNR (4.69 ± 1.37 vs. 4.47 ± 1.22, p = 0.013) and lumen SNR (1.94 ± 0.44 vs. 1.71 ± 0.38, p < 0.001) compared to VISTA-nonCS. CNR was not significantly different between VISTA-nonCS and VISTA-CS (2.76 ± 0.89 vs. 2.75 ± 0.97, p = 0.907). In lesion sites, wall SNR did not differ significantly between VISTA-nonCS and VISTA-CS (8.24 ± 3.28 vs. 8.27 ± 3.09, p = 0.878), while lumen SNR was significantly lower in VISTA-nonCS than in VISTA-CS (2.85 ± 1.52 vs. 3.50 ± 1.83, p < 0.001). The CNR was significantly higher in VISTA-nonCS than in VISTA-CS (5.39 ± 2.21 vs. 4.77 ± 2.00, p = 0.003).

Qualitative Analyses
The visual scores of VISTA-nonCS and VISTA-CS from the two readers are presented in Table 4. VISTA-CS and VISTA-nonCS had comparable overall image quality without any significant difference (averaged scores from the readers, 3.12 ± 0.60 vs. 3.06 ± 0.63, p = 0.297). In VISTA-nonCS, 13 patients (18.1%) received grades of 1 and 2 in overall image quality due to MRI artifacts interrupting the VW assessment. The remaining 59 patients (81.9%) had negligible artifacts or some artifacts causing no difficulty in the VW assessment. In VISTA-CS, 7 (9.7%) and 10 (13.9%) patients received grades of 1 and 2 in overall image quality, respectively. Ripple-like artifacts were common with VISTA-nonCS, likely related to SENSE (Fig. 5C). On VISTA-CS, blurred and coarse vessel boundaries and decreased signal in the center of images were common (Fig. 5B). Normal walls were better delineated with VISTA-nonCS than with VISTA-CS (3.51 ± 0.63 vs. 3.38 ± 0.67, p = 0.019). Except for one
Table 3. Comparisons of Volume, SNR and CNR between VISTA-NonCS and VISTA-CS for Normal and Lesion Sites

|              | VISTA-NonCS | VISTA-CS | P   | ICC |
|--------------|-------------|----------|-----|-----|
| Normal site  |             |          |     |     |
| Wall volume  | 9.90 ± 3.04 | 9.79 ± 2.89 | 0.414 | 0.968 |
| Lumen volume | 13.30 ± 7.34 | 13.14 ± 7.45 | 0.227 | 0.964 |
| SNRwall      | 4.47 ± 1.22 | 4.69 ± 1.37 | 0.013 |     |
| SNRlumen     | 1.71 ± 0.38 | 1.94 ± 0.44 | < 0.001 |     |
| CNRwall-lumen| 2.76 ± 0.89 | 2.79 ± 0.97 | 0.907 |     |
| Lesion site  |             |          |     |     |
| Wall volume  | 10.87 ± 8.55 | 10.94 ± 8.66 | 0.613 | 0.997 |
| Lumen volume | 6.54 ± 5.86  | 6.45 ± 5.73  | 0.422 | 0.996 |
| SNRwall      | 8.24 ± 3.28  | 8.27 ± 3.09  | 0.878 |     |
| SNRlumen     | 2.85 ± 1.52  | 3.50 ± 1.83  | < 0.001 |     |
| CNRwall-lumen| 5.39 ± 2.21  | 4.77 ± 2.00  | 0.003 |     |

Data is presented with mean ± SD. CNR = contrast-to-noise ratio, ICC = interclass correlation coefficient, SNR = signal-to-noise ratio.

Fig. 6. Bland-Altman plots for volumes of wall and lumen from VISTA-nonCS and VISTA-CS in normal sites (A) and lesion sites (B) suggest excellent reproducibility between two sequences. SD = standard deviation.
In the current study, we compared VISTA-CS and VISTA-nonCS, hoping to increase scan coverage to cover both anterior and posterior circulations with the use of CS. While increasing scan coverage, CS showed or trended towards significantly higher SNRs when applied to VISTA in the VW-MRI. CNR was significantly lower with CS in the lesion sites, while there were no significant differences in the normal sites. VISTA-CS provided comparable overall image quality and normal wall delineation to VISTA-nonCS, but lesion walls were better delineated with VISTA-nonCS. We also observed that similar proportions of VISTA-CS and VISTA-nonCS provided acceptable quality images. Thus, we concluded that CS might be useful when applied to the post-contrast images in intracranial VW-MRIs since it facilitates more scan coverage with slightly shorter scan times without significantly sacrificing image quality.

Several prior studies reported that CS achieved substantial scan time reduction while providing similar or superior SNR, CNR, VW delineation, and image quality in intracranial VW-MRIs (13, 14). Those studies enrolled mostly healthy volunteers; therefore, validation in large patient cohorts was necessary to use CS in clinical practice. Another recent study investigated CS in joint intracranial and carotid VW-
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MRIs in 28 subjects, including 20 stroke patients, and concluded that CS reduced scan time by almost half without compromising wall thickness measurements or diagnostic wall visualizations (22). This study, however, used CS only in non-contrast images and the sample size was also small. Here, we applied CS to post-contrast images in a relatively large number of patients who underwent intracranial VW-MRIs for specific reasons and performed both VISTA-CS and VISTA-nonCS. Post-contrast images, which are an essential part of VW-MRIs since they help detect lesions and define the cause of luminal narrowing (2, 16), were chosen for analysis in our study. Instead of reducing scan acquisition time with CS, as seen in previous studies, we increased scan coverage while maintaining a similar acquisition time since larger scan coverage has strengths in clinical practice. Before CS was applied, images were usually limited to a target region covering either anterior or posterior circulation or known stenotic lesions to achieve high resolution in a reasonable scan time. Since scan coverage was increased to cover anterior and posterior circulations by applying CS, we were able to find incidental lesions located in the non-target vessels that might have been missed in VW-MRIs without CS. Especially in patients with intracranial atherosclerosis, atherosclerotic plaques are frequently found in multiple arterial segments (23, 24), necessitating broad imaging coverage of the brain. In addition, radiologists should confirm the scan region in each VW-MRI when CS is not applied (3). This is unnecessary when scan coverage is increased with CS and VW-MRI can be performed without asking radiologists to determine the scan site and without imaging delay. In addition to the benefits from more scan coverage, we found that CS still achieved acceptable image quality that did not significantly affect radiological interpretations; therefore, CS could be a useful method for intracranial VW-MRIs.

A greater acceleration factor introduces more artifacts, blurring, and image degradation (7, 25, 26). Prior investigations of CS in VW-MRI have used acceleration factors ranging from 1.5 to 10 (12-14, 26, 27). For carotid wall evaluation, images obtained with two-fold to five-fold acceleration achieved acceptable image quality that was not significantly different from full acquisition (12, 27). CS also provided robust and repeatable results with an acceleration factor of 1.5 (15). Further, CS with an acceleration factor of 3–5 applied to intracranial VW-MRIs yielded higher or comparable image quality to marked time reductions (13, 14). Here, we used an extra reduction factor of 1.5 in VISTA-CS, which did not significantly affect image quality.

Several CS-related artifacts were reported in previous studies. For instance, Sharma et al. (25) described a global ringing artifact and fine detail blurring in the brain. Worters et al. (28) reported that CS images looked slightly blurrier compared to the original images, possibly due to a denoising effect in the CS reconstruction. In our study, we noted blurring in VISTA-CS as seen previously. This may have affected normal and lesion wall delineation and, thus, led to score reductions in the visual assessment; however, most VISTA-CS images allowed for VW assessments similar to VISTA-nonCS images in the present study. Furthermore, coarse, ripple-like artifacts potentially related to SENSE with limited coverage were more prominent in VISTA-nonCS than in VISTA-CS. Thus, we concluded that artifacts of different characteristics exist in VISTA-CS images without significant impact on intracranial VW-MRI assessments.

VISTA-CS elicited superior SNR from the lumen of both normal and lesion sites and the wall of normal sites in the present study, which was similar to previous studies’ results (13). The SNR increases observed in CS may result from the denoising process in CS reconstruction (8). Furthermore, increased scan coverage in VISTA-CS may increase SNR while reducing aliasing and SENSE-related artifacts. A SNR increase in the lumen, however, might not be desirable since it has a negative effect on the contrast between the lumen and VW wall, affecting the radiologists’ VW assessment. Despite a higher SNR in VISTA-CS, the qualitative analysis by two neuroradiologists revealed that normal and lesion wall delineations were superior in VISTA-nonCS. Meanwhile, wall SNR of lesion sites did not differ significantly between the two sequences. This might be associated with differing time intervals between contrast injection and scan acquisition. Since VISTA-nonCS was obtained before VISTA-CS, delayed enhancing lesions may be more enhanced with VISTA-nonCS than VISTA-CS, increasing the SNR in VISTA-nonCS. This phenomenon might cancel out the SNR increase in the VISTA-CS. Since VISTA-CS revealed comparable wall SNR and higher lumen SNR than VISTA-nonCS, CNR was subsequently lower than with VISTA-nonCS at lesion sites; however, the CNR of the lesion site with VISTA-CS was high (more than 4) and most VISTA-CS images provided acceptable quality images for lesion wall delineation. Moreover, the volumes of enhancing lesions did not differ significantly between VISTA-nonCS and VISTA-CS. Thus, VW assessments may not be affected with CS in intracranial VW-MRIs.

There are several study limitations to be addressed.
First, this is a retrospective study, which might introduce selection bias to our results. To mitigate selection bias, we searched all VW-MRIs performed in a certain time period, but future studies with prospective design are needed to validate our study results. Second, VISTA-nonCS was obtained after VISTA-CS in all patients. This potentially introduced differences in the degree of enhancement between the two scans. The volumes of lesion walls, which were calculated based on the VOIs drawn on the enhancing portions, did not differ significantly between VISTA-nonCS and VISTA-CS; therefore, the differences in the degree of enhancement between the two sequences might not significantly affect the image interpretation. Still, further studies performing contrast-enhanced images with and without CS in a randomized order are required to validate our study results. Third, scan coverage was increased when CS was applied, altering imaging protocols between VISTA-nonCS and VISTA-CS; however, all other scan parameters were the same between VISTA-nonCS and VISTA-CS, so we consider the two protocols to be comparable. Fourth, completely blinded interpretations of VISTA-nonCS and VISTA-CS were not possible since the two sequences provided different coverages and specific patterns of artifacts, which were easily distinguishable by the reviewers.

In conclusion, CS achieved comparable SNR, CNR, image quality, and wall delineation when applied to post-contrast images of intracranial VW-MRIs, while simultaneously increasing scan coverage and maintaining a similar scan acquisition time. Thus, CS may be useful when applied to intracranial VW-MRIs.

Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

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