





Citation: Yamamoto T, Okada T, Fushimi Y, Yamamoto A, Fujimoto K, Okuchi S, et al. (2018) Magnetic resonance angiography with compressed sensing: An evaluation of moyamoya disease. PLoS ONE 13(1): e0189493. https://doi.org/ 10.1371/journal.pone.0189493

Editor: Stephan Meckel, Universitatsklinikum Freiburg, GERMANY

Received: September 6, 2017

Accepted: November 27, 2017

Published: January 19, 2018

Copyright: © 2018 Yamamoto et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Data Availability Statement: Free access to medical image data is ethically restricted by the Japanese governmental organization "Personal Infromation Protection Comission JAPAN" (https://www.ppc.go.jp/en/) per restrictions on Specific Personal Information (https://www.ppc.go.jp/en/legal/). Interested researchers may send data accession requests to the Kyoto University Graduate School and Faculty of Medicine, Kyoto University Hospital Ethics Committee to the following email address: ethcom@kuhp.kyoto-u.ac.

RESEARCH ARTICLE

Magnetic resonance angiography with compressed sensing: An evaluation of moyamoya disease

Takayuki Yamamoto¹, Tomohisa Okada¹*, Yasutaka Fushimi¹, Akira Yamamoto¹, Koji Fujimoto¹, Sachi Okuchi¹, Hikaru Fukutomi¹, Jun C. Takahashi^{1,2}, Takeshi Funaki³, Susumu Miyamoto³, Aurélien F. Stalder⁴, Yutaka Natsuaki⁵, Peter Speier⁶, Kaori Togashi¹

- Department of Diagnostic Imaging and Nuclear Medicine, Graduate School of Medicine, Kyoto University, Kyoto, Japan,
 Department of Neurosurgery, National Cerebral and Cardiovascular Center, Suita, Japan,
 Department of Neurosurgery, Graduate School of Medicine, Kyoto University, Kyoto, Japan,
 Siemens Itd. China, Shanghai, China,
 Siemens Medical Solutions USA, Inc., Huntington Beach, California, United States of America,
 Siemens Healthcare, Erlangen, Germany
- * tomokada@kuhp.kyoto-u.ac.jp

Abstract

Compressed sensing (CS) reconstructions of under-sampled measurements generate missing data based on assumptions of image sparsity. Non-contrast time-of-flight MR angiography (TOF-MRA) is a good candidate for CS based acceleration, as MRA images feature bright trees of sparse vessels over a well-suppressed anatomical background signal. A short scan time derived from CS is beneficial for patients of moyamoya disease (MMD) because of the frequency of MR scans. The purpose of this study was to investigate the reliability of TOF-MRA with CS in the evaluation of MMD. Twenty-two patients were examined using TOF-MRA with CS (CS-TOF) and parallel imaging (PI-TOF). The acceleration factors were 3 (CS3) and 5 (CS5) for CS-TOF, and 3 (PI3) for PI-TOF. Two neuroradiologists evaluated the MMD grading according to stenosis/occlusion scores using the modified Houkin's system, and the visibility of moyamoya vessels (MMVs) using a 3-point scale. Concordance was calculated with Cohen's κ . The numbers of MMVs in the basal ganglia were compared using Bland-Altman analysis and Wilcoxon's signed-rank tests. MRA scan times were 4:07, 3:53, and 2:42 for PI3, CS3, and CS5, respectively. CS-reconstruction completed within 10 minutes. MMD grading and MMV visibility scales showed excellent correlation ($\kappa > .966$). Although the number of MMVs was significantly higher in CS3 than in PI3 (p < .0001) and CS5 (p < .0001), Bland-Altman analysis showed a good agreement between PI3, CS3, and CS5. Compressed sensing can accelerate TOF-MRA with improved visualization of small collaterals in equivalent time (CS3) or equivalent results in a shorter scan time (CS5).

Introduction

Compressed sensing (CS) has been vigorously studied in recent years because of its potential to reduce scan time[1] or to improve spatio-temporal resolution while maintaining the same



Funding: This work was supported by a Grant-in-Aid for Scientific Research on Innovative Areas "Initiative for High-Dimensional Data-Driven Science through Deepening of Sparse Modeling" of The Ministry of Education, Culture, Sports, Science and Technology, Japan was provided to K.T. (MEXT grant number 25120002) (http://sparsemodeling.jp/). The funder had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: Three of the authors (A.S., Y. N., P.S.) are employees of Siemens Healthcare. These authors developed and provided the prototype sequence and reconstruction method, but were not involved in the data acquisition and evaluation, and did not have direct control of the data. This does not alter our adherence to PLOS ONE policies on sharing data and materials.

scan time[2]. The goal of CS is to approximate the image quality of a fully sampled measurement from under-sampled *k*-space data, by exploiting the intrinsic sparsity in the imaged structures in a non-linear iterative regularized reconstruction. Several studies[3,4] on contrast enhanced MRA with CS have reported on its excellent speed (e.g. only 10 seconds were needed to cover the whole brain[5]). Non-contrast time-of-flight (TOF) MRA is also a good candidate for CS based acceleration, as images feature a bright trees of sparse vessels over a well suppressed anatomical background signal[6–10]. In a preceding study on normal volunteers[11], the scan time of cerebrovascular non-contrast TOF-MRA with CS (CS-TOF) was reduced to approximately half of that of TOF-MRA with parallel imaging (PI-TOF), the current clinical standard for accelerating acquisition[12–14], while maintaining the image quality.

CS reconstructions of accelerated (i.e., under-sampled) measurements generate the missing data by making assumptions about the image sparsity after a certain transformation; in our case, magnitude images were sparsified in the wavelet domain of the magnitude image. Therefore, the diagnostic capability of CS-TOF requires validation, especially in regard to whether or not it introduces an over- or under-estimation of vascular pathological changes in comparison with clinically established methods. For cerebrovascular aneurysms, images acquired with CS-TOF have been successfully evaluated [15,16]. Our aim here is to extend this validation to vascular stenosis and small vessels.

Moyamoya disease (MMD) is an uncommon cerebrovascular disorder characterized by progressive stenosis and occlusion of the supra-clinoid internal carotid artery and its main branches within the circle of Willis. These changes result in the formation of a fine vascular network, known as the moyamoya vessels (MMVs), and other collateral circulations such as superficial temporal artery. Visualization of these is directly linked to the surgical treatments [17]. Although digital subtraction angiography (DSA) still remains the gold standard for the diagnosis and staging of MMD, TOF-MRA is now considered to be a reliable alternative with high sensitivity and specificity[18–20]. TOF-MRA is a non-invasive modality that can be readily applied to asymptomatic MMD patients and repeatedly used for follow-up examinations, even in children.

Therefore, we hypothesized that CS-TOF can be used to assess MMD, and examined the grading and visualization of MMVs in a prospective MMD cohort, comparing the results from CS-TOF with those from conventional PI-TOF, with the CS-TOF using the same or a higher acquisition acceleration factor than the PI-TOF.

Materials and methods

Study population

This study was approved by the local institutional review board (Kyoto University Graduate School Ethics committee), and written informed consent was obtained from all subjects. From December 2014 through to October 2015, 22 patients with MMD were enrolled (mean age, 43.1 years; age range, 29–65 years; 14 women and 8 men). All subjects were scanned using both CS-TOF and PI-TOF in the same session. Diagnosis of MMD was confirmed by DSA or conventional MRA (PI-TOF). Ten of the patients had undergone a previous operation for direct bypass from the superficial temporal artery to the middle cerebral artery. Exclusion criteria were failure to acquire the whole imaging sets, as well as large motion and susceptibility artifacts.

Acquisition of MR angiography

CS-TOF and PI-TOF were acquired using a 3T MR scanner (MAGNETOM Skyra, Siemens Healthcare, Erlangen, Germany) with a 32-channel head coil (Fig 1). A research sequence and

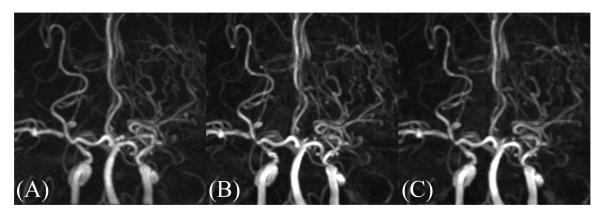


Fig 1. Coronal views of MIP images. (A) PI3, (B) CS3, (C) CS5. Occlusion of the left MCA and the development of MMVs are equivalently visualized in these images. PI3, conventional TOF MRA with an acceleration factor of 3; CS3, compressed sensing TOF MRA with an acceleration factor of 5; MCA, middle cerebral artery.

https://doi.org/10.1371/journal.pone.0189493.g001

reconstruction prototype provided by Siemens Healthcare GmbH (Erlangen, Germany) was used for CS-TOF. Common acquisition parameters were as follows: TR/TE = 20/3.69 ms, FA = 18° , 4 slabs, TONE ramp, 70%; and a reconstructed voxel size of $0.38 \times 0.38 \times 0.38 \times 0.38$ mm. For CS-TOF, the FOV was $220 \times 200 \times 63$ mm and the acquired matrix was $288 \times 264 \times 82$ (interpolated to $576 \times 528 \times 164$), whereas for PI-TOF, the FOV was $240 \times 200 \times 62$ mm, and the matrix was $320 \times 270 \times 81$ (interpolated to $640 \times 540 \times 162$).

For CS-TOF data acquisition, a conventional 3D TOF gradient-echo sequence was extended to facilitate incoherent sampling of k-space using a variable-density Poisson disk sampling pattern in the ky-kz encoding plane on a Cartesian grid[21] (Fig 2). After data acquisition, the image was reconstructed from the under-sampled data by nonlinear iterative SENSE-type reconstruction[14]. The reconstruction was implemented inline on the scanner. The following minimization problem is solved using a modified Fast Iterative Shrinkage-Thresholding Algorithm (mFISTA) [22]:

$$\min_{x} \left| \sum_{j}^{N} y_{j} - F_{u} S_{j} x \right|_{2}^{2} + \lambda |Wx|_{1}$$

where x is the image to reconstruct, y_j and S_j are the k-space data and coil sensitivity for j-th coil element, Fu is the Fourier under-sampling operator, W is the redundant Haar wavelet transform, and λ is the normalized regularization parameter (set to 0.008). This equation and

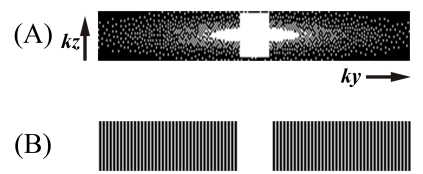


Fig 2. Sampling patterns of the slab. Ky-Kz plane of k-space. (A) Compressed Sensing, (B) Parallel Imaging.

https://doi.org/10.1371/journal.pone.0189493.g002



the parameter was the same and optimized in our previous study[11]. The image reconstruction was fully integrated with the scanner software and was performed inline.

The acceleration rates for CS-TOF were 3 (CS3) and 5 (CS5), while that of PI-TOF was 3 (PI3). PI3 (acceleration in phase encoding direction only) was selected as the reference standard, as it is the acceleration factor in general use in our institution. PI-MRA was reconstructed with the scanner's product implementation of GRAPPA (Generalized Auto-Calibrating Partially Parallel Acquisition) [13,23].

Image analysis

Firstly, two neuroradiologists (T.O. and Y.F., with 26 and 19 years of experience, respectively) graded the stenotic changes at Circle of Willis on both CS-TOF and PI-TOF. MMD grading was assessed according to the modified Houkin's grading system[24]. This classifies the progress of the disease into 4 grades based on the total stenosis/occlusion scores of internal carotid artery, middle cerebral artery, anterior cerebral artery, and posterior cerebral artery.

Secondly, MMV visibility was scored using a 3-point scale: 1, few or no MMVs; 2, a moderate amount of MMVs (weak visualization of vessels, but sufficient for diagnosis); 3, a large amount of MMVs (vessels clearly visualized). Examples are presented in Fig 3.

Thirdly, high-intensity spots recognized as vessels (perforating arteries or MMVs in the basal ganglia) were manually counted on an original 2D slice at the level of the basal ganglia by two neuroradiologists (T.Y. and S.O., with 11 and 11 years' experience, respectively). For each patient, this was performed separately for the left and right side, with the vessels being tracked through upper and lower slices to differentiate vessels from noises. Definite vessels were counted as 1, and probable vessels as 0.5.

Finally, the visibility of bypass vessels was also subjectively evaluated in the 10 patients who had undergone previous bypass surgery.

All of the analysis above was performed independently, and the sequence was blinded for the raters.

Statistical analysis

The Cohen's κ statistic (quadratic weighted) was used to analyzed for the MMD grading and for the MMV visibility on maximum intensity projection (MIP) images, facilitating comparisons

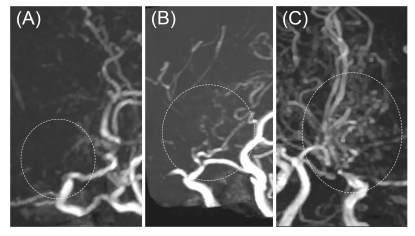


Fig 3. Moyamoya vessels (MMVs) on MIP images. (A) Few or no MMVs, (B) a moderate amount of MMVs (weak vessels visibility, but sufficient for diagnosis), (C) a large amount of MMVs (vessels are clearly visualized). All images are compressed sensing TOF-MRA with an acceleration factor of 3.

https://doi.org/10.1371/journal.pone.0189493.g003



between CS3, CS5, and PI3 (slight agreement, $\kappa < 0.2$; fair agreement, 0.2–0.4; moderate agreement, 0.4–0.6; substantial agreement, 0.6–0.8; almost-perfect agreement, \geq 0.8). The MMV numbers of CS3 and CS5 were compared with PI3 using the Bland-Altman analysis and the Wilcoxon signed rank test. Inter-rater agreement in the MMV numbers was also evaluated by intraclass correlation coefficient (ICC). Statistical analyses were performed using R (version 3.3.1, https://www.r-project.org/), with a p value < .05 being considered as statistically significant.

Results

MRA scan times were 4:07, 3:53 and 2:42 for the PI3, CS3 and CS5 acquisitions, respectively (Fig 2). The CS-TOF reconstruction was started at the end of the acquisition and finished within 10 minutes. No patients were excluded from the analysis.

The modified Houkin's MMD grading of CS3 and CS5 showed a nearly complete agreement with those of PI3 (κ = .993 and .996, respectively; see <u>Table 1</u>), thereby demonstrating equivalent grading capabilities. Inter-rater agreements were substantial (<u>Table 2</u>).

MMV visibility on CS-TOF MIP images was also equivalent to that on PI-TOF images (Table 3). Excellent agreements were obtained between PI3 and CS3 (κ = .992), and between PI3 and CS5 (κ = .967). Inter-rater agreements were substantial (Table 4).

The median number of MMVs at the basal ganglia level was 3 for PI3 (1st–3rd quartile: 2.0–5.3), 4 for CS3 (3.0–6.6), and 3.5 for CS5 (2.0–5.0). The Bland-Altman analysis for the number of MMVs also showed a good agreement. The number of MMVs on CS3 was significantly higher than that on PI3 (p < .0001) or that for CS5 (p < .0001) (Fig 4). No significant difference was observed between CS5 and PI3 for the number of MMVs. The inter-rater ICC for counting of the number of MMVs showed an excellent correlation (.894).

Table 1. The results of the MRA grading of moyamoya disease with PI3, CS3, and CS5.

CS3 (CS5)	PI3			
	Stage 1	Stage 2	Stage 3	Stage 4
Stage 1	8 (8)			
Stage 2		26 (24)		
Stage 3		0 (2)	32 (31)	
Stage 4			1 (2)	21 (21)

Two observers independently assessed 44 sides of 22 patients, thus a total of 88 results are illustrated. The κ value was 0.993 between PI3 and CS3, and 0.966 between PI3 and CS5. PI3, conventional TOF MRA with an acceleration factor of 3; CS5, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.t001

Table 2. Inter-rater agreement in the grading score of moyamoya disease observed as Cohen's κ statistic.

		Rater 2		
		PI3	CS3	CS5
Rater 1	PI3	.755	.751	.765
	CS3	.748	.748	.758
	CS5	.745	.741	.757

Substantial agreement was shown for each combination of PI3, CS3, and CS5.

PI3, conventional TOF MRA with an acceleration factor of 3; CS3, compressed sensing TOF MRA with an acceleration factor of 3; CS5, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.t002



Table 3. The visibility of moyamoya vessels on MIP images from PI3, CS3, and CS5.

	PI3		
CS3 (CS5)	Grade 1	Grade 2	Grade 3
Grade 1	33 (33)	1 (1)	
Grade 2	0 (2)	27 (25)	0(1)
Grade 3			27 (26)

The κ value was 0.992 between PI3 and CS3, and 0.967 between PI3 and CS5.

PI3, conventional TOF MRA with an acceleration factor of 3; CS3, compressed sensing TOF MRA with an acceleration factor of 3; CS5, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.t003

Table 4. Inter-rater agreement on the visibility of moyamoya vessels observed as Cohen's κ statistic.

		Rater 2		
		PI3	CS3	CS5
Rater 1	PI3	.608	.608	.608
	CS3	.631	.631	.631
	CS5	.608	.608	.608

Substantial agreement was shown for each combination of PI3, CS3, and CS5.

PI3, conventional TOF MRA with acceleration factor of 3; CS3, compressed sensing TOF MRA with acceleration factor of 3; CS5, compressed sensing TOF MRA with acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.t004

All of the bypass vessels of the patients who had undergone operations were well recognized in all three acquisitions. The superficial temporal arteries were also satisfyingly depicted in the other patients.

Discussion

We investigated the diagnostic quality of CS-TOF by evaluating MMD grading (which include stenosis/occlusion scores) and MMV visibility on both MIP images and 2D slices at the level of the basal ganglia. CS-TOF better visualized MMVs that work as small collaterals in an equivalent scan time, and attained equivalent visualization in reduced scan time of less than 3 minutes.

The choice of an acceleration rate of 3 or 5 for CS-TOF had little effect on the visual aspect and the grading of MMD, suggesting the absence of either over- or under-estimation due to irregular undersampling and nonlinear iterative reconstruction. In 4 cases (9.1%), MMD grading based on CS5 resulted in an over-diagnosis (2 cases, stage 2 as 3; 2 cases, stage 3 as 4; see Table 1), but we did not observe any cases of under-diagnosis. Thus, if CS-TOF shows a severe stenosis, a confirmation of the finding using contrast-enhanced angiography (MR, CT, or DSA) would be worth considering for the initial evaluation, but is not required for follow-up observations focusing on relative changes. The inter-rater correlation for the stenosis/occlusion scores was substantial; however, the intra-rater agreement was almost perfect. Therefore, the difference in scoring is considered to reflect variations in evaluation due to the individual interpreting the images, and this may have contributed to the over diagnoses in the CS5 cases.

The presence and quantity of MMVs is also important for the evaluation of MMD[25–27], although it is not easy to evaluate it using TOF-MRA[28]. The visibility of MMVs on MIP images was almost the same on PI-TOF and CS-TOF, whereas the number of MMVs counted on the CS3 images was significantly higher than that on the PI3 images. This evaluation had

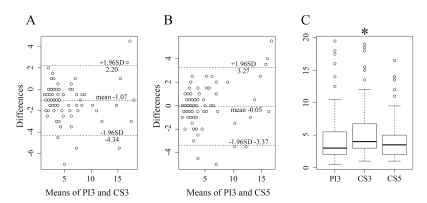


Fig 4. Bland-Altman plots and Box-and-whisker plot of the numbers of high-intensities, which are considered as vessels, observed in the basal ganglia. (A, B) Bland-Altman plots. The Y-axis unit (differences) is numerical. (A) No significant bias was found comparing PI3 with CS3. (B) Hardly any differences were detected when comparing PI3 with CS5. (C) Box-and-whisker plot. The number for CS3 was significantly higher than that of PI3 (p < .0001) and CS5 (p < .0001). PI3, conventional TOF MRA with an acceleration factor of 3; CS3, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.g004

high intra-observer agreement, with inter-observer reproducibility showing substantial agreement, probably because evaluation on the MIP images is tended to be influenced by observer subjectivity, as in the evaluation of stenosis/occlusion. The evaluation of the number of MMVs in basal ganglia helps to understand the difference in characteristics between CS-TOF and PI-TOF. CS3 showed more MMVs than the others; this is likely to be due to the inherent denoising effect of CS[1]. CS reconstruction enhances the contrast of vessels by counteracting background noise, while this denoising effect may also simultaneously diminish signal from vessels. The previous study reported that the apparent SNR of CS-MRA is higher than that of PI-MRA with more than 3-fold acceleration[11]. The high apparent SNR is considered to result in more moyamoya vessels being visualized in CS. In CS3, the positive effect was considered to exceed the negative effect, while even in CS5, the numbers of MMVs were comparable with those on PI3. A previous report supports this argument, as it showed that attenuation of small vessel signals is strong on CS-TOF in comparison with PI-TOF[11].

Fig 5A shows representative images for all three acquisitions. The example PI3 image in Fig 4A shows prominent speckled noise, especially in the center[14,29]; this is due to the g-factor increase in the center. The example CS images of Fig 5B and 5C show highly reduced noises throughout the image; however, curved stripe patterns consistent with ghosts originating from the skull boundaries are apparent, with these becoming most prominent at the center. The intensity of this artifact varied from slice to slice, but the direction of the stripe patterns always aligned with the phase encoding direction. We thus speculate that these artifacts are residual aliasing artifacts originating from the sparse undersampling, possibly derived from the outer edge of the skull. We evaluated the incoherence by the point spread function of sampling patterns of CS (Fig 6), and the ratio of the main peak of point spread function to the standard deviation of the pseudo-noise (incoherent artifacts) is 74.2 for CS3, and 35.1 for CS5. The limited slab size (28 slices, 16.7% oversampling) might constraint the incoherence. Slight patient movements are also a conceivable reason. Increasing the regularization parameters used in the reconstruction can mitigate this artifact, but a strong increase may obscure small vessels. However, this artifact is hardly noticeable on the MIP images (Fig 1), and did not affect the detection of the vessels in the basal ganglia in this study.

The total reconstruction time for CS was approximately 10 minutes, which is too long for clinical use. CS-TOF is scanned in 4 slabs in this study, and slab-wise reconstruction after the

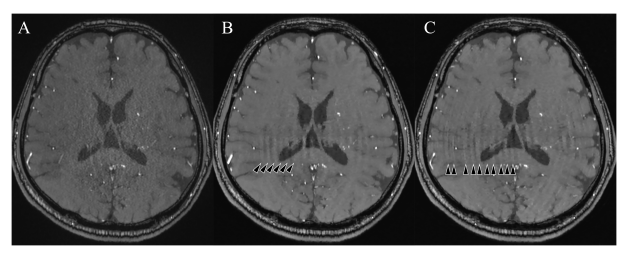


Fig 5. The 2D slice images at the level of the basal ganglia for (A) PI3, (B) CS3, and (C) CS5. On CS images, some artifacts such as curved stripe patterns (arrowheads) were seen. These artifacts were not observable on the MIP images. PI3, conventional TOF MRA with an acceleration factor of 3; CS3, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.g005

acquisition of each slab cloud reduce the actual waiting time. Moreover, the reconstruction time could be drastically shortened to clinically acceptable times by using a graphics processing unit implementation, which was not available in this study.

There are several limitations to this study. Firstly, CS and PI were separately evaluated for comparison in this study, however, CS and PI are not essentially exclusive technique, but they can be used synergistically for further acceleration[30]. Secondly, the acquisition matrix was not the same between CS-MRA and PI-MRA (voxel size was adjusted to be the same). Setting the same values for both acquisition matrix and spatial resolution was not feasible due to the limitation of the sequences at the moment we collected the data. Thirdly, not all patients underwent DSA. Only 8 subjects had DSA within 6 months before and after examination of CS-MRA. DSA is invasive, and in the situation where MRI/MRA can depict time-course changes and offer reliable MMD grading, it is not considered essential. Furthermore, several studies[31,32] have verified a good correlation between MRA (Houkin's criteria) and DSA (Suzuki's criteria[33]). Fourthly, a relatively small number of patients were assessed, but it is considered sufficient to elucidate the capability of CS-TOF. Fifthly, only two CS-TOF acceleration rates (3 and 5) were evaluated. The research scans were added to the routine scan, and further additions were considered ethically inappropriate because of the burden on the patients.

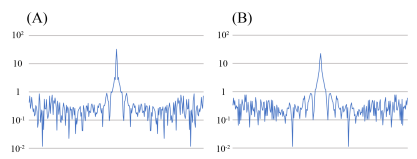


Fig 6. Point spread functions of the sampling pattern. (A) CS3, (B) CS5. The ratio of the main peak to the standard deviation of the pseudo-noise (incoherent artifacts) is 74.2 for CS3, and 35.1 for CS5. CS3, compressed sensing TOF MRA with an acceleration factor of 3; CS5, compressed sensing TOF MRA with an acceleration factor of 5.

https://doi.org/10.1371/journal.pone.0189493.g006



In conclusion, compressed sensing can accelerate TOF-MRA, while preserving the diagnostic capability for MMD. CS-TOF better visualized collateral MMVs in the equivalent acceleration. Moreover, five-fold accelerated CS-TOF was considered as acceptable for the proper diagnosis of MMD, allowing detection of steno-occlusive changes and MMVs. Shortening the scan time without reducing the clinically required coverage and spatial resolution brings benefits to patients, especially those who need to receive repeated life-time follow-up examinations. Further research is needed to investigate whether even higher acceleration rates can be used reliably in clinical practice.

Author Contributions

Conceptualization: Takayuki Yamamoto, Tomohisa Okada, Yasutaka Fushimi, Kaori Togashi.

Data curation: Tomohisa Okada, Yasutaka Fushimi, Akira Yamamoto, Takeshi Funaki.

Formal analysis: Takayuki Yamamoto, Tomohisa Okada, Yasutaka Fushimi, Koji Fujimoto, Sachi Okuchi, Hikaru Fukutomi.

Funding acquisition: Kaori Togashi.

Investigation: Takayuki Yamamoto, Yasutaka Fushimi, Akira Yamamoto, Koji Fujimoto, Sachi Okuchi, Hikaru Fukutomi, Jun C. Takahashi, Takeshi Funaki, Susumu Miyamoto.

Methodology: Takayuki Yamamoto, Yasutaka Fushimi, Akira Yamamoto, Koji Fujimoto, Aurélien F. Stalder, Yutaka Natsuaki, Peter Speier.

Project administration: Tomohisa Okada, Akira Yamamoto, Kaori Togashi.

Resources: Koji Fujimoto, Jun C. Takahashi, Takeshi Funaki, Susumu Miyamoto, Aurélien F. Stalder, Yutaka Natsuaki, Peter Speier, Kaori Togashi.

Software: Aurélien F. Stalder, Yutaka Natsuaki, Peter Speier.

Supervision: Tomohisa Okada, Akira Yamamoto, Susumu Miyamoto, Kaori Togashi.

Validation: Akira Yamamoto, Koji Fujimoto, Jun C. Takahashi, Takeshi Funaki, Susumu Miyamoto.

Visualization: Takayuki Yamamoto.

Writing – original draft: Takayuki Yamamoto, Tomohisa Okada, Yasutaka Fushimi, Akira Yamamoto, Koji Fujimoto, Sachi Okuchi, Hikaru Fukutomi.

Writing – review & editing: Jun C. Takahashi, Takeshi Funaki, Susumu Miyamoto, Aurélien F. Stalder, Yutaka Natsuaki, Peter Speier, Kaori Togashi.

References

- Lustig M, Donoho D, Pauly JM. Sparse MRI: The application of compressed sensing for rapid MR imaging. Magn Reson Med. Wiley Subscription Services, Inc., A Wiley Company; 2007; 58: 1182–1195. https://doi.org/10.1002/mrm.21391 PMID: 17969013
- Vincenti G, Monney P, Chaptinel J, Rutz T, Coppo S, Zenge MO, et al. Compressed Sensing Single– Breath-Hold CMR for Fast Quantification of LV Function, Volumes, and Mass. JACC Cardiovasc Imaging. 2014; 7: 882–892. https://doi.org/10.1016/j.jcmg.2014.04.016 PMID: 25129517
- Rapacchi S, Natsuaki Y, Plotnik A, Gabriel S, Laub G, Finn JP, et al. Reducing view-sharing using compressed sensing in time-resolved contrast-enhanced magnetic resonance angiography. Magn Reson Med. 2014; 74: 474–481. https://doi.org/10.1002/mrm.25414 PMID: 25157749
- Addy NO, Ingle RR, Wu HH, Hu BS, Nishimura DG. High-resolution variable-density 3D cones coronary MRA. Magn Reson Med. 2015; 74: 614–621. https://doi.org/10.1002/mrm.25803 PMID: 26172829



- Gratz M, Schlamann M, Goericke S, Maderwald S, Quick HH. Evaluation of fast highly undersampled contrast-enhanced MR angiography (sparse CE-MRA) in intracranial applications—initial study. Eur Radiol. 2017; 27: 1004–1011. https://doi.org/10.1007/s00330-016-4398-z PMID: 27300194
- Çukur T, Lustig M, Nishimura DG. Improving non-contrast-enhanced steady-state free precession angiography with compressed sensing. Magn Reson Med. 2009; 61: 1122–1131. https://doi.org/10.1002/mrm.21907 PMID: 19230013
- Hutter J, Grimm R, Forman C, Hornegger J, Schmitt P. Highly undersampled peripheral Time-of-Flight magnetic resonance angiography: optimized data acquisition and iterative image reconstruction. MAGMA. 2015; 28: 437–446. https://doi.org/10.1007/s10334-014-0477-9 PMID: 25605300
- Moghari MH, Annese D, Geva T, Powell AJ. Three-dimensional heart locator and compressed sensing for whole-heart MR angiography. Magn Reson Med. 2015; 75: 2086–2093. https://doi.org/10.1002/ mrm.25800 PMID: 26069182
- Kang C-K, Son Y-D, Kim H-K. Energy Spectrum-based Variable-Density Sampling Distribution Optimized for MR Angiography at Compressed Sensing Technique. Applied Magnetic Resonance. Springer Vienna; 2015; 47: 201–210. https://doi.org/10.1007/s00723-015-0742-8
- Ilicak E, Cetin S, Bulut E, Oguz KK, Saritas EU, Unal G, et al. Targeted vessel reconstruction in noncontrast-enhanced steady-state free precession angiography. NMR Biomed. 2016; 29: 532–544. https://doi.org/10.1002/nbm.3497 PMID: 26854004
- Yamamoto T, Fujimoto K, Okada T, Fushimi Y, Stalder AF, Natsuaki Y, et al. Time-of-Flight Magnetic Resonance Angiography With Sparse Undersampling and Iterative Reconstruction. Invest Radiol. 2016; 51: 372–378. https://doi.org/10.1097/RLI.000000000000221 PMID: 26561046
- Sodickson DK, Manning WJ. Simultaneous acquisition of spatial harmonics (SMASH): fast imaging with radiofrequency coil arrays. Magn Reson Med. 1997; 38: 591–603. PMID: 9324327
- Griswold MA, Jakob PM, Heidemann RM, Nittka M, Jellus V, Wang J, et al. Generalized autocalibrating partially parallel acquisitions (GRAPPA). Magn Reson Med. Wiley Subscription Services, Inc., A Wiley Company; 2002; 47: 1202–1210. https://doi.org/10.1002/mrm.10171 PMID: 12111967
- Pruessmann KP, Weiger M, Scheidegger MB, Boesiger P. SENSE: sensitivity encoding for fast MRI. Magn Reson Med. 1999; 42: 952–962. https://doi.org/10.1002/(SICI)1522-2594(199911)42:5<952:: AID-MRM16>3.0.CO;2-S PMID: 10542355
- Fushimi Y, Fujimoto K, Okada T, Yamamoto A, TANAKA T, Kikuchi T, et al. Compressed Sensing 3-Dimensional Time-of-Flight Magnetic Resonance Angiography for Cerebral Aneurysms. Invest Radiol. 2016; 51: 228–235. https://doi.org/10.1097/RLI.00000000000226 PMID: 26606551
- Fushimi Y, Okada T, Kikuchi T, Yamamoto A, Okada T, Yamamoto T, et al. Clinical evaluation of timeof-flight MR angiography with sparse undersampling and iterative reconstruction for cerebral aneurysms. NMR Biomed. 2017; 30. https://doi.org/10.1002/nbm.3774 PMID: 28796397
- Kikuta K-I, Takagi Y, Fushimi Y, Ishizu K, Okada T, Hanakawa T, et al. "Target bypass": a method for preoperative targeting of a recipient artery in superficial temporal artery-to-middle cerebral artery anastomoses. Neurosurgery. Oxford University Press; 2008; 62: 1434–1441. https://doi.org/10.1227/01. neu.0000333807.64530.39 PMID: 18695562
- Yamada I, Matsushima Y, Suzuki S. Moyamoya disease: diagnosis with three-dimensional time-of-flight MR angiography. Radiology. 1992; 184: 773–778. https://doi.org/10.1148/radiology.184.3.1509066
 PMID: 1509066
- Saeki N, Silva MN, Kubota M, Takanashi J, Sugita K, Nakazaki S, et al. Comparative performance of magnetic resonance angiography and conventional angiography in moyamoya disease. Journal of Clinical Neuroscience. 2000; 7: 112–115. https://doi.org/10.1054/jocn.1999.0160 PMID: 10844793
- Cho B-K, Tominaga T, editors. Moyamoya Disease Update. 1st ed. Tokyo: Springer Japan; 2010. https://doi.org/10.1007/978-4-431-99703-0
- 21. Stalder AF, Natsuaki Y, Schmidt M, Bi X, Zenge MO, Nadar MS, et al. Accelerating TOF MRA in Clinical Practice using Sparse MRI with Variable Poisson Density Sampling. Proceedings of the 23rd Annual Meeting of the ISMRM, Tronto, Ontario, Canada; 2015. p. 3606.
- 22. Beck A, Teboulle M. A Fast Iterative Shrinkage-Thresholding Algorithm for Linear Inverse Problems. SIAM J Imaging Sci. 2009; 2: 183–202. https://doi.org/10.1137/080716542
- 23. Wang J, Kluge T, Nittka M, Jellus V, Kuehn B, Kiefer B. Parallel acquisition techniques with modified SENSE reconstruction mSENSE. Proceedings of the first Wurzburg workshop on parallel imaging basics and clinical applications. Wurzburg, Germany; 2001. p. 89.
- **24.** Houkin K, Nakayama N, Kuroda S, Nonaka T, Shonai T, Yoshimoto T. Novel magnetic resonance angiography stage grading for moyamoya disease. Cerebrovasc Dis. 2005; 20: 347–354. https://doi.org/10.1159/000087935 PMID: 16131804



- 25. Research Committee on the Pathology and Treatment of Spontaneous Occlusion of the Circle of Willis, Health Labour Sciences Research Grant for Research on Measures for Infractable Diseases. Guidelines for diagnosis and treatment of moyamoya disease (spontaneous occlusion of the circle of Willis). Neurologia medico-chirurgica. 2012. pp. 245–266. PMID: 22870528
- Sawada T, Yamamoto A, Miki Y, Kikuta K-I, Okada T, Kanagaki M, et al. Diagnosis of moyamoya disease using 3-T MRI and MRA: value of cisternal moyamoya vessels. Neuroradiology. 2nd ed. 2012; 54: 1089–1097. https://doi.org/10.1007/s00234-012-1020-1 PMID: 22349623
- 27. Mikami T, Sugino T, Ohtaki S, Houkin K, Mikuni N. Diagnosis of Moyamoya Disease on Magnetic Resonance Imaging: Are Flow Voids in the Basal Ganglia an Essential Criterion for Definitive Diagnosis? Journal of Stroke and Cerebrovascular Diseases. Elsevier Ltd; 2013; 22: 862–868. https://doi.org/10.1016/j.jstrokecerebrovasdis.2012.07.010 PMID: 22939199
- 28. Okuchi S, Okada T, Fujimoto K, Fushimi Y, Kido A, Yamamoto A, et al. Visualization of Lenticulostriate Arteries at 3T: Optimization of Slice-selective Off-resonance Sinc Pulse-prepared TOF-MRA and Its Comparison with Flow-sensitive Black-blood MRA. Acad Radiol. 2014; 21: 812–816. https://doi.org/10.1016/j.acra.2014.03.007 PMID: 24809322
- Okada T, Kanao S, Ninomiya A, Sato S, Kuhara S, Kamae T, et al. Whole-heart coronary magnetic resonance angiography with parallel imaging: Comparison of acceleration in one-dimension vs. two-dimensions. European Journal of Radiology. 2009; 71: 486–491. https://doi.org/10.1016/j.ejrad.2008.06.005
 PMID: 18640801
- Seo N, Park M-S, Han K, Kim D, King KF, Choi J-Y, et al. Feasibility of 3D navigator-triggered magnetic resonance cholangiopancreatography with combined parallel imaging and compressed sensing reconstruction at 3T. J Magn Reson Imaging. 2017; 46: 1289–1297. https://doi.org/10.1002/jmri.25672 PMID: 28295827
- Kuroda S, Houkin K. Moyamoya disease: current concepts and future perspectives. The Lancet Neurology. Elsevier Ltd; 2008; 7: 1056–1066. https://doi.org/10.1016/S1474-4422(08)70240-0
- Jin Q, Noguchi T, Irie H, Kawashima M, Nishihara M, Takase Y, et al. Assessment of Moyamoya disease with 3.0-T magnetic resonance angiography and magnetic resonance imaging versus conventional angiography. Neurol Med Chir(Tokyo). 2011; 51: 195–200. https://doi.org/10.2176/nmc.51.195
- Suzuki J, Takaku A. Cerebrovascular "moyamoya" disease. Disease showing abnormal net-like vessels in base of brain. Arch Neurol. 1969; 20: 288–299. PMID: 5775283