INTRODUCTION

Time-of-flight (TOF) magnetic resonance angiography (MRA) is a non-invasive, non-contrast-enhanced MRA technique that provides contrast between vessels and stationary tissues by inducing blood inflow effects (1-4). Three-dimensional (3D) TOF MRA with multiple slabs is performed widely to depict the cerebral vasculature in the diagnosis of cerebral vascular pathologies such as intracranial occlusions, intracranial aneurysm, and arteriovenous malformations (5,6). However, conventional TOF techniques suffer from relatively long acquisition times.
for images with high spatial resolution and large coverage, which often require longer than 5 minutes. Parallel imaging techniques such as generalized autocalibrating partially parallel acquisitions (GRAPPA) and modified sensitivity encoding (mSENSE) allow imaging acceleration by a factor of 2–3, but further reduction of the acquisition time is challenging because of the reduced signal-to-noise ratio (SNR) (7, 8).

To overcome this limitation, sparse methods, which exploit known sparsity properties of the data in transform domains, in combination with incoherent k-space sampling and non-linear reconstruction algorithms were recently proposed to further accelerate the acquisition speed in magnetic resonance imaging (MRI) (9-11). Because of the intrinsic sparsity of the pixel domain and high contrast-to-noise ratio, contrast-enhanced and non-contrast-enhanced MRA are excellent candidates for sparse methods (12-14). However, because of the long reconstruction times and often off-line reconstructions, the clinical application and evaluation of sparse methods remain limited (12, 15).

In this context, a sparse TOF MRA using sparse methods with inline iterative reconstruction on a standard clinical system has been suggested recently (16, 17) and applied to cerebral aneurysms (18).

The purpose of this study was to prospectively compare the image quality of 3D sparse TOF MRA with incoherent sampling and iterative reconstruction with that of conventional TOF MRA in a clinical setting and on a standard clinical MR system.

**MATERIALS AND METHODS**

**Subjects**

This study was approved by the local Institutional Review Board and written informed consent was obtained from all patients. A total number of 56 patients with dizziness (33 males and 23 females; mean age, 53.4 ± 18.6 years) were enrolled in this study between March 2017 and July 2017.

**MRI**

The MRI examination was performed on a 3T magnetic resonance (MR) scanner (MAGNETOM Skyra, Siemens Healthineers, Erlangen, Germany) with a 20-channel phased array head-neck coil for evaluation of the intracranial arteries. To compare the image qualities of sparse and conventional TOF MRA, all patients were divided into two groups (Table 1) to be examined by prototype sparse and conventional TOF sequences with 1) nearly matched acquisition times and 2) nearly matched spatial resolutions using the acquisition parameters shown in Table 2.

| Table 1. Clinical Characteristic of Subjects |
|---------------------------------------------|
| Parameters | TOF and Sparse TOF with Same Resolution | TOF and Sparse TOF with Same Scan Time |
| Number of patients | 27 | 29 |
| Age (years) | 48.6 ± 18.3 | 57.8 ± 18.0 |
| Sex (male/female) | 16/11 | 17/12 |

Sparse TOF = TOF with sparse undersampling and iterative reconstruction, TOF = time-of-flight

| Table 2. MRI Parameters of TOF and Sparse TOF |
|---------------------------------------------|
| Parameter | TOF and Sparse TOF with Same Resolution | TOF and Sparse TOF with Same Scan Time |
| TE/TR (ms) | 3.43/21 | 3.43/21 | 3.43/21 | 3.43/21 |
| FOV (mm²) | 200 x 180 | 200 x 180 | 200 x 180 | 200 x 180 |
| Voxel size (interpolated, mm³) | 0.3 x 0.3 x 0.4 | 0.3 x 0.3 x 0.4 | 0.3 x 0.3 x 0.6 | 0.3 x 0.3 x 0.4 |
| Slices per slab | 56 | 56 | 36 | 60 |
| Slabs | 4 | 4 | 4 | 4 |
| Slab distance factor (%) | -19.6 | -19.6 | -16.7 | -16.7 |
| Acceleration method | GRAPPA | Sparse sampling | GRAPPA | Sparse sampling |
| Acceleration factor | 2 | 3.5 | 3 | 5 |
| Partial fourier | 7/8 | NA | 7/8 | NA |
| TA (min) | 6:32 | 4:10 | 3:44 | 3:52 |

FOV = field of view, GRAPPA = generalized autocalibrating partially parallel acquisition, NA = not applicable, TA = acquisition time, TE = echo time, TR = repetition time
Image Sampling and Reconstruction

For the conventional TOF sequence, a GRAPPA parallel imaging technique with an accelerator factor of 2 or 3 was applied along the phase-encoding (left-right) direction. A slice partial Fourier value of 7/8 was used in the image acquisition. For the sparse TOF prototype sequence, each slab was acquired using a variable-density Poisson disk sampling pattern in the $k_x - k_z$ phase-encoding plane (17-22) with acceleration factors of 3.5 and 5.

Image reconstruction was performed using a redundant Haar wavelet-based L1-regularized iterative SENSE algorithm with fast iterative shrinkage and a thresholding algorithm (FISTA) integrated inline on the clinical scanner (17, 23). More specifically, it was achieved by solving the following minimization problem:

$$\min_x \left\{ \frac{1}{2} \| y - Fx \|_2^2 + \lambda \| Wx \|_1 \right\},$$

where $x$ is the reconstructed image, $y$ is the undersampled k-space data, $F$ is the undersampled Fourier operator, and $W$ and $\lambda$ represent the redundant Haar wavelet transform and normalized regularization weighting factor, respectively. The number of FISTA iterations was set to 10 and the regularization weighting factor $\lambda$ was set to 0.008 for sparse TOF acquisition according to a previous report (12). The total image reconstruction time was about 5 minutes. Afterward, sparse TOF images acquired with net acceleration factors of 3.5 and 5 were compared with conventional TOF images acquired using GRAPPA acceleration factors of 2 and 3, respectively. Both the image sampling and reconstruction were carried out on the scanner.

Image Analysis

Image evaluation was performed on a standard post-processing workstation by two neuroradiologists (with 5 and 26 years of experience) who were blinded to the acquisition parameters and patients’ information. For the qualitative analysis, the delineation of the vessels was assessed using a five-point grading scale (5-the display of vessels was excellent and with sharp borders; 4-the display of vessels was good and with clear borders; 3-the display of vessels was moderate and with visible borders; 2-the display of vessels was poor with dim borders and affected the diagnosis; and 1-the display of vessels was unclear and could not be diagnosed). The vessels that were used for evaluation included the main branch, primary sub-branch, and secondary sub-branch of the anterior cerebral artery (ACA), middle cerebral artery (MCA), posterior cerebral artery (PCA), anterior communicating artery (ACoA), posterior communicating artery (PCoA), internal carotid artery (ICA), and basilar artery (BA). The artifacts, overall vessel visualization, and overall image quality were also assessed using a five-point grading scale. The detailed evaluation standards are listed as follows:

1) Artifacts: 5-no artifact; 4-slight aliasing/blurring artifacts that did not affect the diagnosis; 3-a few aliasing/blurring artifacts and slight motion artifacts (the images could be diagnosed); 2-some aliasing/blurring artifacts and significant motion artifacts (artifacts affecting the diagnosis moderately); and 1-severe artifacts (the images could not be diagnosed).

2) Overall vessel visualization: 5-the display of all above-mentioned vessels was excellent; 4-the display of the main and primary branches was good and the secondary branches were visible; 3-the display of the main and primary branches was good but the secondary branches were barely visible; 2-the display of the main branches was good but the primary vessels were barely visible; and 1-the main branches were visible but the primary branches were invisible.

3) Overall image quality: 5-the display of vessels was excellent, sharp, and without artifacts; 4-the display of vessels was good with clear borders and slight artifacts; 3-the display of vessels was moderate with visible borders and a few artifacts, (diagnostic image quality); 2-the display of vessels was poor with blurry borders and some artifacts (image quality moderately affected the diagnosis); and 1-the display of vessels was unclear with severe artifacts (non-diagnostic image quality).

For the quantitative measurement, the signal intensities (SIs) of the vessels and static background tissue were measured on original images and maximum intensity projection (MIP) images to calculate the contrast ratio (CR) as

$$CR = (SI_{vessel} - SI_{background}) / (SI_{vessel} + SI_{background})$$

where $SI_{vessel}$ is the SI of the vessels and $SI_{background}$ is the SI of the static background tissue (24). Regions of interest (ROIs) were placed in the same position on sparse and conventional TOF MRA images from each patient on the proximal left M1 segment of the MCA and the adjacent gray matter. The sizes of the ROIs varied slightly to account for anatomical variations and to maximize ROI size without including voxels from other tissues.
| Parameter | TOF and Sparse TOF with Same Resolution | TOF and Sparse TOF with Same Scan Time |
|-----------|----------------------------------------|----------------------------------------|
|           | TOF | Sparse TOF | TOF | Sparse TOF |
| ACA       |     |            |     |            |
| Main branch |     |            |     |            |
| R1        | 4.9 ± 0.4 | 4.9 ± 0.4 | 4.8 ± 0.4 | 5.0 ± 0.0* |
| R2        | 4.8 ± 0.5 | 4.9 ± 0.5 | 4.8 ± 0.4 | 5.0 ± 0.0* |
| κ         | 0.84 | 0.65       | 0.89 | 1.00       |
| Primary   |     |            |     |            |
| R1        | 4.7 ± 0.7 | 4.7 ± 0.5 | 4.5 ± 0.6 | 4.9 ± 0.3** |
| R2        | 4.6 ± 0.7 | 4.8 ± 0.5 | 4.4 ± 0.6 | 4.9 ± 0.3** |
| κ         | 0.91 | 0.89       | 0.87 | 0.78       |
| Secondary |     |            |     |            |
| R1        | 3.6 ± 1.0 | 3.3 ± 0.9 | 2.8 ± 1.2 | 3.8 ± 1.4** |
| R2        | 3.6 ± 0.9 | 3.4 ± 0.9 | 2.7 ± 1.0 | 3.9 ± 1.4** |
| κ         | 0.84 | 0.89       | 0.81 | 0.89       |
| MCA       |     |            |     |            |
| Main branch |     |            |     |            |
| R1        | 4.9 ± 0.6 | 4.9 ± 0.3 | 4.9 ± 0.3 | 5.0 ± 0.2 |
| R2        | 4.8 ± 0.6 | 4.9 ± 0.3 | 4.9 ± 0.3 | 5.0 ± 0.2 |
| κ         | 0.84 | 0.78       | 0.78 | 1.00       |
| Primary   |     |            |     |            |
| R1        | 4.9 ± 0.6 | 4.9 ± 0.4 | 4.9 ± 0.3 | 4.9 ± 0.3 |
| R2        | 4.8 ± 0.6 | 4.9 ± 0.5 | 4.8 ± 0.4 | 5.0 ± 0.2 |
| κ         | 0.79 | 0.65       | 0.87 | 0.65       |
| Secondary |     |            |     |            |
| R1        | 4.3 ± 0.9 | 4.3 ± 0.8 | 3.5 ± 0.8 | 4.3 ± 0.7** |
| R2        | 4.2 ± 0.9 | 4.3 ± 0.8 | 3.5 ± 0.8 | 4.3 ± 0.7** |
| κ         | 0.94 | 0.94       | 0.95 | 0.89       |
| PCA       |     |            |     |            |
| Main branch |     |            |     |            |
| R1        | 4.9 ± 0.6 | 5.0 ± 0.2 | 4.6 ± 0.6 | 4.8 ± 0.4 |
| R2        | 4.8 ± 0.8 | 4.9 ± 0.3 | 4.7 ± 0.5 | 4.9 ± 0.3 |
| κ         | 0.65 | 0.65       | 0.78 | 0.87       |
| Primary   |     |            |     |            |
| R1        | 4.8 ± 0.6 | 4.7 ± 0.4 | 3.3 ± 1.0 | 4.5 ± 0.7** |
| R2        | 4.7 ± 0.7 | 4.8 ± 0.4 | 3.3 ± 1.0 | 4.5 ± 0.6** |
| κ         | 0.86 | 0.90       | 0.94 | 0.94       |
| Secondary |     |            |     |            |
| R1        | 3.9 ± 0.8 | 3.9 ± 0.8 | 2.0 ± 1.0 | 3.2 ± 1.3** |
| R2        | 3.9 ± 0.7 | 3.9 ± 0.8 | 2.0 ± 1.0 | 3.1 ± 1.3** |
| κ         | 0.87 | 0.88       | 0.90 | 0.95       |
| ACoA      |     |            |     |            |
| R1        | 3.5 ± 1.4 | 3.4 ± 1.5 | 3.3 ± 1.4 | 3.5 ± 1.4* |
| R2        | 3.5 ± 1.4 | 3.4 ± 1.5 | 3.2 ± 1.3 | 3.5 ± 1.5* |
| κ         | 0.90 | 0.91       | 0.87 | 0.86       |
| PCoA      |     |            |     |            |
| R1        | 3.6 ± 1.1 | 3.7 ± 1.0 | 3.5 ± 1.0 | 4.0 ± 0.9** |
| R2        | 3.6 ± 1.1 | 3.7 ± 1.0 | 3.4 ± 1.0 | 4.1 ± 0.9** |
| κ         | 0.89 | 0.95       | 0.90 | 0.89       |
Statistical Analysis

The statistical analysis was performed by using SPSS 19.0 (IBM Corp., Armonk, NY, USA). Numerical data are presented as the mean ± the standard deviation. The non-parametric Wilcoxon signed-rank test was used for qualitative comparisons between sparse and conventional TOF MRA. The paired t test was used for comparisons of CRs. Interobserver agreements for the delineation of arterial vessel segments,

Table 3. Averaged Readers’ Scores of TOF and Sparse TOF (continued)

| Parameter                  | TOF and Sparse TOF with Same Resolution | TOF and Sparse TOF with Same Scan Time |
|----------------------------|----------------------------------------|----------------------------------------|
|                            | TOF                                   | Sparse TOF                             | TOF                                   | Sparse TOF                             |
| ICA                        |                                        |                                        |                                       |
| R1                         | 5.0 ± 0.0                              | 5.0 ± 0.0                              | 4.9 ± 0.3                              | 5.0 ± 0.0*                             |
| R2                         | 5.0 ± 0.0                              | 5.0 ± 0.0                              | 4.8 ± 0.4                              | 5.0 ± 0.2*                             |
| \(\kappa\)                 | 1.00                                   | 1.00                                   | 0.87                                   | 0.86                                   |
| BA                         |                                        |                                        |                                       |
| R1                         | 5.0 ± 0.0                              | 5.0 ± 0.0                              | 4.9 ± 0.3                              | 5.0 ± 0.0                             |
| R2                         | 5.0 ± 0.0                              | 5.0 ± 0.0                              | 4.9 ± 0.3                              | 5.0 ± 0.0                             |
| \(\kappa\)                 | 1.00                                   | 1.00                                   | 0.84                                   | 1.00                                   |
| Artifacts                  |                                        |                                        |                                       |
| R1                         | 4.6 ± 0.8                              | 4.7 ± 0.6                              | 4.2 ± 0.5                              | 4.8 ± 0.5**                           |
| R2                         | 4.6 ± 0.8                              | 4.7 ± 0.6                              | 4.2 ± 0.5                              | 4.8 ± 0.5**                           |
| \(\kappa\)                 | 0.92                                   | 0.89                                   | 0.93                                   | 0.89                                   |
| Overall vessel visualization|                                        |                                        |                                       |
| R1                         | 5.0 ± 0.2                              | 4.9 ± 0.6                              | 4.4 ± 0.5                              | 4.9 ± 0.3**                           |
| R2                         | 4.9 ± 0.3                              | 4.8 ± 0.6                              | 4.3 ± 0.5                              | 4.8 ± 0.4**                           |
| \(\kappa\)                 | 0.78                                   | 0.79                                   | 0.85                                   | 0.87                                   |
| Overall image quality      |                                        |                                        |                                       |
| R1                         | 4.7 ± 0.6                              | 4.9 ± 0.4                              | 4.1 ± 0.5                              | 4.8 ± 0.4**                           |
| R2                         | 4.7 ± 0.7                              | 4.8 ± 0.5                              | 4.1 ± 0.5                              | 4.7 ± 0.4**                           |
| \(\kappa\)                 | 0.89                                   | 0.84                                   | 0.84                                   | 0.91                                   |

Sparse TOF compared with TOF (*\(p < 0.05\), **\(p < 0.01\)). ACA = anterior cerebral artery, ACoA = anterior communicating artery, BA = basilar artery, ICA = internal carotid artery, MCA = middle cerebral artery, PCA = posterior cerebral artery, PCoA = posterior communicating artery

Fig. 1. Axial MIP images from conventional TOF (A) and sparse TOF (B) using same spatial resolution in 60-year-old man. Display of lesion in left MCA is poorer on conventional TOF (A) because of motion artifacts (arrow). MCA = middle cerebral artery, MIP = maximum intensity projection, sparse TOF = TOF with sparse undersampling and iterative reconstruction, TOF = time-of-flight
artifacts, overall vessel visualization, and overall image quality scores were calculated by using the kappa test (κ values of 0.00–0.20 were considered to indicate poor agreement; κ values of 0.21–0.40, fair agreement; κ values of 0.41–0.60, moderate agreement; κ values of 0.61–0.80, good agreement; and κ values of 0.81–1.00, excellent agreement). A p value less than 0.05 was considered to indicate statistical significance.

RESULTS

The acquisition and image reconstruction were successful for all 56 patients. Among them, five were diagnosed as having vessel stenoses, while the others presented healthy intracranial vasculatures.

When comparing sparse and conventional TOF MRA with similar spatial resolutions, sparse TOF showed comparable image quality regarding the delineation of vessel segments, artifacts, overall vessel visualization, and overall image quality compared with conventional TOF (Table 3). Even
in patients with poor compliance, motion artifacts were reduced in sparse TOF MRA (Fig. 1). In the quantitative analysis, the CRs showed no difference on the original images, but a slight increase was observed on MIP images for sparse TOF compared with conventional TOF (0.64 ± 0.05 vs. 0.61 ± 0.05, \( p < 0.001 \)) (Fig. 2A, B).

As shown in Figure 3 for two representative datasets, the presence of artifacts, overall vessel visualization, and overall image quality were superior for sparse TOF when compared with conventional TOF with a similar scanning time. The delineation of vessel segments was also significantly increased in the main branches of the ACA, ACoA, PCoA, and ICA, the primary sub-branches of the ACA and PCA, and the secondary sub-branches of the ACA, MCA, and PCA (Table 3). Comparing sparse TOF with conventional TOF, the CR increased from 0.58 ± 0.06 to 0.60 ± 0.06 on the original images (\( p < 0.05 \)) and from 0.59 ± 0.04 to 0.64 ± 0.04 on the MIP images (\( p < 0.001 \)) (Fig. 2C, D).

Fig. 3. Axial MIP images from conventional TOF (A, C) and sparse TOF (B, D) using same scanning time in two patients. A and B are from 46-year-old woman and C and D are from 70-year-old man with left MCA stenosis. Delineation of fine vessels is better with sparse TOF than conventional TOF (arrowheads). Moreover, display of vessel stenosis is clearer with sparse TOF than with conventional TOF (arrows).
Good-to-excellent interobserver agreement (κ values: 0.65–1.00) was obtained between the two radiologists regarding the delineation of arterial vessel segments, artifacts, overall vessel visualization, and overall image quality scores (Table 3).

DISCUSSION

In the present work, we evaluated a sparse TOF technique for the acceleration of data acquisition in the context of non-contrast-enhanced MRA integrated into a clinical scanner. The performance of sparse TOF with a net acceleration factor of up to 5 was evaluated and compared with that of conventional TOF. Based on comparisons with protocols using either a matched spatial resolution or matched scanning time, we found that the sparse TOF technique reduced the acquisition time by 40% while maintaining comparable image quality and achieved superior results in cases with matched scanning times between sparse and conventional TOF.

High spatial resolution is essential for the detection of small-vessel disease and aids doctors in discovering more vascular lesions and developing the treatment plan. The major drawback of conventional TOF MRA with high resolution is the long acquisition time, which also renders the resulting images susceptible to artifacts due to any type of motion. Conventional parallel imaging techniques, such as mSENSE and GRAPPA, can reduce the acquisition time by reducing the number of phase encoding steps. It is well known that higher acceleration factors can be achieved with more coil elements because the parallel imaging technique needs the coil elements’ phase information to reconstruct the final images. In addition, a higher SNR can be obtained with more elements (25). Therefore, it is useful to perform TOF MRA with multiple coil elements. However, an acceleration factor of 2–3 is feasible with conventional TOF. A further increase in parallel factors results in significant SNR loss (26, 27). Compared with conventional TOF, the acceleration factor of sparse TOF was affected by the undersampling factor. Furthermore, the SNRs of reconstructed images from sparse TOF were also associated with undersampling and the iteration steps in the image reconstruction (12). Therefore, compared with conventional TOF, sparse TOF can use a higher acceleration factor with the same phased array coils. In our study, as conventional TOF and sparse TOF were obtained using the same 20-channel head coil, the effect of the coil elements on the SNR would be the same, but the acceleration factor can be set higher in sparse TOF than in conventional TOF.

Sparse MRI approaches matching the basic requirement for both high acceleration factors and high SNR have been reported previously for contrast-enhanced MRA, whole-heart coronary MRA, peripheral TOF MRA (9, 12, 14, 15, 28), as well as intracranial TOF MRA (16, 17). However, because of the long reconstruction times and/or limited availability of inline reconstruction algorithms at present, the clinical evaluation of compressed-sensing MRI techniques remains limited. In this work, we used a recently reported sparse TOF technique that was fully integrated on a clinical scanner for the clinical evaluation of 56 patients.

Our results demonstrated that with a net acceleration factor of 5, sparse TOF could acquire MRA images with a voxel size of 0.3 x 0.3 x 0.4 mm³ within 4 minutes with much higher image quality than that of conventional TOF with a similar scanning time. The reconstruction time for sparse TOF was 4–5 minutes for each patient on our MRI scanner. By using more advanced graphics processing units, a reduction in reconstruction time is possible. In our protocol, T₁-weighted imaging, T₂-weighted imaging, T₂ fluid-attenuated inversion recovery, and diffusion-weighted imaging were acquired for clinical diagnosis in addition to TOF MRA. These sequences can be performed while the MRI scanner is simultaneously reconstructing the sparse TOF MRA. Therefore, it does not increase the total scanning time.

By using the same spatial resolution for both sequences, our results demonstrated that sparse TOF presented comparable image quality to that of conventional TOF with a remarkable reduction in scanning time. The overall image quality, overall vessel visualization, and presence of artifacts were similar for both sequences, and no significant differences were shown in the delineation of vessel segments. These results are consistent with those of previous reports (18), which found that cerebral aneurysms were visible on sparse TOF accelerated by a factor of 5 with equivalent quality to that of traditional TOF, while the scanning time of sparse TOF accelerated by a factor of 5 was shorter than that of conventional TOF. In our study, the CR in the M1 segment of the MCA showed no significant differences between the two techniques on the original images but showed a slight increase on the MIP images. This was caused mainly by motion artifacts in patients with poor compliance when using conventional TOF MRA as the acquisition time was longer.

To match the scanning time, the slice thickness was
increased for conventional TOF compared with sparse TOF with a higher acceleration factor, as the acceleration factor could not be increased to guarantee an adequate SNR. Considering this condition, thicker slices were needed for conventional MRA to obtain the same coverage and in-plane resolution as those of sparse TOF MRA. The scores of large-diameter vessels, including the main branches of the MCA, PCA, and BA, did not show any difference between the sequences. However, the delineation of small vessels was improved when using sparse TOF, which might be due to the volume averaging effect and iterative reconstruction. Another important reason might be a result of the iterative reconstruction. The process of image reconstruction using an iterative reconstruction algorithm can also be considered for image denoising. With an optimized regularization weighting factor, a good balance between the noise level and image details can be achieved (12), resulting in increased contrast of the small vessels. This result highlighted the usefulness of sparse TOF in the diagnoses of stenosis and small-vessel occlusion. Although the ratings of the main branches did not show any difference, a more detailed analysis using the CR showed that the CR of vessels increased significantly on the original and MIP images.

Our study has limitations. First, the acceleration factor, regularization, and number of iterations of the protocols were fixed to compare the two methods. Further assessment of image quality with regard to these parameters should be addressed. Second, the subjects in this study were mostly patients without lesions in vessels; thus, diagnostic accuracies were not compared in the current study.

In this study, sparse TOF with a variable-density Poisson disk sampling pattern and iterative reconstruction was successfully integrated on a standard clinical MR scanner. The total acquisition time was significantly reduced without any tradeoff in image quality.

Conflicts of Interest
The authors have no financial conflicts of interest.

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