3D Time-Resolved MR Angiography (MRA) of the Carotid Arteries with Time-Resolved Imaging with Stochastic Trajectories: Comparison with 3D Contrast-Enhanced Bolus-Chase MRA and 3D Time-Of-Flight MRA

BACKGROUND AND PURPOSE: Time-resolved MR angiography (MRA) offers the combined advantage of large anatomic coverage and hemodynamic flow information. We applied parallel imaging and time-resolved imaging with stochastic trajectories (TWIST), which uses a spiral trajectory to undersample k-space, to perform time-resolved MRA of the extracranial internal carotid arteries and compare it to time-of-flight (TOF) and high-resolution contrast-enhanced (HR) MRA.

MATERIALS AND METHODS: A retrospective review of 31 patients who underwent carotid MRA at 1.5T using TOF, time-resolved and HR MRA was performed. Images were evaluated for the presence and degree of ICA stenosis, reader confidence, and number of pure arterial frames attained with the TWIST technique.

RESULTS: With a consensus interpretation of all sequences as the reference standard, accuracy for identifying stenosis was 90.3% for TWIST MRA, compared with 96.0% and 88.7% for HR MRA and TOF MRA, respectively. HR MRA was significantly more accurate than the other techniques (P < .05). TWIST MRA yielded datasets with high in-plane spatial resolution and distinct arterial and venous phases. It provided dynamic information not otherwise available. Mean diagnostic confidence was satisfactory or greater for TWIST in all patients.

CONCLUSION: The TWIST technique consistently obtained pure arterial phase images while providing dynamic information. It is rapid, uses a low dose of contrast, and may be useful in specific circumstances, such as in the acute stroke setting. However, it does not yet have spatial resolution comparable with standard contrast-enhanced MRA.

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stroke has an estimated prevalence of 5.7 million (2.6%) adults in the United States, where it is the third most common cause of mortality.1 Atherosclerotic carotid artery disease is an important risk factor for anterior circulation ischemic stroke. Imaging evaluation is essential for optimal management and stroke prevention, as demonstrated in studies of symptomatic2,3 and asymptomatic populations.4,5 Carotid dissection is another potential cause of anterior circulation ischemia, which also requires high-spatial-resolution imaging for definitive diagnosis.

Digital subtraction angiography (DSA) remains the gold standard for assessment of the cervical vasculature,6 with excellent spatial and temporal resolution. However, risks include vascular injury, intracerebral complications, contrast nephrotoxicity, and exposure to ionizing radiation. Therefore, noninvasive techniques are typically used initially. Duplex Doppler sonography, CT angiography (CTA), and MR angiography (MRA) all have high but varying degrees of sensitivity and specificity, ranging from 70% to 99% for carotid stenosis detection.7 Although sonography provides excellent dynamic information and spatial resolution, insonation window limitations restrict anatomic coverage. Similarly, long imaging times with time-of-flight (TOF) MRA also limit anatomic coverage. CTA and conventional arterial contrast-enhanced MRA offer extensive coverage but provide no dynamic information and are dependent on accurate timing for optimal visualization of the arterial tree.

3D time-resolved contrast-enhanced MR angiography (TR MRA) has been previously described.8 TR MRA offers combined anatomic and hemodynamic information and obtains pure arterial and venous phase images consistently and rapidly without a timing run. More recently, the technique has been specifically applied to the extracranial carotid arteries using parallel imaging or keyhole imaging techniques.9,10 In this study, we report the diagnostic accuracy of TR MRA of the extracranial internal carotid artery (ICA) acquired with a combination of parallel imaging (generalized autocalibrating partially parallel acquisition [GRAPPA])11 and time-resolved imaging with stochastic trajectories (TWIST), a new view-sharing technique, which undersamples the periphery of k-space depending on the radial distance from the center of k-space.12,13 We compared TR MRA with 3D high-resolution contrast enhanced MRA (HR MRA) and 3D TOF MRA.

Materials and Methods

Patient Population

The study was institutional review board–approved, and a waiver of consent was obtained for this Health Insurance Portability and Ac-
countability Act–compliant retrospective study. Thirty-two consecutive patients were identified from the radiology data base who underwent carotid MRA on a 1.5 T magnet (Avanto; Siemens Medical Solutions, Erlangen, Germany) from November 2006 to March 2007. One patient was excluded secondary to extravasation of contrast during acquisition. Of the final cohort, there were 16 men and 15 women, with an age range of 21–89 years (mean age, 59.9 years). Clinical indications for MRA included transient ischemic attack or stroke (n = 18), known cervical atherosclerotic disease (n = 5), and dissection evaluation/follow-up (n = 8).

**MR Imaging**

Patients underwent carotid MRA with 3 MRA sequences: 3D TOF MRA, contrast-enhanced 3D TWIST MRA, and 3D HR MRA. GRAPPA\(^1\) was used for all sequences, with an acceleration factor of 2 used for the TWIST sequence. An 8-channel neurovascular array coil was used.

**TOF**

TOF coverage was limited to the cervical common carotid artery and carotid bifurcation, acquired axially, whereas the other 2 sequences acquired a sagittal oblique slab from the aortic arch to the circle of Willis. Imaging parameters were as follows: TR, 25 ms; TE, 7.2 ms; flip angle, 25\(^\circ\); FOV, 230 × 230 mm; matrix, 314 × 448; 90 partitions; voxel size after zero-interpolation, 0.7 × 0.5 × 0.9 mm\(^3\) (true voxel size, 0.7 × 0.5 × 1.4 mm); acceleration factor, 3; acquisition time, 2:47 minutes.

**Time-Resolved Imaging with TWIST**

A 6-mL bolus of gadopentetate dimeglumine (Gd-DTPA, Magnevist; Bayer, Wayne, NJ) was administered at 2 mL/s, followed by a 15-mL saline flush. Imaging parameters were the following: TR, 3.3 ms; TE, 1.3 ms; flip angle, 25\(^\circ\); rectangular field of view (rFOV), 246 × 375 mm; matrix, 210 × 320; 88 partitions; voxel size after zero-interpolation, 1.2 × 1.2 × 2.0 mm\(^3\) (true voxel size, 1.2 × 1.2 × 2.0 mm\(^3\)); acceleration factor, 2. Although detailed physics of the TWIST sequence are beyond the scope of this article, a brief description is provided. TWIST relies on partial k-space undersampling, with emphasis on more frequent sampling of the center of k-space (designated as region A), relative to the periphery of k-space (designated as region B). Region A governs image contrast and is updated more rapidly than region B, which governs fine image detail. (Missing data points in B for each sequential dataset are obtained from adjacent B intervals as shown in Fig 1. K-space sampling occurs in a continuous spiral fashion that allows a full range of k-space coverage from the center to the periphery and updating of high-frequency at the same rate as low-frequency information, though not all points in B are sampled during each acquisition.

The user may select the percentage of the center of k-space (A), compared with the entire k-space volume that is sampled with every acquisition (controlling image contrast), and the percentage of data points in the periphery of k-space (B) sampled per acquisition, with subsequent savings in acquisition time, at the expense of fine detail. Values of A = 10% and B = 15% were used for TWIST MRA in our series. The sequence had a temporal resolution of 3.9 seconds, interpolated to 1.95 seconds, with 9 sequential non-breath-hold datasets obtained (1 precontrast and 8 postcontrast). Automated subtraction and coronal and sagittal maximal intensity projection (MIP) images were scanner-generated for the TWIST sequence.

**Image Evaluation**

Two radiologists (with 1 year and 5 years’ experience) interpreted studies in random order, blinded to patient history and identity, on a dedicated imaging workstation (Leonardo; Siemens Medical Solutions), where source, subtraction images and 3D multplanar reformatting and MIP were available. A quantitative assessment of the greatest percentage of stenosis was made in each cervical ICA, by using North American Symptomatic Carotid Endarterectomy Trial guidelines\(^1\) and by comparing the diameter of the most stenotic segment with the closest distal segment judged to be normal in caliber. Stenosis measurements were made from source data viewed in multplanar reconstruction mode, with reference to subtraction images for TWIST and HR sequences as required. Stenoses were subsequently characterized into clinically relevant categories: mild (0%–49%), moderate (50%–69%), severe (70%–99%), or occluded (100%), for data analysis. Sequences were evaluated for diagnostic confidence on a 3-point scale (1 = poor, 2 = satisfactory, 3 = excellent). The TWIST sequence was evaluated for the number of arterial frames in which contrast was present at the carotid bifurcation before venous opacification.

Plaque surface morphology (1 = smooth, 2 = ulcerated) and dissection, if present, were recorded. Other vascular pathology affecting the vertebral, common carotid, and external carotid arteries was recorded, along with pathology of the aortic arch for the contrast-enhanced sequences. ARTifacts hindering interpretation were noted. The reference standard was a consensus reading by both radiologists of all
sequences and DSA correlation, where available (n = 2, within 1 and 69 days of MRA without interval intervention).

A radiologist with 2 years’ experience evaluated signal-intensity-to-noise ratio (SNR) and contrast-to-noise ratio (CNR) for each sequence (Appendix).

Statistical Analysis
Because all 3 sequences were used to derive data for each subject, statistical techniques appropriate for the analysis of correlated data were used to evaluate and compare the sequences. Specifically, the sequences were compared in terms of average reader confidence, SNR, and CNR by using a Wilcoxon matched-pairs signed rank test (ie, a Wilcoxon signed rank test applied to the within-subject differences between the sequences being compared). Logistic regression for correlated data (generalized estimating equations based on a binary logistic regression model) was used to compare sequences for diagnostic accuracy relative to consensus opinion as the reference standard. For these comparisons, the stenosis grade for specific sequences was declared accurate only when in exact agreement with the consensus characterization. Stenoses outside the TOF MRA FOV were recorded as incorrect assessments for TOF MRA, by using the rationale that failure to diagnose a distal stenosis could negatively impact patient management.

Correlation between TWIST and TOF MRA and TWIST and HR MRA was calculated by using mixed-model regression analysis to account for statistical dependencies among observations provided by each reader for the same patient. Interobserver agreement for each sequence was measured by using the Cohen \( \kappa \) coefficient with linear weights. SAS Version 9.0 software (SAS Institute, Cary, NC) was used for all statistical computations. All reported \( P \) values are 2-sided and were declared statistically significant when \(< .05."

Results
Of 62 extracranial internal carotid arteries evaluated, 4 (6.5%) had 50%–69% stenosis, 1 (1.6%) had 70%–99% stenosis, and 2 (3.2%) were occluded according to the reference standard evaluation. This included 3 carotid dissections, where true lumen diameter was moderately stenotic to occluded (Fig 2). Ulcerated plaque was identified in 2 patients (Fig 3), whereas all other plaque was smooth.

The TWIST technique yielded datasets with distinct arte-
effects, because the proximal ICA is approaching the axial imaging plane in its orientation. MRA (32% and 35%). TOF stenosis overestimation was attributed to in-plane saturation (2, respectively) and was accurately categorized for TWIST MRA (31% and 47%) and HR MRA (32% and 35%). TOF stenosis overestimation was attributed to in-plane saturation effects, because the proximal ICA is approaching the axial imaging plane in its orientation. Note the clear arterial phase image obtained by using the TWIST technique and the presence of venous contamination on the HR image.

Stenosis Evaluation

Overall, accuracy for stenosis assessment was greatest for HR MRA (96.0%), followed by TWIST (90.3%) and TOF MRA (88.7%), with significantly superior accuracy for HR MRA compared with both TWIST and TOF MRA (P < .05). There was no significant difference in accuracy between TWIST and TOF MRA (P = .59). On-line Table 1 summarizes results when data were divided into clinically relevant categories, with the highest accuracy of all individual sequences observed with mild stenosis and occlusion. High correlation between TWIST and TOF MRA (r = 0.78, P < .001) and TWIST and HR MRA (r = 0.82, P < .001) stenosis assessments was observed for all patients, and statistically significant correlation was also observed when analysis was restricted to patients with ≥50% stenosis (TWIST versus TOF MRA, r = 0.70, P < .05; TWIST versus HR MRA, r = 0.73, P < .04).

Severe Stenosis

The single case of severe stenosis was correctly identified with HR MRA and TOF MRA. The less experienced reader underestimated the severe stenosis with the TWIST technique. Severe stenosis was caused by a proximal ICA dissection and an adjacent intramural T1 hyperintense hematoma.

Moderate Stenosis

All sequences had relatively poor accuracy for moderate stenosis. For the TWIST technique, there were 3 incorrect readings. One was classified as incorrect because the stenosis was borderline moderate (69% at consensus interpretation, with the “incorrect” reading of 71% falling into the severe category). The other 2 were the following: 1) an overestimation of stenosis in a patient with remote dissection and recanalization of a complex dysplastic ICA; and 2) a moderate petrous left ICA stenosis missed by the less experienced reader, which was only identified by the second reader on the TWIST sequence and at the consensus reading. The stenosis was not identified by either reader at HR MRA due to venous contamination. For HR MRA, similar accuracy in the moderate stenosis group was demonstrated, with the petrous ICA stenosis described previously not identified by either reader due to adjacent venous signal intensity (SI) and 1 overestimation by 16% of a moderate stenosis by the less experienced reader. TOF MRA performed poorly (accuracy 12.5%) in the moderate stenosis group, with 2 stenoses outside the imaging FOV and an overestimation of stenosis in 1 case in which susceptibility artifact secondary to surgical clips was present.

Mild Stenosis

TWIST led to overestimation of mild stenoses in 5 cases. Cases in which discordant stenosis readings were recorded are detailed in On-line Table 2.

The 3 ICA dissections were identified on TWIST and HR MRA sequences in all cases. TOF MRA depicted the 2 proximal ICA dissections, with 1 dissection not seen because it was in the distal ICA, outside the imaging FOV.

Mean diagnostic confidence was satisfactory or greater for all 3 sequences: 2.61 ± 0.48, 2.29 ± 0.36, and 2.84 ± 0.37 for TOF, TWIST, and HR MRA respectively. Confidence with HR MRA was significantly higher compared with the other 2 sequences (P < .04) and, for TOF, was also significantly higher than that in TWIST (P = .003). SNR and CNR values were highest for HR MRA (which used 14 mL of Gd-DTPA), followed by TOF and then TWIST (6 mL of Gd-DTPA), as shown in On-line Table 3.

TWIST MRA also identified hemodynamic changes, with delayed filling of the carotid bulb in carotid dissection (Fig 2) and delayed opacification of a distal cervical internal carotid dissection following opacification of the circle of Willis (Fig 4), demonstrated with time-resolved MRA. Retrograde filling of a vertebral artery distal to an occlusion at its origin was also incidentally noted in 1 case. These findings were not apparent at HR MRA or TOF MRA.

There was greatest reader concordance for identifying each category of stenosis with the HR sequence (κ coefficient = 0.92), with substantial agreement also calculated for the TWIST and TOF MRA sequences (κ = 0.73 and 0.76, respectively).

Vessel-margin blurring was recorded for 30 of 31 patients for the TWIST sequence. In no case was motion artifact or venous contamination recorded. For TOF MRA, motion artifact (n = 15), wrap in the phase-encoding direction (n = 9), artifact related to flow turbulence (n = 11), and susceptibility artifact secondary to surgical clips (n = 1) were the factors hindering interpretation. For HR MRA, venous contamination was recorded in 16 patients, though diagnostic confidence remained satisfactory to excellent in 15 of these patients. Motion artifact (n = 4) and wrap in the phase-encoding direction (n = 1) also hindered interpretation of HR MRA.

Other pathology included vertebral artery steno-occlusive disease (n = 5) and severe (≥70%) common carotid artery
stenosis (n = 2). Common carotid artery stenosis was detected with high confidence in all sequences in both cases, and similarly, all vertebral artery steno-occlusive disease was detected with HR and TWIST MRA sequences. TOF MRA detected 60% (3/5) of vertebral artery disease, with 1 distal vertebral artery occlusion outside the imaging FOV and 1 proximal vertebral occlusion not detected, with TOF MRA images degraded by motion and wrap artifact in this patient.

**DSA Correlation**

DSA correlation was available in 2 patients. One patient had a complex dysplastic midcervical left ICA, recorded as a dissection causing severe stenosis on all 3 sequences and diagnosed as a remote dissection with recanalization versus a developmentally dysplastic artery at DSA. The second patient had a distal cervical left ICA dissection causing a moderate stenosis, remote bilateral vertebral artery dissections, and an incidental right ophthalmic artery aneurysm diagnosed only at conventional angiography. Both readers identified and accurately graded the ICA dissection on both TWIST and HR MRA sequences (Fig 4).

**Discussion**

Time-resolved MRA of the extracranial carotid arteries has been reported by using a fast low-angle shot 3D gradient-echo sequence with full sampling of k-space.\(^{16,17}\) Time-resolved imaging has also been performed with spatial undersampling by using parallel imaging\(^{9,18,19}\) and temporal undersampling techniques, such as time-resolved imaging of contrast kinetics (TRICKS),\(^8\) or other alternatives\(^{20,21}\) to achieve better temporal resolution.

A time-resolved sequence is beneficial in that it can be useful when there are differential rates of filling between normal and severely diseased cervical vessels or where there is collateral or retrograde flow in the setting of occlusion. The technique has potential value in the setting of carotid stent placement for assessment of altered hemodynamics, when stent-related susceptibility artifact may limit anatomic evaluation, and can also be useful to assess subclavian steal, hypervascular tumors, arteriovenous malformations, arteriovenous fistulas, or extracranial–intracranial bypasses.\(^{22,23}\) Previously, only conventional angiography was available to provide a large FOV dynamic assessment. To our knowledge, this is the first description using the TWIST method of temporal undersam-
sampling in combination with parallel imaging to assess the carotid circulation in symptomatic patients.

TWIST differs in the trajectory of k-space sampling compared with other temporal undersampling techniques described in the literature, such as TRICKS and time-resolved echo-shared angiographic technique (TREAT). Although all use a keyhole imaging approach to sample the center of k-space more frequently compared with high-spatial-frequency information peripherally, with TRICKS, k-space is divided into 1 central and several peripheral sections, with every iteration sampling the central section and 1 of the peripheral sections. TREAT differs in that alternating lines rather than different discrete sections of the periphery of k-space are sampled with each iteration, avoiding jumps between central and peripheral k-space data acquisition. TWIST uses a spiral and pseudostochastic rather than a linear trajectory to traverse a full range of k-space with every iteration, from center to periphery, on the basis of radial distance from the center of k-space (Fig 1), with theoretic advantages in image quality and reduction of artifacts. Although there have been reports in the literature studying different temporal undersampling techniques in this imaging region, no direct comparisons of the techniques are available.

Our study showed that TWIST MRA has good accuracy at 1.5T with high correlation for stenosis estimation compared with established techniques. The transit time from the common carotid artery to the internal jugular vein is approximately 6–10 seconds. Using a combination of spatial undersampling (GRAPPA) and temporal undersampling (TWIST) techniques, we achieved a temporal resolution of 3.9 seconds, allowing clear arterial datasets in every patient without a timing run. Another benefit of rapid acquisition time was that motion artifact, which hindered interpretation of both TOF and HR MRA sequences in our series, was almost completely eliminated. Additionally, near-isotropic datasets were obtained, allowing multiplanar reconstruction of images at consecutive timeframes.

The technique also demonstrated hemodynamic alterations, with delayed filling of the carotid bulb and distal cervical ICA in 2 cases of carotid dissection and retrograde filling of a proximally occluded vertebral artery evident only with time-resolved MRA. In contrast, TOF and conventional HR MRA only captures a single time point. Because subtraction images were available for interpretation, background SI could be eliminated and the small contrast dose used for this sequence still allowed satisfactory diagnostic confidence, which becomes of increasing importance with the advent of three-dimensional and multidetector CT.

The low dose of contrast used for the TWIST sequence made comparison of SNR and CNR between sequences problematic and may also have decreased small-vessel conspicuity in comparison with the higher dose HR MRA. Further, the short duration of the contrast bolus may have contributed to suboptimal vessel conspicuity. A more general limitation was that readers could not be blinded to the sequence they were interpreting and sequences were performed in the same order in all patients. Only a relatively small number of patients in our series had severe disease, with only one 70%–99% stenosis and 2 occlusions, making it difficult to assess the true accuracy of the technique for severe stenosis. A larger prospective study with optimized TWIST parameters would allow further evaluation.

The study was also limited by lack of gold standard correlation, because only 2 of the 31 patients underwent DSA within 3 months of MRA; at our institution, management decisions are often based on sonographic or MRA findings, with was not identified at HR MRA by either reader due to venous contamination. As a result of this missed petrous ICA stenosis and an additional overestimation of TWIST MRA by the less experienced reader, HR MRA had accuracy similar to that of TWIST MRA in the moderate stenosis group. TOF MRA performed poorly in the moderate stenosis group, with factors including its limited FOV and clip susceptibility artifact causing stenosis overestimation in 1 case. The mild stenoses that were overestimated by TWIST MRA were attributed to poor definition of vessel margins, clip artifact, and misclassification of common carotid artery stenosis where plaque was crossing the carotid bifurcation.

It could be argued, however, that a shorter TE than that used in our study might have improved diagnostic accuracy of TOF MRA, because SI loss from turbulent flow may lead to overestimation of stenosis with increasing TE. This likely contributed to the artifact from flow turbulence noted in 11 cases. However, a TE was used in our study that was similar to that reported previously for 3D TOF MRA of the carotid bifurcation. Furthermore, TOF MRA may miss subtle luminal stenosis/plaques and vascular flaps from dissection that may be more apparent on intravenous contrast-enhanced MRA.
DSA reserved for further evaluation or consideration of endovascular management.

It has been reported that there are limitations to the degree of parallel imaging that can be used in combination with temporal undersampling techniques if images are to remain of diagnostic quality at 1.5T. The technique would benefit from imaging at higher field strengths and extracellular contrast agents with higher relaxivity, such as gadobenate dimeglumine. This addition would allow greater SNR in the first instance and greater contrast and vessel conspicuity in the second. Higher parallel imaging factors could be used in conjunction with higher field strengths, where there is increased SI to "spend" on improvements in temporal and/or spatial resolution, as has been described with earlier time-resolved techniques. Specific to carotid imaging, this would again be helpful in differentiating hairline residual patency of the cervical ICA from complete occlusion, with important management implications.

Conclusion

Time-resolved MRA with the TWIST sequence and parallel imaging at 1.5T using a low-contrast dose provides sufficient temporal resolution to assess the anatomy and hemodynamics of the extracranial carotid circulation. It can provide dynamic information noninvasively and rapidly, and delivers consistently pure arterial phase imaging. In its current form, TWIST imaging has potential use as a screening and supplementary tool. It may be particularly useful in the acute stroke setting, where rapid imaging helps limit motion artifact in circumstances where hemodynamic information is essential, such as differentiating critical stenosis from occlusion or when limiting contrast dosage is desirable. However, it does not yet have an accuracy comparable with the HR technique, due to artefacts related to temporal undersampling of k-space. With optimization of k-space sampling parameters, performance at higher field strengths, and use of newer gadolinium contrast agents, further improvements in temporal and spatial resolution are within reach.

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Appendix

SNR was estimated by using the following formula:

$$\text{SNR} = \frac{\text{mean SI vessel region of interest (ROI)}}{\text{SD of homogeneous tissue}}$$

CNR was estimated by using the following formula:

$$\text{CNR with parallel imaging} = \frac{(\text{mean SI vessel ROI} - \text{mean SI homogeneous tissue ROI})}{\text{SD of homogeneous tissue}}$$

Homogeneous tissue (subcutaneous fat) in close proximity to the vessel of interest was used to estimate noise because all 3 sequences used parallel imaging; therefore the use of air to calculate background noise was thought to be less accurate, given spatial variations in SNR with parallel imaging.

In each case, a vessel ROI (terminal common carotid artery just proximal to carotid bifurcation) and subcutaneous fat ROI were obtained for 3 images in each dataset (TOF, arterial phase TWIST, and arterial phase HR MRA) on an imaging workstation (Leonardo, Siemens). SNR and CNR measurements were made, and the average of the 3 SNR and CNR measurements acquired was taken as the final SNR and CNR for each dataset.

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