Simultaneous Multislice for Accelerating Diffusion MRI in Clinical Neuroradiology Protocols

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ABSTRACT

BACKGROUND AND PURPOSE: Diffusion MR imaging sequences essential for clinical neuroradiology imaging protocols may be accelerated with simultaneous multislice acquisitions. We tested whether simultaneous multislice–accelerated diffusion data were clinically equivalent to standard acquisitions.

MATERIALS AND METHODS: In this retrospective study, clinical diffusion sequences obtained before and after implementation of 2-slice simultaneous multislice acceleration and an altered diffusion gradient sampling scheme using the same 3T MRI scanner and 20-channel coil (n = 25 per group) were independently and blindly evaluated by 2 neuroradiologists for perceived quality, artifacts, and overall diagnostic utility. Diffusion tractography was performed in 13 patients both with and without 2-slice simultaneous multislice acceleration (b = 0, 1000, 2000 s/mm²; 60 directions). The corticospinal tract and arcuate fasciculus ipsilateral to the lesion were generated using the same ROIs and then blindly assessed by a neurosurgeon for anatomic fidelity, perceived quality, and impact on surgical management. Tract volumes were compared for spatial overlap.

RESULTS: Two-slice simultaneous multislice diffusion reduced acquisition times from 141 to 45 seconds for routine diffusion and from 7.5 to 5.9 minutes for diffusion tractography using 3T MR imaging. The simultaneous multislice–accelerated diffusion sequence was rated equivalent for diagnostic utility despite reductions to perceived image quality. Simultaneous multislice–accelerated diffusion tractography was rated clinically equivalent. Dice similarity coefficients between routine and simultaneous multislice–generated corticospinal tract and arcuate fasciculus tract volumes were 0.78 (SD, 0.03) and 0.71 (SD, 0.05), respectively.

CONCLUSIONS: Two-slice simultaneous multislice diffusion appeared clinically equivalent for standard acquisitions and diffusion tractography. Simultaneous multislice makes it feasible to acquire higher angular and q-space-resolution diffusion acquisitions required for translating advanced diffusion models into clinical practice.

ABBREVIATIONS: AF = arcuate fasciculus; CST = corticospinal tract; DSC = Dice similarity coefficient; FA = fractional anisotropy; GRAPPA = generalized autocalibrating partially parallel acquisition; SMS = simultaneous multislice.

The diffusion-weighted sequence is essential for clinical protocols in head MR imaging and detecting acute ischemic stroke. Nervous tissue diffusion properties can be informative for other pathologies, including prognosis in lymphoma, detection of skull lesions, or early diagnosis of Creutzfeldt-Jakob disease. Diffusion tractography has improved our understanding of white matter anatomy, like the arcuate fasciculus (AF), the key pathway for language. Diffusion tractography improves neurosurgical planning and reduces perioperative morbidity for patients with brain tumors but requires high-angular-resolution acquisitions that prolong scan time. During the past 15 years, more advanced diffusion models that account for non-Gaussian water diffusion have further improved our understanding of white matter structure but require both high-angular-resolution and high q-space-resolution acquisition strategies; these protocols necessitate relatively long scan times that are much harder for patients to tolerate and thus have not been widely implemented clinically.
Simultaneous multislice (SMS) is a recent innovation that allows concurrently acquiring multiple image slices during each acquisition,13 thereby enabling reduced TR and total scan time. Parallel imaging–based SMS uses controlled aliasing with blipped z-direction gradients14,15 and then reconstructs images individually using a section generalized autocalibrating partially parallel acquisition (GRAPPA) reconstruction16 with little penalty to the image SNR. This technique has become popular in the radiology research community because it supports ever more complex diffusion MR imaging acquisitions for advanced nervous tissue models and representations.17-19 Researchers, however, seldom rely on visual interpretation of SMS-diffusion trace images, nor do they depend on the fidelity of diffusion tractography for high-risk clinical decision-making in individual patients. The goal of this study was to evaluate SMS acceleration for clinical neuroradiology diffusion and tractography applications.

MATERIALS AND METHODS

Initially, volunteers gave consent and were scanned under a prospective, technical development institutional review board (New York University) approval using outpatient 3T MR imaging (Magnetom Skyra; Siemens) with a 20-channel head coil to compare multiple versions of 2-slice SMS-accelerated diffusion sequences to determine an acceptable sequence for common neuroradiology protocols that require trace-weighted and ADC maps, and for a diffusion tractography protocol for patients requiring presurgical planning. The 2-slice SMS-accelerated sequence common to most protocols was then implemented on multiple 1.5T and 3T clinical scanners. Diffusion data for tractography still were obtained with both the original and SMS-accelerated sequences until comparison and approval also could be obtained by the referring neurosurgery service. Data analyzed for this report were collected and analyzed retrospectively from the time period around SMS implementation after institutional review board approval of a Health Insurance Portability and Accountability Act–compliant research study.

Clinical Neuroradiology Head Protocols

Diffusion trace and ADC images from 25 consecutive patients imaged with neuroradiology head protocols using the hospital 3T MR imaging scanner (Magnetom Skyra) with a 20-channel coil were collected from the first day of the month before and after implementation of the 2-slice accelerated SMS diffusion sequence. Consecutive cases were included regardless of reported diagnoses and included inpatients or patients in the emergency department with infarcts, hemorrhage, or brain tumors. The only exclusion criterion was severe motion degradation, but no studies were excluded. There were 2 key changes to the sequence (Online Supplemental Data): First, we used 2-slice SMS acceleration to reduce the TR from 8200 to 3400 ms. Second, the diffusion-sampling scheme was changed by the following: 1) eliminating $b = 500 \text{ s/mm}^2$ sampling, 2) increasing gradient directions from 3 to 6 at $b = 1000 \text{ s/mm}^2$, and 3) reducing averages from 2 to 1 per direction. If we counted $b = 0 \text{ s/mm}^2$ images (2 averages), the total diffusion-weighted images were reduced from 14 to 8 (43% reduction in diffusion volumes acquired). Note that 2-slice SMS TR reductions will always be greater than the resulting total scan time reductions because “dummy” scans are required to reach a steady-state before image acquisition occurs. Overall, these 2 changes reduced scan time from 141 to 45 seconds.

Blinded to the acquisition sequence and alternating side per subject, the senior author used ROIs to extract ADC for the frontal lobe white matter (minor forceps) and frontal horn lateral ventricle CSF (free of the choroid plexus), and to calculate relative SNR for the minor forceps and cerebellar white matter relative to a noise ROI in the same trace-weighted image using previously published methods.20 Next, 2 board-certified neuroradiologists blinded to the acquisition sequence independently rated 3 features of the diffusion-weighted trace and ADC images: artifacts (motion, fat suppression, skull base susceptibility, geometric distortion), image quality (resolution, edge sharpness, noise, gray-white junction), and overall diagnostic utility (ability to interpret). Each rating used the same ordinal scale, where 1 was poor, needs recall; 2 was suboptimal, but interpretable; 3 was average, lower range of normal quality; 4 was good, higher range of normal quality; and 5 was excellent. Sample size per group was determined a priori specifically to detect a 0.5 change in the ordinal rating scale in which the predicted mean $= 4.0 (SD, 0.5), \alpha = .05$, and $\beta \sim 0.95$. ROI and ordinal data were compared by unpaired 2-tailed $t$ tests and Mann-Whitney-Wilcoxon 2-tailed tests, respectively. Confidence intervals and Cohen $\kappa$ coefficients for interrater reliability were calculated.

Diffusion Tractography Protocol

Thirteen consecutive brain-mapping cases obtained for neurosurgical planning had both conventional and SMS acquisitions for DTI; this group included 9 glial tumors, 2 seizure foci, 1 cavernous malformation, and 1 epidermoid cyst (mean age, 44 [SD, 36] years, 5 woman. No potential subjects were excluded. Distance documented in the original radiology report between the lesion margin and the visualized corticospinal tract (CST) and/or AF in the ipsilateral hemisphere was 0 cm ($n = 4$), $<$1 cm ($n = 5$), $<$2.5 cm ($n = 3$), or $\sim$4 cm ($n = 1$). In this latter patient, tractography was requested because the CST was in closer proximity to the planned surgical corridor than the lesion. One patient had a repeat diffusion scan with SMS acceleration 30 days later for a second procedure.

The shared sequence parameters included $54 \times 3 \text{ mm}$ thick slices with 2.8-mm in-plane resolution (200-cm FOV, 88 $\times$ 88 matrix) with identical diffusion parameters (1 average, $b$-values = 0, 1000, 2000 s/mm$^2$, 30 directions per $b$-value, bipolar gradients). Higher $b$-value data are part of our institutional protocol but may not be considered necessary at other institutions. TR and TE were 7100/95 and 4700/95 ms for standard and 2-slice SMS acquisitions, respectively. Shorter TEs (with resultant shorter TRs and overall scan time) are possible with monopolar diffusion gradients but may be more affected by eddy currents. Volumetric T1, diffusion, and SMS diffusion acquisitions were coregistered in third-party brain mapping software (Brainlab). Two ROIs per tract were drawn on the direction-encoded color fractional anisotropy (FA) maps derived from the standard diffusion acquisitions to generate CST and AF tracts ipsilateral to the lesion21 (Online Supplemental Data).

Diffusion data with and without SMS were used separately to generate the tracts while keeping the ROIs constant, thus eliminating manual drawing biases. The minimum tract lengths were 80 and 50 mm for CST and AF, respectively. The lower limit of FA was 0.15 and 0.12 for CST and AF, respectively. Tractography
Table 1: Quantitative and qualitative comparison of diffusion trace and ADC parameter maps for clinical patients in the emergency department [single single 3T MRI scanner]

| Data | Standard (n = 25) | SMS (n = 25) | 95% CI for Difference | P Value |
|------|------------------|-------------|-----------------------|---------|
| Age (yr) | 61.5 [SD, 19.5] | 63.2 [SD, 15.4] | NA | .734<sup>b</sup> |
| Woman (No.) (%) | 52% [13/25] | 56% [14/25] | NA | NA |
| ADC, minor forceps (×10⁻³ mm²/s) | 0.833 [SD, 0.077] | 0.816 [SD, 0.073] | −0.025 to +0.059 | .427<sup>b</sup> |
| ADC, CSF (×10⁻³ mm²/s) | 3.012 [SD, 0.099] | 3.015 [SD, 0.080] | −0.033 to +0.067 | .508<sup>b</sup> |
| SNR, minor forceps | 21.4 [SD, 14.6] | 22.5 [SD, 9.6] | −7.9 to +5.7 | .754<sup>b</sup> |
| SNR, cerebellum | 32.6 [SD, 10.9] | 27.3 [SD, 14.8] | −19.0 to +12.5 | .156<sup>b</sup> |
| Artifacts<sup>c</sup> | 3.5 [SD, 0.5] | 3.7 [SD, 0.5] | −0.5 to +0.1 | .070<sup>d</sup> |
| Image quality<sup>c</sup> | 4.0 [SD, 0.7] | 3.3 [SD, 0.5] | −0.4 to +1.0 | .001<sup>d</sup> |
| Diagnostic utility<sup>c</sup> | 3.8 [SD, 0.4] | 3.7 [SD, 0.5] | −0.2 to +0.4 | .187<sup>d</sup> |

Note: NA indicates not applicable

<sup>a</sup>Data are mean [SD]. (No.) (%) or 95% CI.

<sup>b</sup>Unpaired, 2-tailed t test.

<sup>c</sup>Average of independent, blinded assessment using an ordinal scale 1–5 by 2 board-certified neuroradiologists (no adjudication performed).

<sup>d</sup>Mann-Whitney-Wilcoxon 2-tailed test.

FIG 1. Axial diffusion images from standard [A] and SMS diffusion acquisitions [B] in a clinical patient with headaches. There is no appreciable difference in diagnostic utility, even with a reduction in scan time using SMS. An SMS arc-like scalp artifact (C, arrow) is seen in another clinical patient due to poor fat saturation. With SMS, unsaturated fat signal associated with echo-planar acquisitions can alias into all simultaneously acquired slices.

RESULTS

Clinical Diffusion Trace

At 3T, 2-slice SMS acceleration with an altered diffusion-sampling scheme allowed us to reduce the TR by 59% (from 8.2 to 3.4 seconds), reduce diffusion volumes acquired 43% (from 14 to 8), and reduce overall scan time by 70% (from 141 to 45 seconds). There was an 18% decrease in perceived image quality of the SMS-accelerated scans ($P = .001$). No difference in artifacts or diagnostic utility was detected with 95% confidence interval that true differences, if present, represented $<0.5$ decrement on the 5-point ordinal scale. Representative clinical cases are provided in Figs 1 and 2 for readers to compare sequence results. Raters perceived subtle, lower SNR in the posterior fossa on SMS images (16% absolute quantitative reduction, $P = .156$, with a 95% confidence interval to detect an SNR reduction of $\approx 38\%$), but this remained acceptable for diagnostic use in ratings and subsequent clinical cases (Fig 2). Both readers noted a fat-saturation artifact in 1 subject undergoing SMS (Fig 1C); however, the diagnostic utility of this scan was not compromised. Raters did not observe increased T1-weighting to the SMS diffusion trace images; the TR of 3400 ms remained estimated a priori to require approximately 16 to detect a quantitative difference roughly equal to the anticipated variance of the data (ie, a Cohen D value of 1) with 95% statistical power. Confidence intervals also were calculated. The Dice similarity coefficient (DSC) was used to measure the spatial overlap between standard (A) and SMS (B) tract volumes, where $DSC(A, B) = 2(A \cap B) / (A + B)$ and $\cap$ is the intersection between the 2 volumes. Spatial misregistration from skull base geometric distortion was characterized by measuring the distance from the anterior margin of the conventional and SMS-derived CST tracts to the anterior margin of the cervicomedullary junction on coregistered volumetric T1 at the foramen magnum.

seed density and turning angle limits have fixed values in the clinical software used. While normally part of our clinical practice, here potential spurious tracts were not edited, nor were FA thresholds adjusted for individual cases, to avoid introducing individual bias into the comparisons.

The tract volumes were superimposed on the T1-weighted, volumetric sequences and then retrospectively reviewed in random order separately using Brainlab software by a neurosurgeon with 5 years of experience using tractography for surgical planning (blinded for all assessments). The neurosurgeon evaluated anatomic correctness and preference and whether the tract would alter surgical management. “Anatomically correct” implied that the tract matched known anatomy from previous surgical experience and reports of postmortem white matter dissection. “Different surgical management” implied that a tract changed the surgical approach or planned extent of resection. Finally, in blinded side-by-side comparison, the neurosurgeon determined whether he preferred 1 version of the tract.

The AF and CST length, FA, and volumes were extracted from the tract objects for standard and SMS diffusion tractography and then compared using paired-sample 2-tailed t tests. Note that sample size was determined by practical considerations but was
the reported T1-relaxation times for gray and white matter at 3T.\textsuperscript{24} There was no difference in the ADC values of the minor forceps or frontal horn CSF of standard compared with SMS diffusion images (95% confidence to exclude no greater than a 7% difference). There were no callbacks or nondiagnostic diffusion scans attributed to SMS during the subsequent 10 months after widespread adoption.

Rater agreement for artifacts, image quality, and diagnostic utility were 60%, 62%, and 76%, respectively. The Cohen \( \kappa \) values for interrater reliability were 0.216, 0.363, and 0.413, respectively; these may underestimate agreement due to the first kappa paradox\textsuperscript{25} because ratings of 3 or 4 represented most scores. Reader agreement for artifacts, image quality, and diagnostic utility became 100%, 88%, and 96% respectively if 3 and 4 ordinal scores from raters were collapsed to 1 value. While it was not part of the rater assessment, a similar 2-slice SMS-accelerated diffusion sequence was implemented on a newer 1.5T MRI scanner with a 20-channel coil (Magnetom Aera; Siemens) at the same hospital, reducing scan time from 165 to 65 ms (Online Supplemental Data); for reader interest, additional clinical examples of SMS-accelerated diffusion images at both field strengths are provided in the Online Supplemental Data. Note that full protocol parameters for 1.5T and 3T MR imaging scanners will be provided to interested readers by contacting the communicating author.

**Diffusion Tractography**

2-slice SMS acceleration reduced the TR 34% to 4.7 seconds and reduced overall acquisition time 22% to 5.9 minutes without visual differences in tractography (Fig 3) or quantitative tractography differences for either the CST or AF ipsilateral to clinical lesions (Table 2). The mean DSCs between standard and SMS tract volumes were 0.78 (SD, 0.03) and 0.71 (SD, 0.05) for the CST and AF tracts, respectively ($n=13$ subjects). The SMS-acquired CST demonstrated similar spatial misregistration at the skull base (Table 2). The Online Supplemental Data demonstrate reproducibility for CST and AF tracts in the same subject scanned twice 30 days apart.

The neurosurgeon rated 25 of 26 (96%) SMS-generated tracts to be anatomically correct. The exception was the AF in case 11; however, both the conventional and SMS tracts were deemed equally nonanatomic in this case. This outcome was attributed to a porencephalic cavity, which made the standardized 2-ROI tractography approach ineffective and not from any inherent deficiency of the SMS acquisition. Tractography from SMS diffusion was considered equivalent or

**FIG 3.** Overlap of diffusion tractography between standard and SMS-acquired data in 2 selected patients. **Upper Row:** A 56-year-old man with a multifocal right-cerebral hemisphere glioma. The right corticospinal tract volumes are shown in blue (conventional) and red (SMS) at the levels of the cerebral peduncle (A), posterior limb of the internal capsule (B), and precentral gyrus (C). The DSC between standard and SMS tract volumes is 0.76. **Lower Row:** A 36-year-old man with a left opercular cavernous malformation. The left-arcuate fasciculus volumes are shown in purple (conventional) and yellow (SMS) at the levels of the anterior and posterior frontal projections (D and E) and genu (F). There are slight differences in the edge of the visualized tracts abutting the superior border of the cavernous malformation (F), but these differences would not affect selection of the surgical corridor, and in both situations, the neurosurgeon would be cautious in approaching the superior margin of the lesion. The DSC between standard and SMS tract volumes is 0.75.

3× the reported T1-relaxation times for gray and white matter at 3T.\textsuperscript{24} There was no difference in the ADC values of the minor forceps or frontal horn CSF of standard compared with SMS diffusion images (95% confidence to exclude no greater than a 7% difference). There were no callbacks or nondiagnostic diffusion scans attributed to SMS during the subsequent 10 months after widespread adoption.

**FIG 2.** Examples of 2-slice SMS-accelerated diffusion trace and ADC maps of 4 different clinical patients. Large acute left-occipital infarct (A) contrasted with a small subacute left-thalamic infarct (B) (arrows). Raters suggested subtle SNR reductions for the posterior fossa using SMS (Table 1); however, small infarcts in the brainstem (C) and cerebellum (D) remain well-visualized.
better in 18 of 26 (69%) tracts generated. Reasons for preferring one tract over its counterpart included the following: more robust tract, less spurious fibers, less discontinuities in volume, and less spurious extension into the extracranial soft tissues (removing the latter is normally routine, but not in this investigation to avoid introducing bias). All CST pairs did not change the surgical management (approach and aggressiveness). All AF pairs had no change in surgical approach. Eleven of 13 (85%) AF pairs would have equivalent aggressiveness of resection. The standard AF tract in case 13 was considered a better anatomic representation of projections of the left-frontal operculum adjacent to a left-frontal oligodendroglioma that would result in a less aggressive resection. The SMS AF tract in case 5 was considered a more robust tract adjacent to the cavernous malformation, which would result in a less aggressive resection.

**DISCUSSION**

SMS-accelerated neuroradiology protocol diffusion acquisitions appear safe and clinically feasible without large differences in image quality when assessed by 2 neuroradiologists. Raters perceived mild decreases in the SNR of diffusion trace images, particularly in the posterior fossa structures, but characterization of pathologies of different sizes and in different anatomic locations appeared similar between standard and SMS-accelerated diffusion. Aggressive 2-mm isotropic resolution of clinical diffusion MR imaging protocols; this potential was not emphasized in the current study but could be the subject of future work. SMS-accelerated diffusion images for the current study still had TRs (3200 or 4700 ms) more than 3× longer than our current data. Given that diffusion imaging is ubiquitous to all neuroradiology protocols, the 1.5-minute time savings obtained here would return sufficient time theoretically for an additional 3 scans per day per magnet at our primary outpatient facility (estimation based on ~67.5 minutes saved; the facility is open 15 hours per day, with 20-minute slots and 100% neuroradiology head protocols). A motion-free 45-second diffusion scan should facilitate more rapid decision-making in acute stroke, decrease motion degradation in patients who are less MR imaging-tolerant, and create more time for additional MR imaging sequences. An additional motivation for our center was to create more time within the scheduled slots to clean the magnet between patients during the early stages of the coronavirus 19 pandemic.

SMS-accelerated diffusion tractography was considered anatomically equivalent by an experienced neurosurgeon without altering intended surgical management or the aggressiveness of resection. The tractogram volume agreement was good: mean DSCs of 0.78 and 0.71 for CST and AF, respectively. An imperfect DSC may occur due to low spatial resolution (~3-mm isotropic voxels), SNR differences, patient movement (or other causes of misregistration), and/or different spurious tracts not actively edited from the data. In our clinical practice, neuroradiologists freely and subjectively manipulate the tractography parameters more on an individual basis than permitted in this study design to accomplish the goals of presurgical planning. Yokota et al reported that CST tractography appeared similar to radiologists with or without 2-slice SMS acceleration, but most of the data were from nonsurgical patients and not blindly evaluated by a neurosurgeon experienced using tractography. Here, SMS acquisitions in preoperative cases could reduce tractography scan times by ~2 minutes; these studies tend to be long due to the need for coregistered fMRI, tractography, and 3D structural MR imaging. Even more time savings might be possible with 3- or 4-fold slice acceleration using array coils with more receive elements (eg, a 64-channel coil), less angular or q-space resolution than in our current institutional protocol, or improved monopolar imaging gradients that reduce TE (and subsequently scan time).

Besides increased throughput on busy clinical MR imaging scanners, SMS acceleration can also be used to increase spatial resolution of clinical diffusion MR imaging protocols; this potential was not emphasized in the current study but could be the subject of future work. SMS-accelerated diffusion images for the current study still had TRs (3200 or 4700 ms) more than 3× the T1-relaxation times of gray and white matter at 3T. More aggressive reductions of TR are possible, particularly with 3- or 4-slice acceleration, which may introduce T1-weighting to diffusion trace images. In preliminary experiments, shortening the TR below 2500 ms also introduced more substantial signal loss and increased the frequency of peripheral nerve stimulation reported by volunteer subjects. With SMS, unsaturated fat signal associated with echo-planar acquisitions can alias into all simultaneously acquired slices and result in spurious tractography. While SMS diffusion is now FDA-approved, readers

| Table 2: Quantitative comparison of DTI tractography for the corticospinal tract and arcuate fasciculus ipsilateral to the lesion in 13 subjects* |
| --- | --- | --- | --- | --- |
| Diffusion Technique | Standard | SMS | Mean Individual Differenceb | 95% CI for Difference | P Value |
| **Corticospinal tract** | | | | | |
| Volume (mL) | 30.7 [SD, 9.1] | 32.9 [SD, 9.7] | −2.3 [SD, 4.0] | −9.4 to +5.0 | .062c |
| Length (mm) | 125.2 [SD, 13.1] | 124.0 [SD, 15.0] | 1.2 [SD, 7.0] | −9.6 to +12.0 | .537c |
| Mean FA | 0.45 [SD, 0.04] | 0.46 [SD, 0.04] | −0.01 [SD, 0.02] | −0.04 to +0.02 | .076c |
| **Arcuate fasciculus** | | | | | |
| Volume (mL) | 25.3 [SD, 5.3] | 26.0 [SD, 6.7] | −0.7 [SD, 3.1] | −5.3 to +3.9 | .438c |
| Length (mm) | 77.4 [SD, 7.1] | 76.5 [SD, 6.7] | 0.9 [SD, 4.6] | −4.4 to +6.2 | .482c |
| Mean FA | 0.36 [SD, 0.04] | 0.36 [SD, 0.04] | 0.00 [SD, 0.01] | −0.03 to +0.03 | 1.000c |
| Spatial misregistration (mm)d | 3.4 [SD, 1.6] | 2.8 [SD, 1.0] | 0.6 [SD, 1.3] | −0.4 to +1.6 | .335c |

*Data are mean (SD) or 95% CI.

bDifference between measurements in the same individual (standard value minus SMS value).

cPaired-sample 2-tailed t test.

dDifference between the anterior margin of the cervicomedullary junction at the foramen magnum depicted by volumetric T1 MR imaging versus CST diffusion tractography.
should carefully adapt SMS acquisitions with or without acceleration to their local diffusion protocols.

This study has limitations. Due to practical limitations on the time required for detailed assessment by radiologists and a neurosurgeon, the study is underpowered to exclude smaller quantitative differences between sequences (see confidence intervals in Tables 1 and 2). The SMS-accelerated diffusion sequences used here are derived from a single large academic center (New York University) but may differ from other radiology practice locations and settings. Scan time reductions were not just from 2-slice SMS acceleration but also the elimination of the b=0 500 s/mm² images. The external validity of our results is supported by another report of SMS-accelerated diffusion equivalence, albeit with less time savings, using a different sequence (2-mm isotropic resolution, b=0 and 1000 s/mm² with 10 directions, a final time of 128 seconds, which was 41% faster than the previous protocol). Hence changes in time savings, perceived image quality, artifacts, and diagnostic utility will vary depending on the local accepted protocol and how it is modified. Another limitation was that tractography results were reviewed retrospectively so that the position of the SMS-derived tracts could not be correlated with operative findings (the standard diffusion data were instead used as an index standard). Intraoperative verification of tractography with or without SMS acceleration in patients requiring surgery is not trivial to obtain, but it could be the subject of a future study. There are additional methods for accelerating diffusion acquisitions that are being translated into clinical practice, including compressed sensing and image denoising or without deep learning. These techniques are not readily available for clinical implementation, but direct comparison of these diffusion-acceleration techniques and/or their combination also could be the subject of future study.

The characterization of nervous tissue microstructure with advanced diffusion models in clinical patients has been hampered by the time-intensive high-angular and q-space acquisition requirements that SMS acceleration helps overcome. For instance, as part of an ongoing National Institutes of Health–funded study, we have used SMS to acquire high-spatial and -angular resolution, multi-b-shell diffusion data from 137 patients with multiple sclerosis. We also have used SMS-accelerated acquisitions to generate probabilistic tractography–derived contrast of internal brainstem anatomy. SMS diffusion can complement other means of accelerating clinical neuroradiology MR imaging protocols like compressed sensing, synthetic MR imaging, or MR fingerprinting that have, thus far, been more limited at producing clinically acceptable accelerations or substitutions for conventional diffusion-weighted sequences.

CONCLUSIONS

Using standard 3T MR imaging hardware and coils, 2-slice SMS acceleration can reduce routine diffusion MR imaging and diffusion tractography acquisition times in clinical patients without compromising image quality, clinical interpretation, or surgical planning. SMS acceleration also should increase the feasibility of translating high spatial and angular resolution diffusion acquisitions to clinical patients.

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