ABSTRACT

BACKGROUND AND PURPOSE: The number of centers currently performing 3D fat-suppressed isotropic imaging is limited. If the angular orientations of the major lumbosacral plexus nerves on 3D isotropic MR neurography could be determined, similar planes could be prescribed during acquisition of 2D or 3D nonisotropic techniques for optimal depiction of various nerves. Our aim was to determine oblique sagittal and coronal angular measurements for longitudinal depiction of lumbosacral plexus nerves. Interobserver and intraobserver performance and mean calibers of sciatic and femoral nerves were also determined.

MATERIALS AND METHODS: A consecutive series of lumbosacral plexus MR neurography examinations with 3D nerve-selective imaging performed during a 10-month period on a 3T scanner were evaluated. Two observers performed reconstructions and angular measurements. Sciatic and femoral nerve diameters were measured. Descriptive statistics and intraclass correlation coefficient correlations were used.

RESULTS: There were 52 subjects, 11 men and 41 women. Mean sagittal thecal sac angles for coronal demonstration of lumbosacral plexus nerve roots from L1 to S1 for 2 independent observers measured 13.58° ± 2.87° and 13.61° ± 2.18°. Mean sagittal femoral nerve angles were 27.78° ± 4.81° and 28.94° ± 4.49°, and mean sagittal sciatic nerve angles were −10.7° ± 3.75° and −11.82° ± 2.87°. Coronal angular measurements of the femoral and sciatic nerves were similar. The intraclass correlation coefficient was moderate (0.582–0.671) for interobserver performance. For intraobserver performance among various angular measurements, the intraclass correlation coefficient was moderate to good (0.586–0.788). Femoral nerve caliber on MR imaging was almost half that of the sciatic nerve. Mean right femoral nerve thickness was 4.52 ± 1.11 mm and 4.85 ± 0.64 mm for the 2 observers, and mean left femoral nerve thickness was 4.48 ± 0.97 mm and 4.94 ± 0.57 mm. Mean right sciatic nerve thickness was 9.71 ± 1.76 mm and 9.94 ± 0.83 mm, and mean left sciatic nerve thickness was 10.03 ± 1.71 mm and 9.98 ± 0.99 mm.

CONCLUSIONS: Angular lumbosacral plexus measurements aid in the prescription of different planes on MR imaging for the optimal longitudinal demonstration of nerves.

T he lumbosacral (LS) plexus is a complex network of nerves, which provides both motor and sensory innervation to most structures of the pelvis and lower extremities. Most anatomy illustrations in the text books and Internet Web sources depict coronal views of the LS plexus and its branch nerves to demonstrate their longitudinal extent. Some surgeons and physicians prefer to visualize the LS plexus in the coronal plane, and this plane may be beneficial for patient consultations. MR imaging provides the best soft-tissue contrast for the evaluation of deep soft-tissue structures, including the nerves.1 With increasing frequency, MR neurography is playing a prominent role in the diagnosis, characterization, localization, and determination of the extent of pathology in patients with symptoms of lumbar plexopathy.2,3 Due to the complex anatomy and oblique course of the plexus branches, however, the longitudinal extent of pathology is difficult to determine on direct coronal or sagittal MR images. 3D isotropic MR images can be reconstructed in various arbitrary planes to depict the peripheral nerves in their entirety.4 However, the number of centers currently performing 3D fat-suppressed isotropic imaging is limited due to various hardware and software limitations. If the angular orientations of the major LS plexus nerves
on 3D isotropic MR neurography could be determined, similar planes could be prescribed during acquisition of 2D or 3D nonisotropic techniques for optimal depiction of various nerves, depending on the clinical suspicion of different neuropathies. This added oblique imaging plane/sequence would help the radiologists and referring physicians precisely assess the location and extent of the pathology while facilitating multidisciplinary communication.

The primary aim of the study was to determine oblique sagittal and coronal angular measurements for the optimal longitudinal depiction of lumbosacral plexus nerve roots and sciatic and femoral nerves from their origin to the most distal extent without a break in continuity. As secondary aims, interobserver and intraobserver performance was assessed and the average calibers of sciatic and femoral nerves were measured to generate normative data.

**MATERIALS AND METHODS**

Informed consent for this retrospective evaluation was waived, and anonymized data were evaluated. A consecutive series of LS plexus MR neurography examinations for pelvic pain performed during a 10-month period (November 2013 to August 2014) on a 3T scanner (Achieva; Philips Healthcare, Best, the Netherlands) were evaluated. The inclusion criteria were consecutive examinations with no distortion of images by metal, motion, or poor fat suppression. The exclusion criteria were incomplete imaging or evidence of any major nerve pathology, which could cause abnormal thickening in the evaluated nerves. All the MR imaging examinations were performed on nerve-SHeath signal increased with INKed rest-tissue rarE Imaging (SHINKEI; Philips Healthcare) acquisitions, which were obtained as part of the LS plexus MR neurography examination. This sequence uses adiabatic inversion recovery fat suppression for uniform fat saturation and a motion-sensitive driven equilibrium pulse for vascular signal suppression for selective demonstration of the LS plexus nerves. The SENSE XL Torso coil (Philips Healthcare) combined with spine elements was used for imaging. The parameters of the sequence included the following: TR, 2000 ms; TE, 78 ms; voxels, 1.5 mm isotropic; acquisition time, 7–8 minutes; fat suppression, spectral adiabatic inversion recovery. The source data were manipulated on independent Aquarius iNtuition software (TeraRecon, San Mateo, California). Two observers (E.B. and R.J., second- and third-year radiology residents) performed the reconstructions and measurements independently, following training on an initial set of 10 cases. Sagittal and coronal oblique 20-mm-thick-slab maximum-intensity reconstructions were performed, which allowed maximum longitudinal visualization of the LS plexus nerve roots and sciatic and femoral nerves from their origin to termination without a break in continuity (i.e., the distal-to-inguinal ligament in the case of femoral nerves). Thinner slab MIP reconstructions did not show the maximum longitu-
dinal course of the nerves, while thicker slab MIP reconstructions resulted in blurring of the nerves due to overlapping of adjacent structures. In our initial testing, 20-mm-thick slabs provided the optimal longitudinal depiction of the nerves; MIP parameters would not be expected to change substantially in patients with conditions such as chronic inflammatory demyelinating polyneuropathy, or neurofibromatosis, though parameters could be optimized for each patient.

The sagittal plane measurements for the angle ventral to the thecal sac were denoted by positive numbers and dorsal to the thecal sac, by negative numbers. Femoral and sciatic nerve calibers were measured in their midportions in the abdomen and pelvis, respectively, at the site of optimal nerve visualization and maximum thickness (ie, immediately below the piriformis muscle and sciatic notch for the sciatic nerve and adjacent to the iliopsoas crotch for the femoral nerve). Descriptive statistics were performed, and data were expressed as mean ± SD. Interobserver variance was evaluated by using the intraclass correlation coefficient (ICC), with values of ≥0.65 considered good; <0.65 to ≥0.50, moderate; <0.50 to ≥0.40, fair; and <0.40, poor correlations. Intraobserver performance was also assessed by using the ICC for the reader E.B., who obtained repeat measurements approximately 25 weeks following the initial measurements.

RESULTS

There were 62 examinations with full abdomen and pelvis MR neurography imaging, though 10 were excluded due to hardware artifacts (n = 3) and motion and suboptimal fat suppression and incomplete imaging (n = 7). Final samples included 52 subjects, 11 men (mean age, 45.73 ± 18.4 years) and 41 women (mean age, 48.85 ± 12.65 years). All reconstructions were successfully obtained in <7 minutes.

Mean sagittal thecal sac angles for the coronal demonstration of LS plexus nerve roots from L1 to S1 for the 2 independent observers measured 13.58° ± 2.87° and 13.61° ± 2.18° (Fig 1). For complete longitudinal nerve course depiction bilaterally in the coronal plane, the mean sagittal angular measurements of the femoral nerve were 27.78° ± 4.81° and 28.94° ± 4.49° (Fig 2). Similarly, mean sagittal sciatic nerve angle measurements for coronal depiction were 9.6° ± 3.75° and 11.82° ± 2.87° (Fig 3).

The mean coronal angular measurements for the sagittal long axis demonstration of the right femoral nerve were 33.54° ± 4.55° and 33.87° ± 3.57°; and for the left femoral nerve, 32.85° ± 4.24° and 34.29° ± 3.75°, respectively (Fig 4). The mean coronal angular measurements for the sagittal long axis demonstration of the right sciatic nerve were 33.01° ± 4.41° and 34.53° ± 3.8° and, for the left sciatic nerve, 32.75° ± 3.72° and 34.37° ± 3.5°, respectively (Fig 5). The interobserver ICC was moderate (0.582–0.671) for various angular measurements. For intraobserver performance among various angular measurements, the ICC was moderate to good (0.586–0.788).

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the sciatic nerve. Mean right femoral nerve thickness was 4.52 ± 1.11 mm and 4.85 ± 0.64 mm for the 2 observers, and mean left femoral nerve thickness was 4.48 ± 0.97 mm and 4.94 ± 0.57 mm, respectively. Mean right sciatic nerve thickness was 9.71 ± 1.76 mm and 9.94 ± 0.83 mm, and mean left sciatic nerve thickness was 10.03 ± 1.71 mm and 9.98 ± 0.99 mm, respectively (Fig 6). The interobserver ICC was moderate to good (0.557– 0.737) for the above thicknesses except for the right femoral nerve thickness, which was poor (0.343). Intraobserver performance for nerve thickness was moderate to good (0.646 – 0.756), except for the left sciatic nerve thickness, which was poor (0.392) (Table).

**DISCUSSION**

3D SHINKEI allows excellent vascular and fat signal suppression, leading to selective nerve depiction.\(^6\)\(^,\)\(^7\) This sequence uses adiabatic inversion recovery for uniform fat suppression and a motion-sensitive driven equilibrium pulse for vascular signal suppression, thereby enabling selective demonstration of the LS plexus nerves.\(^5\) The variable turbo spin-echo component limits the acquisition time to 7–8 minutes.

This imaging technique allows ready depiction of the nerves in various arbitrary planes. Thus, the angular prescriptions could be easily and confidently measured\(^8\),\(^9\); these results are reflected in nearly good interobserver and intraobserver performance in angular measurements.

The femoral nerve calibers were ~4–5 mm, the sciatic nerve calibers were ~8–11 mm, and the interobserver performance was moderate to good except for the right femoral nerve, due to the relatively smaller size of the femoral nerve. The normative data can be used for identification of pathologies that frequently cause abnormal nerve thickening, such as chronic demyelinating polyneuropathy, diabetic polyneuropathy, perineurioma, or Charcot-Marie-Tooth disease and so forth.\(^10\)\(^–\)\(^12\) The nerve-caliber measurements are also reproducible, as shown by moderate-to-good ICC results for intraobserver performance, except for the left sciatic nerve caliber, which was poor (0.392).

Although this is the first study depicting angular and nerve-caliber measurements, there are some limitations. We did not obtain surgical or cadaveric correlation because it would not be practical to obtain such correlations for a large sample. In addi-
tion, no correlations with diffusion imaging were obtained, which is also shown to produce nerve-selective images. The observers felt confident in nerve identification on 3D SHINKEI during the evaluation of the initial training set due to excellent vascular signal suppression and nerve depiction by this technique. It was therefore decided that diffusion imaging evaluation can be avoided and left as the subject of another article. Another limitation is that we evaluated only patients with normal nerves. Patients with significant nerve pathology/known mass lesions were excluded from our patient population. Patients with significant nerve pathology may require different angles to optimize the longitudinal display of the nerves, due to distortion of nerve course. Nerve caliber in these patients would also be expected to differ from that of patients with normal nerves. Finally, we did not measure obturator nerve angles. Because this nerve has been shown to travel in a straight coronal plane, it was thought that further study would not yield a significant angle. 

CONCLUSIONS
The study fills a gap in the literature regarding angular LS plexus nerve measurements for their maximum length depiction and nerve thickness. It will aid in prescription of predictable nerve planes on MR imaging for their optimal longitudinal demonstration.

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