High-Resolution Diffusion-Weighted MR imaging of the Human Lumbosacral Plexus and Its Branches Based on a Steady-State Free Precession Imaging Technique at 3T

SUMMARY: 3D diffusion-weighted steady-state free precession imaging (3D DW-SSFP) with isotropic resolution was performed to delineate structures of the human lumbosacral plexus (LSP). 3D DW-SSFP clearly revealed detailed anatomy of the LSP and its branches. Our data suggest that the sequence based on 3D DW-SSFP can be used for high-resolution MR imaging of the peripheral nervous system.

MR imaging evaluation of the normal peripheral nerve anatomy and diseases is mainly dependent on 2D MR imaging techniques, including T1-weighted spin-echo, T2-weighted fast spin-echo, and inversion-recovery sequences with fat saturation. Although these techniques can produce excellent-quality images, they have limitations in describing deliberate orientations of the targeted nerves, in that section gaps in these techniques lead to lower anatomic coverage and less quantitative information. Furthermore, T2-weighted spin-echo techniques cannot image smaller nerves in the periphery because the nerves cannot be distinguished from blood vessels on T2-weighted spin-echo images. A combination of diffusion-weighted imaging (DWI) with fat-suppressed T2-weighted sequences has been proposed to overcome these technical limitations. Additionally, DWI based on spin-echo with an echo-planar readout has been used to evaluate the anatomy of the peripheral nervous system. However, these conventional techniques have limited spatial resolution or low signal-to-noise ratio and often produce severe image distortion.

It is difficult to reveal the detailed anatomy of the lumbosacral plexus (LSP) by using conventional MR imaging in that many important branches of the LSP are small, and contrast of the various anatomic structures is not sufficient to exhibit their discrete identification. The purpose of this study is to describe a high-spatial-resolution 3D diffusion-weighted steady-state free precession (3D DW-SSFP) sequence and prospectively evaluate its feasibility in human LSP imaging at 3T.

Materials and Methods
Twenty-four healthy subjects (10 men, 14 women; mean age, 42.2 years; range, 24–69 years) underwent high-spatial-resolution T2-weighted 3D DW-SSFP imaging with a generalized autocalibrating partially parallel acquisition (GRAPPA) technique on a clinical 3T MR imager (Trio; Siemens Medical Solutions, Erlangen, Germany) with 2-element circularly polarized (CP) phased-array body coils combined with CP phased-array spinal coils.

After optimization of the sequence in 3 healthy subjects (1 woman and 2 men, 30–36 years of age), we used the following parameters: TR/TE, 14.16/4.61 ms; flip angle, 30°; FOV, 360 mm; matrix, 512 × 512; bandwidth, 184 Hz per pixel; fat suppression, water excitation technique; diffusion moments, 90 mT/m (*)msec; 3D slab thickness, 256 mm; voxel dimension, 0.8 × 0.8 × 0.8 mm (320 partitions); number of acquisitions, 1; acquisition time, 10 minutes 49 seconds. The diagram of the 3D DW-SSFP sequence used in this study is shown in Fig 1.

The postprocessing techniques, including maximum intensity projection and multiplanar reformation, were used to generate images of the LSP with the commercial software on the console of the MR imaging scanner. The datasets from 3D DW-SSFP sequences in all subjects were reconstructed in arbitrary planes oriented in relation to the course of the LSP and its branches. All images were analyzed by 2 experienced neuroradiologists. Each radiologist made initial evalu-

Fig 1. The 3D DW-SSFP sequence diagram used in the study. Diffusion gradients in the section direction were applied to reduce the signal intensity of fluid along the axis, and flow-compensation gradients in the read direction were applied to reduce motion sensitivity along this axis. DG indicates diffusion gradient; ACQ, acquisition; RF, radio-frequency; Gs, section-selection gradient; Gp, phase-encoding gradient; Gr, read-out gradient.
ations independently, and any disagreements regarding final conclusions were resolved by consensus. The certainty of identifying the LSP and main branches was individually determined in each patient by using a 3-score scale system (2, identified with certainty; 1, most probable identification; 0, not identified).

The study protocol was approved by the local ethics committee and the review board of our department, and informed consent was obtained from all subjects before MR imaging examination.

Results
In arbitrary planes, the nerve roots from L1 to S4, the spinal nerve ganglion, and the proximal branches of the LSP were identified (score of 1 or 2) in 48 (100%); the main branches of the lumbar plexus including the obturator nerves and the femoral and lateral femoral cutaneous nerves within the psoas major muscle were identified (score of 1 or 2) in 22 (100%), 2 (8%), and 16 (67%), respectively. The main branches of the sacral plexus including the sciatic nerves, the posterior femoral cutaneous nerve, and the gluteal nerves were identified (score of 1 or 2) in 24 (100%), 20 (83%), and 20 (83%), respectively. Typical imaging results with the 3D DW-SSFP sequence are shown in Figs 2–4.

Discussion
The 3D DW-SSFP sequence in this study is a high-resolution diffusion-weighted T2-weighted method of steady-state gradient-echo imaging combined with fat-suppression and flow-compensation techniques. Our sequence has several advantages. First, the signal intensity of the sequence was formed by a spin-echo mechanism with characteristics of a spin-echo sequence and thus with dominating T2 contrast as compared with other spoiled or refocused gradient-echo techniques because the echo of the sequence is formed by a radio-frequency excitation that is unusual for gradient-echo sequences. Therefore, the influence of the local magnetic field in homogeneities can be reduced on the spin relaxation.¹²
Second, the detection of small nerve structures, especially those located near large vessels, is largely improved in that the sequence can completely eliminate the signal intensity from the vessels. The SSFP signal intensity heavily depends on the steady-state condition, and vessels do not produce signal intensity because it cannot construct the steady-state condition.\textsuperscript{13} In addition, the image contrast is further improved by using the DWI technique combined with the SSFP sequence, which completely suppresses the high signal intensity of the stationary fluid, such as urine and synovia. As a result, the signal intensity of stationary fluid is very low in contrast to the high signal intensity of nerves and the intermediate signal intensity of adjacent pelvic parenchymal structures on DWI images. Furthermore, the fat signal intensity is completely suppressed, and the chemical shift effect and motion-related phase errors are reduced by using a spectral spatial water-excitation pulse. With this technique, only water is excited by using section-selective composite pulses, whereas lipid spins are left in equilibrium, thereby producing no signal intensity.\textsuperscript{14}

Besides, this sequence is compatible with the GRAPPA technique at 3T. The combination of this sequence with GRAPPA allows performance of high-resolution imaging within a reasonable acquisition time, and image disturbances due to B0 field inhomogeneity in a high-field MR imager can be further decreased by the technique. Given that the thin contiguous sections and high isotropic resolution can depict tiny structures or lesions, any images can be reconstructed in arbitrary orientation by the multiplanar reconstructive technique without image degradation.

The limitations of this study include the motion, pulsation, and ghost artifacts in the lower abdomen. These artifacts can degrade or even disturb the quality of the images, especially the quality of images on the nerves close to the abdominal wall.

**Conclusion**

In the present study, 3D DW-SSFP clearly revealed detailed anatomy of the LSP and its branches. These data suggest that the sequence based on 3D DW-SSFP can be used for high-resolution imaging of the peripheral nervous system.

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