Dear Reader,

I am pleased to edit this exciting MAGNETOM Flash issue on Magnetic Resonance Neurography (MRN). Incorporation in the RSNA edition of the magazine also makes it special as it will have a wider audience and multispecialty reach.

This edition has three articles covering different aspects of the peripheral nervous system from spinal cord to peripheral nerves. Latest 2D and 3D techniques in practice and future directions in both anatomic and functional domains of peripheral nerve imaging are addressed in detail.

Current high field technology depicts high resolution intra-neural fascicular detail in both normal and abnormal states as described in the article by Philipp Bäumer et al. (Heidelberg University Hospital, Germany). The high resolution imaging is important for diagnosis and surgical planning of nerve injuries, entrapments and for further evaluation of diffuse or polyneuropathies. Various technological developments, interpretation criteria, and future directions towards technique improvement are highlighted in my article from the work I have performed at Johns Hopkins University Hospital. Finally, there is an important article on diffusion tensor imaging improvement using syngo RESOLVE technique for functional spinal cord and brain imaging.

A bright future lies ahead for MRN with rapid deployment of this technology in current clinical setting for a variety of neuropathy diagnoses encompassing referrals from a number of providers, including neurosurgeons, neurologists, peripheral nerve surgeons, hand or foot and ankle specialists, plastic surgeons, and rehabilitation specialists. We will see further improvements in coming years in both anatomic and functional techniques, while user friendly analytic software becomes available.

Please enjoy reading the latest and inspiring RSNA issue of MAGNETOM Flash!

Avneesh Chhabra, M.D.
The information presented in MAGNETOM Flash is for illustration only and is not intended to be relied upon by the reader for instruction as to the practice of medicine. Any health care practitioner reading this information is reminded that they must use their own learning, training and expertise in dealing with their individual patients. This material does not substitute for that duty and is not intended by Siemens Medical Solutions to be used for any purpose in that regard. The treating physician bears the sole responsibility for the diagnosis and treatment of patients, including drugs and doses prescribed in connection with such use. The Operating Instructions must always be strictly followed when operating the MR System. The source for the technical data is the corresponding data sheets. The MRI restrictions (if any) of the metal implant must be considered prior to patient undergoing MRI exam. MR imaging of patients with metallic implants brings specific risks. However, certain implants are approved by the governing regulatory bodies to be MR conditionally safe. For such implants, the previously mentioned warning may not be applicable. Please contact the implant manufacturer for the specific conditional information. The conditions for MR safety are the responsibility of the implant manufacturer, not of Siemens.

1The software is pending 510(k) clearance, and is not yet commercially available in the United States and in other countries.
Magnetic Resonance Neurography – Techniques and Interpretation

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Introduction

Magnetic resonance neurography (MRN), akin to angiography, is an ever-advancing technology for multiplanar depiction of normal and abnormal peripheral nerves. This article will highlight various 2D and 3D pulse sequences available for non-selective and selective nerve visualization as well as their functional evaluation. Related interpretation pearls and pitfalls are discussed.

Techniques

MRN is best performed on 3 Tesla (T) scanners with dedicated extremity or wrap around flex coils as they can provide higher signal-to-noise ratio (SNR), which is essential to enhance the smallest structure in the neurovascular bundle, namely the peripheral nerves [1]. On current 1.5T scanners (MAGNETOM Aera, MAGNETOM Avanto), 2D (dimensional) imaging can be performed with near-similar resolution as on 3T scanners, however 3D imaging is often limited, especially if smaller voxels are used or fat suppression is applied. On the other hand, if there is metal in the field-of-view, in order to mitigate susceptibility artifacts and for superior nerve visualization, 1.5T imaging is often favored.

2D pulse sequences include high resolution (base resolution 256 or higher, in plane resolution 0.3–0.4 mm) T1-weighted and fat suppressed T2w images. Uniform fat suppression is essential to avoid artifactual increase in nerve signal intensity. Options include frequency selective fat suppression (fsT2w), short tau inversion recovery (STIR), and spectral adiabatic inversion recovery (SPAIR), or 2-point or 3-point Dixon techniques. fsT2w is often limited due to loss of fat saturation along the curvatures of extremities, especially in large subjects. STIR works well for 3D imaging with excellent fat suppression, however for 2D imaging it is often marred by poor SNR, excessive SAR (specific absorption rate) deposition and pulsation artifacts. SPAIR works well as it provides better fat suppression than fsT2w and higher SNR than STIR. It has virtually no pulsation artifacts and is also relatively SAR favorable (Fig. 1). It comes in weak and strong contrast versions. The authors prefer the strong version as it provides isointense signal in normal nerves [2].

1 Meralgia Paresthetica. Young woman with right anterolateral thigh pain and suspected lateral femoral cutaneous (LFCN) nerve abnormality. Axial T2 SPAIR image shows uniform fat suppression and abnormally hyperintense right LFCN (arrow) in keeping with clinical diagnosis of meralgia paresthetica.

2 T2 SPACE. Coronal non-fat suppressed T2 SPACE image through the pelvis shows bilaterally split sciatic nerves (arrows) in this patient with no symptoms of sciatica.
Dixon technique applied with T2w imaging gives separate water and fat images and also provides excellent fat suppression with higher SNR and contrast-to-noise ratio than STIR or SPAIR.

3D pulse sequences can be divided into anatomic and functional techniques. Anatomic techniques are further divided into nerve non-selective and nerve selective sequences. Nerve non-selective techniques include T1w imaging, namely VIBE (volume interpolated breathhold examination, T1 3DGRE) or MPRAGE (3D GRE); and T2w multislab acquisition, namely SPACE (sampling perfection with application optimized contrasts using varying flip angle evolutions) [3]. SPACE is heavily used in MRN examinations as isotropic spine echo type imaging can obtained using SPACE in a variety of contrasts (T1, PD, T2, STIR, and SPAIR) with constant or variable echo times (Fig. 2). The authors use VIBE for pre- and post-contrast imaging, and otherwise mostly use 3D imaging with fluid sensitive contrast. 3D STIR SPACE provides best fat suppression for the brachial and L5 plexus while 3D SPAIR SPACE provides higher SNR along with good fat suppression for extremity imaging (Figs. 3, 4). Dixon T2w imaging also provides excellent images of the brachial plexus, however currently can be acquired in 2D mode only (Fig. 5). Others such as 3D PD SPACE with variable echo time can produce similar image quality and allows multiplanar reconstruction (Fig. 6). Currently, 3D STIR SPACE and SPAIR SPACE sequences are most widely used and are time tested. Maximum intensity projections (MIP) of acquired images or their curved planar reformats produce excellent quality nerve images along their long axis. It remains to be seen if other techniques, such as Dixon with SPACE, can be obtained within acceptable time periods and provide the necessary isotropic spatial and good contrast resolution.

3D nerve selective imaging includes diffusion-weighting (DW) with a small b-value (80–200 s/mm²) to suppress flowing blood. It is a fine balance as one adds diffusion gradient to the 3D imaging, as it reduces SNR and can degrade image quality while providing the benefit of vascular flow signal suppression for selec-
Clinical Neurology

3D DW PSIF. Young woman with persistent carpal tunnel symptoms following a prior transverse carpal ligament release. Notice normal median (large arrow) and ulnar nerves (medium arrow) proximal to the carpal tunnel. A neuroma in continuity is seen along the distal aspect of the carpal tunnel (smallest arrow). Notice selective nerve depiction with excellent vascular signal suppression.

3D DW SPACE. MIP image from coronal DW 3D STIR SPACE sequence shows selective depiction of normal bilateral brachial plexuses (arrows) in a healthy volunteer. However note decreased SNR due to added diffusion component.

DTI. MIP coronal tensor image with inverted grey scale from axial DTI sequence shows enlarged ulnar nerve with abnormally low FA and high ADC values in this case of re-entrapped ulnar nerve at the site of prior anterior transposition (large arrows). Notice normal median nerve (small arrows).

Functional imaging of the peripheral nerves primarily includes diffusion tensor imaging (DTI). It has been exploited in various extremity and plexus peripheral nerves in the last few years and continuous refinements are being made. Single shot echo planar imaging (EPI) is the technique used in most centers and multiple diffusion moments (b-values) are applied to obtain functional parameter of apparent diffusion coefficient (ADC) [6]. At least 6 directions of interrogation are needed for DTI, although most authors have used 12–20 directions to obtain reproducible data. The authors use 3 diffusion moments (0, 800, 1000 s/mm²) and 12 directions of interrogation. Tight echo spacing, frequency selective fat suppression, auto-shimming before image acquisition and no motion degradation
Table 1: The imaging protocol employed in the magnetic resonance neurography examination for the LS plexus.

<table>
<thead>
<tr>
<th>MR sequence</th>
<th>Slice thickness (mm)</th>
<th>TR / TE (ms)</th>
<th>TF</th>
<th>Base resolution (pixels)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axial T1</td>
<td>4</td>
<td>800 / 12</td>
<td>6</td>
<td>832</td>
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<tr>
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<td>4</td>
<td>960 / 12</td>
<td>5</td>
<td>384</td>
</tr>
<tr>
<td>Axial T2 SPAIR</td>
<td>4</td>
<td>4890 / 80</td>
<td>22</td>
<td>256</td>
</tr>
<tr>
<td>Sagittal T2 3D SPACE</td>
<td>1</td>
<td>1000 / 97</td>
<td>81</td>
<td>256</td>
</tr>
<tr>
<td>Sagittal STIR</td>
<td>4</td>
<td>3700 / 18</td>
<td>22</td>
<td>256</td>
</tr>
<tr>
<td>Coronal STIR 3D SPACE</td>
<td>1.5</td>
<td>1500 / 91</td>
<td>41</td>
<td>256</td>
</tr>
</tbody>
</table>

are essential to obtain good and reproducible DTI data. Axial images obtained with 4–5 mm slice thickness with 0 gap can be reconstructed in multiple planes without artifacts (Fig. 9). These images then allow accurate tensor calculation, fractional anisotropy (FA) measurements and tractography. DTI has proven useful in non-invasive pre- and post-operative evaluation of carpal tunnel syndrome patients and peripheral nerve sheath tumors (PNST) as the involved nerves show reduced FA values that improve over time with treatment [7]. The benign PNSTs show higher ADC values than their malignant counterparts and tractography differences exist among different tumor types depending upon internal fascicular involvement or mere displacement [8] (Fig. 10). Further investigations are under way to evaluate the role of DTI in other types of neuropathies.

**Interpretation pearls and pitfalls**

While image generation is getting easier with high field MR techniques, the radiologists should learn to correctly interpret these high quality examinations. There is a steep learning curve for those who spend time with attention to detail, since nerve architecture is easily visible to the fascicular and perineurium level. One should learn normal nerve anatomy, variations, diagnostic pearls and pitfalls while obtaining all the information possible from clinical findings and available electrodiagnostic test results. Electrodiagnostic tests are also limited by false negative or indeterminate results, especially in deeply located nerves. So, while these results are helpful, one

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<table>
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<tr>
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<td>1500 / 91</td>
<td>41</td>
<td>256</td>
</tr>
</tbody>
</table>
should not get biased by negative results. The reader should follow a stepwise approach to imaging diagnosis:

1. Image quality – is it adequate and is fat suppression uniform? Are the nerves visible adequately and can the nerves be separately evaluated from adjacent vessels or compared side to side if the contralateral portion of body is available, such as in pelvic imaging. Recognize normal nerves and their variations (bifid nerve, split nerve with muscle belly intervening, intramuscular course of the nerve, etc.)

2. Look for orthopedic internal derangements which can mimic similar neuropathic symptoms or are potential cause of traction neuropathy, such as spondylosis, plantar fasciitis, tibialis posterior dysfunction, etc.

3. Look for clues of disseminated or systemic causes of neuropathy: one nerve abnormal over long distance away from entrapment sites or multiple regional nerves abnormal. This may happen in diabetic neuropathy, demyelinating neuropathies, hereditary neuropathy, vasculitis or toxic metabolic conditions. Usually hereditary neuropathy results in symmetric disease as compared to acquired conditions. Clinical findings should be correlated for insights into above diagnoses.

4. Look for focal area of nerve abnormality, abnormal T2 hyperintensity (approaching adjacent venous signal intensity) and / or fascicular abnormality (enlargement / effacement from edema / discontinuity) indicating entrapment or injury in the correct clinical scenario. If the nerve is really abnormal, the signal intensity will persist over few to many sections along its length versus signal change from a magic angle artifact. The nerve further enlarges with worsening neuropathy forming a pseudoneuroma in entrapment and neuroma in injury (lost partial or complete fascicular continuity with heterogeneous appearance) [9]. The nerve abnormality is generally worst at the site of insult and it fades gradually proximally and distally. Abrupt change in nerve intensity from bright-black-bright signal (Trip B sign) usually means severe focal neuropathy and a potential surgical case. Painful neuroma in continuity and nerve discontinuity in functionally important nerves also require surgical repair/reconstruction. Long standing neuropathy, such as in diabetes, can lead to atrophic appearance of the nerve with fascicular atrophy and intra-epeineurial fatty proliferation / replacement.

5. Evaluate regional muscles. As a rule, the muscle denervation changes are distal to the site of insult. If muscle changes are patchy or widespread in different nerve territories or associated with fascial edema, the diagnosis could be myopathy / myositis rather than denervation change. The diagnosis can be made with confidence if regional nerves are normal.

6. In case of a mass lesion, further characterize the lesion into neural or perineural masses. Age, clinical findings and anatomic MRN plus DTI are useful in the imaging evaluation of neural masses. Perineural lesions are further evaluated based on anatomic imaging and contrast evaluation into lipoma, ganglion cyst, hematoma, and abscess etc.

7. Finally, look for prior local surgical changes or nerve repair/reconstruction changes. The signal alteration may persist but in successful cases, the signal decreases within the nerves and denervation changes in the muscle resolve. In worsening nerve degeneration cases, the nerve signal approaches fluid signal and persists till the nerve atrophy starts, while the regional muscles undergo continued fatty replacement and atrophy [10]. Correlation with prior imaging studies is essential in these cases.

To conclude, magnetic resonance neurography is an exciting imaging technique that affords multiplanar anatomic and functional depiction of peripheral nerves and their related lesions. Appropriate imaging and accurate interpretation are essential components of successful performance of this ever advancing technique.

References

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MR Neurography – Diagnostic Criteria to Determine Lesions of Peripheral Nerves

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I) Introduction with potential clinical indications

Clinical examination and electrodiagnostic testing comprise the traditional methods of diagnostic investigation in peripheral nerve disease. These are excellent tools to assess function of peripheral nerves but can be limited in localizing a lesion. Recent developments in MR scanner and coil technology and the refinement of pulse sequences for increasing structural resolution have allowed imaging of fine details in healthy and diseased peripheral nerve [1, 2]. MR Neurography (MRN) at high magnetic field strength of 3 Tesla allows for the visualization of peripheral nerves at the fascicular level – the fascicle being the first order subunit of a peripheral nerve. The major limitation of clinical and electrophysiological examination is precisely the strength of MRN: the exact localization of nerve lesions. Lesion localization and spatial lesion patterns in the periph-

1 T2-weighted sequence with fat saturation at high resolution allows for imaging of peripheral nerves at the fascicular level. The sciatic nerve is shown here in hereditary NF2 polyneuropathy with various lesions ranging in size from pretumorous microlesions affecting only individual fascicles with no significant caliber change (lower row) to severe caliber thickening and enlargement of the entire nerve cross-section as in true, full-blown schwannomas (upper row, right panel). Modified from Baumer P. et al. Accumulation of non-compressive fascicular lesions underlies NF2 polyneuropathy. J Neurol. 2012, with permission.
eral nervous system are, as in many
diseases of the CNS, arguably among
the most important pieces of diagnostic
information.
One typical example for the use of MRN
is the differentiation between spinal
nerve root compression (radiculopathy),
plexus affection and peripheral nerve
lesion (peripheral neuropathy). Another
frequent and important situation is the
differentiation of a focal mononeuropa-
thy, which is potentially surgically ame-
nable, from neuropathies with involve-
ment of more than one nerve. These
oligo- or polyneuropathies almost exclu-
sively exhibit a disseminated lesion pat-
tern stemming from an inflammatory,
immune-mediated, metabolic or heredi-
tary origin, so that nerve surgery is not a
primary therapeutic option.
In traumatic nerve injuries, exact lesion
localization is the central diagnostic step
in pre-surgical work-up. The assessment
of nerve continuity and the identification
of potential scar tissue within and around
the nerve yield essential diagnostic infor-
mation. In cases of true neuroma with
discontinuity of the nerve, early indica-
tion for reconstructive nerve surgery by
MRN can improve outcome. However,
in instances where MRN shows a neu-
roma in continuity, surgery may become
unnecessary since spontaneous recovery
may occur under clinical monitoring.

II) How I do it
a) MRN protocol
A magnetic field strength of 3 Tesla is
advisable to achieve high-resolution
images at the fascicular level of peripheral
nerves. We perform clinical examinations
on a Siemens MAGNETOM Verio 3 Tesla
MR System (Siemens Healthcare, Erlan-
gen, Germany) which allows comfortable
positioning due to its wide bore diame-
ter. The MRN protocol varies depending
on the body region of interest and the
clinical-diagnostic question. In the vast
majority of patients, T2-weighted fat-
saturated sequences are acquired, which
show high contrast of nerve lesions to

Typical findings in cubital tunnel syndrome, a common entrapment neuropathy. The upper row depicts a typical compressive focal ulnar
nerve lesion at the elbow from proximal (2A) to distal (2C). Nerve T2w signal and caliber are severely increased compared with healthy control
2011;260(1):199-206, with permission.
Inferring the site of nerve lesion from the comprehensive evaluation of muscle denervation patterns. (3A) The denervation pattern in a patient with an L5 lesion additionally includes the posterior tibial muscle and the popliteal muscle. (3B) Patient with a lesion of the common peroneal nerve (NPC). The denervation is confined to the extensor compartment (here shown the anterior tibial muscle and extensor digitorum muscle) and the peroneal compartment (long and short peroneal muscle). Both compartments are known to harbor exactly the target muscles of the common peroneal nerve.

The magic angle effect occurs in tissue which is densely composed of collagen, such as in tendons, but also peripheral nerve. On the left the increase in T2 relaxation time can be seen as a function of angular deviation of nerve from the main static magnetic field (B0). The first maximum of this cosine type function is reached at 55°. The T2 signal increase is visually apparent (zoomed inserts above graph) at the maximum magic angle position of 55° in comparison with the neutral position (0°). However, true nerve lesions in neuropathies of various origins can usually be discriminated easily by the much stronger T2 signal increase (right column, traumatic, compressive and inflammatory true nerve lesions). Modified from Koestel et al. AJNR Am J Neuroradiol. 2011 May;32(5):821-7, with permission.

Artificial T2 signal increase (Magic Angle Effect)

True T2 signal increase (Lesions)

Trauma

Entrapment

Inflammatory

Angle dependent increase of nerve T2 signal

surrounding tissue (Fig. 1). This may be complemented by T1-weighted sequences, with and without contrast administration, and T1-VIBE or TrueFISP for 3D reconstruction. T2-weighted SPACE STIR sequences at isotropic voxel size allow for imaging at submillimeter resolution of brachial and lumbosacral plexus structures. CISS may be added for imaging of intradural filaments in cases of trauma. Choosing appropriate radiofrequency coils is crucial for image quality and diagnostic performance. For extremities, we use an 8-channel knee coil (Siemens Healthcare, Erlangen, Germany) for most regions. For brachial plexus structures, we additionally acquire high-resolution sequences using dedicated surface coils (multichannel phased-array 2 element coil, NORAS GmbH, Höchberg, Germany). The applicability of diffusion tensor imaging (DTI) by echo-planar-imaging (EPI) has been investigated for many years [3]. It is currently not used in routine clinical examinations and its diagnostic accuracy has so far only been studied for carpal tunnel syndrome [4]. Various technical issues may limit the clinical significance of application of EPI DTI in the near future, most importantly its structural resolution, which is significantly inferior to T2-weighted sequences and constrains its use to large peripheral nerves of the extremities. Segmented EPI could help to overcome these limitations. Extent of coverage of an MRN examination is decided on a case-by-case basis. In the setting of focal nerve damage, the examination can be targeted to one region by different sequences. Occasionally, larger coverage of an entire extrem-
Neurology

ity including the plexus and at the same time high resolution may be useful. One such example is the hereditary polyneuropathy in neurofibromatosis type II [5] where a large number of microlesions may lead to motor and sensory loss but a single, surgically removable macroleesion must be ruled out. Also, large coverage MRN is able to characterize the spatial pattern of polyneuropathies such as diabetic polyneuropathy [6].

b) Diagnostic criteria

Nerve T2 signal is a highly sensitive and specific diagnostic sign for the presence of neuropathy [7]. Increased T2 signal as a parameter of high pathomorphological contrast is readily assessed by qualitative visual evaluation. Nerve caliber is a second important diagnostic sign and, when clearly increased, usually indicates the presence of a more severe neuropathy. Looking beyond the nerves, the denervation pattern of muscles adds crucial information [8]. It can help establish the site of a nerve lesion, as in L5 radiculopathy compared to common peroneal neuropathy (Fig. 3). Contrast enhancement of peripheral nerves is observed in cases of peripheral nerve sheath tumors such as neurofibromas, schwannomas and perineuriomas. Whether contrast uptake can discriminate between different tumor entities and dignities has not been formally investigated. In cases of potential nerve trauma, nerve continuity is a straightforward but essential criterion when a ‘true’ neuroma has to be discriminated from a neuroma-in-continuity.

c) Pitfalls

The magic angle effect causes an increase in T2 of fibrous tissue densely packed with collagen which results in artificially bright signal in T2w images. The magic angle effect depends on the angular orientation of the main longitudinal axis of the investigated structure (e.g. a tendon or peripheral nerve) relative to the field direction of the main static magnetic field (B0) and reaches its maximum at 55°. It is related to the interaction of collagen bound protons and can also be observed in the peripheral nervous system [9, 10]. Below an angle of < 30° the artifact becomes negligible so that correct positioning of limbs in an MRN examination can usually avoid its occurrence [10]. There are two regions where an oblique anatomical orientation of nerves cannot be avoided in the scan: the supraclavicular brachial plexus and the proximal common peroneal nerve. Hence, in these regions, the magic angle must always be considered in image interpretation.

A second pitfall in MRN image interpretation is the presence of small epineural (outside the epineurium as the outer nerve sheath), epineurial (inside the epineurium), and intraneurial vessels (within the interfascicular space). Small veins

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5A Traumatic lumbosacral plexus injury. (5A) Coronal SPACE STIR gives an overview of lumbosacral plexus and proximal sciatic nerve. Pseudomeningocele of L4 – S 2/3 indicate root avulsions on the right side. (5B) Additional CISS of the lumbosacral spinal canal in coronal reconstruction confirms roots avulsions of L5 – S2. (5C) Axial reconstruction of CISS. Dorsal portions of the right L4 root appear intact at least in their intra-spinal course. (5D) Coronal SPACE STIR maximum intensity projection (MIP) of a normal lumbosacral plexus including the proximal sciatic nerve.
and nerve lesions can have a similar T2 signal intensity. In most cases, small veins are readily discriminated from nerve lesions due to their winded course, strong and relatively homogeneous T2w hyperintense signal, and their eventual anatomical entry or exit into the nerve via its epineural surface. When in doubt, one proposed method to discriminate the two is to acquire an SSFP sequence [11, 12]. Another approach is to acquire T1-weighted images with fat saturation after administration of contrast media since nerve lesions do not enhance as strongly as vessels.

III) Useful applications of MR Neurography

a) Focal compressive and traumatic lesions

Typical frequent referrals for an MRN exam include suspected compressive neuropathies such as ulnar neuropathy at the elbow or at the wrist. MRN has a high diagnostic accuracy in confirming or excluding these [7, 13]. Main diagnostic criteria are T2 signal and caliber increase. Other frequent focal neuropathies are traumatic nerve injuries. For patients in whom the region of the nerve lesion is known, MRN is most useful in determining whether continuity of the nerve is intact as well as the precise localization of neuroma (Fig. 6).

In other cases, the exact lesion localization and pattern can only be established by MRI. This is often the case in plexus injuries where electrodiagnostics are limited (Fig. 5). Precise determination of lesion pattern and lesion localization by MRN allows targeted surgery.
b) Disseminated inflammatory lesions
Known or suspected disseminated nerve lesions require an examination tailored to the needs of the individual patient. In suspected inflammatory neuropathy, the precise spatial pattern of involvement of nerves is diagnostically important for classification and treatment. MRN is an excellent method to investigate this even in the most complex nerve structures like the brachial plexus. Figure 7 illustrates the precision of the method.

c) Distal symmetric polyneuropathies such as diabetes mellitus associated PNP
In addition to its manifold clinical applications, MRN contains immense potential for understanding and monitoring the evolution of peripheral nerve disease. In diabetic neuropathy for example, as a typical symmetric polyneuropathy with mainly distal symptoms, the MRN findings of a largely proximal lesion pattern are enhancing our understanding of the pathophysiology of the disease. MRN could for the first time document proximal nerve lesions in-vivo [6]. These findings promise a useful research purpose for MRN: to monitor the evolution of nerve lesions in diffuse polyneuropathies and better understand if proximal nerve lesions cumulate to cause distal symptoms, an intriguing pathomechanisms which so far has not been proven by any other clinical-diagnostic or scientific method of investigating nerve pathology.

References

Table 1: Exemplary MRN sequences.

<table>
<thead>
<tr>
<th>Sequence</th>
<th>TR</th>
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<td>180</td>
<td>0.8</td>
<td>0.781 x 0.781</td>
<td>250</td>
<td>320/314</td>
<td>72</td>
<td>2</td>
<td>neck (Siemens)</td>
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<tr>
<td>T2 SPAIR sagittal-oblique</td>
<td>5530</td>
<td>45</td>
<td></td>
<td>3.0</td>
<td>0.469 x 0.469</td>
<td>150</td>
<td>320/198</td>
<td>51</td>
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<td>dedicated surface coil (NORAS GmbH)</td>
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<td>3.0</td>
<td>0.300 x 0.300</td>
<td>130</td>
<td>512/358</td>
<td>45</td>
<td>3</td>
<td>knee 8-channel phased-array (Siemens)</td>
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Abstract
In this paper we present some applications of the syngo RESOLVE sequence that enable high-resolution diffusion-weighted imaging. The sequence is based on a readout-segmented EPI strategy, allowing susceptibility distortions and $T_2^*$ blurring to be minimized. The RESOLVE sequence can be combined with other acquisition strategies such as reduced field-of-view (FOV) and parallel imaging, to provide state-of-the-art tractography of the full brain and cervical spinal cord. The RESOLVE sequence could be of particular interest for ultra high field systems where artifacts due to susceptibility and reduced $T_2$ values are more severe. At all field strengths, the sequence promises to be useful in a number of clinical applications to characterize the diffusion properties of pathology with high resolution and a low level of image artifact.

1. Introduction
1.1. Diffusion-weighted imaging
Diffusion-weighted MRI makes it possible to map white matter architecture in the central nervous system based on the measurement of water diffusion [11]. The technique works by using MRI sequences that are sensitized to the microscopic motion of water molecules, which are in constant motion in biological tissues (Brownian motion). Using the well-known pulse sequence introduced by Stejskal and Tanner in 1965 [29], it is possible to quantify the extent of water displacement in a given direction. This sequence consists of magnetic field gradient pulses (diffusion-encoding gradients) that are applied before and after a $180^\circ$ radiofrequency (RF) refocusing pulse. The first gradient pulse dephases the precessing nuclear spins that generate the signal in MRI. In the theoretical case of static molecules, the second gradient pulse completely rephases the spins and there is no attenuation due to the application of the gradients. However, if water molecules move during the application of this pair of gradients, spins are dephased and signal decreases as a function of the magnitude of the displacement, leading to a so-called diffusion-weighted signal. The magnitude by which the signal is weighted by diffusion is dictated by the so-called $b$-value, which depends on the length and amplitude of the applied diffusion-encoding gradients, as well as the duration between the first and the second gradient pulse (also called diffusion time). By applying diffusion gradients in various directions (e.g., 20 directions equally sampled on a sphere), it is possible to estimate the rate and direction of water diffusion. For example in pure water, molecules diffuse equally in all directions, hence the diffusion is described as isotropic. Conversely, in mesenchymal structures such as the white matter or muscles, water diffuses preferentially along the direction of the fiber. In such a case, the diffusion is anisotropic [3].
1.2. DTI and tractography

After acquiring a set of diffusion-weighted data, it is possible to compute a parametric model representing the main diffusion direction at each voxel [2]. This type of model is described by a tensor and is usually represented graphically as a 3D ellipsoid. This technique is known as Diffusion Tensor Imaging (DTI) and makes it possible to derive metrics such as fractional anisotropy (FA), mean diffusivity and axial and radial diffusivities. These metrics are relevant as they have been shown to correlate with demyelination and axonal degeneration [28] and can therefore be used to obtain markers of demyelination in multiple sclerosis [26], spinal cord injury [7] or amyotrophic lateral sclerosis [8].

Having modeled the diffusion tensor, it is possible to reconstruct global axonal pathways by linking every voxel with a similar eigenvector (principal axis of the diffusion tensor) to its neighbors. This procedure is called fiber tracking or tractography [18]. It should be mentioned however that the reconstructed fiber bundles do not represent real axonal tracts but rather represent the path where water diffuses preferentially. It is therefore an indirect measure of axon orientation. Hence, tractography results should be handled with care since they may not represent the real pathway of axons [14]. For instance, false negatives could be induced by the presence of crossing fibers, which would artificially decrease the FA and stop the tracking procedure. Inversely, false positives could be induced by fibrous structures such as scar tissue, in which water diffusion also has anisotropic properties [27]. In such cases, fiber tracking can become fiber tracking.

To illustrate the need for caution, Figure 1 shows an example of tractography performed in a healthy subject, before and after correction for susceptibility artifacts, which can induce strong image distortions in the phase-encoding direction (antero-posterior in this example). On the ‘raw data’ (1A), a false apparent interruption of the tracts is clearly observed, and can be misinterpreted as an interruption of the white matter tracts. However, this apparent interruption is simply due to susceptibility distortions, as can be seen on the ‘corrected data’ (1B). This result suggests two things: (I) a careful assessment of the image data is essential before further processing is applied and (II) susceptibility-induced distortions should be minimized by an appropriate choice of data acquisition technique. This might ultimately reduce the need for quality assessment, which is not often possible in clinical routine. This brings us to the next section on how to reduce susceptibility artifact during the acquisition.

1.3. Artifacts in diffusion-weighted imaging

In this section we will mostly focus on susceptibility artifacts and T2* blurring. Other artifacts exist in diffusion-weighted imaging [15], but are not in the scope of the presence article. Single-shot echo planar imaging (ss-EPI) is an ultra-fast imaging sequence that makes it possible to acquire a full slice within a single repetition time (TR) [16]. To achieve this, spatial encoding gradients are played out rapidly along two gradient axes (e.g. X and Y) in order to sample all raw data (or k-space) points in a ‘single shot’ after an RF excitation pulse. By applying the EPI readout directly after the excitation pulse, a gradient echo image can be generated. Alternatively, an RF refocusing pulse can be inserted between the excitation and EPI readout to produce a spin echo image, which has reduced susceptibility-induced signal loss, but has the same sensitivity to susceptibility-based distortions. An oscillating readout gradient is applied along one axis (e.g. X), consisting of a series of gradient pulses with alternating polarity and each of these readout gradient pulses is used to sample a single line of k-space. A short phase-encoding gradient or ‘flip’ is applied along an orthogonal gradient axis (e.g. Y), so that each readout gradient corresponds to a different line of k-space in the Y direction. While acquisition of a full line a k-space in the read-
out direction is obtained rapidly (in about 1 ms), the acquisition of k-space data along the phase-encoding direction takes much more time. For example, for a 128x128 matrix, acquisition of the full k-space can take more than 100 ms. During that time, spins experience dephasing due to local $B_0$-inhomogeneities. These undesirable inhomogeneities are typically induced at interfaces between structures with different susceptibility properties, such as air/tissue or bone/muscle. As a result, phase errors accumulate in the phase-encoding direction, causing geometric distortions in the reconstructed image in that direction. EPI sequences are fast and particularly appropriate for diffusion-weighted experiments [31] where multiple datasets need to be acquired in a clinically acceptable time with low sensitivity to motion-induced phase error [17]. However, single-shot EPI also has the disadvantage that it is particularly sensitive to susceptibility-related geometric distortions. Reduction of susceptibility artifacts can be achieved by sampling the k-space faster, which decreases the accumulation of the phase errors. Faster acquisition can be achieved by reducing the echo spacing, which is defined as the total time between the application of two phase-encode blips. Echo spacing can be reduced by (I) increasing the readout bandwidth, (II) reducing the number of phase lines by reducing the FOV (because the sampling density in k-space defines the FOV in the image domain) and (III) combining several subsampled k-space acquisitions (multi-shot). In addition to susceptibility artifacts, T2* blurring can be induced due to the relatively long sampling of k-space, during which T2* relaxation occurs (Figure 2, left). This effect is more pronounced in regions with large $B_0$ inhomogeneities. Another undesirable effect of the long sampling is that it will impose a relatively long echo time (TE) (Figure 2, right). As a result, the recorded signal amplitude will be small due to the relatively short transverse relaxation (T2) in white matter (about 70 ms at 3T). This effect is critical since SNR is proportional to $\exp(-TE/T2)$, therefore even a small increase of TE (several ms) can substantially reduce the signal-to-noise ratio (SNR).

All these limitations become even more problematic at higher field, where T2 is reduced (about 50 ms at 7T) and $B_0$ homogeneity is more difficult to achieve, resulting in a greater level of susceptibility artifact. Figure 3 shows an example of the high-field degradation in image quality when using the standard single-shot EPI technique at 1.5T and 7T.

To summarize, the main issues encountered with diffusion-weighting EPI are:

- **Susceptibility artifacts**, which are caused by an accumulation of phase error during the readout, leading to image distortions.
- **T2* blurring**, which is caused by the long data sampling interval during the readout, leading to a broadening of the point spread function and a blurring of the reconstructed image.
- **Long TE**, which is imposed by a combination of the diffusion-encoding gradients and the long data sampling interval and results in low SNR.

These three arguments provide motivation for a reduction in the sampling time, which will address all these issues, as illustrated in Figure 4.

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3 A comparison of single-shot EPI at 1.5T (3A) and 7T (3B).

4 Benefits of faster sampling for: (I) reducing susceptibility distortions (less accumulation of phase errors), (II) reducing T2* blurring (faster sampling) and (III) enabling shorter TE (due to the faster sampling of k-space, which in turn brings the center of the readout window closer to the 180º pulse).
2. Multi-shot techniques and syngo RESOLVE

As described in the previous section, susceptibility artifacts and T2* blurring can be reduced by decreasing the sampling time (or the effective echo spacing). In EPI, this can be done by skipping lines of k-space in the phase-encoding direction (also called interleaved EPI). Multi-shot techniques consist of acquiring several sub-sampled images and combining them to reconstruct a full k-space data set. An example of a 2-shot interleaved EPI acquisition is illustrated in Figure 5. In this approach, the first image is acquired by only filling even lines of k-space, while the second image is acquired by filling the odd lines. Then, both data sets are combined in order to obtain the full k-space data that is required for image reconstruction. The key aspect is that each of the sub-sampled data sets is acquired twice as fast as in the single-shot case, which reduces the effective echo spacing by a factor two, and hence also reduces the distortions by a factor of 2. However, one disadvantage of the interleaved EPI approach is that the overall acquisition time is longer than single-shot techniques (because multiple TR intervals are required to form an image). Another disadvantage is that phase errors due to motion (from subject or physiology) can occur between the acquisition of different interleaves, which can lead to severe ghosting artifacts [6].

To overcome the mismatch between different data interleaves, one can acquire navigator echoes that monitor phase errors and correct them before image reconstruction. At their introduction, navigator echoes consisted in single line of k-space acquired at the center, which was used to apply a linear phase correction in the readout direction [1, 20]. Later, 2D navigator echoes enabled linear phase correction both in the readout- and in the phase-encoding directions [4]. More recently, non-linear approaches have been introduced [17], which is particularly important for diffusion-weighted imaging of the brain, where cerebrospinal fluid pulsation leads to non-rigid-body brain motion that results in non-linear phase errors if the motion occurs during the application of the diffusion-encoding gradients. 2D non-linear phase correction is most easily applied when each shot in the multi-shot data acquisition samples a contiguous set of data points from a sub-region of k-space. In this case, the Nyquist sampling condition is fulfilled and the phase correction can be applied as a complex multiplication in image space. The interleaved EPI sampling scheme does not fulfill this local Nyquist sampling condition and results in aliased signal contributions that complicate the phase correction procedure.

2.1. Readout-segmented EPI

An alternative to the interleaved approach is to perform readout-segmented EPI [25]. In readout-segmented EPI, k-space is filled with a series of concatenated segments in the readout direction (Figure 6), corresponding to shorter readout gradient pulses and an overall reduction in the length of the EPI readout. As with interleaved EPI, the shorter readout time results in a reduction in the level of susceptibility-based distortion.

2.2. Diffusion-weighted readout-segmented EPI

One advantage of readout-segmented EPI compared to phase-segmented (or interleaved) EPI is that each segment consists of a Nyquist-sampled region of k-space that is well suited to the application of 2D non-linear phase correction, as described above. This property of the readout-segmented EPI sampling scheme was first exploited by Porter et al. [24] who proposed the sequence shown in figure 7 for application to diffusion-weighted imaging.
weighted imaging. The sequence consists of an imaging echo, which is used to sample the standard readout-segmented EPI data, and a 2D navigator echo, in which a low resolution EPI readout is used to sample the center of k-space at each shot. The sequence was subsequently modified to support parallel imaging [22] using GRAPPA [10]. This made it possible to further shorten the readout time with a corresponding additional reduction in the level of susceptibility artifact. The resulting sequence has been given the acronym RESOLVE1 (Readout Segmentation Of Long Variable Echo-trains).

### 2.3. Reacquisition technique

In some cases, the level of motion-induced phase error can be too severe for a reliable phase correction due to signal voids in the navigator images. Although physiological gating strategies can be employed to mitigate these effects, they do not address all sources of motion and can significantly increase acquisition time. To avoid the residual artifacts associated with these occasional extreme phase errors, the RESOLVE1 sequence uses a reacquisition strategy to identify and re-measure corrupt readout segments that cannot be reliably phase corrected. This method is based on an original concept that was proposed for interleaved EPI and 1D navigator echoes [19], but uses an adapted real-time algorithm, which has been optimized for the combination of 2D navigator data and the readout-segmented EPI sampling scheme [21].

For a comprehensive description of the RESOLVE1 sequence, including the adaptations for parallel imaging and data reacquisition, the reader is referred to [23].

### 3. Application of the RESOLVE1 sequence

To illustrate the benefits of the RESOLVE1 sequence, Figure 8 shows a comparison between the conventional single-shot EPI (8A) and RESOLVE1 (8B). Acquisition parameters were:

**RESOLVE**: Axial slices, TR 3800 ms, TE 80 ms, 4 nex, 12-4 mm thick slices with 0.4 mm gap, 320 × 256 matrix, b=700 s/mm² only in slice direction (S-I), GRAPPA x 2, segment 21, FOV 163 mm, acquisition time 6–7 min (depending on reacquisition).

**Single-shot EPI**: Axial slices, TR 3800 ms, TE 102 ms, 32 nex, 12-4 mm thick slices with 0.4 mm gap, 192 × 192 matrix, b=700 s/mm² only in slice direction (S-I), GRAPPA x 2, FOV 220 mm, acquisition time 2 min 15 sec.

All scans were performed at 3T using a 32-channel array head coil.

Images represent an axial mean diffusion-weighted image located in the lower brain region, which is well-known for its susceptibility artifacts due to the sinuses and temporal poles. In the ss-EPI image, strong susceptibility artifacts can be observed as image and signal distortions. For example, the temporal region not only shows image distortion along the phase-encoding direction (antero-posterior), but also shows increase of signal intensity caused by the overlap of distorted voxels (arrow). This spurious signal increase could be falsely interpreted as pathological. Conversely in the RESOLVE1 image, the same region shows a substantial reduction in this susceptibility artifact. In addition, an overall blurring can be observed on the ss-EPI image, which is caused by the relatively long k-space sampling relative to the T2* decay. Reducing the sampling time also reduces this T2* blurring, as can be seen on the RESOLVE1 image, where the image appears sharper, making it possible to delineate white and gray matter with greater precision.

#### 3.1. 1x1x1 mm³ resolution full brain at 7T

With the RESOLVE1 sequence it is possible to acquire high-resolution images without the large increase in susceptibility and T2* artifacts that would affect an ss-EPI acquisition. In one study at 7T, the sequence was used to perform full brain diffusion-weighted imaging at 1x1x1 mm³ resolution [12]. The study was performed on a whole-body scanner (MAGNETOM 7T, Siemens Healthcare, Erlangen, Germany) with a 24-channel RF receive coil (Nova Medical, Wilmington, MA, USA). Acquisition parameters were: 104 axial slices, matrix 220×220, FOV 220 mm, slice thickness 1 mm (no gap), TR 7500 ms, TE 76 ms, 7 readout segments, 30 diffusion-weighting directions at a b-value of 1000 s/mm², acquisition time was 78 min. The diffusion-weighted images were filtered with a two-stage hybrid image restoration procedure, corrected for subject motion and registered to a T1-weighted anatomical image (1 mm resolution) of the same participant. For each voxel, multiple fiber orientations were modeled by constrained spherical deconvolution [30] followed by whole-brain fiber-tracking using...
Figure 9 shows results of the fiber orientation distribution function (ODF) and tractography centered over the pre-motor cortex. The spatial consistency of the fiber ODF demonstrates the high SNR of the data (Figure 9B). Tractography successfully demonstrated intersection of the corpus callosum (CC, pink), the corona radiata (CR, blue) and the superior longitudinal fasciculus (SLF, green, tracks not shown).

The same data were processed using track-density imaging [5], which makes it possible to reconstruct fiber bundles at 200 µm resolution. Figure 10 shows the results at the three orientations in order to appreciate the 1 mm isotropic resolution. The views are centered over the thalamus. The ultra-high resolution in combination with high SNR and low susceptibility artifacts provide an unprecedented level of detail about the anatomical organization of the thalamus. The ability of the sequence to provide full brain coverage at an isotropic resolution of 1 mm made it possible to reconstruct track density images of the whole brain with no orientation bias.

3.2. Spinal cord DTI at 3T
Another powerful application of the RESOLVE sequence is the spinal cord. Indeed, spinal cord DTI is particularly challenging due to (I) the small cross-sectional size of the spinal cord, which requires high spatial resolution, resulting in low SNR at standard imaging times; (II) susceptibility artifacts, which are caused by inhomogeneous magnetic field in the spinal region due to the close proximity of tissue with very different magnetic susceptibility (tissue, bone, air, cartilage) and (III) physiological-related motion due to respiration, heart beat, cerebrospinal fluid pulsation. The first two arguments (small size and susceptibility artifacts) are particularly well suited for the RESOLVE¹ sequence, as this

³MRtrix is not a Siemens Healthcare product. Siemens bears no responsibility for this product including, but not limited to, its regulatory status. Further information about the software can be found at the website listed.
sequence provides high SNR and enables reduction of susceptibility distortions. The last challenge (physiological motion) is a typical cause of failure for segmented approaches, however the non-linear 2D phase correction combined with the reacquisition scheme enabled in the RESOLVE sequence ensures low sensitivity to physiological motion.

An example of the RESOLVE in the spinal cord is shown in Figure 11. To further improve image quality, the RESOLVE sequence was used with parallel imaging using the GRAPPA algorithm [10] to further reduce susceptibility artifacts and shorten the TE. To maximize the capabilities of parallel imaging, a dedicated 32-channel head/spine coil was used, which enables up to 4× acceleration [9]. Acquisition parameters were: 1.7 mm isotropic, 3 shots, R = 3 acceleration (iPAT3), TE 74 ms, Cardiac gating, b 800 s/mm², 30 directions, acquisition time 8 min. The sequence only used 3 shots (i.e., 3 segments per k-space), therefore the acquisition was relatively short. Although minor distortions are still visible on the b = 0 image at the level of each intervertebral disk, the spinal cord itself is not affected by these intervertebral distortions as assessed by the mean diffusion-weighted image, the fractional anisotropy (FA) map and the FA color map. Note the particularly high spatial resolution (1.7 mm isotropic, 4.9 mm³ voxel volume), which is rarely achieved in spinal cord imaging. One particularly difficult acquisition is when one wants to image the full brain and spinal cord. The challenge of B₀-inhomogeneity can be partially overcome when only the spinal cord is imaged by adjusting the shim volume within an elongated box that nicely encompasses the spinal cord. However, when the FOV includes both the brain and spinal cord, it is a difficult for the shimming algorithm to find appropriate fitting coefficients for ensuring B₀-homogeneity both in the brain and spinal cord, due to the complex geometry and the presence of several susceptibility interfaces (sinuses, temporal poles, brainstem, intervertebral disks and lungs). This type of acquisition is therefore particularly well-suited to the RESOLVE sequence.

Images were acquired with the following parameters: 2.2 mm isotropic resolution, matrix 138 × 104, TR ~ 15000 ms (cardiac gating), TE 66 ms, 3 shots, R = 3 acceleration, b 800 s/mm², 30 diffusion-weighting directions, acquisition time 10:45 min. Images were further corrected for residual susceptibility-distortions in the thoracic region (due to the close proximity with the lungs), using the reversed gradient technique [13]. Tractography was performed using the Diffusion Toolkit¹ (deterministic FACT algorithm) and displayed with TrackVis¹ (http://www.trackvis.org/). Diffusion-weighted images were then registered to the 1 mm isotropic anatomical image (MPRAGE). Results are shown in Figure 12 and demonstrate reduction of susceptibility artifacts, enabling to explore connections within and between the brain, brainstem and cervical spinal cord.

4. Conclusion

¹Diffusion Toolkit and TrackVis are not Siemens Healthcare products. Siemens bears no responsibility for these products including, but not limited to, their regulatory status. Further information about the software can be found at the website listed.
tions in research include the study of white matter architecture in the brain, brainstem and spinal cord, and also the possibility to study fine structure such as the thalamus and the gray nuclei. In clinical application, the reduced level of susceptibility artifact with the RESOLVE sequence can provide better interpretation of abnormalities in white matter pathology. Although acceptable image quality can already be obtained using the conventional single-shot EPI sequence providing shimming is successful, this procedure is strongly user-dependent, subject-dependent and structure-dependent. Because this variability is not acceptable in clinical routine, the RESOLVE sequence has the potential to ensure robust image quality. Moreover, the RESOLVE sequence may be particularly useful in cases where patients have metallic implants, notably in spinal cord injury, where assessing the integrity of spinal pathway would be of tremendous help for patient prognosis.

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The software is pending 510(k) clearance, and is not yet commercially available in the United States and in other countries. The 7T system is a research system only. It cannot be used outside of a research study.

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