Peripheral Nerve Lesions
High-Resolution MR Neurography of Diffuse Peripheral Nerve Lesions

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High-resolution MR imaging of peripheral nerves is becoming more common and practical with the increasing availability of 3T magnets. These magnets provide high SNR, which can be used for a quicker acquisition time as well as higher image contrast and resolution. There have been multiple reports of MR imaging of peripheral nerves in compression and entrapment neuropathies. However, there is a relative paucity of literature on the MRN appearance of diffuse peripheral nerve lesions. These lesions seen on MR imaging present a diagnostic dilemma because a long list of pathologies could be causing them. We attempt to highlight the salient imaging features of myriad diffuse peripheral nerve disorders and to describe a diagnostic approach to these lesions on the basis of the available literature and our experience in this area. Various clinical and pathologically proved relevant examples are illustrated.

Fig 1. Normal (A and B) versus abnormal (C) MRN appearance of the sciatic nerve. Axial T1WI (A) and T2WI (B) sections at the level of the midthigh show low-intermediate nerve signal intensity (circles). T1WI demonstrates fat planes delineating the normal nerve (perineural fat). C, Axial STIR SPACE at the level of the thighs shows an abnormal sciatic nerve. Notice the enlarged size and T2 hyperintensity of the fascicles. The dark rim of perineural fat is also disrupted. Also note posterior compartment denervation muscle edema (arrows).

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High-Resolution MRN Technique

MRN techniques can be broadly divided into T2-based imaging or diffusion-based imaging. T2-based MR imaging of peripheral nerves is generally preferred over diffusion-based imaging because radiologists and technologists are more familiar with T2-based imaging; moreover, the techniques and the pulse sequences are easier to implement. Diffusion-based imaging of peripheral nerves is still in the research phase due to suboptimal SNR obtained in peripheral nerve imaging and at present has not been widely implemented. T2-based MRN imaging relies on obtaining high-resolution (2- to 3-mm section thickness) T1WI SE/T1WI FLAIR images and T2WIFS SE/SPAIR/STIR images in multiple planes (Fig 1). The abundant fat around the fascicles and the nerve itself makes these structures clearly visible on T1WI. STIR works best at 1.5T, while SPAIR with varying flip angles is the favored fat-suppression technique for 3T magnets because it provides higher SNR than STIR images. T2WIFS may provide inhomogeneous fat suppression, especially for large peripheral areas of the body chosen for evaluation of diffuse nerve lesions. 3D pulse sequences such as SPACE have been reported to perform well at 1.5T and 3T MR imaging (Fig 2). Multiplanar reconstructions of these 3D sequences allow good depiction of peripheral nerves, which commonly course through a variety of obliquities.

Normal and Abnormal Peripheral Nerves on High-Resolution MRN Imaging

The reader should carefully evaluate various characteristics of peripheral nerves on high-resolution MRN imaging (Table 1), such as size, signal intensity, fascicular pattern (Fig 1), and course (Fig 2). Various imaging criteria are helpful in differentiating normal and abnormal nerves.

<table>
<thead>
<tr>
<th>Table 2: Etiologic classification of diffuse lesions that may be seen on MR imaging of peripheral nerves</th>
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<tr>
<td>Classification</td>
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<tr>
<td>Traumatic</td>
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<tr>
<td>Inflammatory</td>
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<tr>
<td>Brachial plexitis</td>
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<tr>
<td>CIDP</td>
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<tr>
<td>MMN</td>
</tr>
<tr>
<td>Vasculitis</td>
</tr>
<tr>
<td>Infectious</td>
</tr>
<tr>
<td>Pyogenic, viral, leprosy, others</td>
</tr>
<tr>
<td>Hereditary</td>
</tr>
<tr>
<td>CMT/HSMN</td>
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<tr>
<td>Radiation plexopathy/neuropathy</td>
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<td>Neoplastic and tumor variants</td>
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<tr>
<td>Neurofibromatosis</td>
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<tr>
<td>NL</td>
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<tr>
<td>Perineuroma</td>
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<tr>
<td>FLH</td>
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<td>Amyloidosis</td>
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Table 3: Summary of MR imaging characteristics in diffuse peripheral nerve lesions

<table>
<thead>
<tr>
<th>Etiology</th>
<th>Enlargement</th>
<th>T2 Hyperintensity</th>
<th>Fascicular Pattern</th>
<th>Course</th>
<th>Enhancement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trauma</td>
<td>±</td>
<td>+</td>
<td>Minimally effaced/disrupted</td>
<td>May be altered due to hematoma/fracture</td>
<td>–</td>
</tr>
<tr>
<td>Acute inflammation</td>
<td>±</td>
<td>++</td>
<td>Effaced/preserved</td>
<td>Not altered</td>
<td>+</td>
</tr>
<tr>
<td>CIDP</td>
<td>++</td>
<td>++</td>
<td>Effaced/enlarged</td>
<td>Not altered</td>
<td>++</td>
</tr>
<tr>
<td>Infectious</td>
<td>++</td>
<td>+</td>
<td>Effaced/disrupted</td>
<td>May be altered by abscess/granulation tissue</td>
<td>+</td>
</tr>
<tr>
<td>CMT/HSMN</td>
<td>++</td>
<td>++</td>
<td>Preserved ↑ fatty infiltration around atrophic fascicles</td>
<td>Normal</td>
<td>±</td>
</tr>
</tbody>
</table>

Radiation ± (linear; geographic distribution of radiation field) + | Preserved/effaced | Altered in subacute-chronic stages due to developing fibrosis | + |

NL ++ | ++ | Effaced/disrupted | Altered with focal masses | ++ |

NF1 ++ | ++ | Preserved | Altered with focal masses | ++/++ |

MPNST in NF1 ++ | ++ | Disrupted | Altered due to mass-effect | ++ (heterogeneous) |

Perineuroma ++ | ++ | Preserved/effaced focally | Altered with focal masses | ++ |

FLH ++ | ++ | Preserved (coaxial cable/spaghetti) | Altered with thick tortuous nerve | – |

Amyloid ++ | ++ | Cutaneous induration with subcutaneous stranding | Preserved/disrupted focally | Altered with focal masses | ± |

* ± indicates variable; +, mild; ++, moderate; ++++, marked; –, absent.

Fig 3. Traumatic neuropathy. A 46-year-old man had a posterior shoulder dislocation during a seizure episode. After the postictal phase, he had weakness of shoulder abduction and sensory loss over the deltoid area. Coronal STIR image shows minimal diffuse nerve enlargement and T2 hyperintensity of the terminal branch of the brachial plexus (arrow) as a result of direct trauma of the recent shoulder dislocation.

Fig 4. Brachial plexus neuritis. A 19-year-old man presented with acute onset of right shoulder pain without any history of trauma. A week later the pain subsided, but weakness was noted on movement of the right shoulder. On examination, atrophy and severe weakness of abduction of the right shoulder (2/5, Medical Research Council scale for muscle power) were noted. Coronal T2WI shows T2 hyperintense and minimally enlarged right-sided C5 and C6 nerve roots (long arrow) and upper trunks (short arrow), in keeping with acute neuritis. Acute denervation changes were seen in the deltoid and suprascapularis muscles (image not shown).

entiating a normal peripheral nerve from a diseased nerve (Fig 1).\(^{1,9}\) Intravenous administration of gadolinium is also helpful in detecting abnormal nerve and denervated muscle enhancement in patients suspected of various inflammatory, infectious, and neoplastic neuropathies.

Etiology of Diffuse Peripheral Nerve Lesions

Diffuse peripheral nerve lesions may be divided into traumatic, inflammatory, infectious, hereditary, radiation-induced, neoplastic, and tumor variants (Table 2).\(^{10–18}\) Relevant examples of cases from each category with their MRN imaging characteristics are illustrated (Table 3).

Traumatic Neuropathy

Direct trauma to the peripheral nerve may be caused by stretching or laceration during a traumatic event. Other mechanisms of injury include acute or chronic compression related to hematoma, bony fracture, and ischemia and, rarely, intramuscular injections.\(^{10}\) Superficial nerves and fascicles that lie...
Fig 5. CIDP. A 55-year-old woman presented with a 1-year history of progressive pain in the hands and feet and weakness on finger extension of both hands. On examination, sensory loss and severe weakness in the bilateral distal radial nerve distribution were demonstrated. A nerve-conduction study found severe asymmetric sensory-motor peripheral neuropathy with demyelination (multifocal partial motor conduction block and reduced velocities). A, Coronal contrast-enhanced T1WIFS of the right upper extremity shows an enlarged and enhancing radial nerve (ellipse). The radial, median, and ulnar nerves were enlarged and T2 hyperintense bilaterally with contrast enhancement (images not shown). The diagnosis of CIDP was proved by biopsy in this case. B, Axial STIR image of the right upper extremity shows enlarged and hyperintense radial (white circle), median (gray circle), and ulnar (black circle) nerves. The changes are most pronounced in the radial nerve.

Fig 6. CIDP. A 46-year-old man had an acute episode of pain 2 years before imaging. The pain subsided, but he developed bilateral distal arm paresthesia. Clinical examination at the time of imaging showed distal sensory loss and motor weakness in both arms. Coronal T1 FLAIR (A) and coronal STIR (B) images show T2 hyperintense and symmetrically enlarged brachial plexus nerve roots and trunks (ellipses). The diagnosis of CIDP was proved by biopsy in this case.

Fig 7. Infectious neuropathy (viral): Hansen disease (leprosy) in peripheral nerves. A 40-year-old man developed multiple small areas of lighter skin on the upper extremity 2 years before imaging. Subsequently, he developed weakness and decreased sensation on the arms and hands. T2WIFS (A) and a proton-attenuation axial (B) image show an enlarged hyperintense median nerve with prominent fascicles continuing into the carpal tunnel (arrows). The diagnosis of leprosy was proved by biopsy.
along the periphery of the nerve may be affected more severely in direct trauma, such as greater involvement of the peroneal component of the sciatic nerve by external trauma or compression. MRN may not only show the abnormal nerve (Fig 3), thereby confirming the clinical suspicion, but may also demonstrate the underlying causative lesion.

Inflammatory Neuritis
Peripheral nerves may be affected by a variety of inflammatory processes, and the clinically suspected pathologic nerves can be visualized in detail with high-resolution MRN imaging. Brachial plexus neuritis (plexitis) is a common occurrence. It is possibly immune-mediated and inflammatory, as is the form furste of acute GBS. The usual presentation is sudden onset of pain, followed by weakness and paraesthesia. There have been possible associations with viral or bacterial infection, vaccination, trauma, or surgery. This condition is self-

Fig 8. Infectious neuropathy (pyogenic). A 30-year-old man with a history of a long-standing right ischial decubitus ulcer and leg pain developed fever and chills with slow onset of sciatic neuropathy. MR imaging was performed for clinical suspicion of osteomyelitis. Axial T2WIFS shows an abnormally enlarged T2 hyperintense right sciatic nerve (thin arrow). Note the right ischial decubitus ulcer and osteomyelitis (thick arrow). The sciatic nerve demonstrated enhancement on postcontrast images (not shown).

Fig 9. CMT. A 49-year-old man with progressive left hand and arm weakness and numbness, which started approximately 5 years before the current imaging. Coronal T1 FLAIR (A) and coronal STIR (B) images show hyperintense and enlarged brachial plexus nerve roots and trunks with a preserved fascicular pattern (arrows). There is asymmetric involvement of the brachial plexus in this case, with the left side more severely affected than the right side.

Fig 10. Radiation neuropathy. A 69-year-old woman with a history of breast carcinoma and external beam radiation therapy 2 years before imaging presented with worsening right-arm sensory deficits. Coronal (A) and axial (B) 3D STIR SPACE images show a T2 hyperintense right-sided brachial plexus (arrow) with minimal linear enlargement in the area of the radiation field. No focal mass is visualized.

Fig 11. Neurofibromatosis with MPNST. A 57-year-old man with neurofibromatosis presented with persistent left-leg pain and weakness. A, Coronal STIR image shows enlarged lobulated masses (arrow), which represent MPNST in this case. B, Coronal contrast-enhanced T1WIFS shows a heterogeneously enhancing lobulated mass with areas of necrosis along the left femoral nerve, demonstrating MPNST.
limiting (4–6 weeks), and treatment is conservative. MRN plays a crucial role in its management because it may confirm the diagnosis of brachial plexitis (Fig 4) as well as distinguish it from cervical radiculopathy. Surgery is needed for the latter diagnosis. Brachial neuritis involves multiple nerves, unlike the fixed nerve distribution of cervical spondylosis and degenerative disk disease.

**CIDP**
CIDP is rare and is considered the chronic counterpart of GBS; the peak of illness in GBS is <4 weeks.\(^\text{12,21}\) The clinical features include a chronic course (>2 months); and proximal and distal muscle weakness with sensory involvement, areflexia, and albuminocytologic dissociation (Figs 5 and 6).

**Infectious Neuropathy**
Involvement of peripheral nerves in leprosy (Hansen disease, Fig 7) is well-known, and specific findings such as nerve calcifications in subacute-chronic phases may be seen.\(^\text{13,22}\) Pyogenic infections are rarer and may involve the nerve by direct extension (Fig 8).

**Hereditary Neuropathy**
CMT is a rare disease;\(^\text{14}\) it is also known as hereditary motor and sensory neuropathy. A positive family history may be found in as many as 80% of cases.\(^\text{23}\) The usual clinical presentation is chronic degeneration of peripheral nerves and roots, with muscle atrophy and sensory impairment in a distal distribution.\(^\text{15,24}\) Clinical findings include atrophy and sensory loss affecting all extremities, ataxia, areflexia, palpably enlarged peripheral nerves, pes cavus, and hammer toes. Bilateral lumbosacral plexus and peripheral nerve involvement are common. The enlarged nerves are frequently related to increased fatty interfascicular epineurium with demyelinating atrophic fascicles (Fig 9).

**Radiation Neuropathy**
There may be a long and variable interval (weeks to years) between radiation therapy and the onset of symptoms.\(^\text{16}\) It is essentially a diagnosis of exclusion. Neoplastic disease (primary or metastatic), infectious, and other etiologies should be ruled out first (Fig 10). The severity is dose-dependent, and radiation neuropathy is seen more commonly with doses of >60 Gy.\(^\text{25}\)
Neoplasia and Tumor Variants

A variety of tumors and tumor variants may involve the peripheral nerves. The role of MR imaging has been widely reported and is well-defined in neurofibromatosis (Figs 11 and 12). Multiple signs, such as a bag-of-worms appearance in plexiform neurofibroma, “split fat” sign on T1WI, and target and tail signs on T2WI, have been described.26,27 Other signs that may help to differentiate malignant from benign nerve sheath tumors are size > 5 cm, necrosis or heterogeneous enhancement, and invasion of surrounding structures.26 NL or lymphomatous infiltration of the peripheral nerves has been reported in ≤40% of patients dying of lymphoma.18 The typ-
ical presentation is progressive sensorimotor neuropathy, and the diagnosis is usually confirmed by a histopathologic examination (nerve biopsy or autopsy) demonstrating lymphomatous infiltration of nerves (Fig 13).

Perineuroma is a benign tumor of neoplastic perineural cells (Fig 14). It may present as mononeuropathy or plexopathy. Motor symptoms with extensive chronic muscle denervation changes are more commonly seen than sensory symptoms. 28,29 FLH is a rare benign lesion, which has distinctive features on MR imaging. 30,31 The median nerve is affected most commonly; however, it can also be seen at other sites (Fig 15). A variety of proteins may be deposited in nerves with systemic amyloidosis, resulting in severe progressive mixed autonomic dysfunction. 17,32 The MR imaging appearance of amyloidosis of nerves has also been described (Fig 16). 33

Summary

Diffuse abnormalities of peripheral nerves may be seen with numerous pathoetiologies. Currently, biopsy and pathologic examination of the nerves remain the only definitive ways of achieving a final diagnosis. However, a high-resolution MRN examination may be used to confirm the clinical suspicion and exclude other etiologies, such as a compressive mass lesion; in the correct clinical setting, it may aid in diagnosis and management of these lesions. Future research is needed to further validate the outlined differentiating MR imaging features of diffusely diseased peripheral nerves.

References