MRI characteristics of neuromyelitis optica spectrum disorder
An international update

Abstract
Since its initial reports in the 19th century, neuromyelitis optica (NMO) had been thought to involve only the optic nerves and spinal cord. However, the discovery of highly specific anti-aquaporin-4 antibody diagnostic biomarker for NMO enabled recognition of more diverse clinical spectrum of manifestations. Brain MRI abnormalities in patients seropositive for anti-aquaporin-4 antibody are common and some may be relatively unique by virtue of localization and configuration. Some seropositive patients present with brain involvement during their first attack and/or continue to relapse in the same location without optic nerve and spinal cord involvement. Thus, characteristics of brain abnormalities in such patients have become of increased interest. In this regard, MRI has an increasingly important role in the differential diagnosis of NMO and its spectrum disorder (NMOSD), particularly from multiple sclerosis. Differentiating these conditions is of prime importance because early initiation of effective immunosuppressive therapy is the key to preventing attack-related disability in NMOSD, whereas some disease-modifying drugs for multiple sclerosis may exacerbate the disease. Therefore, identifying the MRI features suggestive of NMOSD has diagnostic and prognostic implications. We herein review the brain, optic nerve, and spinal cord MRI findings of NMOSD. Neurology® 2015;84:1165–1173

Glossary
- AQP4 = aquaporin-4
- IgG = immunoglobulin G
- LETM = longitudinally extensive transverse myelitis
- MOG = myelin-oligodendrocyte glycoprotein
- MS = multiple sclerosis
- NMO = neuromyelitis optica
- NMOSD = neuromyelitis optica spectrum disorder
- ON = optic neuritis.

Neuromyelitis optica (NMO) is an inflammatory disease of the CNS that is characterized by severe attacks of optic neuritis (ON) and longitudinally extensive transverse myelitis (LETM). The past decade has witnessed dramatic advances in our understanding of NMO. Such advances were initiated by the discovery of the disease-specific autoantibody, NMO–immunoglobulin G (NMO-IgG), and subsequent identification of the main target autoantigen, aquaporin-4 (AQP4), which has distinguished NMO as a distinct disease from multiple sclerosis (MS).

Current diagnostic criteria, however, still require both ON and myelitis for an NMO diagnosis. Nevertheless, the identification of anti-AQP4 antibodies beyond the current diagnostic criteria of NMO indicates a broader clinical phenotype of this disorder, so-called “NMO spectrum disorder” (NMOSD). The NMOSD encompasses anti-AQP4 antibody seropositive patients with limited or inaugural forms of NMO and with specific brain abnormalities. It also includes anti-AQP4 antibody seropositive patients with other autoimmune disorders such as systemic lupus erythematosus, sarcoidosis, and other autoimmune diseases.

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lupus erythematosus and Sjögren syndrome. In this regard, MRI has an increasingly important role in differentiating NMOSD from other inflammatory disorders of the CNS, particularly from MS. Differentiating these conditions is critical because treatments are distinct. Furthermore, recent advanced MRI techniques are detecting additional specific markers and help elucidate the underlying mechanisms of tissue damage in NMOSD.

We herein summarize the MRI findings of NMOSD and discuss their diagnostic and prognostic implications.

**BRAIN MRI FINDINGS IN NMOSD** Since the early studies using brain MRI in NMO, unexplained clinically silent and nonspecific white matter abnormalities were found in some patients. With the advent of AQP4-IgG assays, it became clear that a high proportion of patients with NMOSD harbored brain MRI abnormalities, frequently located in areas associated with high AQP4 expression. However, brain abnormalities also occurred in areas where AQP4 expression is not particularly high. Although nonspecific small dots and patches of hyperintensity in subcortical and deep white matter on T2-weighted or fluid-attenuated inversion recovery sequences are the most common findings in NMOSD, certain lesions have a location or appearance characteristic for NMOSD.

Before the discovery of anti-AQP4 antibody, brain MRI abnormalities were reported in only 13% to 46% of patients with NMO. However, when excluding the brain MRI criteria, the incidence of brain MRI abnormalities increased to 50% to 85% using the revised 2006 NMO diagnostic criteria and to 51% to 89% in seropositive patients with NMOSD. Furthermore, brain MRI abnormalities at onset have been reported in 43% to 70% of patients with NMOSD. One of the explanations for discrepancies in frequency between studies may be that brain MRI abnormalities become more frequent with duration of disease. In a published series of 88 seropositive children, brain abnormalities were observed in 68% of the children with available MRI studies, and were predominantly located within periventricular regions of the third (diencephalic) and fourth ventricles (brainstem), supratentorial and infratentorial white matter, midbrain, and cerebellum. This is consistent with the observation that 45% to 55% of children with NMOSD show episodic cerebral symptoms, including ophthalmoparesis, intractable vomiting and hiccups, altered consciousness, severe behavioral changes, narcolepsy, ataxia, and seizures.

**Classification of brain MRI findings seen in NMOSD.**

- **Periependymal lesions surrounding the ventricular system.**
  - Diencephalic lesions surrounding the third ventricles and cerebral aqueduct. Diencephalic lesions surrounding the third ventricles and cerebral aqueduct, which include the thalamus, hypothalamus, and anterior border of the midbrain have been reported in NMOSD (figure 1A). These lesions frequently are asymptomatic, but some patients may present with a syndrome of inappropriate antidiuretic hormone secretion, narcolepsy, hypothermia, hypotension, hyperpnea, obesity, hypothyroidism, hyperprolactinemia, secondary amenorrhea, galactorrhea, and behavioral changes.

  Dorsal brainstem lesions adjacent to the fourth ventricle. One of the most specific brain MRI abnormalities in patients with NMOSD is a lesion in the dorsal brainstem adjacent to the fourth ventricle including the area postrema and the nucleus tracts solitarius. Such lesions are highly associated with intractable hiccups, nausea, and vomiting, and have been reported in 7% to 46% of patients with NMOSD. This area, the emetic reflex center, has a less restrictive blood-brain barrier, making it more accessible to AQP4-IgG attack. The MRI as well as clinical evidence support the notion that area postrema is an important point of attack in patients with NMOSD and further suggests that this area is a portal for entry of circulating IgG into the CNS. Pathologic abnormalities were noted in this region in 40% of patients with NMO, but there was no obvious neuronal, axonal, or myelin loss. Medullary lesions are often contiguous with cervical cord lesion, usually taking a linear shape (figure 1B). These lesions may be associated with the first symptoms of the disease or herald acute exacerbation. Various symptoms corresponding to a brainstem lesion may develop, such as nystagmus, dysarthria, dysphagia, ataxia, or ophthalmoplegia.

  Periependymal lesions surrounding the lateral ventricle. Lesions in the corpus callosum have been described in 12% to 40% of patients with NMOSD. Because both NMO and MS frequently have callosal lesions, location by itself is not a unique finding that differentiates NMOSD from MS. However, while the callosal lesions in MS are discrete, ovoid, and perpendicular to the ventricles and involve inferior aspects of the corpus callosum (figure 2A), NMOSD lesions are located immediately next to the lateral ventricles, following the ependymal lining (figure 1C). The acute callosal lesions in NMOSD are often edematous and heterogeneous, creating a “marbled pattern” and sometimes involving the complete thickness of splenium in a unique “arch bridge pattern” (figure 1, Cb and Cc). Sometime, the callosal lesions extend into the cerebral...
hemisphere, forming an extensive and confluent white matter lesion. In the chronic phase of NMOSD, the callosal lesions tend to reduce in size and intensity and may even disappear; however, cystic changes and atrophy of the corpus callosum have been described. Certain clinical symptoms, such as dysfunctions of cognition and motor coordination, may be attributed to callosal lesions, but they have not been well evaluated yet.

**Hemispheric white matter lesions.** Extensive and confluent hemispheric white matter lesions are often tumefactive (>3 cm in longest diameter) or have long spindle-like or radial-shape following white matter tracts (figure 1D). Mass effect is usually absent. Increased lesion diffusivity on apparent diffusion coefficient maps suggests vasogenic edema in association with acute inflammation (figure 1D.c), occasionally mimicking posterior reversible encephalopathy syndrome or Baló lesions. These extensive lesions have been found more frequently in anti-AQP4 antibody seropositive than seronegative patients. In the chronic phase, these large lesions tend to shrink and even disappear, but in some cases, cystic-like or cavitary changes are revealed (figure 1D.d). These lesions may cause various symptoms such as hemiparesis, encephalopathy, and visual field defects depending on the area they involve. Large confluent hemispheric white matter lesions are not uncommon in children with NMOSD. Tumefactive lesions with a surrounding zone of edema and variable mass effect may resemble acute disseminated encephalomyelitis or CNS malignancies.

**Lesions involving corticospinal tracts.** Lesions involving the corticospinal tracts can be unilateral or bilateral, and may extend from the deep white matter in the cerebral hemisphere through the posterior limb of the internal capsule to reach the cerebral peduncles of the midbrain or the pons (figure 1E). These lesions are contiguous and often longitudinally extensive, following the pyramidal tracts (figure 1E.c). Corticospinal tract lesions have been found in 23% to 44% in some cohorts of patients with NMOSD and have occasionally been reported in other cohorts. It is of interest that, unlike circumventricular areas, corticospinal tracts are not the areas where the AQP4 is highly expressed; it is unknown why these regions are also frequently involved in NMOSD.

**Nonspecific lesions:** Not unique, but most common. Non-specific punctate or small (<3 mm) dots or patches of hyperintensities on T2-weighted or fluid-attenuated inversion recovery sequences in the subcortical or deep white matter have been described most frequently on brain imaging studies of NMOSD (35%–84%) and are usually asymptomatic.

**Enhancing lesions.** Although the exact frequency is unclear, previous studies have described a variable...
percentage of gadolinium-enhancing brain lesions (9%–36%) in patients with NMOSD. Most of the enhancement was displayed in a poorly marginated, subtle, and multiple patchy pattern, a so-called “cloud-like” enhancement (figure 1F.a). These cloud-like enhanced lesions differ from the ovoid or open-ring gadolinium-enhancing lesions with well-defined borders that are more typical of MS (figure 2). A linear enhancement of the ependymal surface of the lateral ventricles (pencil-thin lesion) has also been described in NMOSD (figure 1F.b). Rarely, well-marginated nodular enhancement or meningeal enhancement has been reported in NMOSD (figure 1F.c).

**OPTIC NERVE MRI FINDINGS IN NMOSD**

MRI studies have reported nonspecific optic nerve sheath thickening, optic nerve hyperintensities on T2-weighted sequences, and gadolinium enhancement on T1-weighted sequences in acute ON of NMOSD. However, as similar findings also have been described in ON of MS, these findings are not considered diagnostic of NMOSD. Recent studies have looked at the differential MRI features of the optic nerve lesion between MS and NMOSD. A trend to more posterior involvement of the optic nerve including chiasm, and simultaneous bilateral disease, has been observed in NMOSD (figure 3). Thus, long-segment inflammation of the optic nerve, particularly when simultaneous bilateral and extending posteriorly into the optic chiasm, should lead us to suspect the diagnosis of NMOSD in the appropriate clinical context.

**SPINAL CORD MRI FINDINGS IN NMOSD**

The inflammatory process of NMOSD in spinal cord MRI is characterized by hyperintensity on T2-weighted sequences and by hypointensity on T1-weighted sequences. These abnormalities in the spinal cord MRI have been reported to be, in general, more frequently present in the cervical and the upper thoracic spinal cord segments than the lower thoracic and lumbar regions with a preferential involvement in the central gray matter. In the spinal cord, AQP4 is abundant in the gray matter and in glial cell processes adjacent to the ependymal cells of the central canal and to a lesser degree in the white matter of the spinal cord.

The most distinct manifestation of NMO is LETM, defined as a lesion that spans over 3 or more contiguous vertebral segments and predominantly involves central gray matter on the spinal cord MRI (figure 4). However, not all LETM is NMOSD and several studies of patients with LETM have observed significant differences in demographic and clinical features between anti-AQP4 antibody positive compared with negative patients with LETM. LETM seems to be less specific for NMO in children than in adults. LETM is frequently observed in children with acute disseminated encephalomyelitis, but also in 17% of those with MS, and in 67% to 88% of children with monophasic transverse myelitis. Therefore, it is important to bear in mind that numerous other differential diagnoses than NMOSD need to be considered when a patient presents with LETM.

Spinal cord lesions during follow-up of NMOSD. MRI changes of LETM have been observed over the course of NMOSD and MRI data indicate that LETM lesions may evolve into multiple shorter lesions during remission or after treatment with high-dose steroids. In addition, spinal cord atrophy as a consequence of recurrent myelitis has been reported and may correlate with neurologic disability. Consequently, the timing of MRI may be important for the demonstration of LETM.

**COMPARING THE IMAGING OF NMOSD WITH MS**

In clinical practice, the main differential diagnosis of NMO is MS, particularly disease limited to the optic nerves and spinal cord. Differentiating these conditions is of prime importance because of differences in prognosis and therapy, as some MS therapies can exacerbate NMO.
improve the methods and analysis by which to distinguish these conditions to facilitate early and accurate diagnosis. Contrasting features between the 2 conditions may further improve our understanding of the different pathogenic processes.

Whereas it is possible to select patients with NMOSD using the specific marker (serum anti-AQP4 antibodies), there is no corresponding specific biomarker for MS. Studies contrasting NMO and MS have often used different selection criteria, particularly whether they have restricted the NMO inclusion criteria to patients positive for anti-AQP4 antibody or not, and this may influence the results. Conflicting data may also be partly explained by the use of various assays for anti-AQP4 antibodies, which differ in sensitivity and are confounded by differences in the duration of follow-up.

As previously described, the most important imaging hallmark of NMO is the LETM, but a few patients may have centrally located short myelitis. Other MRI features of the spinal cord lesion that appear to differ between NMOSD and MS are summarized in the table.

The 2006 NMO diagnostic criteria include a brain MRI that is nondiagnostic for MS (using the Paty criteria) at onset as support for NMO. However, it is now known that MS-like lesions may appear in 10% to 12.5% of cases, and 5% to 42% of patients with NMO fulfill the Barkhof criteria. A recent report showed that 13% and 9% of patients with NMOSD, respectively, met Barkhof and the European Magnetic Imaging in MS diagnostic criteria for MS on brain MRI at onset. Lesion probability maps have not found statistically significant lesion locations in patients positive for anti-AQP4 antibody over those with MS. However, distinguishing features were identified on MS brain MRI that were sensitive and specific, such as the presence of a lateral ventricle and inferior temporal lobe lesion, Dawson fingers, or an S-shaped U-fiber lesion, to classify the patient as MS. Imaging sensitive to cortical lesions has revealed their absence in NMO (excluding one Japanese study of NMO pathology), whereas they are seen in the majority of patients with MS. Characteristic MS brain lesions surround central venule in >80% on high-strength MRI. In NMO lesions, this is less frequent, reported in 9% to 35% of cases and likely indicates the different pathogenic mechanisms of the disease.

The frequency of silent lesion formation appears to differ between the 2 diseases. Patients with NMOSD are less likely to develop clinically silent MRI lesions than patients with MS. However, new silent MRI lesions do occur in a small proportion of patients with NMOSD. In addition, most studies show that nonlesional tissue damage as measured on nonconventional imaging such as diffusion tensor imaging is well recognized in MS and may not occur in NMO except in the connecting tracts up and downstream of lesions. Collectively, these findings support the clinical observation that NMO, in contrast to MS, may be a lesion-dependent disease that produces relapses without more generalized neurodegenerative pathology, and hence the lack of a progressive phase.

The differences noted between NMO and MS may relate to the CNS-specific antibody-mediated pathology against astrocytes rather than a T-cell– predominant inflammatory condition targeting myelin. In support of this possibility, a marker of astrocytic function, myo-inositol was reduced in
cervical cord lesions of patients positive for anti-AQP4 antibody, but not in patients with MS. In contrast, \(N\)-acetyl-aspartate, a marker of myelin- and neurofilament-specific injury, was significantly reduced in patients with MS compared with controls and nonsignificantly reduced in the patients positive for anti-AQP4 antibody.55

Important comparisons between NMOSD and MS scans are summarized in the table. Because long-term systematic imaging studies in NMO have not yet been performed, the reported cross-sectional differences compared with MS require further confirmation. Developing algorithms using the brain criteria described by Matthews et al.6 in combination with spinal cord and optic nerve imaging features and possibly nonconventional imaging may further improve the sensitivity and specificity.

**PROGNOSTIC IMPLICATION OF MRI ABNORMALITIES** Anti-AQP4 antibody positivity is established as a prognostic marker, and its positivity indicates a high risk of further relapses of ON and myelitis.56,57 Because of the presence of many imaging features suggesting severe damage of spinal cord, such as T1 hypointensity with edema or cavitation and atrophy, patients with NMOSD are more likely to have a poor recovery, refractory pain,28 and a high risk of permanent disability. In addition, patients with NMOSD who have lesions in the upper cervical region extending to the brainstem may be at risk of respiratory failure.

High levels of glial fibrillary acidic protein in the CSF of patients with NMOSD during acute attacks correlated with length of MRI spinal cord lesion and Expanded Disability Status Scale score 6 months after those attacks. This correlation suggests that imaging findings may be proportional to the amount of astrocyte damage and have potential prognostic implications.e29 The presence of extensive brain lesions might predict a higher rate of relapse and increased disability at follow-up.560 Longitudinal follow-up studies are required to confirm whether patients with brain lesions have a worse prognosis than those without brain lesions. At this point, there are no individual MRI parameters that can predict the prognosis of NMO.

More recently, antibodies against myelin-oligodendrocyte glycoprotein (MOG) have been found in some patients with clinical features of NMOSD, but who lack anti-AQP4 antibodies.59 Patients exhibiting the anti-MOG–positive and anti-AQP4–negative serotype have been suggested to have fewer attacks, bilateral ON, more caudal myelitis, and recover better than patients with anti-AQP4 antibodies and those who are seronegative for both antibodies.59,60 Therefore, patients presenting with an NMOSD phenotype with anti-MOG antibodies may have a distinct underlying disease mechanism presumably with a better prognosis than those with anti-AQP4 antibodies, although this needs to be confirmed by further studies.

**OUTLOOK: MRI FINDINGS IN THE CONTEXT OF NMO DIAGNOSTIC CRITERIA** The notion that brain MRI abnormalities are frequent in patients with NMOSD refutes the older doctrine that a normal brain MRI is a prerequisite for a diagnosis of NMO. Herein, we have reviewed the advances in our knowledge on the spectrum of imaging findings in NMOSD. However, sensitivity and specificity of these imaging features for NMOSD have not been systematically investigated in a prospective manner.
and none of the findings can be considered pathognomonic or evidentiary for NMOSD. Therefore, as with other inflammatory CNS conditions, imaging findings should prompt broad differential diagnostic consideration—a topic that is beyond the scope of this review. The actual utility of lesion probability maps to distinguish NMO and MS is limited by an unclear definition of some traditional criteria for MS—suggestive findings, such as “Dawson fingers.” The picture is further complicated by recent observations of patients with anti-MOG antibodies that some, but not all, have considered part of the NMO spectrum. Some commonalities but also differences in clinical presentation, epidemiology, and imaging have been reported between these 2 conditions, suggesting that NMOSD may not be a homogeneous nosologic entity. In addition, because NMOSD can coexist with other autoimmune diseases and antibodies to other CNS antigens such as anti-NMDA receptor antibodies may be present in patients seropositive for anti-AQP4 antibody, it is possible that autoimmunity against multiple CNS autoantigens may participate in the formation of inflammatory lesions of NMOSD. The emerging heterogeneity of NMOSD is mirrored by the broad range of neuroimaging findings summarized in this article. Areas for improved imaging that may facilitate more specific diagnostic, prognostic, therapeutic efficacy or other patient care benefit include higher-resolution imaging methods, 3-dimensional imaging of site-specific lesions, and potential computationally guided analysis of images for quantitative comparisons. International collaborative efforts are now under way that will permit accrual of sufficiently large, carefully characterized NMO/NMOSD patients to better understand the frequency of brain involvement and to more thoroughly appreciate the implications of MRI abnormalities in clinical diagnosis and prognosis.

AUTHOR CONTRIBUTIONS

Dr. H.J. Kim conceived and designed the work, analyzed the literature, wrote the manuscript, critically reviewed and approved the final manuscript. Dr. F. Paul conceived the work, analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript. Dr. N. Asgari analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript. Dr. S.-H. Kim analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript. Dr. N. Asgari analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript. Dr. M.A. Lana-Peixoto analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript. Dr. A. Saiz critically reviewed and revised the manuscript, and approved the final manuscript. Dr. P. Villoslada critically reviewed and revised the manuscript, and approved the final manuscript. Dr. A. Sait critically reviewed and revised the manuscript, and approved the final manuscript. Dr. K. Fujihara critically reviewed and revised the manuscript, and approved the final manuscript. Dr. S.-H. Kim analyzed the literature, wrote the manuscript, critically reviewed and revised the manuscript, and approved the final manuscript.

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