Radiology update in neuro-ophthalmology
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Introduction
Imaging techniques for the evaluation of neuro-ophthalmologic disorders include both conventional and specialized types of studies (Table 1). MRI is often the imaging test of choice, but computed tomography has advantages in certain situations, such as in detecting nerve head buried drusen, in the evaluation for orbital decompression in thyroid eye disease, and in detecting bony fracture after orbital trauma or traumatic optic neuropathy.

The language of MRI can be confusing, especially with the variety of possible sequences, but indications for some of these sequences are important to know (Table 2). Intensity is the term used to describe brightness on MRI. T1-weighted images are hyperintense when they reflect water molecules bound to fat or cell membranes, but are hypointense when they reflect free water or tightly bound water (e.g. muscle). T2-weighted images are hyperintense when they represent free water such as cerebrospinal fluid (CSF), but become progressively more hypointense as the water becomes more tightly bound. For more detail, an excellent explanation of the physics of MRI from an ophthalmologist’s perspective was recently published as a monograph supported by the American Academy of Ophthalmology [1].

Optic neuritis and multiple sclerosis
Typical MRI features of multiple sclerosis (MS) include disseminated periventricular, callosal, and subcortical white matter T2 signal hyperintensities (Fig. 1). The McDonald MRI criteria for disease dissemination in space were based on standard Tesla (1.5 T) field strength. Higher field strength MRI magnets (e.g. 3 T or greater) are becoming increasingly used. These higher-field MRIs may influence the diagnosis of MS because they detect more lesions and these lesions are more conspicuous with higher field strength MRI [2]. In contrast, higher field strength magnets can also decrease T1 signal contrast. If available, ophthalmologists can request a 3 T MRI for patients with optic neuritis at risk for MS, but it is not mandatory that these higher strengths be used.

Other nonconventional MRI techniques are becoming more valuable for imaging MS and optic neuritis. Diffusion tensor MRI (DTMRI) is one such modality. DTMRI applies the direction of water diffusion through tissues to map out neural pathways in the brain, such as white matter tracts. This situation allows imaging based on functional systems (such as the visual system), and can provide insights into rostral changes in the visual neuraxis after damage to first order neurons. For example, DTMRI performed 1 year after optic neuritis showed changes downstream in the optic radiations, not only in the affected optic nerve [3].

Another MRI technique called magnetization transfer ratio (MTR) can be used to study the central nervous system. MTR images, unlike standard MRI, reflect not only protons on water, but also protons on macromolecules, such as brain tissue. MTR may better reflect the structural
integrity of tissue. MTR was studied in optic neuritis, using optical coherence tomography (OCT) and multifocal visual evoked potential (mfVEP) amplitude to differentiate optic nerves with and without axonal loss. When there was MTR reduction in the optic nerve, the OCT retinal nerve fiber layer thickness was decreased and the mfVEP showed decreased amplitude, suggesting that MTR reduction reflects axonal damage rather than demyelination [4]. Standard MRI techniques do not always differentiate which T2 MS lesions represent pure demyelination and which reflect axonal loss.

Although DTMRI and MTR hold promise for future study in optic neuritis and MS, ophthalmologists will generally use standard MRI sequences. For the patient with a clinically isolated demyelinating syndrome (CIS) such as optic neuritis, the presence and number of disseminated brain lesions on T2-weighted MRI are the most important factors to predict future risk of demyelinating attacks. In contrast, neither retinal nerve fiber layer thinning of the unaffected eye on OCT nor a decrease in normalized brain volume on MRI were present in patients after a first attack of optic neuritis [5], suggesting that the first demyelinating event may be too early to find evidence of diffuse axonal degeneration.

The location of MRI lesions in patients with CIS is also important, as having even one brainstem lesion is predictive of an increased risk of conversion to MS and a risk of higher disability compared with patients without brainstem lesions [6]. This finding holds true mainly when there are at least nine lesions. For patients with ophthalmoplegia and CIS who have a visible demyelinating brainstem lesion on MRI, evidence suggests that the ophthalmoplegia improves, and usually resolves, independent of progression to MS [7].

### Neuromyelitis optica

Neuromyelitis optica (NMO) is characterized by unilateral or bilateral optic neuritis and longitudinally extensive transverse myelopathy. Figure 2 shows bilateral optic nerve and chiasm involvement in NMO (Fig. 2). It is usually associated with antibodies to the aquaporin-4 water channel, the NMO antibody. NMO was originally considered as being restricted to the optic nerves and spinal cord, but radiographic brain lesions are now more recognized as the spectrum of NMO expands. In fact, one study showed that 10% of MRIs from NMO patients had lesions similar in appearance to those of patients with MS [8].

The aquaporin-4 antibody is specific for NMO. A retrospective study in patients with NMO AB positivity showed 79% had an abnormal MRI, 45% of which were symptomatic, and approximately 15% of which enhanced. These lesions were usually nonspecific, but often involve the corticospinal tracts, either in the internal capsule or cerebral peduncle. These structures do not usually show direct damage in MS. MRI lesions in NMO can surround the ventricles. Although periventricular involvement is common in MS, the periventricular involvement in NMO is distinct, appearing as an extensive lesion involving the entire thickness of the corpus callosum, in contrast to the Dawson’s fingers of MS [9]. Extensive callosal atrophy was similarly reported in a patient with NMO of at least 18 years’ duration [10]. Extensive callosal atrophy was similarly reported in a patient with NMO of at least 18 years’ duration [10]. MRI lesions in the area postrema, at the floor of the forth ventricle, may also be characteristic of NMO, but not MS [11]. Lesions of the area postrema result in nausea, vomiting, and hiccups, which occur in NMO, but are very rare in MS.

### Key points

- Both conventional and nonconventional imaging techniques are used in neuro-ophthalmologic disorders.
- MRI is the most commonly used imaging modality in neuro-ophthalmology.
- Particular MRI sequences each have their own advantages in discrimination of different types of lesions.

| Imaging studies used in neuro-ophthalmology | Advantages | Limitations |
|--------------------------------------------|------------|-------------|
| MRI | High resolution | Contraindicated with pacemaker and certain metallic implants |
| CT | Visualization of bony anatomy | Insensitive for some types of lesions compared to MRI |
| fMRI | Areas normal on MRI may show metabolic abnormalities not found on fMRI | Mainly used as a research tool |
| MRS | Measures biochemical changes in tissue | Results of spectra are nonspecific and must be correlated with MRI appearance |
| MTR | May better reflect the structural integrity of tissues | Limited clinical use at this time |
| Diffusion Tensor MRI | Mapping out functional white matter tracts | Less helpful for grey matter or nuclear structures |
| PET | Areas of high or low metabolism can be located anywhere in the body | Expensive |

CT, computed tomography; fMRI, functional MRI; MRS, magnetic resonance spectroscopy; MTR, magnetisation transfer ratio.
Magnetic resonance spectroscopy (MRS) is a magnetic resonance technique that measures different brain metabolites, including N-acetylaspartate, lactate, choline, and creatine, depending on the tissue being studied. These metabolites increase or decrease from expected norms based on tissue integrity. For example, lactate is elevated in tissue necrosis. MRS was recently studied in NMO. Normal appearing grey and white matter had normal MRS markers in NMO. This finding contrasts to the abnormal MRS values found in MS and marks MRS as a possible future differentiating diagnostic test for NMO against MS [12/C15].

Functional MRI (fMRI) is another nonconventional radiographic test, which measures neural activity by measuring changes in cerebral blood flow. Resting state fMRI measurements were different in NMO versus normal controls [13]. This difference in functional activity may explain why NMO patients have more cognitive problems than people without NMO [14].

Other optic neuropathies
Imaging can detect optic nerve abnormalities in many different neuro-ophthalmologic conditions. An example of typical inflammatory optic nerve enhancement is seen in Fig. 3. Some familiar optic nerve disorders may demonstrate unusual radiographic features on conventional MRI. For example, neuroretinitis can affect the optic pathway beyond the optic nerve head, and both intraocular optic nerve and optic nerve sheath enhancement (optic perineuritis) have been reported [15/C15]. Similarly, although optic nerve enhancement on MRI

| Table 2 Conventional MRI protocols |
|-----------------------------------|
| Principle of use | Examples |
| T1 weighted | Good quality anatomic detail. Bright T1 signal occurs from proteinous tissue, fat, calcification and marrow signal | Evaluation of midline structure anatomy; visualizing 'black holes' from chronic demyelination |
| T2 weighted | Water is bright on T2. Better to identify certain lesions compared with T1, especially those in the posterior fossa | Demyelinating infratentorial MS plaques are hyperintense |
| FLAIR | T2 with nullification of CSF signal, which increases conspicuity of periventricular lesions | Demyelinating supratentorial MS plaques are hyperintense |
| Gadolinium-enhanced | Evaluates areas of breakdown of the blood–brain barrier | Enhancing lesions include tumor and abscess. The ring of enhancement from demyelination is often incomplete |
| DWI | Identifies restricted diffusion | Most commonly used to identify acute infarct. Restricted diffusion can also occur in abscesses, densely cellular tumors and epidermoids |
| GRE | Keenly sensitive in visualizing hemosiderin deposition related to hemorrhage | Old hemorrhage related to hypertension, cerebral amyloid angiopathy and cavernous malformations may appear hypointense |
| Fat-sat | Suppresses fat signal to better visualize enhancement of orbital structures such as the optic nerve | Orbital MRI for evaluation of optic neuritis should use fat-sat |

CSF, cerebrospinal fluid; DWI, Diffusion weighted; FLAIR, fluid attenuated inversion recovery; GRE, gradient echo; MS, multiple sclerosis.

Figure 1 Multiple sclerosis

Typical MRI features of multiple sclerosis. Clockwise from the bottom: sagittal FLAIR MRI showing periventricular ovoid 'Dawson's fingers' plaques; T1 axial black holes; T1 postgadolinium areas of enhancement; sagittal cervical spine MRI showing nonlongitudinally extensive demyelinating lesions.

Figure 2 Neuromyelitis optica

Arrow on the left image shows a normal nonenhancing chiasm on T1 postgadolinium orbital fat-sat MRI. Arrow on the right image shows contrast enhancement to the optic nerves and chiasm of a patient with neuromyelitis optica.
usually denotes an inflammatory optic neuropathy, contrast enhancement was also seen in a case of posterior ischemic optic neuropathy due to internal carotid dissection [16].

Using fMRI to study cortical reorganization after nonarteritic anterior ischemic optic neuropathy (AION), a correlation was found between visual acuity and activation in the occipital visual areas, suggesting that afferent nerve pathway integrity is related to primary visual cortical activation. Over time, there is greater activity in the occipital regions when the non-AION eye was stimulated, which speaks to the plasticity of the rostral visual systems after an optic nerve injury. fMRI also showed impaired activation in visual pathways after Leber's hereditary optic neuropathy (LHON). Interestingly, radiographic abnormalities in LHON were not restricted to the visual cortex, as the auditory network was also affected [17].

DTMRI may assist in diagnosis of septo-optic dysplasia. Standard T1- and T2-weighted MRI are adequate to show hypoplasia of the optic nerves and abnormalities in midline brain structures. However, by measuring the entire visual tract pathway, DTMRI at higher tesla has the advantage of providing a more comprehensive measurement of the entire visual pathway, and may also allow for earlier radiographic diagnosis of septo-optic dysplasia [18].

Ophthalmoplegic migraine is one example of an isolated ocular motor neuropathy, which may have MRI changes, and because it often affects young people, will often be imaged. In these cases, the third nerve may enhance and enlarge during symptomatic periods, with radiographic resolution during periods of remission [22].

Trochlear nerve palsy can be congenital or acquired, unilateral or bilateral. Because the trochlear nerve is difficult to visualize [19], it has been historically challenging to confirm that congenital superior oblique palsy is a hypoplastic/aplastic cranial neuropathy, as is the case in congenital third nerve palsy or Duane’s syndrome. By using a higher field 3 Tesla MRI, patients with congenital superior oblique palsy (SOP) were indeed found to lack a trochlear nerve on the affected side, whereas visualization of the nerve was present on the contralateral, unaffected side [23]. This finding suggests the classification of SOP as a dysinnervation disorder. On MRI, the superior oblique muscle is also usually aplastic or hypoplastic in congenital SOP. Controls have almost complete symmetry of the two muscles on MRI, whereas a wide range of ratios of the thickness of the paretic side

The ocular motor system

Ferreira et al. [19] produced a succinct yet thorough review of the anatomy, pathology and normal MRI characteristics of the ocular motor nerves. The brainstem, cisternal, interdural, cavernous, and intraorbital segments are each considered separately. Among the pearls in this review, it is mentioned that the ocular motor nerve nuclei and fascicles cannot be directly visualized on MRI, and their intracavernous segments are prone to artifact.

To better detect cranial nerve abnormalities in the cavernous sinus, an MRI technique called contrast enhanced three-dimensional constructive interference in steady state (CISS) can be used. With superior contrast and resolution, CISS MRI can show smaller structures within the CSF, which makes it ideal for study of cranial neuropathies in the cavernous sinus [20]. CISS is not usually included with standard MRI acquisition series, however.

Which ocular motor cases need to be imaged? Savino’s group reviewed 93 patients who were older than 50 with no history of cancer and an isolated CN III, IV or VI palsy, and found that only one case had a lesion on MRI to explain the defect [21]. The cost and testing exposure to patients does not seem to justify imaging in all patients with ocular motor mononeuropathy. However, patients still need to be imaged if the palsy is progressive or persistent.

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to contralateral superior oblique muscle was found in patients with congenital SOP, from complete aplasia to equal thicknesses [24\(^\ast\)]. This variability is also found in familial forms of superior oblique hypoplasia [25\(^\ast\)].

Fig. 4 shows a cross-section of the ocular motor muscles in the orbit. The presence of supernumerary extraocular muscles (EOMs) has been documented both in cadavers and in vivo, but is difficult to confirm with imaging. When accessory orbital structures are seen on MRI if they are indeed muscles, they should be isointense to the EOMs. Because Grave’s orbitopathy (GO) increases the size of these accessory muscles, they may become more obvious on imaging during this clinical state. Because they act as other muscles do, they may enlarge in GO and decrease after treatment [26\(^\ast\)]. Another study concluded that approximately 2\% of humans have deep orbital bands consistent with supernumerary EOMs, which are sometimes significant contributors to restrictive strabismus. Thus, MRI may be helpful in guiding operative management of strabismus [27\(^\ast\)].

**Pseudotumor cerebri**

Imaging of the venous sinuses in pseudotumor cerebri is discussed elsewhere in this edition of *Current Opinion in Ophthalmology*. The radiologic findings of pseudotumor cerebri (idiopathic intracranial hypertension; IIH) in fact require that brain imaging does not show another cause of increased intracranial pressure. That being said, there are some radiologic features of IIH, which deserve mention. The commonly described ‘slit-like ventricles’ are not actually helpful in diagnosing IIH [28]. Figure 5 illustrates certain features of IIH that are supportive of the diagnosis, which include an enlarged optic nerve sheath, optic nerve tortuosity, protrusion of the optic nerve head, and concavity or flattening of the posterior globes [29]. Two additional MRI features of IIH have now been described: a narrowing of Meckel’s cave and a reduction in the diameter of the cavernous sinuses. Narrowing of Meckel’s (the trigeminal) cave, which normally appears as CSF signal lateral to the posterior cavernous sinus and clivus on axial sections, is an especially sensitive MRI marker of IIH [30\(^\ast\)].

**Posterior reversible encephalopathy syndrome**

Posterior reversible encephalopathy syndrome (PRES) may present to the ophthalmologist as cortical blindness, often associated with mental status changes or seizures. It occurs mainly in the setting of eclampsia, hypertension, chemotherapeutic agents, or sepsis, as a vasculopathy with characteristic MRI abnormalities of bilateral cortical vasogenic edema (Fig. 6). Because the diseases associated with PRES also can cause coagulopathy, it is rational that one report found the incidence of subarachnoid and intraparenchymal hemorrhage in PRES nears 20\%. The subarachnoid blood appears within the sulci, with sparing of the basal cisterns. This distribution is not found commonly in aneurysmal rupture, and extensive workup for aneurysm in patients with PRES with this pattern of hemorrhage is probably unnecessary [31\(^\ast\)].

**Migraine**

PET scanning is another type of functional imaging, which uses radiolabeling to measure regional blood flow. PET is used more often in the clinical setting because the whole body can be imaged for hypermetabolic or hypometabolic activity. Single-photon emission computed tomography is similar to PET, although it
measures glucose metabolism rather than blood flow, and tissue resolution is not as favorable [11]. PET scanning was used to measure cortical activity as a response to light stimulation in migraineurs. PET imaging found more cortical activation after a light stimulus during migraine headaches compared with between migraine attacks [32]. This response on PET suggests that the photophobia of migraine may be linked to impaired sensory modulation and cortical hyperexcitability.

**Conclusion**

Although practicing clinicians are not expected to possess the knowledge of a radiologist with regards to imaging modalities used in neuro-ophthalmologic conditions, it is important that they remain up to date on both conventional and nonconventional radiographic techniques.

Critical review of the limits and advantages of imaging allows the ophthalmologist to make informed choices about when to order imaging, which type of scan to order, and how to interpret the results.

**Acknowledgements**

**Conflicts of interest**

There are no conflicts of interest.

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**References and recommended reading**

Papers of particular interest, published within the annual period of review, have been highlighted as:
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Additional references related to this topic can also be found in the Current World Literature section in this issue (pp. 523–524).

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