Magnetic resonance imaging of optic nerve

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Abstract

Optic nerves are the second pair of cranial nerves and are unique as they represent an extension of the central nervous system. Apart from clinical and ophthalmoscopic evaluation, imaging, especially magnetic resonance imaging (MRI), plays an important role in the complete evaluation of optic nerve and the entire visual pathway. In this pictorial essay, the authors describe segmental anatomy of the optic nerve and review the imaging findings of various conditions affecting the optic nerves. MRI allows excellent depiction of the intricate anatomy of optic nerves due to its excellent soft tissue contrast without exposure to ionizing radiation, better delineation of the entire visual pathway, and accurate evaluation of associated intracranial pathologies.

Key words: Magnetic resonance imaging; optic nerve; visual pathway

Introduction

Optic nerves are the second pair of cranial nerves and are unique as they represent an extension of the central nervous system and, hence, are myelinated by oligodendrocytes.[1] They are the connection between eyes and brain, i.e. they relay visual impulses from retina to brain. Hence, most often, optic nerve abnormalities are associated with brain abnormalities.

This article describes the various common pathologies affecting optic nerves, such as demyelinating and inflammatory processes to less common optic nerve tumors and very rare congenital optic nerve abnormalities. Magnetic resonance imaging (MRI) is extremely useful in promptly identifying some of these pathologies to avoid permanent visual loss. In some cases, especially hereditary and congenital pathologies, unnecessary additional work-up can be avoided as it helps in better counseling of patients and their families.

Anatomy of Optic Nerve

The first sensory bipolar cell body is located in the inner nuclear layer of retina. The central processes of bipolar cells synapse on ganglion cells in the ganglion cell layer of retina, and the central processes of the ganglion cells form the optic nerve proper. The axons leave the globe at optic disc to form the optic nerves, which then relay in lateral geniculate bodies.

Optic nerve is approximately 50 mm long and is divided into four segments [Figure 1].[2] These are:

- Intraocular (1 mm) - when it emerges through the scleral opening
- Intraorbital (25 mm) - the longest segment and communication between subarachnoid space around the optic nerve with that in suprasellar cistern
- Intracanalicular (9 mm) - as it passes through bony optic canal along with ophthalmic artery (OA)
- Prechiasmatic (16 mm) - intracranial segment in suprasellar cistern.

It joins the contralateral optic nerve to form optic chiasma, where the nasal fibers from each optic nerve decussate and temporal fibers do not decussate.
Optic chiasma lies typically about 10 mm above the pituitary gland, separated by the suprasellar cistern. In about 80% of the population, it rests directly above the sella. However, it may rest above tuberculum sellae in 10% (prefixed chiasm) and above dorsum sellae in the remaining 10% of the population (postfixed chiasm). The prefixed chiasm has short optic nerves and long optic tracts, whereas the postfixed chiasm has long optic nerves and short optic tracts.\(^3\) Pituitary stalk angle is 90° or more in prefixed optic chiasm and acute angle for normal or postfixed chiasm.\(^4\)

From the optic chiasma, optic tracts course posterolaterally along the cerebral peduncles to synapse at lateral geniculate bodies. From the lateral geniculate nuclei, optic radiations fan out as optic radiations and reach the primary visual cortex in the occipital lobes.

**MRI Protocol**

MRI is the imaging modality of choice for evaluation of optic nerve pathway. The examinations were performed on 3 T MRI system (Achieva; Philips Medical Systems, Best, the Netherlands or HD.xt TwinSpeed; GE Healthcare, Milwaukee, Wisconsin). Head coil is commonly used; however, surface coil, if applied, improves the signal to noise ratio (SNR) of the globe.\(^5\) Imaging at 3 T MRI scanners offers superior SNR as well as spacial resolution to evaluate orbital structures as well as intracranial pathologies, as compared with 1.5 T scanners. Thin-section high-resolution spin-echo T2-weighted images at 3 T depict the optic nerves and orbital anatomy much better than 1.5 T scanner.\(^6\)

Thin coronal and axial T1 and short tau inversion recovery (STIR)/T2 fat-saturated sequences, as well as sagittal T2 fat-saturated sequences are obtained for optic nerve evaluation. A section thickness of 3-4 mm is preferred with an interslice gap of 0-1 mm. The entire examination takes about 30 min, and the patient is asked to refrain from eye movement during the scan.

### Pathologies

The etiological spectrum ranges from congenital, demyelination, inflammatory, neoplastic, and ischemic to miscellaneous pathologies [Table 1]. Pathologies based on onset of symptoms are also described [Table 2].
Congenital and Hereditary Optic Neuropathy

Optic nerve aplasia
It includes complete lack of the optic nerve, optic disc, retinal ganglion and nerve fiber layer, and optic nerve vessels. It can be unilateral or bilateral [Figure 2]. Other ocular abnormalities like micro-ophtalmia, cataract, iris hypoplasia, anterior coloboma, persistent primary hyperplastic vitreous, cranial and systemic abnormalities are often associated.[7,8]

Mutations in PAX6 (paired box 6) gene (11p13) have been associated with optic nerve malformations, including optic nerve aplasia/hypoplasia, coloboma, morning glory disc anomaly, aniridia, and persistent hyperplastic primary vitreous. PAX6 is involved in ocular morphogenesis as well as in the development of Rathke’s pouch and early anterior pituitary gland.[9]

Optic nerve colobomas
These are characterized by focal defect in the posterior globe at the optic nerve head insertion [Figures 3 and 4]. The defect can be of variable size - typically small, less than the size of the optic nerve head. It is caused by incomplete closure of the embryonic fissure and may be associated with microphthalmia and retrobulbar colobomatous cyst. Many brain abnormalities like gyration abnormalities, lateral ventricular dilatation, dilatation of the Virchow-Robin and subarachnoid spaces, white matter signal and corpus callosal abnormalities are associated.[10]

Morning glory syndrome is an optic disc anomaly characterized by three primary features - excavated optic disc with central glial tuft, the defect is surrounded by elevated rind of peripapillary pigment, and blood vessels traversing radially from the disc margin. It is named for its fundoscopic resemblance to morning glory flower.[8]

It can be isolated or associated with wide spectrum of congenital abnormalities like anterior midline craniofacial defects, pituitary abnormalities, corpus callosum agenesis/dysgenesis, basal encephalocele, and vascular abnormalities including Moyamoya syndrome.[11]

Intraorbital MRI findings include funnel-shaped morphology of the optic nerve head, elevated adjacent retinal surface showing T1-hyperintense signal corresponding to peripapillary pigment, discontinuity of T2-hypointense line, and enhancement of posterior chorioid-lamina cribrosa, abnormal tissue associated with the distal intraorbital segment of the optic nerve causing effacement of the subarachnoid spaces of the distal optic nerve sheath. Other findings include distal optic nerve enhancement, fatty infiltration of distal optic nerve sheath, small size of globe, asymmetry in optic nerve and chiasma,
and optic glioma. Ocular associations include retinal detachment, congenital cataract, persistent hyperplastic primary vitreous, drusen, and eyelid hemangioma.

Intracranial findings include a range of vascular abnormalities from hypoplasia/segmental agenesis of vessels to progressive vasculopathy with Moyamoya syndrome. Basal encephaloceles including persistence of craniopharyngeal canal with herniation of pituitary gland may be seen. Agenesis/dysgenesis of corpus callosum is also well seen on MRI.

Morning glory disc anomaly is distinct from optic disc coloboma. It occurs sporadically as compared to optic disc coloboma which is commonly familial and associated with multisystem congenital malformation syndromes. Morning glory disc anomaly is also associated with PHACE syndrome (posterior fossa malformation, hemangiomas, arterial anomalies including coarctation of aorta, cardiac anomalies, and ocular anomalies), when additional finding of sternal clefting is present, it is called PHACES syndrome, CHARGE syndrome (coloboma, heart defects, atresia of the choanae, retarded growth, ear anomalies), Okihiro syndrome (upper limb, ocular anomalies, deafness, and in some renal anomalies), and neurofibromatosis type 2. Various syndromes such as Aicardi [Figure 3] and Meckel syndrome are associated with colobomas.

Optic nerve hypoplasia

It is characterized by small optic disc affecting one or both eyes [Figures 3-6]. It can occur in isolation or in combination with endocrine (pituitary and hypothalamic dysfunction) and brain abnormalities [ventricles or white or gray matter development abnormalities, septo-optic dysplasia (SOD) [Figure 5], hydrocephalus, and corpus callosal abnormalities].

SOD is a heterogeneous condition diagnosed when two or more features of the classical triad, i.e. i) Optic nerve hypoplasia, ii) pituitary hormone abnormalities, and iii) midline brain defects – agenesis of septum pellucidum and/or corpus callosum, are present. Clinical presentations include hypopituitarism, the commonest being growth hormone deficiency presenting as short stature, failure to thrive, hypoglycemia, developmental delay, and visual impairment.

It usually occurs sporadically; however, a number of familial cases have been described. Mutations in homeobox *HESX1* gene are associated with sporadic cases and mutations of both *SOX2* and *SOX3* in the etiology of variants of SOD. Mutations have also been identified in *SOX2* in association with severe bilateral eye abnormalities (anophthalmia, microphthalmia) and defects of the corpus callosum with anterior pituitary hypoplasia. Additional features described in association with *SOX2* mutations include developmental delay, short stature, esophageal atresia, male genital tract abnormalities, and sensorineural hearing loss.

MRI shows hypoplastic optic nerves and chiasma, and various other findings of hypothalamic–pituitary dysfunction, i.e. anterior pituitary hypoplasia, ectopic posterior pituitary, absent/hypoplastic infundibulum, partial or complete absence of septum pellucidum and corpus callosum. Other findings are schizencephaly, hydrocephalus, and Chiari II malformation.

Sakoda complex includes sphenethmoidal encephalomeningocele, agenesis of the corpus callosum, and cleft lip and/or palate [Figure 6]. Associated optic disc dysplasia, microphthalmia, cortical dysgenesis, mental retardation, and epilepsy may be seen. Hereditary optic neuropathies include dominant optic atrophy and Leber hereditary optic neuropathy. Dominant optic atrophy is autosomal dominant, whereas Leber hereditary optic neuropathy has mitochondrial inheritance. These are characterized by painless loss of central vision, manifesting in childhood or adulthood. The diagnosis is based on ophthalmoscopic findings. MRI reveals diffuse bilateral optic nerve atrophy.

Wolfram's syndrome is a rare hereditary genetic disorder characterized by combination of diabetes insipidus, diabetes mellitus, optic atrophy, and deafness (DIDMOAD). It can be autosomal recessive or can have mitochondrial...
Optic nerve hypertrophy

Hypertrophy of prechiasmatic optic nerves and optic chiasma has well been described in Krabbe’s syndrome [Figure 8] which is an autosomal recessive lysosomal disorder caused due to deficiency of galactocerebroside β-galactosidase resulting in accumulation of globoid cells containing psychosine (galactosylsphingosine). Other MRI findings are white matter abnormalities, which are symmetric, patchy, and confluent involving periventricular and cerebellar white matter. High density on non-contrast CT scan is noted in basal ganglia, posterolateral thalami, and posterior limb of the internal capsule. T2-hypointense signal is seen in basal ganglia and dentate nucleus. Brain atrophy occurs later in the disease course. Enhancement of multiple cranial nerves and spinal roots has also been described. This occurs secondary to myelin breakdown and associated inflammation.

Differential diagnosis of optic nerve hypertrophy is broad [Table 3] and includes optic nerve glioma with dural ectasia, neurofibromatosis 1 (NF1), optic nerve sheath meningioma (ONSM), histiocytic or granulomatous infiltration of the optic nerves, juvenile xanthogranuloma, medulloepithelioma of the optic nerve, leukemia, and orbital pseudotumor.
NF1 is much more common than Krabbe’s disease. Both optic nerve glioma and optic nerve sheath ectasia may occur in NF1 patients. Orbital imaging findings in NF1 are: Enlargement of optic canal due to optic nerve glioma, dilatation of CSF within optic nerve sheath, retinal astrocytomas, infiltrating plexiform neurofibromas causing soft tissue alterations in the globe and extraocular muscles and also causing enlargement of various foramina (foramen ovale, rotundum, vidian canal, infraorbital canal, etc.), secondary sphenoid wing dysplasia which is characterized by remodeling or decalcification of sphenoid wings with anteroposterior enlargement of middle cranial fossa.

<table>
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<th>Diseases</th>
<th>Key features/orbital findings</th>
<th>Associated brain/Systemic findings</th>
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<tr>
<td>Krabbe’s disease</td>
<td>Autosomal Recessive disease due to deficiency of galactocerebrosidase β-galactosidase enzyme</td>
<td>Non-contrast CT- hyperdense basal ganglia and posterolateral thalami</td>
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<td>Infantile form most common</td>
<td>MRI-Symmetric, patchy confluent periventricular and cerebellar white matter signal, T2 hypointensity in basal ganglia and dentate nuclei, Cerebral and cerebellar atrophy in late stages, Enhancement of multiple cranial nerves and spinal roots</td>
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<td>Thickened optic nerves on MRI</td>
<td>Brain: 1) Hamartomas- T2 hyperintense foci in bilateral basal ganglia, thalami, dentate nuclei and white matter due to spongiform changes</td>
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<td>2) Gliomas</td>
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<td>3) Sphenoid wing dysplasia, enlargement of middle cranial fossa with arachnoid cyst</td>
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<td>4) Arterial abnormalities including moyamoya pattern, aneurysms</td>
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<td></td>
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<td>Others: Neurofibromas along peripheral and spinal nerves, dural ectasia, meningococes, kyposcoliosis, tibial bowing, pseudoarthroses, focal overgrowth of digit or limb</td>
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<td>Neurofibromatosis 1</td>
<td>NF1 is a neurocutaneous condition with autosomal dominant inheritance in 50% due to mutation in gene on chromosome 17. Rest of 50% cases are sporadic. Ocular manifestations: 1) Lisch nodules in iris - 2 or more nodules are included in diagnostic criteria for NF1 2) Buphthalmos 3) Optic sheath dural ectasia- idiopathic expansion of CSF within optic sheath 4) Optic nerve glioma-thickened enlarged optic nerves, unilateral or bilateral, may involve optic chiasma, optic tracts 5) Plexiform neurofibroma-infiltrative lesion mainly in superior orbital soft tissue resulting in proptosis 6) Choroid hamartomas-flat ill-defined lesions usually at posterior pole 6) Retinal tumors-astrocytic hamartomas, retinal hemangiomas</td>
<td>Brain: 1) Hamartomas- T2 hyperintense foci in bilateral basal ganglia, thalami, dentate nuclei and white matter due to spongiform changes 2) Gliomas 3) Sphenoid wing dysplasia, enlargement of middle cranial fossa with arachnoid cyst 4) Arterial abnormalities including moyamoya pattern, aneurysms</td>
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<td>Optic nerve glioma</td>
<td>Usually occurs in children and are associated with NF1 Imaging findings: Fusiform enlargement of nerve with nerve not seen separate from tumor Variable T2 signal and enhancement May have Perineural arachnoidal gliomatosis</td>
<td>Can involve optic nerve, chiasma or optic tracts</td>
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<td>Optic nerve menigioma</td>
<td>Usually occurs in middle aged female and in children associated with NF2 Imaging findings: Tram-tract sign-central non-enhancing optic nerve surrounded by homogenously enhancing tumor Erosion/Hyperostosis of adjacent bone</td>
<td>The intraorbital tumor may be extension from cavernous sinus, clinoide process, pituitary fossa, planum sphenoidale or olfactory groove</td>
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<td>Histiocytic or granulomatous</td>
<td>Histiocytic disease: Along with optic nerve infiltration, there are lesions involving bony orbital wall, associated soft tissue mass may show intracranial extension Granulomatous diseases like sarcoïdosis may affect optic nerves. Associated findings are diffuse lacrimal gland enlargement with extra ocular muscle infiltration, eyelid and periorbital inflammation</td>
<td>Histiocytic disease: Bones commonly involved. Definite diagnosis by biopsy</td>
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<td>infiltration of the optic</td>
<td>Brain findings in sarcoïdosis: Thickening and enhancement of cranial nerves especially optic nerves, nodular leptomeningeval enhancement, intraparenchymal lesions, infundibular thickening with suprasellar mass, dorsal based enhancing T2 hypointense lesions</td>
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<td>nerves</td>
<td>Characteristic skin lesions and Biopsy-granulomatous reaction with presence of foreign body giant cells and touting giant cells and absence of intracytoplasmic granules and lack of staining with S100 (which are present in histiocytosis) help in its diagnosis</td>
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<td>Juvenile xanthogranuloma</td>
<td>Benign self-limiting cutaneous disorder may have rarely systemic manifestations. Ocular manifestations include localized or diffuse iris tumor, unilateral glaucoma, spontaneous hyphaema, red eye with signs of uveitis, iris heterochromia</td>
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<tr>
<td>Medulloepithelioma</td>
<td>Mimics optic nerve glioma. It causes fusiform enlargement of optic nerve</td>
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<td>Leukemia</td>
<td>Both acute and chronic leukemia can present with infiltrating mass involving eyelid, optic nerve, canal and intracranial structures or with leukemic retinopathy. Relapse of disease is known to occur in orbit due to suboptimal penetration of chemotherapeutic drugs</td>
<td>Brain: Extramedulatious lesions, parenchymal lesions or leukemic meningitis</td>
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<td>Orbital pseudotumor</td>
<td>Acute painful diplopia, or other oculomotor symptoms Dramatic improvement with corticosteroids Imaging findings: T1, T2 hypointense enhancing soft tissue intraorbital mass. Optic nerve involved along with retrobulbar fat and extraocular muscles</td>
<td>Extension into orbital fissure and cavernous sinus may be seen-Tolosa Hun syndrome</td>
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</table>
The cause can be observed in ADEM, whereas Intravenous steroids are Disease-modifying drugs such as beta-interferon.

Demyelination of the spinal cord has an appearance with one or two lesions, and 51% in those with three or more lesions.

In the ONTT, the 5-year risk of developing MS was 16% in patients with normal brain MRI findings, 37% in patients with one or two lesions, and 51% in those with three or more lesions.

Hamartomatous lesions are commonly seen in MRI Brain in NF1 patients. These are T2-hyperintense non-enhancing lesions without any mass effect occurring in the brain stem, cerebral peduncles, and optic radiations. Gliomas, mostly low-grade astrocytomas, may also occur in these patients.

**DEMYELINATION**

Optic neuritis and its association with multiple sclerosis and acute disseminated encephalomyelitis

Optic neuritis (ON) literally means inflammation of the optic nerve. However, it is usually reserved for demyelinating events, isolated or with multiple sclerosis (MS) or acute disseminated encephalomyelitis (ADEM) or neuromyelitis optica (NMO). It results in acute monocular vision loss associated with some painful eye movements. It is two times more common in females than males.

The cause is presumed to be an autoimmune reaction that results in a demyelinating inflammation of the nerve. A gradual recovery of visual acuity with time is characteristic of ON. For most patients, even without treatment, visual function improves from 1 week to several weeks after onset. In the Optic Neuritis Treatment Trial (ONTT) study, intravenous steroids improved the vision faster as compared to those treated with oral steroids or placebo. Intravenous steroids showed statistically significant benefit in contrast sensitivity, color vision, and visual field, but not in visual acuity at 6 months. At 1-year follow-up, there was no statistically significant difference in visual function among the groups. However, Plant et al. suggested that high-dose corticosteroids, when administered at the onset of pain, may abort the attack of ON and prevent visual loss.

Intravenous steroids are recommended to reduce 2-year risk of developing MS when three or more signal abnormalities are present in MRI brain. They can also be considered to expedite recovery of vision. Disease-modifying drugs such as beta-interferon or glatiramer acetate have also been used to reduce the development and severity of clinically definite MS.

Visual outcome is better in patients with an isolated episode of ON as compared with patients who develop MS. Up to 75% of female patients and 35% of male patients initially presenting with ON ultimately develop MS.

In the ONTT, the 5-year risk of developing MS was 16% in patients with normal brain MRI findings, 37% in patients with one or two lesions, and 51% in those with three or more lesions.

ADEM may be also associated with ON. It is a rare multifocal, inflammatory demyelinating disorder of the central nervous system (CNS) occurring after vaccination, viral infections, or spontaneously. It is most commonly a non-progressive acute monophasic illness. Encephalopathy is a feature required for the diagnosis of ADEM, which differentiates it from MS. ADEM-associated ON is often bilateral, as compared to MS which frequently has unilateral involvement. MRI can differentiate between these two demyelinating conditions.

ADEM lesions are multiple, large, usually bilateral, but asymmetrical involving subcortical white matter or centrum semiovale. Basal ganglia, thalami, and brain stem may be involved in ADEM.

Neuromyelitis optica

Neuromyelitis optica (NMO) is an autoimmune demyelinating disease induced by a specific autoantibody, NMO-IgG, directed against aquaporin-4 water channels.

**Imaging pearls**

The optic nerve is swollen with hyperintense signal on STIR/T2 fat-saturated images and has intense homogenous enhancement. Occasionally, it reveals peripheral tram-track pattern of enhancement mimicking ONMS. However, ON will show enhancement limited to the nerve, rather than the sheath-like pattern of meningioma; there is absence of significant mass or expansion with clinical features of acute onset visual loss and pain.

Cerebrospinal fluid pleocytosis (greater than or equal to) 50 white blood cells/mm³ can be observed in ADEM, whereas this finding is highly atypical for MS.

Neuromyelitis optica

NMO preferentially affects the optic nerve and the spinal cord. Demyelination of the spinal cord has an appearance similar to that of transverse myelitis, involving over four to seven vertebral segments and the full transverse diameter.

Female are nine times more affected than males.

Brain lesions can occur and often are distinct from those seen in MS, and are around the ventricles due to high concentration of aquaporin-4 water channels. Periependymal lesions surrounding the third ventricle...
and cerebral aqueduct are highly characteristic, with involvement of thalamus, hypothalamus, and midbrain. The dorsal part of brain stem adjacent to the fourth ventricle may be involved in NMO. However, non-specific brain lesions are most commonly seen.

Table 4 shows the various demyelinating diseases affecting the optic nerves.

**Inflammatory**

**Perineuritis**
It is an uncommon variety of orbital inflammatory disease that is distinct from demyelinating ON. It affects the older age group as compared with ON and classically shows sparing of central vision. Optic perineuritis usually is idiopathic; however, few case reports have been described with tuberculosis, syphilis, sarcoidosis, Wegener’s granulomatosis, and giant cell arteritis.

**Imaging pearls**
MRI reveals enhancement around, rather than within, the optic nerve (tram-track sign on axial and doughnut sign on coronal images) and “streaky” fat with or without extraocular muscle enhancement [Figure 13]. Associated MRI findings with secondary etiologies include multiple tuberculomas, enhancement of multiple intracranial nerves with dural-based or intraparenchymal lesions, infundibular thickening and nodular leptomeningeal enhancement in sarcoidosis, wall thickening and enhancement of superficial temporal arteries with adjacent fat stranding in giant cell arteritis. Often ON can be the initial presentation of these diseases.
Optic perineuritis mimics ONSM on MRI, but clinically, it is more likely to be mistaken for acute demyelinating ON. Response to corticosteroids is more dramatic than in patients with ON, and patients are more likely to experience recurrence after stopping treatment. Secondary optic perineuritis patients need specific treatment often with corticosteroids for a longer time, with duration depending on response to treatment. Tuberculous perineuritis is treated with anti-tuberculous treatment for a year, syphilis with antibiotics (penicillin), giant cell arteritis needs high dose corticosteroids, Wegener’s granulomatosis with high dose of corticosteroids may be followed by more complex regimen to induce remission: A combination of rituximab, steroids, and cyclophosphamide. Sarcoïd optic perineuritis is treated with corticosteroids which need slow tapering. In refractory cases, immunosuppressive agents (azathioprine and cyclosporine) and antimetabolites (cyclophosphamide, chlorambucil, and methotrexate) may be given. SLE, keratoconjunctivitis sicca is the most common orbital condition; however, the most dreaded and uncommon is optic nerve involvement and retinal vaso-occlusion. It is caused by an ischemic event followed by axonal loss. Initial visual loss is severe; final outcome is variable. It is necessary to differentiate SLE-associated ON from idiopathic ON because of the severe visual impairment and steroid dependence associated with the SLE-induced ON. Early diagnosis and prompt treatment are important for restoring visual function in these patients.

**Orbital pseudotumor**
An idiopathic orbital inflammatory syndrome or orbital pseudotumor is a nonspecific, non-neoplastic inflammatory process of the orbit.

**Imaging pearls**
Imaging shows diffuse orbital mass, uveoscleral thickening, involvement of optic nerve, extraocular muscles, and lacrimal glands. There may be presence of fluid in Tenon’s capsule. The soft tissue mass is hypointense on T1 and T2W images and shows marked post-contrast enhancement. It involves the muscle bellies and tendons of extraocular muscles.
muscles. Optic perineuritis may be seen when there is involvement of optic nerve sheath with inflammatory soft tissue which also infiltrates adjacent fat. When the inflammatory soft tissue is confined to superior orbital fissure and cavernous sinus, it is known as Tolosa Hunt syndrome. Dramatic improvement with corticosteroids is the hallmark of orbital inflammatory syndrome.[55]

**Sarcoidosis**

This may involve any part of the orbit, i.e. extraocular muscles, orbital fat, lacrimal glands, or globe, and can be indistinguishable from pseudotumor. Cranial nerves are commonly involved in sarcoidosis.[56] Optic nerve involvement can be unilateral or bilateral and is involved in intraorbital segment or at the level of chiasma.[57] The nerves are enlarged, thickened, and show hyperintense signal on STIR/T2 fat-saturated images and reveal post-contrast enhancement. The dural sheath may be involved mimicking optic nerve meningioma.

**Associated orbital findings**

Diffuse lacrimal gland enlargement (50-60%) with extension and involvement of lateral rectus and/or other extraocular muscles’ infiltration, eyelid and periorbital inflammation.

**Associated brain findings**

Nodular leptomeningeal enhancement, intraparenchymal lesions, infundibular thickening with suprasellar mass, dural based enhancing T2-hypointense lesions.[56]

**TUMORS**

**Optic nerve glioma**

These are relatively rare tumors and occur in children; they are often associated with NF1. These are mainly low-grade astrocytomas; however, their biological behavior is unpredictable.[58] They are often multifocal and bilateral in NF1 patients. In NF1 patients, the orbital nerve is the commonest site, followed by optic chiasma and hypothalamus, whereas in non-NF1 patients, chiasma is the commonest site of involvement. In NF1 group, the tumor infrequently extends beyond the optic pathway and the shape of the nerve and chiasma are preserved as compared to that in non-NF1 patients. Cystic component of the tumor is commonly seen in non-NF1 patients. Most of the patients with NF1 show stable tumor volume on follow-up, as compared to non-NF1 patients.[58]

**Imaging pearls**

The tumor is isointense on T1 and iso- to hyperintense on T2W images and shows variable contrast enhancement. On T2W images, it demonstrates central isointensity, with surrounding hyperintensity corresponding to perineural arachnoidal gliomatosis. The perineural arachnoidal gliomatosis may enhance with gadolinium administration. There is resultant fusiform enlargement of the nerve with the nerve indistinguishable from the tumor [Figures 15 and 16].

**Optic nerve sheath meningioma**

ONSM is the term applied either to i) primary meningioma which arises from meningoepithelial cap cells along the optic nerve from the globe to the prechiasmatic segment or to ii) secondary meningioma which is actually an extension from the cavernous sinus, clival process, pituitary fossa, planum sphenoidale, or olfactory groove. These are more common in middle-aged females or in children with neurofibromatosis 2.[59]

![Figure 15: Tumors: Optic nerve glioma in a non-NF1 patient. Axial T2 fat-saturated and post-contrast T1 fat-saturated images show thickening with enhancement of right optic nerve. Perineural arachnoidal gliomatosis is also seen, which also shows enhancement (arrow).](image)

![Figure 16 (A and B): Tumors: Optic nerve glioma in an NF1 patient. Coronal STIR (A) and axial T2W (B) images of a 7-year-old male with NF1 with mild decrease in vision on the right side show thickening and hyperintense signal in right optic nerve (arrow) suggestive of glioma. Hyperintense signal is seen in bilateral globus pallidi (elbow arrows), thalami, dorsal aspect of pons, and brachium pontis (curved arrows) due to spongiform changes seen in NF1](image)
Imaging pearls

These are iso- to hypointense on T1 and iso- to hyperintense on T2W images with intense homogenous enhancement. The central hypointense, non-enhancing optic nerve results in the “tram-tract sign.” These tumors often show calcification [Figure 17] and adjacent bone hyperostosis or erosion. They appear plaque-like and grow linearly along the nerve sheath. They cause diffuse or segmental circumferential thickening of the optic nerve sheath.\[60\]

Optic nerve can be separated from the tumor unlike optic nerve glioma. Posterior optic pathway may show atrophy. Other conditions showing tram-track sign include sarcoidosis, periocular neuritis, orbital pseudotumor, periorbital hemorrhage, metastases and leukemia/lymphoma, and Erdheim–Chester disease [Table 5].\[1,60,61\]

Metastases

Metastases [Figures 18 and 19] to orbits are rare and when they occur, they usually involve the uveal tract. Isolated metastases to optic nerve are extremely uncommon; however, they are reported with carcinomas from breast, lung, gastrointestinal tract, etc. Radiologically, isolated metastatic optic nerve tumors appear similar to primary optic nerve tumors; however, history of previous malignancy is helpful.\[62\]

Retinoblastoma is the most common tumor of globe in childhood. MRI is used to assess intraocular spread including invasion of the optic nerve, extraocular as well as intracranial extension. Inherited forms of retinoblastoma may be associated with primary tumors of the suprasellar or pineal region. Involvement of optic nerve or choroid infiltration is an important predictor of metastases in retinoblastoma [Figure 19]. Involvement of optic nerve beyond lamina cribrosa is associated with poor prognosis.\[63\] MRI shows disruption of linear enhancement at choroidoretinal complex or thickening of enhancing choroidoretinal complex in prelaminar optic nerve involvement. Postlaminar optic nerve involvement is seen as focal optic nerve enhancement and/or thickening.\[63\]
Table 5: Differential diagnosis for Tram-tract appearance of optic nerve

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<th>Associated brain/Systemic findings</th>
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<td>Diffuse or segmental circumferential thickening of optic nerve sheath</td>
<td>Wegener’s granulomatosis: Cavitating lung nodules, masses in nasal cavities eroding nasal septum, cANCA+, Tuberculosis: Tuberculomas, leptomeningeal enhancement, Mantoix test positive Giant cell arteritis: Wall thickening and enhancement of superficial temporal arteries with adjacent fat standing</td>
</tr>
<tr>
<td>Perineuritis</td>
<td>Clinically mimics acute demyelinating optic neuritis, however MRI shows enhancement around rather than within the optic nerve. It can be idiopathic or secondary to tuberculosis, syphilis, sarcoidosis, wegener’s granulomatosis, giant cell arteritis</td>
<td></td>
</tr>
<tr>
<td>Pseudotumor</td>
<td>Acute presentation with proptosis, painful red eye. Along with enhancement of optic nerve sheath, other inflammatory findings of uveoscleral thickening, fat stranding in retrobulbar fat, fluid in tenon’s space, enlargement of extracocular muscles may be seen. Dramatic response to steroid treatment</td>
<td>Extension into orbital fissure and cavernous sinus may be seen-Tolosa Hunt syndrome</td>
</tr>
<tr>
<td>Sarcoïdosis</td>
<td>Associated orbital findings: Diffuse lacrimal gland enlargement with extra ocular muscle infiltration, eyelid and periorbital inflammation</td>
<td>Thckening and enhancement of cranial nerves especially optic nerves, Nodular leptomeningeal enhancement, intraparenchymal lesions, infundibular thickening with suprasellar mass, dural based enhancing T2 hypointense lesions</td>
</tr>
<tr>
<td>Perioptic hemorrhage</td>
<td>Can be spontaneous or traumatic. Acute hemorrhage may not show any enhancement and appears hypodense on CT scan. At later stages, it reveals peripheral enhancement. Clinical history is important</td>
<td></td>
</tr>
<tr>
<td>Leukemia/lymphoma</td>
<td>Other than involvement of optic nerves, leukemia can present with an infiltrating mass involving eyelid, canoral and intraconal structures or with leukemic retinopathy. Relapse occurs due to suboptimal penetration of chemotherapeutic drugs into orbit</td>
<td>Brain: Extradural enhancing lesions, parenchymal lesions or leukemic meningitis</td>
</tr>
<tr>
<td>Metastases</td>
<td>Usually history of primary malignancy is present Carcinomas from breast, lung, and gastrointestinal tract can metastasize to optic nerves Retinoblastoma may directly infiltrate optic nerve</td>
<td>Brain and systemic metastases</td>
</tr>
<tr>
<td>Erdheim Chester disease</td>
<td>Rare non-Langerhans’s cell systemic xanthogranulomatosis. Orbital masses arise from optic nerve sheath</td>
<td>Brain lesions include T2 hypointense enhancing dural masses, intraparenchymal masses involving hypothalamus and pituitary infundibulum resulting in diabetes Insipidus. Other systems involved are appendicular skeleton, kidneys, lungs, retroperitoneum, heart, blood vessels and skin</td>
</tr>
</tbody>
</table>

Table 6: Blood supply of optic nerve

<table>
<thead>
<tr>
<th>Optic nerve segments</th>
<th>Arteries</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optic nerve head</td>
<td>Superficial part: Branches of CRA Region around lamina cribrosa: Short posterior ciliary arteries or via circle of Zinn-Haller</td>
</tr>
<tr>
<td>Intraorbital segment</td>
<td>Intraorbital segment is supplied by posterior ciliary arteries via pial network Collateral circulation by Middle meningeal artery</td>
</tr>
<tr>
<td>(Resistant to ischemia)</td>
<td></td>
</tr>
<tr>
<td>Orbital apex</td>
<td>Collateral circulation by Middle meningeal artery</td>
</tr>
<tr>
<td>Intracanicular segment</td>
<td>Anteriorly from collateral branches of OA and posteriorly from pial branches of ICA and superior hypophyseal arteries</td>
</tr>
<tr>
<td>(Prone to ischemia)</td>
<td>Pial branches of ICA, superior hypophyseal, A1 segment of anterior cerebral and anterior communicating arteries</td>
</tr>
<tr>
<td>Intracranial segment</td>
<td></td>
</tr>
</tbody>
</table>

Ischemic Optic Neuropathy

Blood supply of optic nerve

Blood supply of the optic nerve is different for its four segments [Table 6]. Most of the optic nerve derives its blood supply from the OA, which is the first branch of supraclinoid internal carotid artery (ICA). Occasionally, OA may have its origin from the middle meningeal artery due to enlargement of anastomoses between recurrent branch of lacrimal artery and orbital branch of middle meningeal artery, from the cavernous ICA, and rarely from the middle or anterior cerebral arteries. After its origin, OA enters the optic canal inferolateral to the optic nerve and is separated by a dural sheath. In intraorbital course, it crosses over (in 83%) or under (17%) the optic nerve to lie medial to the nerve. Further in orbit, it runs between the medial rectus and superior oblique muscles and lies close to anterior ethmoid foramen and medial orbital wall. It terminates at the superomedial angle of the orbit into supratrochlear and dorsal nasal branches. There is a wide variation in the branches of OA; however, its important branches in the orbit are two or three posterior ciliary arteries and a central retinal artery (CRA).[64]

The superficial aspect of the optic nerve head is supplied by the branches of CRA, whereas the region around lamina cribrosa gets blood supply directly from the short posterior ciliary arteries or via circle of Zinn–Haller. Intraorbital segment is
supplied by posterior ciliary arteries via pial network. This segment is resistant to ischemia because of a plethora of pial vessels. Middle meningeal artery, branch of the external carotid artery forms collateral circulation at the orbital apex. Intracanalicular segment lies within the watershed zone and, hence, is prone to ischemia and vulnerable to shearing injuries in skull fracture. It derives blood supply anteriorly from the collateral branches of OA and posteriorly from the pial branches of ICA and superior hypophyseal arteries.

Intracranial segment is supplied by the pial branches of ICA, superior hypophyseal, A1 segment of anterior cerebral and anterior communicating arteries.

Anterior ischemic optic neuropathy
Anterior ischemic optic neuropathy (AION) occurs when ischemia/infarction involves optic disc. It results in acute painless monocular vision loss. Diagnosis is mainly clinical based on history, clinical examination, and fundoscopic findings of pallor/edema of the optic disc and peripapillary hemorrhage. Imaging plays no role in its diagnosis. However, MRI may be necessary in cases of signficant pain with eye movement to exclude ON or MS, in patients with atypical course, i.e. with prolonged disc edema or progressive and/ or recurrent visual loss more than 2 months after their initial presentation to exclude inflammatory or compressive lesions.

AION can be:
- Arteritic AION: Occurs due to blood vessel inflammation, most commonly from giant cell arteritis. Other causes include polyarteritis nodosa, Wegener’s granulomatosis, connective tissue diseases such as SLE, Churg‑Strauss syndrome, and rheumatoid arthritis
- Nonarteritic AION: Seen in association with hypertension, diabetes mellitus, myocardial infarction, and hypercholesterolemia.

Posterior ischemic optic neuropathy
Posterior ischemic optic neuropathy (PION) involves the optic nerve and/or optic chiasma. Color vision loss is the presenting symptom. It is seen in the perioperative period following cardiac surgery, spine surgeries, and major abdominal surgeries. It may be associated with hypertension, diabetes mellitus, myocardial infarction, and hypercholesterolemia.

Diagnosis is by MRI, especially DWI, which shows restricted diffusion [Figure 20].

Miscellaneous Causes
Idiopathic intracranial hypertension
It is characterized by elevated CSF pressure and papilledema without focal neurological deficit. It is mostly seen in middle‑aged females and is a diagnosis of exclusion.

Imaging pearls
MRI shows peri optic nerve sheath distention, posterior flattening of globe and optic nerve head protrusion, vertical buckling of the optic nerves, empty sella or partially empty sella, narrowing and decreased flow in dural sinuses (usually in transverse sinuses), which normalize after reduction in CSF pressure. Small meningoceles may often be seen [Figures 21 and 22].

Figure 20 (A and B): Ischemic: DWI (A) and FLAIR (B) images of a 45-year-old female with sudden loss of right vision after cholecystectomy surgery 4 days back show restricted diffusion in right optic nerve (arrow) with hyperintense signal on FLAIR image

Figure 21 (A and B): Idiopathic intracranial hypertension. Axial (A) sagittal and coronal (B) T2 fat-saturated images of a 26-year-old female with blurring of vision and b/l papilledema show peri optic nerve sheath distention, flattening of posterior globe, and optic nerve head protrusion (arrowhead). Partial empty sella (elbow arrow) and meningoencephalocele are seen at left foramen rotundum (star)
Secondary compressive optic atrophy

Tumor

The common tumors compressing prechiasmatic optic nerve and chiasma are: Pituitary macroadenoma [Figure 23], meningioma in adults [Figure 24], craniopharyngioma, and gliomas of visual pathway in children. These cause gradual and progressive loss of vision. It is important to mention whether the optic chiasm is prefixed or postfixed. Pituitary/tuberculum sellae lesions are more likely to compress prefixed chiasm than postfixed chiasm.\(^{[60]}\) Prefixed chiasm or prominent tuberculum sellae limits the access to suprasellar area in transcranial approach.\(^{[60]}\)

Other sellar/parasellar lesions include ICA aneurysm in adults. Intraorbital tumors like lymphangioma [Figure 25], hemangioma, and dermoid cysts can also compress the intraorbital optic nerve resulting in atrophy.

Thyroid orbitopathy

It is characterized by fusiform enlargement of extraocular muscles with sparing of its tendinous insertions and increase in orbital fat volume resulting in compression of optic nerve at the orbital apex [Figure 26]. Inferior and medial recti are most commonly involved followed by lateral and superior recti muscles. Involvement may be symmetrical or asymmetrical, but is usually bilateral. Other findings include proptosis, bony changes in lamina
papyracea with bowing due to muscle pressure, lacrimal gland displacement and enlargement, and superior ophthalmic vein dilatation.\textsuperscript{[71]} Surgical decompression may be needed when the optic nerve is compressed.

**Vascular**

Optic pathway compression can rarely occur due to tortuous vessels, resulting in visual loss and optic atrophy [Figure 27]. The symptoms occur due to direct pressure compression and/or ischemia secondary to occlusion of small arterial supply branches. Visual loss is usually mild and slowly progressive. Most patients are elderly with other forms of vascular disease. It is managed conservatively; however, occasionally surgical intervention may be required when there is rapid progression.\textsuperscript{[72]}

Progressive visual loss due to optic nerve compression can also occur with ophthalmic segment carotid artery aneurysms (carotid-ophthalmic aneurysms). These aneurysms may be asymptomatic or occasionally may cause proptosis.

Ophthalmic segment of ICA gives rise to two branches—OA and superior hypophyseal artery. OA aneurysms arise from ICA, just distal to the origin of OA, point superiorly or superomedially, and typically displace the optic nerve superomedially. Anatomic variations in OA origin as mentioned above are important when embolization or surgery is planned.\textsuperscript{[73]}

Superior hypophyseal artery aneurysms arise above the dural ring from the medial bend of ICA. These aneurysms are located medial to ICA and can extend beneath the chiasm (suprasellar variant) or extend anteriorly beneath anterior clinoid process (paraclinoid variant). Preoperative differentiation helps the surgeon in achieving low operative and visual morbidity.\textsuperscript{[74]}

**Traumatic optic neuropathy**

Injury can be direct when the optic nerve fibers are anatomically disrupted due to penetrating trauma or due to fracture fragments in optic canal or optic nerve sheath hematomas associated with craniofacial trauma. CT scan of the orbit is recommended to visualize small fracture fragments and acute orbital/intracranial hemorrhages. Indirect injury occurs due to transmission of forces to the optic canal from blunt trauma. Road traffic accidents are the most common causes of traumatic optic neuropathy [Figure 28].\textsuperscript{[75]} The condition can sometimes

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**Figure 26:** Compressive optic neuropathy: Thyroid orbitopathy. Coronal STIR and axial T2W images of a 50-year-old female with diplopia show thickened and hyperintense signal in both medial and inferior recti muscles (arrow) resulting in compression of optic nerve at the orbital apex (arrowhead)

**Figure 27:** Compressive optic neuropathy: Vascular. Coronal STIR and MR angiography images of a 59-year-old male with gradual diminishing vision in both eyes show compression of optic chiasma by right A1 segment superiorly (arrow) and by left PCOM from the inferior aspect (dotted arrow) resulting in distortion of chiasma

**Figure 28:** Traumatic optic neuropathy. Coronal STIR images of a 23-year-old male with loss of right vision following trauma show signal in intracanalicular segment of the right optic nerve with severe atrophy of prechiasmatic segment (arrow). Gliosis is also seen in the right temporal lobe (arrowhead)

**Figure 29:** Traumatic optic neuropathy. Preoperative scan showing tuberculum sella meningioma (arrowhead) and elevation of optic chiasma (arrow) in a 68-year-old male with bitemporal hemianopsia. Postoperative scan shows gliosis of optic chiasma (right > left). Postoperatively, the patient had complete right monocular vision loss with severe reduction in left vision
Figure 30 (A and B): Toxic (alcohol) optic neuropathy. Coronal STIR (A) images of a 42-year-old alcoholic male with gradual loss of vision for 6 months showing hyperintense signal with atrophy of both optic nerves (arrow) and chiasma (elbow arrow). Also note on coronal T2W (B) image the prominence of cerebellar fissures (arrowhead) and cortical sulci suggestive of atrophy.

Figure 31: Intraocular pathology resulting in optic nerve atrophy. Axial T2 fat-saturated and coronal STIR images show bupthalmos (arrowhead) due to high myopia with bilateral optic nerve signal and optic atrophy (arrow).

occur secondary to surgery when the optic nerve or chiasma is located close to the tumor [Figure 29].

Toxic optic neuropathy
Various toxins affect the optic nerves resulting in visual problems [Figure 30]. The common toxins include ethanol, methanol, ethylene glycol, tobacco, and drugs like ethambutol, isoniazid, chloroquine, quinine, sulfonamides, linezolid, amiodarone, and digitalis.

The diagnosis is often delayed when optic nerve pallor has already set in and vision is severely affected. These usually cause central or cecocentral scotomas due to selective involvement of maculopapillar bundle. With tobacco-alcohol amblyopia, genetic and nutritional factors like deficiency of vitamin B complex increase the susceptibility of injury to the optic nerves.[9]

Intraocular pathologies resulting in optic nerve atrophy
Glaucoma is associated with elevated intraocular pressure and affects the intraocular segment of the optic nerve; phthisis bulbi, etc., can also result in optic atrophy [Figure 31].

Conclusion
Various pathologies affect the optic nerves, either isolated or associated with intracranial abnormalities. A clear understanding of MR imaging protocols, key anatomical structures, onset of symptoms, along with familiarity with various pathologies is fundamental for a radiologist to arrive at an accurate diagnosis and guide the referring clinician in patient care.

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