Title: **Optical Coherence Tomography (OCT) in Neuro-ophthalmology**

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Abstract
Optical coherence tomography (OCT) is a non-invasive medical imaging technology that is playing an increasing role in the routine assessment and management of patients with neuro-ophthalmic conditions. Its ability to characterise the optic nerve head, peripapillary retinal nerve fibre layer and cellular layers of the macula including the ganglion cell layer enables qualitative and quantitative assessment of optic nerve disease. In this review, we discuss technical features of OCT and OCT-based imaging techniques in the neuro-ophthalmic context, potential pitfalls to be aware of, and specific applications in more common neuro-ophthalmic conditions including demyelinating, inflammatory, ischaemic and compressive optic neuropathies, optic disc drusen and raised intracranial pressure. We also review emerging applications of OCT angiography within neuro-ophthalmology.

Introduction
Optical coherence tomography (OCT), a quick and reproducible imaging technique using low coherence interferometry to produce cross-sectional images of the retina and optic nerve head (ONH), has become one of the most valuable tools employed in the assessment of ophthalmic patients. Alongside advances in technology, its application in the field of neuro-ophthalmology specifically continues to expand.

OCT allows non-invasive visualisation of the anatomy of the most anterior part of the visual pathway, from retina to lamina cribrosa. Lesions involving the pre-laminar area can be assessed with spectral domain (SD-OCT) and enhanced depth imaging OCT (EDI-OCT) of the ONH. Beyond this, afferent visual pathway lesions involving the optic nerve, chiasm or
tracts can lead to visible axonal loss caused by direct retrograde axonal degeneration. OCT imaging can capture and quantify axonal loss through measurements of retinal nerve fibre layer (RNFL) thickness, and neuronal damage through measurements of ganglion cell layer (GCL) or combined ganglion cell layer-inner plexiform layer (GCIPL) thickness. The strict anatomical structure of the retina and maintenance of retinoscopic organisation with the afferent visual pathway (1), as described in table 1, increases the utility of OCT in evaluating central nervous system pathology.

In this review, we discuss the imaging parameters useful in the assessment of neuro-ophthalmic conditions, pitfalls to be aware of, and describe specific neuro-ophthalmic conditions where OCT imaging can help with diagnosis, management and follow-up.

**Imaging the optic nerve head (ONH) and peripapillary retinal nerve fibre layer (pRNFL) thickness**

OCT imaging of the ONH can be undertaken in two or three dimensions (2D or B-scan and 3D or volume scan). The OCT software can construct an ONH topographical map (figure 1) which can be used to quantify parameters such as ONH diameter, depth and diameter of the central cup, and thickness of the neuroretinal rim. Qualitative assessments can also be made of structural abnormalities such as optic nerve pits, tumours, optic disc drusen (ODD) and choroidal neovascularisation (CNV). EDI-OCT is beneficial for analysing the deeper structure of the ONH up to the level of the lamina cribrosa.
The peripapillary retinal nerve fibre (pRNFL) thickness is a measure that can be used to quantify axonal integrity in pathological processes involving the optic nerve. Technical details are outlined in table 2. Figure 2 shows an example of a pRNFL report. Normal global average pRNFL thickness is approximately 105µm with an estimated physiological loss due to aging of about 0.017% per year from age 18 years onwards, equating to a 10- to 20-µm loss over a period of 60 years (2).

**Imaging the macula and macular ganglion cell-inner plexiform layer (mGCIPL) thickness**

Aside from detailed visualisation of macular structure, macular OCT scans offer the possibility to conduct manual and automated segmentation analysis of macular ganglion cells layer (mGCL) and inner plexiform layer (mIPL) densities. Technical details are outlined in table 2. Figure 3 shows an example of a mGCL thickness map report. Both mGCL and combined mGCIPL thickness values have revolutionised the evaluation of optic neuropathies.

**Pitfalls to be aware of in OCT assessment**

As with any technology, there are limitations, pitfalls and potential errors that must be considered, especially when using derived measurements in clinical decision-making. Foremost, different OCT machines have different measurement protocols and so patients must be reviewed on the same machine using the same scanning protocol for accurate longitudinal comparisons to be made. Secondly, all OCT measurements are compared to a normative database. Often these normative databases are made up of Caucasian middle-
aged subjects, and as such this must be considered when evaluating measurements from a patient not in this demographic, for example a child or a patient of different ethnicity.

Interpretation of automated OCT measurements need to take into account potential artefacts. The centre of the fovea on OCT may differ from the retinal locus of fixation (3,4) and deviations of 60 ± 50μm between fixation and the centre of the foveal avascular zone can occur (5). For correct analysis, accurate placing of the measurement area should be checked and amended manually if necessary (6).

OCT segmentation software algorithm automatically detects the pRNFL and mGCIPL. In some cases, particularly in eyes with ocular pathology or in scans of poor image quality, the software can fail. Again, manual correction should be done to improve the accuracy and reproducibility of the measurements (6). This may not be possible with severe retinal pathology. The introduction of eye tracking and retest software has improved reproducibility of RNFL measurements (7).

Repeatability of automated segmentation measurements has been demonstrated to vary across different OCT devices and depends on the specific scan protocol and algorithm software updates (8). As described earlier, patients should be monitored consistently using similar OCT methodologies. The presence of ocular disorders and poor visual acuity may result in fixation errors and induce measurements that appear variable over time (9). Likewise, assessment of retinal thickness should be limited to the area encircling the parafoveal macular rim (10).
The importance of distinguishing artefacts from true pathological changes cannot be understated, not only to ensure optimum patient care but also to avoid invasive and costly investigations.

**Neuro-ophthalmic conditions**

**Multiple Sclerosis (MS) and demyelinating optic neuritis (MSON)**

The earliest use of OCT in neuro-ophthalmology was in 1999 where Parisi and colleagues reported pRNFL thinning on OCT in MS patients with previous MSON (11). In the acute phase of MSON, ONH swelling due to axoplasmic flow stasis in the inflamed optic nerve may be demonstrated on OCT and the elevated pRNFL measurements used to objectively quantify the swelling (12). pRNFL thickening may also be seen in MSON where there is no clinically evident ONH swelling, and this may be useful in detecting optic neuritis flares in clinic (13). Typically, 3 months after the acute episode of MSON, pRNFL atrophy develops which again can be quantified using OCT measurements demonstrating pRNFL thinning, most evidently in the temporal pRNFL region (14).

Due to the potentially elevated pRNFL measurements initially, followed by thinning some weeks to months later, it can be challenging to rely on pRNFL thickness to track structural changes in axonal integrity. Furthermore, there is an OCT floor effect, whereby mean pRNFL values do not generally reduce below 30µm regardless of extent of optic nerve injury, making it difficult to detect appreciable change in severe optic atrophy or recurrent optic neuritis (15). OCT-mGCL or mGCIPL volume and thickness analysis has been shown to be a more sensitive and reliable measure of retrobulbar neuroaxonal injury. mGCIPL thinning
occurs earlier in the time course of acute ON (after about 2 weeks), even whilst the RNFL remains oedematosus, thus providing an early indication of retinal ganglion cell dropout and possible permanent visual dysfunction (16). Whilst lower pRNFL values have been shown to correlate with reduced visual acuity, contrast sensitivity, visual field mean sensitivity and colour vision mean testing scores (17,18), mGCIPL thickness correlates better with these measures of visual dysfunction, as well as quality of life measures, disability and MRI findings (19,20). Overall OCT is helpful for prognostication after MS related relapses.

MS patients can experience substantial subclinical disease activity, and even in the absence of clinical MSON events, can develop progressive pRNFL and mGCIPL thinning (21,22). A meta-analysis comparing 1,667 MSON eyes and 4,109 MS non-ON eyes to 1,697 eyes from healthy control subjects found significant pRNFL and mGCIPL thinning in both MSON eyes (mean difference -20μm and -16μm respectively) and MS non-ON eyes (mean difference -7μm and -6μm respectively) relative to control eyes (23). Without additional MRI or electrodiagnostic data, the meta-analysis cannot conclude whether the finding in the MS non-ON eyes is a manifestation of a primary neurodegenerative component of MS, or whether it is due to retrograde transsynaptic degeneration from pathology of the posterior visual pathway. These data come from smaller sized, multimodal cohort studies and case reports (24). However, the finding does suggest that OCT could be used in the measurement of disease activity in MS, potentially reducing the need for frequent MRIs. This is further supported by the fact that reduced pRNFL and mGCIPL values have been shown to correlate with other surrogate markers used to monitor disease activity such as MRI-measured brain atrophy (25,26), spinal cord lesions (27) and neurological disability
scores (25,26,28-30). Figure 4 demonstrates findings in an MS patient with post-chiasmal demyelination with mGCL density measurements mirroring the visual field defect.

Microcystic macular changes detected on retinal OCT has been a recent area of interest since first reported in patients with demyelinating optic neuropathy in 2012 by Gelfand and colleagues (31). Small hyporeflective spaces within the inner nuclear layer (INL) in the parafoveal region of the fovea are characteristic, with sparing of the fovea itself, which would otherwise be involved in vascular leakage. Microcystic macular changes and thickening of the INL have been shown to be associated with disease activity and worse disability in MS (32). However, microcystic macular change is not unique to MS and can be observed across a variety of optic neuropathies, including non-inflammatory aetiologies such as compressive optic neuropathy (33). With several studies showing no corresponding dye leakage on fluorescein angiography (34,35), it is likely that these changes are due to retrograde axonal degeneration and that loss of retinal Muller cell function plays a role in microcyst formation (36). The literature uses the terms ‘microcystic changes’, ‘retrograde maculopathy’, ‘microcystic macular oedema (MMO)’, and microcystic macular edema (MME). None of these terms distinguish transient from permanent changes. Terminology and understanding of this novel observation is still evolving.

**Neuromyelitis Optica Spectrum Disorder (NMOSD)**

Whilst pRNFL and mGCIPL thinning is typical in optic neuritis generally, the degree and pattern of loss may help with differentiating underlying aetiology. It has been shown that ON related to NMOSD leads to more pronounced thinning of the pRNFL and mGCIPL than in MSON (37,38). The distribution of pRNFL thinning also differs, with NMO-ON preferentially
involving the inferior and superior quadrants, versus the temporal quadrant which is more typically affected in MS (37). Patients with NMO/SD do not seem to suffer the progressive pRNFL atrophy that occurs in MS independent of ON episodes (39), as described earlier, with damage appearing to be attack-related, although some studies have shown the mGCIPL to undergo progressive thinning (40). This seems to be specific for aquaporin4 IgG positive NMO/SD, whereas myelin-oligodendrocyte-glycoprotein (MOG) IgG positive NMSOD patients demonstrate a more similar pattern to MS patients (41). The reason for this is not clear, and certainly the time course in NMO-ON and MOG-ON is less well understood than in MSON, with the relationship between severity of atrophy observed and loss of visual function also being less well correlated. Microcystic macular change is also seen more frequently in NMO/SD patients (20-26%) than in MS patients (1-5%) (37,42), and INL measurements are thicker in NMSOD versus MS patients (43,44). This may be a reflection of the severity of optic neuropathy and the resultant retrograde degeneration. Figure 5 demonstrates progressive pRNFL and mGCL thinning in a patient with NMO after right eye optic neuritis.

Optic disc drusen (ODD)

ODD are acellular intracellular and extracellular deposits located in the optic nerve that may become calcified over time. Whilst superficial ODD can be detected on clinical examination, distinguishing buried optic disc drusen from optic disc swelling remains a clinical challenge. Ancillary tests including fundus autofluorescence, computer tomography, fluorescein angiography and B-scan ultrasonography are used to help distinguish between the two. These do have their limitations, however. B-scan ultrasonography, for example, whilst
traditionally viewed as the ‘gold-standard’ test for detection of buried ODD, is unable to
detect non-calcified drusen or peripapillary hyperreflective ovoid mass-like structures
(PHOMS) (45). Resolution is relatively poor, making it difficult to monitor drusen
progression (45). Furthermore, B-scan ultrasonography does not give information regarding
the neuroaxonal integrity of optic nerve and retinal structures, and as such does not allow
prognostication with regards to visual outcomes in patients with ODD (46).

OCT, specifically EDI-OCT, has earnt its place as not only a useful complement but as a
potential competitor as the gold-standard for diagnosis and analysis of ODD. Prior to
mainstream use of EDI technology, there was much variability in the literature with regards
to ODD morphology on OCT. Early studies focussed on RNFL thickness differences between
ODD and disc oedema (47, 48), but this was found to be unreliable (49). Other OCT features
described in the literature included ODD appearing as highly reflective round sub-retinal
structures with well-defined margins displacing the adjacent tissue (47), and a ‘lazy V’
pattern or ‘lumpy bumpy’ internal contour of the subretinal hyporeflective space helping to
distinguish between ODD and papilloedema (50,51). However, the hyporeflective space was
actually an imaging artefact and a by-product of poor-penetrance inherent to early
generations of OCT (45).

The improved penetrance afforded by EDI-OCT technology now enables quantification of
drusen size, delineation of drusen borders and assessment of the integrity of adjacent
retinal structures. The Optic Disc Drusen Studies Consortium have published guidelines for
the assessment of ODD (52). They define ODD as hyporeflective structures always located
above the lamina cribrosa, with a full or partial hyperreflective margin, often most
prominent superiorly (figure 6). Blood vessels, which appear as elongated tube-like structures, are distinguished from ODD as they lack the hyperreflective signal around them. ODD are often seen as conglomerates of smaller ODD with internal reflectivity within the signal-poor core. Hyperreflective horizontal lines might represent early ODD but should not be diagnosed as ODD.

EDI-OCT has been shown to have significantly higher ODD detection rate than B-scan ultrasonography (53). Furthermore, the size and type of ODD classified by EDI-OCT have been shown to correlate with visual field defects. Specifically, confluent ODD have been associated with worse mean deviation scores on visual fields testing (54). Some studies have suggested that EDI-OCT ODD volumes correlate with structural optic nerve head damage and functional deficits among patients (55). The capability of EDI-OCT to quantify ODD volume, and additional ability of OCT to measure neuroaxonal changes in the retina, will enable the monitoring of ODD progression and potentially provide guidance with regards to risk factors for vision loss in ODD patients.

Peripapillary hyper-reflective ovoid structures (PHOMS)
Peripapillary hyper-reflective ovoid mass-like structures (PHOMS) were originally described in patients with ODD (47-49) and were thought to represent precursors or variants of ODD (54). However, as explained by the Optic Disc Drusen Studies Consortium, PHOMS, unlike ODD, are hyperreflective without a sharp outer margin or hyporeflective core (52) (figure 7). They are neither visible on fundus autofluorescence nor on B-scan ultrasonography, and they can be found on OCT in patients with papilloedema without ODD (52). They are thought to represent disruption of retinal layers caused by local axoplasmic build-up (45),
with the histopathology of papilledema suggesting that PHOMS might correspond to the lateral bulging of distended axons into the peripapillary retina (52). Of note, PHOMS can also be seen in crowded discs, or “pseudo-papilloedema” and therefore are not useful as a distinguishing feature for papilloedema.

**Idiopathic intracranial hypertension and papilloedema**

Idiopathic intracranial hypertension is a condition of unknown aetiology, typically associated with obesity, that can lead to headaches, optic disc swelling and, in some cases, permanent visual loss (56). OCT allows accurate and objective monitoring of this condition, not only with respect to the degree of optic disc swelling present, but also with quantification of resultant optic neuropathy.

The OCT pRNFL measurements can be used to serially monitor the degree of optic disc swelling present over consecutive visits (figure 8), which is a far more reliable and sensitive method of evaluating change than comparisons of clinician-assessed Frisén grades (57, 58). Furthermore, whilst reduction in disc swelling and pRNFL thickness may be a result of treatment success or general improvement, it may also be a result of worsening axonal loss from disease progression. OCT mGCIPL thickness measurements play a critical role here for distinguishing between these two entities. Reduction in pRNFL thickness with preserved mGCIPL thickness indicates treatment success with preservation of the neuroaxonal structure, whereas reduction in both indices indicates worsening optic neuropathy and treatment failure or fulminant IIH.
EDI-OCT may demonstrate PHOMS in papilloedema. EDI-OCT also enables visualisation of Bruch’s membrane, and peripapillary deformations of Bruch’s membrane surrounding the neural canal due to a differential pressure between the retrobulbar optic nerve and vitreous cavity can also be useful in the diagnosis and management of patients with intracranial hypertension. In raised intracranial pressure, there can be an upward deflection of Bruch’s membrane towards the vitreous (59, 60) (figure 9), and after treatment there can be normalisation to a downward deflection of the Bruch’s membrane complex (59, 61). There is anecdotal evidence that this can be seen straight after lumbar-puncture induced lowering of intracranial pressure, much sooner than fundal appearances of papilloedema, and indeed even OCT pRNFL thickness changes (62,63).

OCT is also helpful in differentiating between vision loss in papilloedema due to optic neuropathy, or due to retinopathy such as subretinal macular fluid or choroidal folds. Decreased vision due to the latter two are generally more benign and reversible, and treatment may not need to be as aggressive as is the case in optic neuropathy (64).

Lastly, functional overlay is not uncommon in patients with IIH. OCT analysis of mGCIP thickness as an objective marker of established optic neuropathy provides additional insight, along with the results of other tests such as kinetic perimetry and visual evoked potentials, as to whether functional overlay may play a role in explaining poor visual function.

Anterior ischaemic optic neuropathies (AION)
OCT plays a similar role in the assessment of ischaemic optic neuropathies as it does in assessment of optic neuritis. Acutely, optic disc swelling can be diagnosed and quantified
using pRNFL thickness measurements. Within 2 months, 80% of patients subsequently show pRNFL thinning, with progressive thinning and thus optic disc atrophy occurring between months 2 and 4, with stability typically reached at month 6 (65-67). As with optic neuritis, thinning of the mGCIPL has been shown to occur much earlier than pRNFL thinning. It can occur as early as 2.2 days after onset of symptoms, and thinning of the mGCIPL density has been shown to be present in 62.5% of eyes at presentation (68). Longitudinal assessment of the mGCIPL layer offers a reliable and objective tool in monitoring of these patients (figure 10). Furthermore, the pattern of mGCIPL loss often indicates the degree and pattern of visual field loss (69-71), which in non-arteritic ischaemic optic neuropathy (NA-AION) is often altitudinal.

OCT can also be used to distinguish between a chronic branch retinal artery occlusion and NA-AION. Whilst both can cause an altitudinal visual field defect and corresponding pallor of the optic nerve in the post-acute phase, retinal artery occlusions will cause thinning of the entire inner retina including the INL, whereas in NA-AION the thinning will be limited to the pRNFL and mGCIPL. As retinal artery occlusions require a thromboembolic work-up and NA-AION do not, it is important to clinically distinguish between the two. In clinical practice it is helpful to compare the visual field with the macular OCT. The anatomical defect seen in the mGCIPL thickness should match the functional deficit depicted in the visual field.

**Compressive optic neuropathies**

Compression of the anterior visual pathway most commonly occurs at the level of the optic chiasm, for example by pituitary lesions, but compression of the optic nerve can also occur elsewhere, for example by meningiomas or in the orbit by the enlarged muscles of thyroid
eye disease. OCT plays a role, not only in diagnosis and monitoring of compression, but also in prediction of visual outcomes after decompression surgery.

Compression causing optic nerve damage can be detected on OCT earlier than may be visible on fundoscopy. Both pRNFL and mGCIPL thinning can be detected and quantified on OCT imaging, with mGCIPL analysis being particularly sensitive for detecting compressive optic neuropathy. Not only may mGCIPL thinning occur earlier than pRNFL thinning (72), but in some studies it has also been shown to occur before standard automated perimetry changes (73, 74). A hemianopia on perimetry can present as a hemi-macular atrophy on the OCT.

The pattern of pRNFL and mGCIPL loss may be helpful in detecting some compressive optic neuropathies. In pre-chiasmatic unilateral optic nerve compression, asymmetry in pRNFL and mGCIPL thickness between eyes gives a clue to the location of the compression. Lesions compressing the optic chiasm superiorly or inferiorly predominantly compress the decussating nasal fibres, resulting in retrograde RNFL loss on the nasal and temporal sides of the optic disc. This can be identified clinically as bow tie or band optic atrophy (75, 76). Whilst multiple studies have shown patients with band atrophy to have pRNFL loss in all 4 quadrants around the optic disc, not just the horizontal band (77-81), there is greater proportional thinning nasally and temporally in patients with bitemporal hemianopia from chiasmal compression (78) (figure 11). The pattern of mGCIPL loss is more consistent, with binasal thinning occurring typically in chiasmal compression (73,82). Homonymous mGCIPL thinning is seen in optic tract and lateral geniculate nucleus injuries, not only from compressive lesions but also from vascular ischaemia or demyelination (83). Post-
geniculate lesions can also lead to homonymous mGCIPL thinning through trans-synaptic retrograde degeneration, but this OCT pattern does not occur acutely and may take over a year to develop (84).

OCT can be helpful in differentiating compressive optic neuropathies from normotensive glaucoma. In glaucoma, pRNFL thinning tends to be vertical and the mGCIPL thinning tends to respect the horizontal meridian (82), distinct from that seen in chiasmal compressive lesions.

OCT may potentially play a role in monitoring and management decisions in paediatric intrinsic optic pathway gliomas, which are typically managed conservatively with treatment initiated on documented visual decline. However, reliably measuring visual function using visual acuity and visual fields in young children can be difficult, and optic nerve glioma size on MRI is poorly predictive of visual function (85). OCT pRNFL thinning has been shown to be sensitive and specific for vision loss in this group of patients (86), although this finding has been challenged (87). Further research is needed to fully support the use of longitudinal OCT assessments to aid therapeutic decision-making.

OCT is helpful in assessing visual dysfunction and predicting post-surgical visual outcomes in compressive optic neuropathies. Both pre-operative pRNFL thickness (62, 88) and mGCIPL thickness (74, 89, 90) should be considered, and one parameter is not necessarily superior to the other as prediction of visual outcomes is multifactorial, depending also on lesion location, duration and severity of compression (46). Both have been shown to correlate
with visual field loss and visual outcomes post-surgical decompression, with lower chance of complete visual recovery when there is greater pRNFL and mGCIPL thinning.

**Optical coherence tomography angiography (OCTA) in neuro-ophthalmology**

OCT technology continues to improve and evolve, and the availability of optical coherence tomography angiography (OCTA) is one such advancement. OCTA provides non-invasive imaging of the perfused vascular network of the neurosensory retina and optic nerve head. Technical details are outlined in table 3.

In neuro-ophthalmology, OCTA assessment of the peripapillary radial capillary network, a layer not visible on traditional fluorescein angiography (91), can be useful. Peripapillary radial vessels are located in the RNFL, superficial to the inner retinal plexus. These vessels are susceptible to fluctuations in intraocular pressure and their density has been shown to correlate with RNFL thickness (92). For instance, in glaucomatous optic neuropathy, reduced perfusion, reduced vessel density and pre-laminar blood flow of the optic nerve head correlate with pRNFL, mGCIPL, visual field mean deviation, and visual field index (93).

Reduced peripapillary radial capillary network perfusion has been demonstrated in demyelinating optic neuritis (94) and ischaemic optic neuropathies (95). The difficulty here is that OCTA is not reliable in a swollen disc. Moreover, partial reperfusion of peripapillary vascular flow has been linked to visual acuity improvement in NA-AION (96). The association of absent focal perfusion alongside superficial peripapillary capillary dilation has been advanced as a potential diagnostic sign for arteritic AION (97). In established cases of
Leber hereditary optic neuropathy and autosomal dominant optic atrophy, reduction in peripapillary perfusion is consistently observed (98,99).

**Conclusion**

OCT imaging has revolutionised neuro-ophthalmic practice and with ever advancing technology the scope for further applications will continue to expand. We have described its application in the diagnosis, monitoring and prognostication of a number of more common neuro-ophthalmic conditions, including demyelinating, inflammatory, ischaemic and compressive optic neuropathies, optic disc drusen and raised intracranial pressure. This list is not exhaustive, and there are a multitude of other neuro-ophthalmic conditions that also benefit from the capabilities that OCT imaging affords with respect to diagnosis and management. The use of computer-generated information, however, must always be reviewed for potential inaccuracies or artefacts, correctly and critically interpreted, and integrated into the clinical context. This is an exciting area in both neuro-ophthalmic research and clinical practice, and as OCT technology continues to improve, it will undoubtedly enhance our understanding of the pathophysiology of neuro-ophthalmic diseases in the future.

**Conflicts of Interest**

The authors declare no conflicts of interest.
Table 1: Anatomy of the anterior visual pathway

<table>
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<tr>
<th>Light energy reaching retina is converted by photoreceptors into an electrochemical signal</th>
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<tr>
<td>Signal is relayed to retinal ganglion cells via bipolar, horizontal and amacrine cells</td>
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<tr>
<td>Axons of ganglion cells travel in retinal nerve fibre layer (RNFL) and converge at optic nerve head (ONH):</td>
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<tr>
<td>- Foveal ganglion cell axons travel to temporal aspect of optic nerve in papillomacular bundle (caeco-central projections)</td>
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<tr>
<td>- Temporal ganglion cell axons form arcuate bundles that travel above and below the fovea, crossing the horizontal meridian in a small delta area behind the foveola, finally entering the superotemporal and inferotemporal aspects of the optic nerve</td>
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<tr>
<td>- Nasal ganglion cell axons enter the nasal portion of the optic nerve</td>
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<tr>
<td>Retinal ganglion cell axons travel posteriorly through the lamina cribrosa where they become myelinated and are called the optic nerve</td>
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<tr>
<td>Distal to proximal rearrangement of positions of the ganglion cells axons occurs in the first section of the optic nerve:</td>
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<tr>
<td>- Macular ganglion cell axons move to the optic nerve centre</td>
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<tr>
<td>- Peripheral temporal fibres become positioned more temporally, both superior and inferior to the macular fibres</td>
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<tr>
<td>- Nasal fibres remain in the nasal past of the optic nerve</td>
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<tr>
<td>The two optic nerves reach the optic chiasm, typically located 10 mm above the pituitary gland, in the sella turcica within the sphenoid bone</td>
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<tr>
<td>Decussation of axons from the optic nerves takes place in the chiasm</td>
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<tr>
<td>Each optic tract contains axons from the ipsilateral temporal retina and the contralateral nasal retina</td>
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<tr>
<td>Fibres synapse in the ipsilateral lateral geniculate nucleus (LGN)</td>
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<tr>
<td>Peripapillary retinal nerve fibre (pRNFL) thickness analysis</td>
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<td>-----------------------------------------------------------</td>
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<tr>
<td>Circular cross-sectional retina scan obtained at diameter of 3.4-3.5mm, centred around the optic nerve head (ONH)</td>
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<tr>
<td>Spectralis SD-OCT (Heidelberg Engineering, Heidelberg, Germany): average pRNFL thickness displayed in four quadrants. Temporal, superior, nasal and inferior and sectoral thicknesses are measured at each of the 12 clock-hours or in 16 equal sectors</td>
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<tr>
<td>Cirrus HD-OCT (Carl Zeiss Meditec, Jena, Germany): deviation map generated where RNFL measurements are compared at each pixel with age-matched normative database, and location under lower 95% of normal range highlighted. Quantitative optic nerve head parameters are provided in the centre panel of the scan report</td>
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<tr>
<td>Other devices display the RNFL thickness as colour-coded thickness maps of the entire peripapillary region (useful in the assessment of small, localised areas of thinning outside the sampling location)</td>
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Table 3: Technical details of optical coherence tomography angiography (OCTA)

<table>
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<tr>
<th>OCTA generates motion-contrast images, that arise due to variable backscattering of light from the vascular and neurosensory tissue in the retina</th>
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<tr>
<td>Scans are performed at the same location and temporal changes are measured, permitting the differentiation of moving particles from static tissue. In the retina, movement is largely generated by red blood cells</td>
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<td>Motion detection can be gathered by amplitude decorrelation or phase variance</td>
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<td>Amplitude decorrelation detects differences in amplitude between two different OCT B-scans</td>
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<td>Phase relies on emitted light wave properties and the variation of phase derived from the interception with moving objects</td>
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<td>After Fourier transform, the OCT signal contains amplitude (intensity) and phase information</td>
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<td>Two averaging methods are then applied to reduce background noise from normal eye movements, namely the split spectrum amplitude decorrelation technique and volume averaging</td>
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<td>The images, with a size between 3 mm² to 12 mm², provide information from the superficial retinal plexus (including the superficial radial peripapillary capillaries and inner vascular plexus within the ganglion cell layer), the deep retinal plexus, the outer retina, and the choriocapillaris</td>
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Figure 1:
Horizontal spectral-domain optical coherence tomography (SD-OCT) B-scan images of the optic nerve head (ONH) in a healthy 33 year old female referred for assessment of non-pathological retinal pigmentary changes. The diameter of the optic disc is defined as the distance between the edges of the retinal pigment epithelium (RPE). The perpendicular orientation of the retinal nerve fibre layer (RNFL) is lost as the nerve fibres blend with the optic nerve head. Abbreviations: BM = Bruch’s membrane; EZ = ellipsoid zone; ELM = external limiting membrane; ILM = internal limiting membrane; CC = choriocapillaris; OS = outer segment; IS = inner segment; ONL = outer nuclear layer; OPL = outer plexiform layer; INL = inner nuclear layer; IPL = inner plexiform layer; GCL = ganglion cell layer.
Figure 2:
Spectral-domain optical coherence tomography (SD-OCT) peripapillary retinal nerve fibre layer (pRNFL) report in a healthy asymptomatic 27 year old female. The RNFL thickness falls within normal range in all segments (TMP = temporal; SUP = superior; NAS = nasal; INF = inferior).
Figure 3:
Spectral-domain optical coherence tomography (SD-OCT) thickness map report of the right eye showing macular ganglion cell layer (mGCL) density and thickness analysis (1, 2.22m 3.45 mm volume scan) after segmentation in a healthy asymptomatic 27 year old female.
A 36 year old male with relapsing-remitting multiple sclerosis (MS) presented with visual complaints and was found to have a right inferior incongruous visual field defect. The spectral-domain optical coherence tomography (SD-OCT) macular ganglion cell layer (mGCL) analysis shows a corresponding pattern of left hemi-retinal loss. This is indicative of left-sided post-chiasmal demyelination (left optic tract or radiation), with MRI images confirming left optic radiation volume loss.
Figure 5

A 44 year old female with sero-negative neuromyelitis optica (NMO) presented with right eye optic neuritis. The spectral-domain optical coherence tomography (SD-OCT) imaging demonstrates progressive thinning of the peripapillary retinal nerve fibre layer (pRNFL) and macular ganglion cell layer (mGCL) after presentation, shown here at 3 weeks and 3 months post episode.
Figure 6

Optic disc drusen (ODD) visible on fundus autofluorescence (A) with corresponding morphology demonstrated on enhanced depth imaging spectral-domain optical coherence tomography (EDI SD-OCT) (B). ODD are seen as signal-poor structures with a partial hyperreflective margin.
Figure 7

Enhanced depth imaging spectral-domain optical coherence tomography (EDI SD-OCT) shows a peripapillary hyper-reflective ovoid mass structure (PHOMS) in a 37 year old who also has optic disc drusen (ODD).
Figure 8

A 23 year old female presented with headaches and was found to have bilateral optic disc swelling, and after investigation was diagnosed with idiopathic intracranial hypertension (IIH). Spectral domain optical coherence tomography (SD-OCT) imaging demonstrates bilateral peripapillary retinal nerve fibre layer (pRNFL) thickening at presentation (A). There was improvement in symptoms and pRNFL thickening after commencement of acetazolamide and weight loss strategies at 4 months after diagnosis (B).
Figure 9

A 47 year old male presented with headaches and was found to have bilateral optic disc swelling, after investigation deemed secondary to raised intracranial pressure. Enhanced depth imaging spectral-domain optical coherence tomography (EDI SD-OCT) demonstrates upward deflection of Bruch’s membrane (arrows).
Figure 10

A 70 year old female presented with headache and reduced vision in the right eye. She was diagnosed with right arteritic anterior ischaemic optic neuropathy (A-AION) due to giant cell arteritis (GCA). Spectral domain optical coherence tomography (SD-OCT) imaging demonstrates progressive thinning of the macular ganglion cell layer (mGCL) from onset, shown here at 3 weeks and 8 weeks post presentation.
Figure 11

Spectral-domain optical coherence tomography (SD-OCT) peripapillary retinal nerve fibre layer (pRNFL) report in a 64 year old male with bilateral established optic neuropathy secondary to a pituitary adenoma causing compression of the optic chiasm.

Reference database: European Descent (2009)
References


47. Lee KM, Woo SJ, Hwang JM. Differentiation of optic nerve head drusen and optic
disc edema with spectral domain optical coherence tomography. Ophthalmology
2011;228:971-977

48. Bassi ST, Mohana KP. Optical coherence tomography in papilledema and
pseudopapilledema with and without optic nerve head drusen. Indian J Ophthalmol.
2014;62:1146-1151

49. Kulkarni KM, Pasol J, Rosa PR, Lam BL. Differentiating mild papilledema and buried
optic nerve head drusen using spectral domain optical coherence tomography.
Ophthalmology. 2014;121:959-963

50. Johnson LN, Diehl ML, Hamm CW, Sommerville DN, Petroski GF. Differentiating optic
disc edema from optic nerve head drusen on optical coherence tomography. Arch
Ophthalmol 2009;127:45-49

51. Sarac O, Tasci YY, Gurdal C, Can I. Differentiation of optic disc edema from optic
nerve head drusen with spectral-domain optical coherence tomography. J
Neuroophthalmol 2012;32:207-211

Drusen Studies Consortium Recommendations for Diagnosis of Optic Disc Drusen

imaging optical coherence tomography of optic nerve head drusen. Ophthalmology
2013;120:1409-1414

54. Traber GL, Weber KP, Sabah M, Keane PA, Plant GT. Enhanced Depth Imaging
Optical Coherence Tomography of Optic Nerve Head Drusen: A Comparison of Cases


80. Monteiro ML, Cunha LP, Vessani RM. Comparison of retinal nerve fiber layer measurements using Stratus OCT fast and regular scan protocols in eyes with band atrophy of the optic nerve and normal controls. Arq Bras Oftalmol 2008;71:534-539


