Optical coherence tomography in compressive lesions of the anterior visual pathway

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Introduction

Optical coherence tomography (OCT) has revolutionized the clinical management of retinal and optic nerve disease. The strict anatomical structure of the retina and maintenance of retinotopic organization within the afferent visual pathway means that OCT is also relevant to central nervous system disease.

OCT allows objective segmental analysis of the retinal layers at the macula. The retinal ganglion cell (RGC) layer can be measured at the macula and the RGC axons can be measured as the retinal nerve fiber layer (RNFL) at the optic nerve. These RGC axons form into the optic nerve, traverse the optic chiasm and synapse at the lateral geniculate ganglion whilst strictly conforming to retinotopic organization. Compression of these axons anterior to the geniculate synapses can cause retrograde changes to the RGCs which can be quantified with OCT. As axons are more compact in the anterior visual pathway than the visual cortex, lesions affecting the anterior regions can cause substantial loss of visual function.

Compression of the anterior visual pathway most commonly occurs at the level of the optic chiasm. Mass lesions affecting this area include pituitary macroadenomas, craniopharyngiomas, Rathke’s cysts and aneurysms of the internal carotid and ophthalmic arteries. However, compression can also occur from other lesions including meningiomas of the tuberculum sella (and contiguous structures) and metastases. Within the orbit, compression can occur from infiltrative neoplastic lesions, optic nerve sheath meningiomas, systemic inflammatory conditions (i.e., ANCA positive disease or sarcoid), or the enlarged muscles of thyroid eye disease.

The anatomical changes on OCT and their relationship to visual function in patients with anterior visual pathway compression is explored in this review. The role of OCT as a non-invasive tool to guide diagnostic and treatment decisions for lesions compressing the optic nerve and chiasm to minimize loss of visual function is discussed.
a prognosticator for visual recovery after treatment of these compressive lesions, is also detailed.

**OCT as an ocular imaging technique**

OCT is a non-invasive method of structurally assessing microscopic damage to RGCs and their axons by performing high-resolution cross-sectional imaging of macula and peripapillary retinal tissue. Low-coherence near-infrared light is transmitted to the retina and the magnitude and echo time delay of backscattered light from the retina is measured to construct a cross-sectional, three-dimensional, real-time tomographic retinal image. Time domain OCT has axial resolution up to 10 μm and scanning speeds of 400 A-scans per second. Spectral domain OCT is a more current technology with digital axial resolution of 4–6 μm and scan speeds of 50,000–85,000 A-scans per second. The techniques of OCT are explored in more detail in the ‘Overview of OCT’ in this focused issue.

OCT measurements are compared to a normative database but these comparisons do not necessarily consider age, sex, or race variables as Caucasian middle-aged subjects make up the bulk of the normative population. Ocular variables such as long or short axial length and optic disc area also alter RNFL measurements. Intraocular disease can hinder the accuracy of OCT such as media opacities, coexistent retinal disease, and other optic disc abnormalities like drusen. Machine factors such as segmentation inaccuracies and scan quality need to be scrutinized to minimize measurement error. Different OCT machines such as Spectralis (Heidelberg, Germany), Cirrus (Zeiss, Germany) and RS-3000 (Nidek, Japan) have different measurement protocols, so patients need to be reviewed on the same machine using the same scanning protocol for accurate comparisons over time.

**OCT patterns of change with compressive lesions of the anterior visual pathway**

Based on the specific anatomical course of RGC axons, compressive lesions along the anterior visual pathway can create certain predictable patterns on OCT. These patterns can sometimes detect visual pathway compression that may not be initially apparent on neuroimaging, or by standard ophthalmic assessments.

Axons at the level of the RNFL do not cross the horizontal meridian. The nerve fibers originating in the nasal hemiretina directly enter the nasal optic nerve. The macula, which lies temporal to the optic disc, has the highest density of RGCs which direct their axons through the “papillomacular bundle”. The remaining axons of the temporal retina approach the optic disc by curving around the papillomacular bundle and thus are compressed into the superotemporal and inferotemporal sectors of the optic nerve. Within the optic nerve, the papillomacular fibers travel centrally, therefore external compression will typically cause a loss of peripheral vision. At the optic chiasm the nasal fibers, which supply the temporal visual field, decussate to join the temporal fibers of the contralateral optic nerve to form the optic tract. The optic tract fibers carry visual information from the contralateral hemifield of each eye.

As the RGC axons are unmyelinated anterior to the lamina cribrosa, reduction in RNFL thickness can be attributed to axonal loss (1). The pathophysiological basis of retrograde RGC degeneration and axonal damage from anterior visual pathway compression is not well known. Mechanical axonal destruction, ischemia and metabolic mechanisms have all been suggested (2-4). Damage to the axons may occur through axon loss, axoplasmic stasis, blockage of conduction and/or demyelination.

**OCT assessment of lesions involving the optic chiasm**

The chiasm is a common site for compression. Lesions abutting the optic chiasm superiorly or inferiorly will predominantly compress the decussating nasal fibers which results in retrograde RNFL loss on the nasal and temporal sides of the optic disc, clinically identified as bow tie or band atrophy of the optic disc (5,6). As a clinical sign, band atrophy can often be difficult to detect with ophthalmoscopy, whereas OCT segmentation techniques can quantify patterns of axonal loss objectively based on RNFL thinning.

Multiple studies have confirmed that patients with band atrophy tend to have RNFL loss in all quadrants around the optic disc, not just along the horizontal band (6-11). Monteiro et al. did not detect more RNFL thinning in the temporal and nasal regions compared to the vertical quadrants (12). However, ROC area under the curve analysis by Moura et al. showed preferential nasal and temporal RNFL loss in patients with band atrophy (13). Danesh-Meyer et al. also showed greater proportional thinning nasally and temporally in patients with bitemporal hemianopia from chiasmal compression (8).

Global macular ganglion cell layer (GCL) thickness
is also reduced in patients with anterior visual pathway compression, especially nasally (13-15). Some have compared the sensitivity of peripapillary RNFL versus macular GCL thickness in detecting visual pathway damage from chiasmal compression. Moura et al. found no difference in the degree of thinning between different sectors of the macular GCL and peripapillary RNFL (13). However, some studies have found macular RGC thinning to be more sensitive than RNFL loss in detecting chiasmal compression (16,17). Tieger et al. identified that the nasal RNFL and GCC thinning in patients with chiasm compression were both statistically significant, yet the effect size was much greater for GCC thinning (16). This discrepancy between RNFL and macular GCL thinning may reflect RNFL measurement variability in the peripapillary region due to variables such as optic disc area. Figure 1 demonstrates macular GCL thinning with normal RNFL in a case of left optic nerve compression.

Macular microcystic changes in the inner nuclear layer may also be seen in anterior visual pathway compression.

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**Figure 1** OCT for a patient with left optic nerve compression from an aneurysm (Cirrus HD-OCT 5000, Carl Zeiss Meditec Inc., Germany). This patient had normal visual acuity (6/6 and N5 in both eyes), color vision and visual fields. The RNFL was of normal thickness in both eyes (A,B). However, the macular GCL is thinner in the left eye (D) compared to the right eye (C) due to compression from an intracranial aneurysm. OCT, optical coherence tomography; RNFL, retinal nerve fiber layer; GCL, ganglion cell layer.
Inner nuclear layer thickening has also been reported (18). Figure 2 shows inner nuclear layer microcysts in a case of craniopharyngioma and giant prolactinoma where severe anterograde RNFL/RGC loss developed. Microcysts have also been found in optic neuritis (multiple sclerosis and neuromyelitis optica), glaucoma, Lebers hereditary optic neuropathy, dominant optic atrophy, and optic disc drusen (20-23). This finding has not been shown to be a disease specific biomarker but may rather reflect the severity of RGC loss and retinal thinning (24).

OCT angiography has been increasingly used to study perfusion of the macula and peripapillary retina at the level of the capillary microvasculature. A recent study found reduced circumpapillary and macular vessel density in eyes with band atrophy which correlated with RNFL thinning, GCL thinning and visual field loss (25). A smaller study of four patients with chiasmal compression also found reduced vessel density in peripapillary areas which correlated with visual field defects (26). The true benefit of OCT angiography as a biomarker in Neuro-Ophthalmological testing requires further research.

**OCT and compressive lesions of the pre-chiasmatic optic nerves**

Pre-chiasmal compression can also cause distinctive OCT changes. In unilateral optic nerve compression, asymmetry in RNFL and GCL thickness between the eyes is a major clue to the location of compression. RNFL may be increased from optic disc swelling or reduced in optic atrophy (27). Loo et al. looked at optic nerve compression by anterior visual pathway meningiomas (28). These eyes had significantly thinned peripapillary RNFL in all quadrants except temporally. However, RNFL can be normal in prechiasmal/chiasmal compression as exemplified in more than half the meningioma patients in another study (29). Also, Sibony et al. used OCT to identify an inward deformation of the retinal pigmented epithelial layer in patients with optic nerve sheath meningioma which worsened with larger tumors closer to the globe (27).

Apart from the aforementioned studies, mainly case reports document specific OCT changes in anterior optic nerve compression within the orbit. Convex retinal bowing and chorioretinal folds have been reported in retrobulbar cavernous hemangiomas and these OCT changes can persist even post tumor resection (30-32).

Figure 3 demonstrates a case of unilateral optic nerve sheath meningioma which presented with optic disc swelling. Whilst the OCT RNFL was thickened, GCL analysis showed thinning. In this case, OCT pointed towards a chronic pathogenic process of the anterior optic nerve.

**Use of OCT to differentiate compressive optic neuropathies from glaucoma**

In an Ophthalmology clinic setting, OCT can be an important diagnostic tool in distinguishing patients with compressive lesions from normotensive glaucoma which can both present with an enlarged cup to disc ratio. As band atrophy can be difficult to identify clinically, the predominantly horizontal peripapillary RNFL thinning is an important clue in compressive lesions which contrasts to the vertical RNFL thinning in glaucomatous optic neuropathy (33). Similarly, GCL loss in the nasal and temporal areas of the macula in chiasmal compressive lesions differs from the GCL thinning seen in glaucoma which tends to respect the horizontal meridian (14). Asymmetry in RNFL and GCL thinning between the eyes is also useful in distinguishing optic nerve compression from glaucoma. Figure 4 highlights two cases where OCT was used to identify optic nerve compression in patients who were mistakenly referred for glaucomatous optic disc cupping.

**Relationship between OCT and visual function**

Sellar masses can lead to visual dysfunction by causing progressive and often painless deterioration of visual acuity, color vision, and visual field. OCT augments the analysis of visual field testing in anterior compressive lesions by providing a structural-functional comparison. This is particularly important given the high test-retest variability and poor reliability of perimetry.

RNFL loss has shown to correlate with the severity of visual field loss from chiasmal compression (8). Sectoral analysis has shown strongest correlation between visual field mean deviation and temporal RNFL thinning (34,35). However, it is unwise to rely on RNFL alone as there are reports of severe visual field loss with intact peripapillary RNFL. This may occur in cases with more acute compression as axonal dysfunction may precede structural loss of axons as detected by OCT.

Macular GCL thickness has shown a greater relationship than RNFL thickness with visual field loss. Monteiro et al. found a stronger correlation (R, 0.65–0.78) for macular...
Figure 2 Inner nuclear layer microcysts in patients with chiasmal compression. (A-D) It shows OCT images of a 45-year-old man with a craniopharyngioma (RS-3000, Nidek Co. Ltd., Japan). (E-H) It shows OCT images of a 20-year-old female with a giant prolactinoma (Spectralis, Heidelberg Engineering Inc., Germany). (A) Right macula with microcysts; (B) left macula with microcysts; (C) right and left optic disc RNFL with severe thinning; (D) macula GCL thickness; (E) right macula with microcysts; (F) left macula with microcysts; (G) right optic disc RNFL with severe thinning; (H) left optic disc RNFL with severe thinning. OCT, optical coherence tomography; RNFL, retinal nerve fiber layer; GCL, ganglion cell layer.
Figure 3 OCT of a 59-year-old female with right optic nerve sheath meningioma (Cirrus HD-OCT 5000, Carl Zeiss Meditec Inc., Germany). Initially the right optic disc was swollen with a thickened RNFL (A) but the right macula had an atrophic RGC complex (C). One-year post radiotherapy, the right optic disc became atrophic with severe RNFL thinning especially superotemporally (E) and the macular GCL thinning worsened (G). The right visual field showed inferonasal changes corresponding to the OCT RNFL (I,J). The left eye remained unaffected with a normal RNFL and GCL thickness (B,D,F,H) and normal visual field (K,L). OCT, optical coherence tomography; RNFL, retinal nerve fiber layer; GCL, ganglion cell layer; RGC, retinal ganglion cell.
Figure 4 OCT detection of optic nerve compression in two patients with disc cupping (Cirrus HD-OCT 5000, Carl Zeiss Meditec Inc., Germany). Asymmetry in the RNFL and GCL thickness between optic discs prompted investigations for optic nerve compression instead of glaucoma. Images A-D are from a patient with left optic nerve sheath meningioma. (A) Right eye normal RNFL; (B) left eye thin RNFL; (C) axial T1 MRI of the corresponding left optic nerve sheath meningioma; (D) coronal T1 MRI of the left optic nerve sheath meningioma. Images E-I are of patient with a right frontal meningioma invading the right optic canal. This patient was mistakenly referred for glaucoma management based on disc cupping and ocular hypertension. (E) right eye thin RNFL; (F) left eye normal RNFL; (G) right eye thin GCL; (H) left eye normal GCL; (I) T1 axial MRI of the corresponding right frontal meningioma. OCT, optical coherence tomography; RNFL, retinal nerve fiber layer; GCL, ganglion cell layer.
thickness than RNFL thickness (R, 0.60) with visual field loss (36). Moura et al. identified a correlation between nasal average macular thickness and visual field temporal mean deviation (R²=48%) (13). The stronger correlation with macular GCL thinning is plausible because it represents the central area of the visual field.

Different studies have shown RNFL and macular GCL thinning in patients with chronic mass lesions that radiologically appear to be compressing the anterior visual pathway despite no change in visual acuity or visual field (8,14,16,37). This suggests that the sensitivity of OCT supersedes perimetry testing. Thus, like in pre-perimetric glaucoma, OCT may have a role in identifying visual pathway damage from compressive lesions prior to functional loss.

Electrophysiological testing through visual evoked potentials (VEP), multifocal VEP (mfVEP) and pattern electroretinography (pERG) can quantify dysfunction along the visual pathway in compressive disease which can augment the interpretation of OCT structural analysis. VEP measures electrical signals received by the occipital cortex and the amplitude and velocity of these signals are diminished in lesions compressing any component of the visual pathway. In chiasmal compression, studies show reduced or delayed P100 during temporal hemifield stimulation which reflects dysfunction of the crossed RGC axons (crossed asymmetry distribution) (38). Topographical changes on mfVEP correspond to visual field defects in chiasmal compression (8,39,40). However, a relationship between VEP, mfVEP and OCT has not been shown, most likely because these electrophysiological tests reflect entire visual pathway function whereas OCT specifically focuses on RGC loss (41,42). PERG is thought to reflect RGC function in the central retinal area and in chiasmal compression, the N95 and b-wave can reduce (43). These changes have not shown correlation with OCT RNFL and GCL thinning (15,44), although multifocal pERG has (45).

**OCT and compressive visual pathway lesions in pediatric patients**

The use of OCT in children has been challenging. There is no normative data and data capture is dependent on patient age and cooperation. However, its role in monitoring optic nerve gliomas, the most common pediatric central nervous system tumor, has been increasingly studied.

Anatomical studies have shown that there is a decline in RNFL thickness with an increase in total glioma volume (46). However, the relationship of RNFL thickness with visual function has been less clear. In a study of children and young adults aged from 6 to 21 years with optic nerve glioma, RNFL thickness did not correlate well with visual acuity or visual field damage (46). The same research group has previously found RNFL to be predictive of vision loss (47).

Children with optic nerve gliomas who have normal RNFL, have demonstrated normal visual acuity and visual fields which can help with prognostication but limits the use of OCT in monitoring for progression before functional deficits occur (48,49). Another study showed RNFL thinning in patients with glioma growth regardless of vision change, meaning that OCT may be able to detect subclinical changes (50).

There has been growing interest in the use of OCT to identify choroidal abnormalities in neurofibromatosis associated optic nerve gliomas. These abnormalities are detected with the use of infrared reflectance retinography on spectral domain OCT (51).

**OCT and visual prognostication after surgical intervention**

RGC axonal structural and functional recovery post decompression surgery is not well understood. Visual field recovery likely reflects recovery of dysfunctioning axons. Many factors have been explored as predictors of visual acuity and visual field post-operative recovery such as symptom duration, age, optic disc appearance, severity of preoperative visual field loss, surgical technique, tumor size and volume and pattern electroretinography, however the results have been variable (28,52-56). Pre-operative OCT findings have proven to be the most consistent prognosticator.

Multiple studies, including a meta-analysis, have shown that preoperative RNFL thinning is a predictor of poor visual field recovery (8,29,35,57-59). The pooled odds ratio in the meta-analysis was 15.61 (95% CI: 4.09–59.61) for field recovery in eyes with normal RNFL compared to those with abnormal RNFL (57). In a large prospective study, visual field recovery was better in the patients with normal RNFL compared to those with abnormal RNFL (81% of eyes versus 37% respectively over one year postoperatively) (35). Garcia et al. showed that eyes with greater nasal, rather than global, RNFL thickness had more peripheral visual field recovery postoperatively (60). Jacob et al. demonstrated that temporal RNFL thinning below
the 5th percentile was associated with poorer visual field recovery (59).

Multiple studies have also identified a strong association between preoperative GCL thinning and poorer visual field recovery (16,34,61). Tieger et al. found a greater correlation of visual field mean deviation with GCL than RNFL thickness ($R^2 0.25$ vs. $0.15$ respectively) (16).

In chiasmal decompression, eyes with normal RNFL have greater improvement in visual acuity as well (8). 98% of the eyes with normal RNFL achieved acuities 6/12 or better compared with 88% of the thin RNFL eyes postoperatively. Danesh-Meyer et al. suggested a presurgical threshold RNFL of 75 μm is associated with a worse postoperative visual prognosis. Loo et al. suggested a RNFL threshold of 70 μm but the focus was on optic nerve more than chiasmal compression (28).

RNFL and macular GCL thickness do not change significantly post-operatively despite visual acuity and field recovery over months to years (8). Thus, further thinning of these parameters can be useful in monitoring for recurrent compression of any residual tumor. However, the direct effect of surgery on OCT measurements needs to be considered when monitoring.

**Conclusions**

OCT has an emerging role in the management of patients with space occupying lesions compressing the anterior visual pathway. Peripapillary RNFL and the macular RGC complex have both proven important in diagnosis, preoperative prognostication and post-operative monitoring of these patients. The ease of access, resolution of detail, reliability and clinical objectivity of OCT heighten the need to incorporate it into routine neuro-ophthalmic assessment. Small study populations and tumor heterogeneity limit the external validity of many OCT studies. However, more research and technological advancement in OCT will shed more light into its role in areas such as pediatric visual pathway gliomas and indications for surgical tumor decompression.

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**Footnote**

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