OCT & OPTIC NERVE

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Like many complementary examinations in ophthalmology, optical coherence tomography (OCT) resulted from a collaboration between ophthalmologists and orthoptists: this book is also based on such a collaboration. I want to thank the orthoptics team of Quinze-Vingts Hospital, especially Frédérique Brion, Audrey Payeras, Cybelle Blatrix and Meddy Metref. Thanks to their efforts, I have been able to present most diseases of the optic nerve, and to compare OCT with other complementary examinations, particularly visual field assessment and photography of the optic nerve. I would also like to thank Professor Philippe Denis, President of the French Society of Ophthalmology, for his help, and in particular for providing many interesting examples.

Knowledge has no meaning unless shared, and Laboratoires Théa and Carl Zeiss have agreed to write, publish and distribute this book free of charge. I am very grateful to them.
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**Atypical features of the retina possibly suggesting optical neuropathy**

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Optical Coherence Tomography (OCT) is a relatively new technique that produces images of biological tissue by measuring the reflection of light from the structure being examined. Depending on the wavelength used, the resolution is in the range of 1 to 15 µm, which is at least twice as high as can be achieved with the best conventional methods such as magnetic resonance imaging (MRI) or high-resolution ultrasonography. However, the main limitation of OCT is that it can only be used to examine structures that allow the passage of sufficient light to obtain a reflected image. The eye is clearly an ideal organ in this respect because many ocular structures, including the cornea, lens, vitreous humour, the neurosensory layer of the retina, and the anterior layers of the iris, are mainly or partially transparent. Furthermore, other highly reflective surface structures, such as the retinal pigment epithelium, can also be studied using OCT.

In 10 years, OCT has assumed a key position in diagnosis and follow-up of retinal disorders, particularly those affecting the macula. This is due to the highly informative nature of OCT images, which can provide a level of detail comparable with that of histological examination. In addition, the latest generation of OCT devices can not only reproduce this descriptive aspect, but also provide quantitative measurements of ocular structures such as the retina and its component layers.

Retinal imaging by OCT has thus emerged as a crucial technique in the investigation of glaucomatous disease, as it is now possible to quantify the thickness of the nerve fibre layer (also known as the optical fibre layer or the retinal nerve fibre layer, RNFL), which is predominantly affected in glaucoma.

This quantitative approach can also be applied to the optic nerve, allowing measurement of the area of the neuroretinal rim and cavity volume. Hence, all parts of the eye that are potentially affected in glaucoma have become accessible with OCT. By obtaining quantitative data from glaucomatous eyes, and comparing these with normal controls, the existence of glaucoma can be confirmed and the severity assessed during follow-up.

The iridocorneal angle can also be analysed, albeit less accurately because the back of the pigmented iris epithelium is inaccessible to OCT. In addition to glaucoma, OCT also appears to be useful in assessing other optic nerve diseases and, to some extent, central nervous system pathologies that affect the eye.
As with any new technology, OCT results must be interpreted with care, because data acquisition is complex and does not involve simple photography of the structures studied. It is good practice to examine the quality of the results, not only in general, but also in detail when apparently abnormal findings are obtained. Once these pitfalls have been avoided, a new field of investigation (and, above all, discussion) opens up. What is the clinical meaning of the observed pathology? How should a worsening of optical fibre pathology on OCT be interpreted if other examinations are stable? How can we explain a finding that a particular structure is no longer changing on OCT examination when clinical findings show that the glaucoma is indisputably worsening?

OCT is advancing so quickly that it is impossible to list its applications definitively: any review is certain to become obsolete quite quickly. Already indispensable for retinal analysis, OCT is now becoming essential for glaucoma and other optic neuropathies.

This monograph aims to give an update on the use of OCT in glaucoma and pathologies of the optic nerve. The first chapter covers the basic principles and interpretation of OCT and is intended to help the reader interpret OCT images. Subsequent sections cover glaucoma and other eye diseases, especially neuro-ophthalmological diseases, because these can mimic glaucoma or be a source of confusion if they are comorbid with glaucoma.
OCT works by passing a beam of laser-generated light, with a wavelength in the infrared range (approximately 840 nm), through a tissue and analysing the reflected fraction. Logically, when a beam of light passes through a structure, one part of the light will continue its path (especially if the structure is quite transparent), one part will be absorbed by the structure, one part will be reflected in all directions and the final part will be reflected towards the emission zone.

It is this last part of the light that is analysed in OCT: it corresponds to between one-billionth and one-millionth of the incident light, so it is very weak.

Principles of OCT
Sound waves propagate relatively slowly (300 m/sec in air), and hence the echo time of sound waves can be recorded directly, as in ultrasonography. However, the high speed of light (300,000 km/sec) means that the time taken for incident light to be reflected back to its source is too short to measure, being approximately 30 femtoseconds (30 x 10^-15 sec). OCT uses the principle of interferometry to analyse this delay. An incident wave is divided into two, one part being projected onto a plane mirror and the other onto the eye. The two waves thus created are reflected; the wave projected onto the mirror returns as a single echo, whereas the wave projected on to the eye returns as multiple echoes depending on the structures it has passed through. These waves are compared by an interferometer which measures the coherence between them (hence the term optical coherence tomography). This measurement is used to deduce the thickness of the ocular structures that the light has passed through.

A simple measurement of coherence at a given point and a given depth is called an A-scan. With time domain OCT, the mirror moves repeatedly at different depths to analyse the different layers of the retina (Figure 2).
The fastest time domain scanners can measure 17,000 A-scans per second, and these one-dimensional measurements are then combined in 2 dimensions to obtain a cross-sectional view of the retina at a given location (known as a B-scan). Time domain OCT is limited by the need for a moving mirror, because the eye has to remain relatively fixed and thus an examination cannot reasonably take more than a few seconds.

A very different technology, spectral domain OCT, dramatically improved the quality of OCT images. Instead of measuring the coherence between two waves, this approach measures the interference spectrum between the two reflected beams of broad-spectrum waves, which is analysed mathematically by Fourier transformation (Figure 3). The advantage of this “spectral/Fourier domain” technology is that rays reflected from different depths can be analysed simultaneously, rather than successively at each depth. The reference mirror therefore no longer has to move, which accelerates the process 50–100-fold.

As a result, much more accurate measurements (around 2 µm) can be made during a reasonable examination time. Nevertheless, the examination is not instantaneous because, although the different depths of the retina can be studied at the same time, the laser beam still has to scan the different regions of the retina successively, and hence the quality of the results may still be affected by eye movement.
As described above, OCT measures the reflection of light from different structures in the eye. This reflection is particularly strong when there are sharp edges between two media with different refractive indices, for example, in the cornea. Moreover, reflections are sharper when the structure under investigation is perpendicular to the incident light (rather like a reflection in a window). In general, the nature of the reflection (only a part of which is directed back towards the incident source) is determined by heterogeneous structures, such as the cell membranes, nuclei, cytoplasm, and neuronal axons, that produce small changes in the refractive index. The more reflective a structure is, the more red it appears on OCT imaging. The most reflective structures in the retina are the RNFL, the pigment epithelium and the interplexiform layers. From measurements of the reflected light, OCT measures the thickness of the structure under investigation, taking into account the return time of the incident beam and the refractive index of the structure.

In structures that contain melanin, which is a strong absorber of light, a combination of reflection and absorption results in an exponential reduction in the power of the incident beam. As a result, OCT cannot be used to study structures distal to the retinal pigment epithelium, because the outer layer is highly reflective, whereas the basal region is highly absorbent.

It is important to note that OCT images are not direct images of the retina, but rather a mathematical reconstruction that is subsequently transposed into images of the ocular fundus. This can be understood by looking at the changes in appearance of optical nerve fibres on OCT, for example, when they change direction to exit from the eye. The reflection of light from the RNFL is displayed in red, indicating that the fibres are perpendicular to the incident light, and this colour changes as the fibres change direction on moving towards the optic nerve, becoming more parallel to the incident light (Figure 4). An awareness of such changes can help in interpreting ambiguous OCT findings to determine whether or not structural changes are present.
When OCT is used to analyse a layer of cells, such as the macular ganglion cell complex, it must be remembered that the measurement corresponds to the entire layer (ganglion cells, support cells, interstitial fluid, etc.) This explains why the macular ganglion cell complex does not completely disappear in patients with glaucoma, because some support cells remain even when the condition is advanced.
In this section, we shall review the use of OCT to investigate those structures that are modified in glaucoma and optic nerve pathologies: the optic nerve head, the parapapillary RNFL and the macular ganglion cell complex.

**Optic nerve**

The initial problem when imaging the optic nerve is how to define the contours of the optic nerve, and the start of the cavity. It is important that these are adequately defined because, when the disc is uneven, measurements may be distorted by erroneous identification of the reference plane. For OCT investigations, an arbitrary reference plane is set at 150 µm above the level of the peripapillary pigment epithelium: everything below this plane is considered to be a cavity, whether physiological or not (Figures 5 and 6). This approach allows measurement of papillary dimensions such as:

- the width and area of the neuroretinal rim at different meridians
- the size of the disc and cavity
- the cup/disc ratio.

**Methods for analysing ocular structures with OCT**

The normal configuration of the eye on OCT.

*Figure 5*
Optical fibres around the papilla

OCT examination of the papillary region involves two processes:
- the two most refractive zones of the retina, the front of the RNFL and the pigment epithelium, are identified
- the point at which the reflection of the RNFL decreases sharply is determined, allowing measurement of its thickness.

The speed of the spectral domain OCT allows measurement of a cube around the optic nerve, but the zone to be imaged must be chosen with care. The thickness of the optical fibres is measured 3.4 mm from the centre of the optic disc (Figure 7). This distance was chosen because it offers the best compromise between the thickness of the RNFL and interindividual variability: the thickness of the RNFL is greatest close to the disc, and thus small changes in thickness should be readily measurable, but inter-individual variability in the location of the vessels and the shape of the disc is also greatest in this region. Conversely, measurements made further from the disc are more consistent between individuals, but the fibre layer narrows quickly and small changes become less detectable.

Figure 6:

The papillary region is uneven, and the pink line does not clearly mark the boundary between the retina and the cavity.

OCT findings in an eye with severe myopia.
However, measurements made 3.4 mm from the centre of the disc may not reveal the first signs of disease, and hence follow-up may be sub-optimal with measurements at this point; the zones closest to the disc are likely to provide better measures of disease progression during follow-up.

**Macular ganglion cell complex**

The thickness of each layer of the retina, particularly the ganglion cell layer in the macular region, can be measured using spectral domain OCT. This zone is particularly interesting because it accounts for around 30% of the entire thickness of the retina.

Conversely, the foveal region is of less interest because it contains fewer ganglion cells (Figure 8). Most OCT scanners measure across a range that includes the internal interplexiform layer, ganglion cell bodies, and ganglion cell axons passing above them (Figure 9).
With some devices, such as the Cirrus™ HD-OCT (Carl Zeiss), the measurement can be refined to improve the location of cell damage by eliminating the ganglion cell axon layer and measuring only the internal interplexiform and ganglion cell body layers.
OCT is an objective examination, so is less prone to patient-related errors. However, it is subject to errors relating to either the imaging process or the structure of the eye.

**Signal reception errors**

Although the quality of imaging is clearly crucial in OCT, it is not straightforward because less than one-millionth of the incident light from the laser is reflected towards the sensor. Imaging quality is expressed in terms of the signal strength, which must be greater than 6/10 with the Cirrus™ HD-OCT (Carl Zeiss) or 50 with the Optovue (RTVue). A poor signal is often responsible for underestimation of the RNFL thickness.

**Errors related to head positioning**

As described above, the strength of the reflected light measured in OCT is dependent on small changes in the refractive index of the tissue, and the orientation of the structure under investigation relative to the incident beam. As a result, changes in the position of the head, and thus in the orientation of the axis of the eye, can lead to variable results.

This can be seen in the example shown in Figure 10. With correct presentation, the parapapillary RNFL (in red) is thick, whereas when the depth setting is incorrect (for example, too distant, or when the head is tilted back), this layer seems to decrease steeply, and the thickness of the red layer is decreased.

However, it is important to note that such errors do not always result in inadequate signal quality, and therefore may not be easily detected.
Principles of OCT

Figure 10a

OCT image of the RNFL, with correct head positioning.

Figure 10b

OCT image from the same eye, with the head tilted back. The apparent thickness of the RNFL is reduced.

Figure 10c

OCT image from the same eye, showing poor adjustment in myopia. Again, the thickness of the RNFL is reduced.
Because time domain OCT scanning takes about 1.5 s, artefacts related to eye movements may also contribute to poor quality images (Figure 11). OCT software can correct for this, or even for eye tracking during the examination, but care should nevertheless be taken to avoid such artefacts as far as possible.

Spectral domain OCT may also be subject to artefacts when the eye is incorrectly positioned (Figure 12), or when there are significant variations in retinal depth in patients with severe myopia.

The upper border of the OCT image shows an immediate reflection, and hence no weaker signals can be detected. However, if the eye is not positioned correctly, an even closer image will be detected in the mirror, which will appear completely inverted, with the outer part of the eye appearing at the top and the inner part at the bottom. In this situation, the retinal layers would also be completely inverted.
Errors related to the optics of the eye

The optics of the eye may also affect the quality of OCT images. Opacification of the posterior capsule tends to reduce the apparent thickness of the RNFL. Conversely, the use of multifocal lenses does not affect the results at either macular or parapapillary levels. The size of the optic disc does not affect the thickness of the optical fibres unless they are very small; in this case, signs of disease may be observed irrespective of whether a genuine abnormality is present. Conversely, myopia may reduce the apparent thickness of the optic fibre layer due to measurement errors resulting from an increased axial length. An increased axial length increases the circumference of the circle 3.4 mm from the centre of the optic nerve; as a result, the optical fibres are more dispersed than in a normal eye, leading to a reduction in apparent thickness even though there is no reduction in the number of optical fibres. This phenomenon is not present to any significant degree in the macular ganglion cells, and it may therefore be preferable to measure this region in patients with severe myopia.
Errors related to ghost images

OCT may not always distinguish between adjacent and highly reflective structures, such as the posterior hyaloid and the inner boundary of the retina.

Figure 13
A further cause of potential error is that in some cases the thinning of the RNFL seen in most optic nerve pathologies is masked by structural oedema resulting from other process, such as traction phenomena (Figures 13 and 14).

**Figure 14**

OCT images from a patient with bilateral glaucoma, with visual impairment skimming the central 10°. The right hand image shows damage to the ganglion cell complex (arrow), while the left hand image shows a macular hole. Traction at this level, and the resulting oedema, are masking the thinning of the RNFL, which appears to be normal.
Many different OCT devices are currently available: in the United States, for example, more than 30 devices have obtained a marketing authorisation. It is therefore not possible to give a detailed review of all the available OCT devices.

In France, the most commonly used devices are the Stratus and Cirrus™ HD-OCT (Carl Zeiss). and the Optovue (RTVue), and most of the examples given in this book were obtained using these two instruments. Because the most relevant parts of the eye affected by glaucoma are the papillary and parapapillary, and the macular ganglion cell complex, OCT findings from these regions obtained with these two instruments will be reviewed in detail.
Presentation of the optic nerve head and retinal nerve fibre layer (Optic Disc Cube)

In each of the following examples (Figures 15 and 16), OCT results are presented with the right eye (OD) on the left and the left eye (OS) on the right. For statistical analyses, yellow indicates a significant anomaly at the < 5% level, and red indicates a statistical anomaly at the < 1% level. Where patterns are small and difficult to analyse, these will be enlarged on a separate page.

The values obtained are compared with those of a patient of the same age with an optic disc of the same size. The size limits of the optic disc range from 1.3 mm² to 2.5 mm²; outside these limits, all measured parameters are shown in grey, because they are not compared to a standard. For each value, this number appears on a white background if it is 5% of the best values, and on a green background for 90% of cases.
Patient’s full name. The patient’s date of birth is important because the results are compared with age-dependent standards.

Signal strength: an indication that the results have been correctly acquired. Values higher than 6/10 are required. However, even if the signal is strong, the results may be inaccurate if the eye is incorrectly positioned.

Vertical section of the posterior pole. Imaged elements are identical to the horizontal section.

Position of the retina 3.4 mm from the optical centre, corresponding to the region where the retinal nerve fibres are studied.

Cirrus™ HD-OCT (Carl Zeiss)

Image of the ocular fundus in which the statistical values of anomalies in the optical fibre are coloured (yellow: $p < 0.05$; red: $p < 0.01$). The optic disc is indicated, as well as the zone 3.4 mm from the centre of the disc where the thickness of parapapillary nerve fibres is measured. It is important that this circle is not located in a zone of peripapillary atrophy, because the reflectance of the pigment epithelium would be evaluated incorrectly and the results potentially distorted.

Horizontal section of the posterior pole. It is important to ensure that the retina does not touch the upper part of the square, as this would distort the results. The reflectance of the different layers are shown, as well as the boundaries of the retina and Bruch’s membrane. Two small dots indicate a zone 150 µm above Bruch’s membrane that defines the start of the cavity.

Thickness of the optical fibre layer. Warmer colours indicate thicker layers. A “clock hands” appearance, centred on the disc, is normal.

Figure 15
Average thickness of the optical fibre layer (RNFL);
Symmetry of this layer between the 2 eyes (RNFL Symmetry);
Area of the neuroretinal rim;
Area of the optic disc;
Mean cup/disc ratio;
Mean vertical cup/disc ratio;
Cavity volume.

Regional thickness of the neuroretinal rim (TEMP: temporal; SUP: superior; NAS: nasal; INF: inferior). The right eye is shown as a continuous line, the left eye as a dotted line; normal zones are indicated in green, and abnormal zones in yellow and then red. This colour coding makes it easy to locate a notch.

Regional thickness of the optical fibre layer 3.4 mm from the centre of the optic nerve. The right eye is shown as a continuous line, the left eye as a dotted line; normal zones indicated in green, abnormal zones in yellow and then red. This colour coding makes it easy to locate.

Representation of optical fibre thickness for each eye in sections 6 hr wide. In glaucoma, the most affected zones are the upper and lower zones.

Representations, of optical fibre thickness in each eye, in sections 2 hr wide. In glaucoma, the most affected areas are located at 7 hr for the right eye and 5 hr for the left eye.

Figure 16
Presentation of the ganglion cells (macular cube)

Figure 17 shows typical OCT findings for the ganglion cell layer, with the right eye (OD) on the left and the left eye (OS) on the right. For each value, the number appears on a white background if it is within 5% of the best values, and on a green background for 90% of cases. Yellow indicates an anomaly at the < 5% level, and red indicates an anomaly at the < 1% level.

Signal strength indicates whether the results have been correctly acquired: values should be higher than 6/10. However, even if the signal is strong, the results may be inaccurate if the eye is incorrectly positioned.

Thickness of the RNFL. The thicker this layer is, the warmer the colour. A “clock hands” appearance, centred on the disc, is normal.

Presentation of the macular sector, excluding the fovea, showing the average thickness in each zone.

Mean and minimum values for the thickness of the ganglion cell/internal interplexiform layer, excluding the ganglion cell fibre layer. The minimum value corresponds to the thinnest 1° sector.

Horizontal scan of the macular passing through the fovea.

Statistical values of anomalies in this layer. Yellow: $p < 0.05$; red: $p < 0.01$

Figure 17

OCT imaging of the ganglion cell layer (macular cube).
Presentation of the parapapillary RNFL and the optic nerve

Figure 18 shows the results of an OCT examination of the parapapillary RNFL and optic nerve, performed using the Optovue scanner. The figure presents findings in a single eye, showing the overall results for the optic nerve head and the parapapillary region, with statistical analysis.
General presentation of the macular ganglion cells and parapapillary RNFL

OCT findings can be summarized in a single page, without showing the actual OCT images (Figure 19). This presentation includes most of the elements shown in the previous figures, including:

- RNFL thickness
- statistical analysis of RNFL thickness
- ganglion cell complex
- optic nerve head parameters.

**Statistical representation of the macular ganglion cell complex (optical fibre axons, ganglion cells and internal interplexiform layer).**

**Thicklness of the RNFL in the different regions: the warmer the colour, the greater the thickness.**

**Interocular differences in parapapillary RNFL thickness**

**Regional thickness of the RNFL, with statistical values: normal zones are green, abnormal zones are yellow and then red.**

**Figure 19**

General presentation of OCT (Optovue) findings in the macular ganglion cells and parapapillary RNFL.
Glaucoma is a slowly progressive optic neuropathy that is usually associated with ocular hypertension. Progressive deformation of the optic nerve head resulting from hypertension leads to cavity formation and destruction of the retinal nerve fibres passing through the cribriform lamina. This neuropathy in turn leads to visual field impairment. The progressive nature of glaucoma suggests it should be theoretically possible to detect structural changes in the retina and optic nerve before the condition becomes clinically apparent with visual field impairment. Prior to the introduction of OCT, however, this could not be confirmed with the available techniques.
With the new generation of spectral domain OCT devices, structural changes in the eye can often be detected before the onset of visual impairment. However, in some cases, this may not be the case: visual impairment may become apparent before any structural impairment can be detected. The reason for this is that each eye has between 800,000 and 1.2 million optical fibres, and so in many cases significant structural damage can be tolerated without visual impairment. Conversely, in an eye with a smaller number of fibres, visual impairment may become apparent more quickly.

**Optic nerve**

At the optic nerve head, OCT measurements show successive changes in:
- the vertical thickness of the neuroretinal rim,
- the overall area of this rim,
- the vertical C/D ratio (Figure 20).

Although these parameters are useful for the detection of glaucoma, they are less effective in differentiating early glaucoma from moderate glaucoma.

**RNFL**

The OCT measurements parameters that best differentiate normal subjects from patients with early glaucoma are:
- RNFL thickness in the lower temporal zone,
- RNFL thickness in the lower quadrant,
- average RNFL thickness (Figure 20).

There is some evidence that measurements in the upper temporal sector would be as effective in discriminating between individuals with and without glaucoma as measurements in the lower temporal quadrant.
OCT analysis in glaucoma

Mean RNFL thickness

Optic nerve parameters:
- Vertical thickness of the neuroretinal rim,
- Overall area of the neuroretinal rim
- Vertical C/D ratio

RNFL thickness in the lower quadrant
RNFL thickness in the lower temporal zone

Figure 20
OCT imaging of the optic nerve and RNFL in glaucoma.
Macular ganglion cell complex

Macular ganglion cells can only be studied using spectral domain OCT, and hence imaging of these cells in glaucoma is a more recent innovation. Early glaucoma is mainly characterized by changes in the mean minimum thickness and lower temporal thickness (Figure 21). For OCT examination, the macula is divided into 360 sectors, each one degree wide, and the mean minimum thickness is that of the thinnest sector.

Figure 21
Spectral domain OCT measurements of the macular ganglion cell complex in glaucoma.
Optic nerve or retinal nerve fibres: which should be examined first in glaucoma?

As we have seen, OCT can provide detailed information about a variety of ocular structures, and this raises the question of which parameters are most reliable for the early detection of glaucoma: in particular, is it better to examine the optic nerve head or the parapapillary ganglion fibres? When a large number of optical fibres are present, the thickness of this layer can be easily measured, but as the thickness decreases the posterior edge of the layer becomes more difficult to differentiate from adjacent structures, and hence measurements become more variable. Conversely, in the presence of a thin ganglion cell layer, it would be easier to determine optic nerve parameters: OCT is very effective in measuring optic nerve parameters by analysis of the end of Bruch’s membrane.

However, the appearance of the optic nerve varies greatly between individuals, making statistical comparisons difficult, particularly when the optic nerve is irregular, such as in patients with myopia or dysversion of the optic nerve head.

In general, it is easier to detect preperimetric glaucoma in the RNFL, although in rare cases optic nerve analysis may be preferable. In some patients, analysis of macular ganglion fibres is as effective as RNFL measurement.
Patients with isolated optic nerve impairment

Due to the marked variability in the appearance of the optic nerve between individuals, which is not adequately represented in the standard databases used by OCT software, isolated optic nerve impairment may occur. In this situation, the retinal nerve fibres appear normal at both parapapillary and macular ganglion cell level, and the visual field is also normal (Figure 22).

Optical fibre analysis is preferred in such cases, although it is still desirable to monitor the progressive changes in optic nerve parameters over time in the same patient.
All results are normal, except from an abnormal cavity volume for each eye. Glaucoma is not established. Simple monitoring is recommended in such cases.
The term preperimetric glaucoma denotes glaucoma that is detected by the presence of structural impairment or changes in early visual field detection tests, such as the blue-yellow visual field or the frequency doubling technology (FDT) Matrix, while conventional visual field assessments (e.g. Humphrey-type automated perimetry or Octopus® perimetry) are normal. OCT findings in various presentations of preperimetric glaucoma are shown in the following figures.

Isolated impairment on OCT

Optic nerve OCT and FDT matrix findings in a 60-year-old patient with ocular hypertonia (23 mmHg). OCT and FDT matrix findings are at the limits of normal ranges.
OCT may be the only complementary examination required in such patients. Simple monitoring is usually sufficient, but treatment is a viable alternative, particularly if the OCT deficit is large.
Preperimetric glaucoma: Impairment of the RNFL and the FDT Matrix visual field

Early visual field impairment detected by OCT and FDT Matrix indicates the beginning of glaucoma, particularly if the deficits are consistent. These deficits may precede the development of conventional perimetry deficits by 5 years.

Optic nerve OCT and FDT matrix findings in a 70-year-old patient with ocular hypertonia (24 mmHg). The optic nerve appears normal, but OCT and FDT matrix show matching impairments of the RNFL in the right eye.

Figure 25
Optic nerve OCT and FDT matrix
Figure 25 shows optic nerve OCT and FDT matrix findings from a 74-year-old patient with ocular hypertension (24 mmHg). The optic nerve is normal, but OCT and FDT Matrix show matching impairments in the RNFL of the right eye. In this patient, there was no central impairment, as assessed by macular cube OCT (Figure 26).

Macular cube OCT analysis in the same patient as in Figure 25. There is no central impairment.

Figure 26
Macular cube OCT analysis
Preperimetric glaucoma: impairment of the RNFL layer, the macula and the FDT Matrix visual field

There is often an impairment of both the RNFL layer and the macular ganglion cell complex in OCT. If the deficit is more present in the parapapillary region, we can assume that hypertension likely plays a major role. Otherwise, we will consider more vascular problems.

Figure 27

OCT and FDT matrix findings in a 55-year-old patient with ocular hypertension (26 mmHg). Automated periphery measurements are normal, but FDT matrix shows bilateral impairment. OCT confirms the presence of this impairment, in both the optic nerve and RNFL.
This situation is illustrated in Figure 27, which shows OCT and FDT matrix findings in a 55-year-old patient with ocular hypertonia (26 mmHg). Automated perimetry findings are normal, but bilateral impairment is apparent on Matrix evaluation. OCT confirms this impairment, both in the optic nerve and the RNFL layer. OCT examination of the macular region shows slight macular impairment on the right side and normal results on the left side (Figure 28). This profile would suggest early hypertonic glaucoma.

Macular cube OCT analysis in the same patient shown in Figure 27. There is little impairment on the right side, and normal findings on the left side. The combination of RNFL, macular and FDT matrix impairment suggests early hypertonic glaucoma.

Figure 28
Preperimetric glaucoma: impairment of the FDT Matrix with normal OCT

It is rare to observe normal OCT findings in a patient who already has perimetric visual field impairment, although such cases could hypothetically occur in glaucoma (Figures 29 and 30). These findings should suggest a more central impairment, and further testing is needed to establish the diagnosis.

OCT and FDT matrix findings in a 60-year-old patient with ocular hypertonia (24 mmHg). The matrix visual field shows marked impairment on the left side, although only slight impairment is visible on OCT.

Figure 29
Preperimetric glaucoma

Macular cube OCT analysis and visual field assessment in the same patient as in Figure 29. The macular ganglion cell complex is normal on the left side, and at the limit of normal on the right side, but the matrix central 10° test results are abnormal.

Figure 30
Early open-angle glaucoma

OCT findings are often consistent with visual field assessments in patients with early or moderate glaucoma (Figures 31 and 32). An OCT finding that impairment is greater in the macular region than in the peripapillary region is not indicative of severe glaucoma, provided the deficit remains moderate.

Optic disc OCT and visual field assessments in a 79-year-old patient with ocular pressures of 22 mmHg in the right eye and 26 mmHg in the left eye. In the right eye, the visual field is normal, and OCT shows a lower temporal deficit typical of early glaucoma. In the left eye, the impairment is more marked and extensive.

Figure 31
In young patients with early or moderate glaucoma, OCT impairment is greater than visual field impairment.

Macular cube OCT and central 10° visual fields in the same patient as in Figure 31. Macular impairment is observed in the left eye, with abnormal values for mean and minimum thickness. The central 10° visual fields are normal.
Early open-angle glaucoma: depth of OCT and visual field impairment in early open-angle glaucoma

The impact of a reduction in RNFL thickness on the visual field depends on the severity of the impairment, and hence the thickness must be analysed accurately in the different regions of the retina. Figures 33 and 34 show OCT findings in a 70-year-old patient with an 8-year history of open-angle glaucoma. OCT and visual field impairments are almost concordant, with more marked changes in both the RNFL and the cavity on the left side.
marked changes in both the retinal nerve fibre and the cavity being seen on the left side. Significant macular impairment is present on both sides, but there is no significant impairment of the right visual field. This is due to the fact that the macular ganglion complex in the corresponding zone is still very thick (> 60 µm); by contrast, on the left side, the thickness in the upper sector is reduced to 44 µm, resulting in a deep lower paracentral scotoma. Such findings explain why impairment may not be noticeable in the central region of the visual field despite marked regional thinning of the macular ganglion cell complex. For this reason, both the minimum and mean thickness of the complex should be measured.

Macular cube OCT and central 10° visual fields in the same patient as in Figure 33. Significant macular impairment is present on both sides (arrow), but there is little impact on the right visual field. This is because the macular ganglion complex is still very thick (> 60 µm) in the corresponding zone, whereas on the left side the thickness in the upper sector is reduced to 44 µm, resulting in a deep lower paracentral scotoma.

Figure 34
Early open-angle glaucoma: predominant OCT impairment in early open-angle glaucoma

In younger patients with early open-angle glaucoma, OCT findings may be altered to a greater extent, compared with older patients, before changes in visual fields become apparent (Figures 35 and 36). This is due to the fact that in younger patients there is a greater reserve of receptor fields in the ganglion cells: in each region of the visual field, several ganglion cells process the same zone, and thus greater damage can be tolerated before visual impairment develops. This reserve decreases with age.

Figure 35

OCT findings in a 31-year-old patient with a 4-year history of open-angle glaucoma. All optic nerve parameters, and most RNFL parameters, are abnormal, but there is little impairment of the visual field.
Early open-angle glaucoma

Macular cube OCT in the same patient as in Figure 35. There is significant macular impairment without loss of visual acuity or impairment of the central visual field.

Figure 36
Early open-angle glaucoma

Primary impairment of the optic nerve without impairment of the RNFL

Importantly, the absence of RNFL layer thinning does not completely exclude the possibility of glaucoma. The presence of a cavity on clinical examination or OCT, associated with mild visual field impairment, should be taken as evidence of true glaucoma.

OCT of the RNFL and optic nerve, and visual field assessments, in a 68-year-old patient with ocular hypertension. In the right eye, OCT and visual field findings unequivocally show the presence of glaucoma. However, in the left eye, the RNFL is normal but all optic nerve parameters are abnormal. In this patient, glaucoma is clearly bilateral, and the slightly abnormal visual fields support this diagnosis.

Figure 37
Early open-angle glaucoma

This is illustrated in Figures 37 and 38, which show OCT findings in a 68-year-old patient with ocular hypertonia. OCT findings from the right eye unequivocally indicate the presence of glaucoma. In the left eye, however, the RNFL is normal but all optic nerve parameters are abnormal; in this case, the diagnosis is supported by the presence of abnormal visual fields (Figure 37). In the presence of a normal RNFL, analysis of the macular ganglion cells can differentiate between isolated hypertonia and early optic fibre impairment (Figure 38).

Macular cube OCT and central 10° visual fields in the same patient as in Figure 37. Bilateral impairment of the macular ganglion cell layer is present, but the central visual field is normal.
Comparison between time domain and spectral domain OCT in glaucoma with incipient open angle

Comparisons between time domain OCT (Stratus OCT, Carl Zeiss) and spectral domain OCT (Cirrus™ HD-OCT, Carl Zeiss) show that the two methods are comparable in terms of their ability to identify incipient glaucoma (Figure 39).

The principal difference between them is that spectral domain OCT is more reproducible, and therefore allows better monitoring than time domain OCT.

Figure 39a
However, the results obtained with the two techniques are not identical; in particular, in patients with severe glaucoma there may be 10–20% variability.

Furthermore, the colour codes used in the displayed results from Stratus OCT and Cirrus™ HD-OCT are not interchangeable and, in general, the measured thickness of the RNFL is lower with the Cirrus than with the Stratus device.
Moderate open angle glaucoma: concordance of OCT findings and visual field testing

In patients with moderate open-angle glaucoma, perfect concordance between OCT and visual field testing is rare, because in many cases OCT can detect more serious impairment than perimetry testing.
Results from a patient with moderate glaucoma are shown in Figures 40 and 41. Visual field testing shows a marked deficit, but OCT shows RNFL involvement only in the lower concordant region of the right eye (Figure 40). Analysis of macular ganglion cells (Figure 41) confirms this impairment, which is concordant with the results of visual field testing, but also shows subclinical impairment in the upper region and in the left eye.
Moderate open angle glaucoma: predominant OCT impairment

In patients with moderate glaucoma, OCT impairment is sometimes more significant than would be expected from visual field testing alone. Hence, in addition to confirming perimetric deficits, OCT can be very useful in the investigation of other regions that appear normal in visual field testing.

OCT findings in the RNFL and visual field test results in a patient with moderate glaucoma. RNFL impairment is seen in both the lower and upper concordant regions of the left eye. Both Matrix visual field testing and OCT findings are normal in the right eye.
An example of this situation is shown in Figures 42 and 43. In this patient, OCT reveals RNFL thinning in both the upper and lower regions (Figure 42), together with subclinical impairment of macular ganglion cells in the upper region (Figure 43).
In patients with advanced glaucoma, OCT findings can confirm the presence of structural damage but the diagnosis is based predominantly on visual field testing. Indeed, beyond a certain point in advanced glaucoma, there is no further decrease in the thickness of the RNFL or macular ganglion cells, despite the presence of optical atrophy, because support structures account for the residual thickness of the RNFL.

**Advanced open-angle glaucoma: concordant impairment on OCT and visual field testing**

In advanced glaucoma, the RNFL regions corresponding to regions of absolute deficit on visual field testing are about 50 µm thick (Figures 44 and 45).
Advanced open-angle glaucoma

Macular cube OCT and central 10° visual fields from the same patient as in Figure 44. Moderate impairment is present in the right eye, whereas all macular regions are affected in the left eye (the eye mostly affected by glaucoma). Note that a difference in thickness of 15 µm to 20 µm separates areas of incipient deficit (right eye) and absolute deficit (left eye).

Figure 45
Advanced open-angle glaucoma: predominant impairment of the visual field

In the most advanced stages of glaucoma, it is difficult to rely on measurements of cell layer thickness, either at the papillary or macular level, because only slight reductions are seen in this situation (figures 46 and 47).

OCT findings in an 84-year-old patient with advanced glaucoma and reduced visual acuity in the right eye. On this side, the visual field mean deviation (MD) is -26dB, compared with -11dB in the left eye. OCT differences between the eyes are less pronounced, with a mean RNFL thickness of 54 µm in the right eye and 60 µm in the left eye.

Figure 46
Macular cube OCT and central 10° visual fields from the same patient as in Figure 46. OCT. Again, there is no clear correlation between visual field deficits and the thickness of the ganglion cell complex: the MD is -20dB in the right eye and -9dB in the left eye, and the mean thickness of the macular fibres is 57 µm and 58 µm, respectively.

Figure 47
Advanced open-angle glaucoma: discordance between findings on OCT and visual field testing

In some cases, only moderate impairment can be seen on OCT despite obviously abnormal visual fields. This could be a genuine finding, or a result of measurement errors due to poor signal quality.

OCT findings in a 65-year-old female patient with almost symmetrical bilateral glaucoma. Despite similar visual field abnormalities in both eyes, the RNFL only appears to be abnormal in the right eye. This could be due to measurement errors associated with serious sparkling synchysis of the left eye, which is well visualised by OCT and leads to decreased signal quality.

Figure 48
In the case shown in Figures 48 and 49, for example, abnormalities in the RNFL are seen only in the right eye, despite the presence of almost symmetrical bilateral visual field impairment. This could be due to the formation of crystalline opacities resulting from liquefaction of the vitreous body (synchysis) in the left eye, which decreases the signal quality in OCT.

Macular cube OCT and central 10° visual fields in the same patient as in Figure 48. Macular abnormalities are shown on OCT in both eyes.
Normal-pressure glaucoma

In normal-pressure glaucoma, impairment of the central 10° of the visual field is often observed. Due to the high number of ganglion cells in this central region, this is reflected in a marked reduction in the retinal ring area, and hence in major hollowing (excavation; also known as cupping) of the optic nerve head in the advanced stage. The same characteristics can be seen on OCT. In addition, an isolated or predominant impairment of the macular region may be present in many cases; indeed, this is characteristic of early-stage normal-pressure glaucoma.

OCT findings from a 60-year-old patient, with an ocular pressure of 15 mmHg without treatment. A slight fasciculate deficit, affecting both the optic nerve and the para-papillary fibres, is observed in the left eye; optic nerve parameters are normal. The results for the right eye are normal.
Normal-pressure glaucoma: isolated fascicular deficit

An isolated deficit may be found in patients with normal-pressure glaucoma, which manifests as a notch in the optical nerve head and the RNFL (Figures 50 and 51).
Incipient normal-pressure glaucoma

In the early stages of normal-pressure glaucoma, the mean RNFL is reduced less than in hypertonic glaucoma (Figures 52 and 53), because the affected fasciculi are finer.}

OCT findings in a 68-year-old female patient with an ocular pressure of 16 mmHg without treatment. OCT shows localised impairment in the right eye: the mean RNFL thickness is not reduced, but there are significant changes in optic nerve parameters.

Figure 52
In the patient shown in Figure 52, the mean RNFL thickness is not affected, but the optic nerve parameters do show significant changes. The RNFL thickness in the macular region may be gradually reduced without producing loss of visual field.

Paracentral scotomas in normal-pressure glaucoma are deep and absolute, and there is only a brief relative scotoma phase.

Figure 53

Macular cube OCT and central 10° visual fields in the same patient as in Figure 52. There is very localised lower macular impairment, concordant with the perimetric deficit.
Normal-pressure glaucoma

Moderate normal-pressure glaucoma

Changes in OCT measurements between normal-pressure glaucoma and high pressure glaucoma become apparent during the early stages of the disease (Figures 54 and 55). When glaucoma is more advanced, resulting in more specific impairments of visual fields, these differences gradually disappear.

OCT findings in a 60-year-old patient with normal-pressure glaucoma. The right eye shows the usual indications of open-angle glaucoma, with a nasal step and arcuate scotoma.

The thickness of the RNFL remains relatively unchanged: a small paracentral scotoma can be seen on the left, which only slightly alters the thickness of the RNFL.

Figure 54
Spectral domain OCT analysis of the macular ganglion cell complex can be used to detect the beginning of impairment, which is reflected in extremely localized reduction of the fibres, with less global macular impairment than in primary open-angle glaucoma.

Figure 55
Closed-angle glaucoma or sequelae of acute hypertonia

Immediately after a glaucoma crisis involving angle closure, OCT findings do not change significantly. However, after, 3–9 months, the RNFL is reduced both locally in the upper and lower regions and globally, whereas the perimeter usually remains unchanged (Figures 56 and 57). It might therefore be interesting to compare OCT findings immediately and some time after the acute crisis to determine whether structural impairment occurs as a consequence of acute hypertonia.

OCT and visual field findings in a 58-year-old patient who experienced a closed-angle glaucoma crisis in the left eye a year previously. The crisis was resolved by medical treatment and iridectomy. Intraocular pressure is 16 mmHg on each side. Matrix and OCT findings are normal in the right eye. In the left eye, the visual field appears normal but, upper region impairment is seen on OCT; optic nerve parameters in the left eye are slightly abnormal.
Even in the absence of an acute crisis and with normal pressure, RNFL thickness in the low temporal region is sometimes reduced in patients with narrow angles, particularly Asian patients. This might reflect nocturnal increases in pressure.

Macular cube OCT findings in the patient shown in Figure 56. There is impairment of the macular region in the left eye, and the mean thickness is abnormal.

Figure 57
OCT can be used to detect and monitor the progression of glaucoma, although because spectral domain OCT is a relatively recent technique few long-term longitudinal studies are available: a 4-year patient follow-up has been reported in one study. Such studies have shown that glaucoma may progress in several ways.

- The most common form of progression is an expansion of the deficit area, with increasing encroachment towards the macula.

- Less frequently, the impairment is deepened or appears in a different region. In order to detect such progression, examination at 2 mm from the centre of the optic nerve offers better discrimination of small changes; the region usually selected for examination, 3.4 mm from the centre of the optical nerve, is too peripheral. In more than 50% of cases, such progressive changes in OCT findings are not confirmed by automatic perimetry testing, a finding which underlines the low correlation between these two examinations, at least over short periods. Conversely, in about 20% of cases, isolated progression in the perimeter is confirmed without changes in OCT parameters.

The Guided Progression Analysis (GPA) report has proposed a glaucoma monitoring programme and an analysis of progression, based on Cirrus HD-OCT findings (Figure 58).

This GPA programme, like perimetry testing, is used to detect significant disease progression. However, it does have important limitations. Glaucoma is considered to be progressive if changes seen in a repeated examination...
Monitoring of open-angle glaucoma

**Figure 58**

OCT parameters recommended for monitoring of glaucoma progression, according to the Guided Progression Analysis (GPA) report.

- **Optical fibre thickness**
- **Changes in the thickness of optical fibres** between examinations, when a first change is detected, are drawn in yellow; if confirmed during the next examination, they are in red.
- **Longitudinal changes** in the global mean thickness of the RNFL in the upper and lower regions (3 traces)
- **Changes in mean cup/disc ratio**
- **Fibre thickness profile** in the various regions (TSNIT), using the same colour code in the event of aggravation
- **Probability of stability or progression aggravation** for fibre thickness and TSNIT (focal progression), average thickness (diffused progression), and mean cup/disc ratio
are greater than variability limits determined in healthy individuals, but it is not known whether this variability is different in glaucoma patients. In one study, an increase in the volume of the RNFL was found in 13% of cases, which is difficult to accept because that would indicate an improvement of glaucoma; undoubtedly, this result is attributable to errors in the initial measurements. The same team carried out a similar study 3 years earlier using OCT Stratus, which showed a reduced ability to detect progression with previous generations of OCT devices. Furthermore, progression of structural and functional impairment does not occur in parallel. Indeed, the relationship between optical fibre loss and visual field deterioration is not linear.

These studies clearly show that automatic perimetry testing is essential for glaucoma monitoring, but OCT also appears to be very useful, at least during the incipient and moderate glaucoma phases. When evaluating glaucoma progression by measuring the reduction in RNFL thickness, the glaucoma stage must also be considered:

- During the initial phases of glaucoma, optical fibre loss may increase without major changes in the visual field.
- As the disease progresses, moderate changes in RNFL thickness are reflected in severe deterioration of the visual field (Figure 59).
- During the later stages, the thickness of the RNFL no longer changes significantly, while the visual field continues to worsen. As noted previously, the persistence of a certain RNFL thickness does not mean that a significant number of ganglion cells remain, but rather that the support cells in this layer are still present.

Simultaneous worsening of results from visual field testing and OCT over 6 months. Mean RNFL in the left eye changed during this period from 78 µm to 74 µm: a 3 µm decrease is considered significant.

Figure 59
Can OCT findings improve in glaucoma?

In practice, OCT findings do not improve over time in patients with glaucoma. After a steep reduction in ocular pressure through surgery, OCT may show an increase in RNFL thickness, but this is a temporary phenomenon that only lasts for about 3 months. Most commonly, apparent improvements in OCT parameters result from improved image quality during subsequent measurements following a suboptimal initial measurement.

Comparisons between spectral domain OCT devices in glaucoma patients

Several spectral domain OCT devices are available, including the Cirrus™ HD-OCT (Carl Zeiss), the 100RTVue (Optovue), and the Spectralis (Heidelberg). These use the same operating principle, but vary in terms of acquisition speed, the ability to monitor eye movements and the retinal layer segmentation method used. Although the different instruments produce similar results, the absolute values obtained may differ: for example, the Cirrus HD-OCT gives lower RNFL values than RTvue, due to differences in the measuring zone between the two devices.

For this reason, absolute values obtained using one spectral domain OCT scanner cannot be extrapolated to another.

OCT and cribriform plate analysis

In addition to impairment of optical nerve fibres, glaucoma is known to cause deformation of the cribriform plate. By using specific software (Enhanced Depth Imaging-Optical Coherence Tomography: EDI-OCT), it is possible to view the cribriform plate deforming towards the rear, and also shifting backwards in relation to the sclera in certain meridians. It is not known whether this is a cause or a consequence of impairment of the plate; however, after surgical reduction of ocular pressure the cribriform plate gains anterior adhesion and thickens again. This analysis appears to be an important development, as it offers the potential to visualize a number of deep components of the optic nerve complex. Conversely, there does not appear to be a great benefit from measuring the thickness of the choroids, which changes only slightly in glaucoma and is unchanged in cases of peripapillary atrophy.
Optic neuropathies lead to deterioration of the axons of both parapapillary and macular nerve fibres, which can be detected by OCT. As a result, OCT has become a very important diagnostic tool in neuro-ophthalmology. In many cases, OCT findings can aid the diagnosis and provide information of the severity of the neuropathy. However, in some cases the OCT profile may resemble that of glaucoma, and hence it is important to understand the OCT findings characteristic of different neuropathies.

Non-glaucomatous optic neuropathies
Analysis of the retinal ganglion fibre layer is of particular interest in patients with multiple sclerosis (MS) because the thickness of this layer correlates better with patients’ functional symptoms (decreased lower visual acuity or reduced visual field) than other examinations, such as magnetic resonance imaging (MRI) 27.

**Acute neuropathy in MS**

During the acute phase of optic neuropathy in MS, the RNFL is sometimes thicker than normal, a finding which reflects slight, sub-clinical, papillary oedema. This oedema is present even when the demyelinating plaque is located in the posterior part of the optic nerve.

Thinning of the RNFL occurs 1–3 months after an acute crisis, and stabilises after approximately 6 months. If no further crises occur, the RNFL subsequently remains stable after 6–8 months. The most marked changes in OCT findings are found in the macular area, where the ganglion cell layer accounts for approximately 34% of the retinal volume. MS-related optic neuropathy is characterized by a reduction of about 35–45% in the thickness of the macular optical fibre layer (i.e. a 20–40 µm reduction from a normal thickness of 110–120 µm) 28. The contralateral eye is also affected in most cases, albeit to a lesser degree (20% reduction in thickness). Importantly, defects in perimetry testing do not become apparent until the macular layer is reduced by approximately 75 µm, and hence it is essential that OCT findings are evaluated according to the dynamics of neuropathy 29: during the acute phase, the absence of layer thinning may provide false reassurance.

Both the papilla and macula are affected by MS. As noted above, during the initial phases the RNFL may be artificially enlarged by papillary oedema, which is seen in macular ganglion cells. However, as the disease progresses, changes in the two structures develop in parallel 30.
Correlations between OCT findings and visual field testing in MS

In patients with minimal evidence of neuropathy on OCT, the visual field and visual acuity generally remain normal after an acute crisis. By contrast, if the thickness of the peripapillary RNFL decreases to approximately 75 µm, decreases in the visual field become apparent (Figures 60 and 61).

OCT and visual field findings in a patient with multiple sclerosis (MS) after an acute attack. OCT predominantly shows impairment in the temporal sectors of each eye, which is not seen with Goldmann’s kinetic perimetry testing. The optic nerves are normal. Temporal impairment, as shown here, is typical of non-glaucomatous neuropathies such as MS-related optic neuropathy.

Figure 60
Location of optic neuropathy in MS

In patients with MS, optic neuropathy is most commonly seen in the temporal quadrant (Figures 60 and 61). This contrasts with the situation in glaucoma, in which impairment is generally seen in the superior or inferior quadrants.
Multiple sclerosis

Important sequelae of MS neuropathy

The progression of MS is associated with progressive deterioration in both visual field and OCT findings (Figures 62 and 63). However, these may not always occur in parallel: in some cases, changes in OCT findings may be greater than changes in visual field testing.

OCT and visual field findings in a female patient with MS and several acute attacks of optic neuropathy. The visual field of the left eye is severely affected: OCT confirms this impairment and also reveals an almost identical change in the right eye. The optic nerves show bilateral optical atrophy, which is more marked on the left. There is no pathological excavation of the optic nerve head.

Figure 62
Multiple sclerosis

Macular cube OCT from the same patient as in Figure 62. Ganglion cell impairment is severe on both sides.

Figure 63
Detection of subclinical MS neuropathy with OCT

Impairment of the visual pathways is almost always present in MS, and hence OCT is a useful tool for the detection of subclinical impairment (Figures 64 and 65) and for monitoring progressive neurological changes 31.

Figure 64

OCT and visual field findings in an MS patient with no known episodes of optic neuropathy. Marked pathology is seen on OCT, while the visual field is normal.
In addition, it could be used to monitor the effects of treatment on the course of MS.

Macular cube OCT from the same patient as in Figure 64. Diffuse impairment of the macular ganglion cells is visible, but visual acuity is normal.

Figure 65
Ophthalmologists are often confronted with acute optic neuropathy of unknown origin. In this situation, it is essential to consider the clinical context when trying to establish whether the pathology is demyelinating. Demyelinating optic neuropathy is characterized by:

- occurrence between the ages of 20 and 50 years
- ocular pain, particularly during eye movement
- strictly unilateral presentation
- progressive deterioration over about 1 week, followed by recovery after 1 month.

OCT and visual field test results from a 30-year-old patient with bilateral blurred vision, without reduced acuity. MRI results did not indicate MS. The ocular fundus shows greater papillary oedema on the left than on the right; this finding is confirmed by OCT, which shows a marked increase in parapapillary RNFL thickness on both sides.
Unfortunately, OCT profile cannot specifically identify demyelinating neuropathologies, which manifest as an initial increase in RNFL thickness resulting from oedema, followed by progressive reduction of the RNFL.

Figures 66 and 67 show OCT findings from a 30-year-old patient with bilateral blurred vision, in whom MRI findings did not indicate MS. OCT showed significant papillary oedema, particularly in the left eye (Figure 66), with severe ganglion cell impairment (Figure 67).
Toxic or nutritional optic neuropathies are characterized by bilateral, symmetrical and painless impairment, which is classically associated with centrocaecal scotoma (Figures 68 and 69). In such cases, OCT often reveals bilateral and gradual impairment of the RNFL. Impairment of peripapillary optical fibres is confined to the temporal region, with relative preservation of the upper and lower sectors.

**Apparent toxic optic neuropathy without papillary impairment**

In the presence of papillary oedema, the parapapillary fibres may appear normal on OCT because the thickness of the cell layer is artificially increased by oedema.
In this situation, there may be a marked contrast on OCT between normal parapapillary optical fibres and major impairment of the ganglion cell complex.
Toxic optic neuropathy with papillary and macular impairment

OCT and visual field test results from a 42-year-old patient with optic neuropathy associated with alcohol and nicotine. Goldmann’s visual fields show distortion of the blind spot on the right and a normal result on the left. OCT reveals bilateral RNFL impairment in the temporal region: the upper and lower sectors are not affected. The optic nerves are normal, and there is no pathological excavation.

Figure 70
In patients with incipient toxic optic neuropathies, impairment may be more evident in the macular ganglion cells, even when visual acuity is preserved (Figures 70 and 71). Such impairment is generally diffuse and symmetrical.
It is sometimes difficult, to distinguish between remote sequelae of anterior ischaemic optic neuropathy (AION) and glaucoma on the basis of OCT findings. Impairment of the RNFL is temporal in both cases 33 but the deficit in AION is more altitudinal than in glaucoma, affecting not only the lower temporal sector (7hr for the right eye, 5hr for the left eye), but also affecting the upper quadrant (Figures 72 and 73) 34.

Figure 72
Macular cube OCT findings in the same patient as in Figure 72, showing global impairment of the left eye. This presentation is perhaps atypical, because a more altitudinal deficit would be expected in a patient with AION.

In addition, the excavation volume is likely to be smaller in AION than in glaucoma. During the acute phase of AION, papillary oedema causes an initial increase in parapapillary fibre thickness, while the ganglion cell layer is unaffected.
Left unilateral uveopapillitis, probably of toxoplasmic origin, in a 19-year-old patient. OCT can determine the severity of papillary and subretinal oedema, and determine the proximity of oedema to the fovea.

In this situation, OCT can be used to detect oedema and intra- and subretinal location. An example is shown in Figure 74.
As we have seen, papillary oedema, of any cause, results in a marked increase in the thickness of the RNFL, which will be reversed as the oedema regresses. The rate of development of sequelae to papillary oedema depends on the cause of oedema, and hence it is important to monitor the time course of changes in the RNFL (Figure 75).

OCT and visual field test findings in a 20-year-old patient with intracranial hypertension due to an expansive intracranial process. OCT reveals increases in the volume of the optic nerve and RNFL thickness; parameters of the RNFL are measured over time to monitor the evolution of papillary oedema.

Figure 75
Isolated papillary oedema without retinal diffusion

In cases of isolated oedema, OCT shows an increase in the volume of the optic nerve, while the RNFL remains normal (Figures 76 and 77). Because the RNFL may appear normal in patients with oedema associated with incipient optic atrophy, long-term assessment is necessary.

OCT and visual field test findings in a 54-year-old patient with intracranial hypertension. OCT reveals an increase in the volume of the optical nerve without associated retinal impairment.
An absence of optic fibre impairment indicates a good prognosis in patients with macular oedema, and re-assessment of the acute episode will confirm this.

Macular cube OCT in the patient shown in Figure 76. Macular OCT findings are normal.
Papillary oedema with subretinal oedema

In the absence of subretinal oedema, it can be anticipated that visual function will improve during regression of papillary oedema. By contrast, the presence of subretinal oedema associated with papillary oedema indicates a less favourable prognosis. OCT can readily distinguish between intra- and subretinal oedema in the parapapillary region (Figures 78 and 79).

OCT and visual field test findings in a 23-year-old patient with decreased visual acuity of unexplained origin. Bilateral papillary oedema is present in the ocular fundus, and the visual fields show enlargement of the blind spot on both sides. OCT reveals an increase in RNFL thickness due to oedema, associated with an absence of optical fibre impairment.

Figure 78
Macular cube OCT and central 10° visual fields in the same patient as in Figure 78. In the macular region, the ganglion cell complex is already altered in the right eye, whereas in the left eye this impairment is hidden by oedema (arrow).

Figure 79
Optic nerve compression is sometimes difficult to identify because atrophy of the optic nerve may be accompanied by secondary excavation. As a result, this condition is sometimes misdiagnosed as glaucoma.
Optic nerve compression is usually unilateral, and hence normal findings in the contralateral eye can help to establish the diagnosis (Figures 80 and 81).

Macular cube OCT from the same patient as in Figure 80, showing diffuse and isolated macular impairment in the left eye.

Figure 81
OCT changes and visual field loss are seen in patients with chiasma impairment (Figures 82 and 83). A favourable prognostic factor for functional recovery after surgery, is the maintenance of an RNFL thickness of at least 80 µm in the affected zone, although postoperative improvement is also seen in patients in whom the RNFL is thinner than this. Thus, in compressive neuropathies, the thickness of the RNFL is an indicator of the probability of clinical improvement after surgery: the thicker the fibres preoperatively, the better the likely recovery.
Importantly, the RNFL is not always affected during early stages of optic nerve or chiasma compression: in some cases, changes in the visual field are the first signs of impairment. This therefore constitutes one of the rare cases where the absence of RNFL thinning does not indicate optical neuropathology.
Amblyopia is characterized by decreased visual acuity in the affected eye. Ganglion cells are present, but non-functional. OCT parameters are only slightly altered (Figures 84 and 85).

OCT and visual field test results in a 48-year-old patient with amblyopia in the left eye (visual acuity 2/10th). Visual field testing shows a nasal deficit of unexplained origin. However, OCT findings are normal, both in the peripapillary region and the optic nerve.

Figure 84
Macular cube OCT and central 10° visual fields in the patient shown in Figure 86. The central visual field of the left eye shows a diffused deficit consistent with a visual acuity of 2/10th. OCT findings are normal.
Perinatal impairment of the central nervous system

Although it might be expected that decreases in the thickness of the RNFL would be seen only when cell bodies or axons in this layer are directly affected by disease, there is evidence that even isolated lesions in the occipital cortex can manifest as a decrease in RNFL thickness on OCT. This situation was first documented in cases of congenital or perinatal pathologies (Figures 86 and 87), but it is not certain whether it also applies to acquired impairments.

Figure 86

OCT and visual field test results in a 42-year-old patient with sequela of epilepsy and slight mental retardation associated with perinatal injury. The visual field is irregular with lower homonymous lateral deficit, but the optic nerves are normal. Although the underlying pathology is central in origin, thinning of the RNFL is seen on OCT.
Macular cube OCT and central 10° visual fields in the same patient as in Figure 86. The central visual field is irregular, and the macular OCT shows bilateral impairment.
Impairment of the central nervous system in adults

Isolated impairment of the central nervous system in adults is reflected in much greater changes in visual fields than in OCT findings (Figures 88 and 89), although OCT parameters may be affected by trans-synaptic degeneration from the occipital lobe to the optic nerve 38.

OCT and visual field test results in a 49-year-old patient. The visual field tests show an unexplained, relatively symmetrical, bilateral deficit in the upper field. The OCT findings are normal, and the optic nerves also appear clinically normal.

Figure 88
Normal OCT findings in a patient with decreased visual acuity or impairment of the visual field can thus be helpful in confirming a neuro-ophthalmological cause of the symptoms.  

Macular cube OCT and central 10° visual fields in the same patient as in Figure 88. There is minor impairment of the ganglion cell complex, whereas there is a very marked perimetric deficit. The underlying cause of these deficits is probably central in origin.
In many situations it is difficult to establish the organic origin of functional symptoms. Examples of such situations include:
- post-cranial trauma
- perimetric deficits of hysteric origin
- ill-defined pathologies.

In such cases, OCT can be used to determine the organic or non-organic nature of the functional deficits (Figures 90 and 91).
Determination of the organic nature of visual impairment

Macular cube OCT and central 10° visual fields in the same patient as in Figure 90. The ganglion cell complex is significantly impaired, confirming the organic nature of the patient’s complaint.

Figure 91
In more general pathologies such as Alzheimer's disease or Parkinson, we also observe a reduction of optical fibres in OCT. In certain cases, this reduction is diffuse and sometimes located in the superior region 39.

In Alzheimer's disease, a reduction of the RNFL and the macular ganglion cell layer is observed, but this is not directly related with the level of cognitive functions.

In Parkinson's disease, a certain relationship between reduction of the optical fibre layer and the functional repercussions of Parkinson's disease has been found.
Atypical features of the optic nerve that can suggest optic neuropathy in OCT

Numerous atypical features of the optic nerve can lead to changes in papillary OCT parameters. In some cases, faulty measurement techniques can result in incorrect diagnoses. OCT measures excavation based on a reference plane, which is arbitrarily set at 150 µm above the level of the peripapillary pigment epithelium: if the pigment epithelium is not analysed correctly, all the results may be wrong. For this reason, papilla sections, indicating the theoretical start of the excavation with a red point, are displayed, enabling the ophthalmologist to determine whether or not the points are placed coherently.
Moderate myopia does not cause changes in OCT parameters (Figures 92 and 93). In the event of peripapillary atrophy, it should be ensured that the measurement 3.4 mm from the centre of the optic nerve is located outside the atrophic region, in order to obtain good quality measurements.

OCT and visual field test results from a patient with myopia (6 dioptres) and mild lower temporal dysversion. There are irregularities in the matrix visual field test results, but OCT findings are normal.
Macular cube OCT in the patient shown in Figure 92. There are ganglion cell complex abnormalities in the right eye. However, this result indicates only that a pathological process is developing.

Figure 93
Papillary dysversion

Moderate dysversion of the papilla does not lead to changes in OCT parameters, particularly if the temporal axis is involved.

However, in some cases a slight nasal deficit may be observed, a feature that does not suggest glaucoma.

Conversely, more severe impairment, particularly affecting the nasal quadrant, can lead to significant changes in OCT parameters, which may be confused with those seen in glaucoma (Figures 94 and 95).

OCT and visual field test results in a patient with bilateral dysversion. The visual field is affected mainly on the right, with changes in RNFL thickness suggestive of glaucoma. No excavation is visible on OCT.

Figure 94
The macular ganglion cell complex is easier to evaluate than the RNFL in patients with dysversion, because it is further from the optic nerve. This can reveal the presence of optical fibre impairment (Figure 95).
Physiological excavation

The large papillae, which are responsible for physiological excavations, are reflected in the preservation of the RNFL and the macular ganglion cell complex on OCT (Figures 96 and 97).

Optic nerve OCT parameters are not usually compared with normative values, because the databases used do not generally include patients with physiological excavation.

OCT and visual field test results in a patient with bilateral physiological excavation. The RNFL is normal. The optic nerve parameters (shown in grey) have not been analysed statistically because the device does not include patients with such large disc areas in its database.
Physiological excavation

In patients with deep, and possibly physiological, excavation, optical fibre parameters should be considered in the diagnosis to a greater extent than optic nerve parameters.

Macular cube OCT from the same patient as in Figure 96. The ganglion cell complex is normal.

Figure 97
In patients with papillary colobomas, the changes seen on OCT affect the opposite side to the coloboma. Thus, the sectors usually affected in glaucoma, particularly the lower temporal region, are not affected if this region is the seat of the coloboma (Figures 98 and 99).
Macular cube OCT in the same patient as in Figure 98. The ganglion cell complex is normal.

Figure 99
Atypical features of the retina possibly suggesting optical neuropathy

Certain macular or retinal pathologies can affect the optical fibre layer at both the macular and peripapillary levels. As a result, OCT changes in these regions do not necessarily indicate direct impairment of the optic nerve. If the initial retinal impairment mainly affects the macula, only changes in the macular ganglion cell complex may be seen in many cases. By contrast, if the impairment is more extensive, OCT changes in all structures may be seen. Two examples - vein occlusion sequelae and pigmentary retinopathy - are discussed here, but the ophthalmologist should be aware that diverse retinal pathologies may be suggestive of optic neuropathy.
Vein occlusion results in destruction of the ganglion cell layer in addition to impairment of the optic nerve. OCT findings will depend on the clinical context.

Figures 100 and 101 show OCT findings in a patient who had experienced upper temporal branch occlusion in the right eye 3 years before examination.
Macular cube OCT and central 10° visual fields in the same patient as in Figure 100. There is diffuse and unilateral impairment of the macular ganglion cell complex.

Figure 101
Impairment of the macular ganglion cell complex is very common in patients with retinal pathologies. It may also be present when the initial impairment is not in the RNFL, for example, in patients pigmentary retinopathy (Figures 102 and 103).
All other sources of macular injury can lead to impairment of the macular ganglion cell complex, including:

- epiretinal membrane
- age-related macular degeneration
- macular oedema of any origin
- acquired or constitutional maculopathies.

Macular cube OCT in the patient shown in Figure 102. There is diffuse and bilateral impairment of the macular ganglion cell complex.

Figure 103
Conclusion

OCT is developing constantly: in the last year, more than 300 articles on “Optic nerve and OCT” have been published in international journals, underlining the innovative character of this technology and its diverse applications in ophthalmology.

This accumulating experience has raised a number of questions. For example, should OCT be used as a basic examination in the context of glaucoma monitoring, or simply to supplement visual field testing and ocular pressure measurement? Can we ignore visual field testing, a restrictive examination that is poorly accepted by many patients? What should be done if OCT findings worsen significantly without changes in the visual field? Conversely, what should be done if the visual field deteriorates but this is not confirmed on OCT? At present, it seems reasonable to conclude that OCT has become a key examination in the investigation of glaucoma, but that it cannot replace visual field testing (particularly automated perimetry testing).

Similar questions are being asked about other optic neuropathies. Should OCT eventually replace visual field testing? And what should be done if OCT reveals signs of impairment in the absence of clinical signs?

OCT is still developing in terms of image precision and quality, and at the same time the technology is being extended to the investigation of new structures, such as the cribriform plate. Other developments are possible, in particular the ability to analyse both layer thickness and content, can be foreseen. This would be very useful, for example in counting the number of residual ganglion cells in patients with optic nerve diseases while excluding the support cells responsible for the persistence of this layer even after total destruction of the visual fibres.

Similarly, long wavelength (≥ 1000 nm) laser beams, such as those used in swept source-OCT, can be used to study structures beyond the pigmentary epithelium of the retina, albeit at the cost of poorer image quality. This approach would allow visualization of the choroid and the region beyond the cribriform plate of the papilla. In the future, these technologies will provide an understanding of complex pathologies that involve both modification of peripapillary cladding structures and cribriform plate impairment.


