Spectral-domain Optical Coherence Tomography Imaging of Age-related Macular Degeneration

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Abstract
Spectral-domain optical coherence tomography (SD-OCT) high-speed, high-resolution imaging of the macula has become an essential tool for evaluating dry and wet age-related macular degeneration (AMD). This high-speed, high-resolution imaging strategy, combined with new innovative algorithms, permits reproducible measurements of the anatomical changes associated with AMD, which include drusen, geographic atrophy (GA) and choroidal neovascularisation (CNV). To visualise drusen and larger retinal pigment epithelial detachments, an algorithm was developed for Cirrus HD-OCT (Carl Zeiss Meditec, Dublin, CA) to detect elevations in the retinal pigment epithelium (RPE). To visualise GA, an algorithm was developed to provide en face visualisation of the macula, which easily identifies and measures areas where the RPE has been lost. To visualise CNV and the associated macular fluid, an algorithm was developed to measure the retinal thickness between the internal limiting membrane and the RPE. No other imaging modality is capable of qualitatively and quantitatively following patients at all stages of AMD, which makes SD-OCT the ideal instrument for following disease progression and the effect of therapies.

Keywords
Age-related macular degeneration, drusen, geographic atrophy, retinal pigment epithelium, retinal pigment epithelial detachment, choroidal neovascularisation; spectral-domain optical coherence tomography

Age-related macular degeneration (AMD) is a common cause of irreversible vision loss among the elderly worldwide. It is estimated that approximately 30 % of adults older than 75 years have some sign of AMD and that approximately 10 % of these patients have advanced stages of the disease. AMD can be classified in two forms: non-neovascular (dry) and neovascular (wet or exudative). The non-neovascular form accounts for 80–90 % of cases, while the neovascular form accounts for 10–20 % of cases, but the neovascular form is responsible for the majority of the cases with severe vision loss (80–90 %). Time-domain optical coherence tomography (OCT) is a non-contact optical technique that images the retina and the macular area. Spectral-domain OCT (SD-OCT), also known as Fourier-domain or high-definition OCT, is a higher-speed, higher-resolution technique that permits even better visualisation of the retina, in particular the photoreceptor and retinal pigment epithelium (RPE) layers, as well as changes associated with disease progression. SD-OCT instruments acquire images at a speed of at least 20,000 A-scans per second, compared with 400 A-scans per second for time-domain OCT. The higher speed and higher resolution of SD-OCT imaging result in the ability to cover a much larger area of the macula in greater detail and, with the use of new algorithms, these data sets can be reconstructed into 3D images of the macula. These images permit visualisation of the real retinal geometry that is less affected by eye movements. Different scan patterns can be used depending on the information desired. For example, the high speed of SD-OCT permits the acquisition of scan patterns with a large number of lower density B-scans. Segmentation algorithms have been developed to extract quantitative information from these SD-OCT data sets, which includes the retinal thickness map, the OCT fundus image (OFI), and an RPE elevation map. The retinal thickness map is ideally suited to show the accumulation of fluid in the macula or the increased thickness in the macula resulting from traction, the OFI is ideally suited to show the boundaries of geographic atrophy (GA) and the RPE elevation map is ideally suited to show drusen and retinal pigment epithelial detachments (PEDs). The ability to perform averaged B-scans increases image quality and can be used to evaluate subtle changes in retinal anatomy. The purpose of this article is to review the use of SD-OCT for the diagnosis, treatment and follow-up of patients with AMD.
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Non-neovascular (dry) AMD is characterised by abnormalities of the RPE and the basement membrane between the RPE and choriocapillaris. Deposits develop under the RPE and within Bruch’s membrane. These deposits can be seen ophthalmoscopically as drusen, which can be scattered throughout the macula and posterior pole. Deposits can also be seen on top of the RPE and these are known as subretinal drusenoid deposits. Increased pigment clumping at the level of the RPE is another characteristic sign of AMD, followed by focal atrophy of the RPE. These abnormalities may be asymptomatic or accompanied by compromised vision and are considered to be the precursors of GA and choroidal neovascularisation (CNV).13–15

Drusen appear clinically as focal, white-yellow excrescences deep to the retina. They vary in number, size, shape and distribution. Several grading strategies have been developed to image drusen using colour fundus photography.16,17 Although colour fundus photography is useful for assessing the appearance of drusen, these images only provide 2D area information on the geometry of the drusen and cannot be used to measure quantitative properties such as drusen volume. SD-OCT can provide a 3D geometric assessment of drusen.

The high-definition averaged B-scans are useful for assessing the ultrastructure of drusen and to examine adjacent retinal layers that can be compromised by the disease process. Schuman et al. used SD-OCT to examine the retinal layers overlying drusen. A thinning in the photoreceptor layer was observed in 97 % of the cases, with an average photoreceptor layer thickness reduced by 27 % compared with age-matched control eyes. The inner retinal layers remained unchanged. They observed a correlation between the decrease of photoreceptor layer thickness and the height of drusen. These findings suggest a degenerative process, with photoreceptor loss leading to visual impairment.18

The acquisition of raster scans comprising a large number of lower-density B-scans, combined with the use of segmentation...
Imaging

Figure 3: Right Eye of a 72-year-old Man with Drusen that Increased in Volume during 16 Months of Follow-up

A–D: Colour fundus images, retinal pigment epithelium (RPE) map, RPE elevation map and horizontal B-scan from the baseline visit; E–H: Colour fundus images after 16 months of follow-up. The volume within a 3 mm circle increased from 0.085 to 0.108 mm³ (E and G). Some areas of drusen disappeared during the follow-up period (B and F, white arrow) while the total drusen volume increased over time.

An example of the segmentation algorithm used to measure drusen is shown in Figure 1. In this figure, the 6 x 6 mm scan area (white box) was superimposed on a colour fundus image of a macula containing drusen (see Figure 1A). A representative B-scan is shown with the boundaries of the actual RPE segmentation and the interpolated RPE floor identified in red and yellow, respectively (see Figures 1B and 1C). In Figure 1D, the OFI is shown, which represents the summation of the reflected light from each A-scan when viewed en face. This topic will be discussed later in this article. A 3D RPE segmentation map is shown along with the RPE elevation map, which represents the difference between the actual RPE and the RPE floor for each B-scan in the data set and this map is used to generate the area and volume of the drusen (see Figures 1E and 1F).

Figures 2 and 3 represent the clinical use of SD-OCT when observing patients with dry AMD. Figure 2 shows an example of an eye with significant regression of drusen as demonstrated using the algorithm. The colour fundus image, a representative B-scan over the drusen, the RPE segmentation map and the RPE elevation map with a 3 mm circle at baseline (see Figures 2A–2D), at six months (see Figures 2E–2H) and at one year (see Figures 2I–2L) of follow-up, respectively, are shown. The drusen area and volume decreased over time as shown most convincingly in the RPE segmentation and elevation maps. Drusen volume decreased from 0.125 mm³ at baseline to 0.067 mm³ at six months and to 0.039 mm³ at one year of follow-up.

Figure 3 shows an example of an eye with increasing drusen volume and area over 16 months. The volume within the 3 mm circle increases from 0.085 to 0.108 mm³. It is interesting to note that some areas of drusen disappear during the follow-up period (Figures 3B and 3F, white arrow) while the total drusen volume increases over time. This example demonstrates that drusen are dynamic.

Geographic Atrophy

The formation of GA impairs visual function, impacts the quality of life and may result in blindness.20 GA is seen clinically as one or more well-demarcated areas of hypopigmentation or depigmentation due to the absence or severe attenuation of the underlying RPE. The larger, deeper choroidal vessels are more readily visualised through the atrophic patches, which also lack photoreceptors and choriocapillaris. The natural history of GA has been described as a progressive condition that evolves through stages with loss of vision occurring over years.8,21,22 The initial size and configuration of the atrophy appear to influence its progression rate. Average linear rates of growth of 140–200 μm/year from the GA margin have been reported and the area of GA may enlarge by up to 3 mm² or more annually.23–25 Multiple imaging modalities have been used to document and quantify the area of GA. Historically, colour fundus photography was used to image GA; however, the use of colour photos can be challenging due to the reported difficulty in detecting and accurately delineating GA.15,16 Other imaging modalities, such as fluorescein angiography (FA), fundus autofluorescence (FAF) and SD-OCT are now used to evaluate and quantify GA. Although these imaging modalities...
provide different information, none has been shown to be superior to another.

SD-OCT was shown to be a useful tool for imaging and measuring GA.8,9,28 A wide spectrum of morphological alterations can be observed when evaluating eyes with GA using high-definition B-scans. The loss of the RPE and photoreceptors is easily observed in these B-scans. Bearelly et al. reported that photoreceptor loss occurred most frequently in a bridging fashion across the margin of GA.28

GA is currently imaged using SD-OCT by using the OFI, which represents a virtual fundus image resulting from the en face summation of the reflected light from each A-scan. This en face OFI identifies GA as a bright area, due to the increased penetration of light into the choroid where atrophy has occurred in the macula. The absence of the RPE and choriocapillaris are responsible for this increased penetration of light associated with GA.8,9 The OFI was shown to correlate well with the GA seen on clinical examination, colour fundus imaging and autofluorescence imaging.10,28,29

More recently, a newer algorithm developed by Carl Zeiss Meditec, Inc., provides an enhanced OFI, which is the summation of the reflected light from beneath the RPE. In addition, this new algorithm automatically measures the area of GA. A study comparing the measurements of GA area with the OFI and the enhanced OFI...
showed excellent correlation between the different modalities (Pearson correlation 0.999) and a good correlation between the automated algorithm and the manual grading (Pearson correlation 0.795) (Yehoshua Z et al. Personal communication).

Figure 4 shows an eye with GA secondary to AMD. Figures 4A and 4B show the colour fundus image and FAF image of the right eye. The entire area of GA is difficult to visualise using the colour fundus image, but more easily seen using FAF. The SD-OCT B-scan demonstrates the increased light penetration into the choroid in the area where the RPE is absent (see Figure 4C). At the border of GA, where there is a transition between intact and atrophic RPE, there is a marked difference in light penetration into the choroid (arrow). This transition in the penetration of light is responsible for creating the border of GA seen with the en face imaging. Within the GA, the area of RPE atrophy appears brighter on the en face image because of the increased penetration of light into the choroid. The summation of the reflected light from the sub-RPE layers is used to compose the of GA seen with the en face imaging. Within the GA, the area of light penetration into the choroid is responsible for creating the border of GA seen with the en face imaging. Within the GA, the area of RPE atrophy appears brighter on the en face image because of the increased penetration of light into the choroid. The summation of the reflected light from the sub-RPE layers is used to compose the OFI (see Figure 4D, area between red lines). A good correlation can be observed between the OFI (see Figure 4D) and the enhanced OFI (see Figure 4E). The manual measurements of the GA areas were 14.52 and 14.59 mm² for the OFI and the enhanced OFI, respectively.

The advantage of the enhanced OFI is that the lesions usually appear brighter than in the OFI, which facilitates identification of the lesion’s boundary. Another advantage is the fact that the OFI represents the light reflected from all the retinal layers and the presence of other macular pathologies may interfere with the identification of GA.

Spectral-domain Optical Coherence Tomography in Wet Age-related Macular Degeneration

The neovascular (wet) form of AMD is responsible for the majority of the cases with severe vision loss. It is characterised by the growth of abnormal vessels in the macular region induced by the overproduction of vascular endothelial growth factor (VEGF). These vessels may arise from thechoroidal circulation and penetrate Bruch’s membrane to form a fibrovascular tissue external to the RPE, or they may arise primarily from the retinal circulation. In either case, the presence of VEGF and abnormal vessels leads to anatomical changes in the retina and choroid with the accumulation of fluid in the subretinal space, within the retina, or under the RPE.

Since the advent of drugs that inhibit VEGF, the ideal strategy for following eyes with wet AMD has been to use OCT to determine whether the treatment is effective in resolving the macular fluid. The macular fluid can be identified by examining the B-scans and by reviewing the retinal thickness maps, which calculate the retinal thickness between the internal limiting membrane and the RPE. The effect of anti-VEGF therapy can then be assessed based on the qualitative appearance of the B-scans and the qualitative, as well as quantitative, changes in the retinal thickness maps. In addition, the same algorithm used to measure drusen can also be used to measure retinal pigment epithelial detachments (PEDs), since both involve the elevation of the RPE. This algorithm will measure both the area and volume of a PED. Penha et al. showed excellent reproducibility when measuring the area and volume of PEDs in patients with wet AMD. Moreover, Penha et al. have shown that changes in these quantitative measurements of PEDs may be useful in making retreatment decisions (unpublished data).

Figure 5 shows the case of a 76-year-old woman with wet AMD before treatment (upper row) and after treatment (lower row) with a single injection of an anti-VEGF drug. In Figure 5A the colour fundus image shows the lesion and in Figure 5B the B-scan shows the macular fluid thickness measured by the enhanced OCT (see Figure 5C). The PED volume decreased to 0.171 mm³ after treatment. In Figure 5D–5F the PED volume decreased, as shown in the colour fundus image (see Figure 5F), the B-scan (see Figure 5G) and the retinal thickness map (see Figure 5H). The PED volume decreases, as shown in Figures 5I and 5J. The PED volume decreased to 0.171 mm³ after treatment. This case demonstrates the usefulness of SD-OCT in the management of patients with wet AMD.

Summary

The recent advances in OCT technology, combined with the development of new algorithms capable of identifying RPE elevations and GA, provide quantitative tools for following patients with both wet and dry AMD. This imaging approach provides one-stop shopping for clinicians interested in managing patients with AMD. The advantage of SD-OCT over other imaging modalities for AMD is that the same scan pattern can be used to image the progression from drusen to GA and CNV. No other imaging modality is able to quantitatively assess all forms of AMD, so now clinicians can reliably follow the normal disease progression of their patients and their response to therapy. Although SD-OCT has changed the way we image AMD, the future of OCT holds even more promise with the use of longer-wavelength light sources for deeper choroidal penetration, faster scan times and higher image resolution.
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