

INTRODUCTION

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The term choroid is derived from the Greek words for “membrane” and “form.” Histologic investigations of this tissue have been performed since the 17th century. The choroid is a vascularized and pigmented tissue that extends from the ora serrata anteriorly to the optic nerve posteriorly. According to histopathologic examination, it is 0.22 mm thick posteriorly and 0.10 mm to 0.15 mm thick anteriorly.⁽¹⁾

A structurally and functionally normal choroidal vasculature is essential for retinal function: abnormal choroidal blood volume and/or compromised flow can result in photoreceptor dysfunction and death.⁽²⁾ Consequently, the choroid plays a vital role in the pathophysiology of many conditions, such as central serous chorioretinopathy (CSC),⁽³⁾ age-related macular degeneration (AMD),⁽⁴⁾ choroidal melanoma,⁽⁵⁾ Vogt–Koyanagi–Harada (VKH),⁽⁶⁾ and others.

A precise clinical understanding of choroidal changes showed proves to be critical for an accurate assessment of many posterior segment diseases. Until recently, the choroid could only be evaluated by indocyanine green (ICG) angiography,^(7,8) laser doppler flowmetry,⁽⁹⁾ and ultrasound.

Indocyanine green angiography enables visualization of the choroidal vessels and blood flow below the retinal pigment epithelium (RPE) from the posterior pole to the periphery as well as the vortex veins. Indocyanine green angiography may reveal greater detail of a choroidal neovascularization and detect choroidal polyps, versus fluorescein angiogram (FA). The specific properties of ICG with longer wavelength fluorescence and limited diffusion within the choriocapillaris enhance the visualization of structures beneath blood, exudates, or RPE detachments in greater detail.⁽¹⁰⁻¹³⁾

Laser Doppler flowmetry is a technique that allows noninvasive measurement of hemodynamic parameters of the optic nerve head (ONH), iris, and subfoveal choroidal circulation. Using laser Doppler flowmetry, the mean speed of the erythrocytes in the sampling volume and the number of moving erythrocytes in this volume can be determined. Several studies with laser Doppler flowmetry showed a decrease in volume in the choroidal circulation in various diseases like diabetic retinopathy, AMD, and retinitis pigmentosa.⁽¹⁴⁻¹⁶⁾

Ultrasound also plays a role in the diagnosis and management of a variety of vitreoretinal pathologies, especially in the presence of opaque media. Additionally, it can detect and characterize tumors and other thickenings in the choroid and retina. However, the image resolution is low, which makes the detection of small changes in the choroid difficult.⁽¹⁷⁾

Although these techniques are useful for determining vessel abnormalities or changes in the choroidal blood flow, they do not provide three-dimensional anatomical information about the RPE or the choroidal layers. The development of optical coherence tomography (OCT) makes it possible to have high-quality, cross-sectional images of the macula or ONH, analogous to ultrasonography but with greater resolution. However, adequate morphologic examination of the choroid using OCT has not been possible until recently, owing to its posterior location and the presence of pigmented cells in the RPE that attenuate the incident light. Recent reports demonstrated successful examination and measurement of choroidal thickness in normal and pathologic states using spectral-domain optical coherence tomography (SDOCT) instruments.^(4,18-22)

Improved optical coherence tomography technology

Optical coherence tomography uses light waves to obtain a reflectivity versus depth profile of the tissue under investigation.⁽²³⁾ Light from a broadband light source is divided into a reference beam traveling a known path and a sample beam that is directed onto the subject's retina. Light backscattered by retinal structures interferes with light from the reference beam that has traveled a known path delay. This interference is used to measure the light echoes versus delay or depth. Earlier OCT systems, such as the Stratus OCT (Carl Zeiss Meditec, Inc, Dublin, CA), used time-domain detection, in which the reference mirror position and delay are mechanically scanned to sequentially measure echoes from different depths and produce axial scans (A-scans). Scan rates of 400 A-scans per second with an axial resolution of 8 μm to 10 μm in the eye are achieved with the Stratus OCT.⁽²⁴⁾

More recently, spectral domain OCT (SDOCT), a type of Fourier domain detection, which uses an interferometer with a high-speed spectrometer to measure light echoes from all time delays simultaneously, has become commercially available in multiple OCT platforms. In SDOCT, the reference mirror does not require mechanical scanning and light echoes are detected simultaneously by measuring and Fourier transforming the interference spectrum. Increased sensitivity enables dramatic improvements in image acquisition speed and signal-to-noise ratio.^{25,26} Commercially available SDOCT devices, coupled with improvements in light sources, achieve axial scanning speeds of 20,000 to 52,000 A-scans per second with an axial resolution of 5 μm to 7 μm in the eye.⁽²⁷⁾

The choroid cannot usually be well visualized using time-domain OCT. The RPE is highly scattering and attenuates the OCT signal from the choroid. Because of the relatively low signal-to-noise ratio of time-domain OCT compared with SDOCT, there is less signal and image information detected from the deeper layers of the choroid. The pixel density of time-domain OCT, which is limited by the number of axial scans in the OCT image, also makes visualization of fine detail difficult in any part of the retina, including the choroid.⁽²⁸⁾

The outer limit of the choroid and the sclera cannot usually be reliably identified using standard SDOCT. In SDOCT, depth information is encoded as different frequencies of the interference spectrum. With increasing depth into tissue, echoes occur further from the point of detection, which is known to be the "zero delay line." These echoes are more difficult to discern from one another by the spectrometer because they have a higher frequency modulation than echoes found closer to the zero delay line. Two ways to overcome this is to increase the sensitivity of the spectrometer to higher frequency modulation and to increase pixel number in the line scan camera. The sensitivity to frequency modulation of a spectrometer does not have an impact on the resolution of the SDOCT imaging system because axial resolution is determined by the bandwidth of the light source and not by the sensitivity in depth.⁽²⁹⁾

Therefore, retinal structures that are closer to the zero delay line have higher signals than structures that are farther from zero delay line. By convention, commercial SDOCT machines operate with the inner retina closest to the zero delay line to maximize the sensitivity from the retina and vitreoretinal interface. In this way, the retina image is improved but it leaves the outer choroid farther from the zero delay line with subsequent diminished signal. Imaging the choroid using SDOCT requires specific imaging protocols that have been described recently. The most important technique to visualize the choroid uses the multiple scans obtained from the same retinal location that are subsequently averaged together by the OCT software. Between scans, the signal should remain constant while any noise is variable. When the subsequent images are averaged, the software

reduces the “speckle,” which enhances the continuity and sharper the tissue features. This technique also results in images with improved continuity of retinal features (Figure 1). Averaging typically increases signal-to-noise ratio in proportion to the square root of the number of images averaged. Most commercially available SDOCT devices have image averaging as an option. The different commercial available softwares average 8 to 100 images. At present, there is no consensus about the optimal number of images that should be averaged to adequately visualize the choroid. In addition to image averaging, several devices have mechanical eye tracking as a feature to ensure that all the images to be averaged are taken from the same retinal location. This also increases signaling by reducing the motion artifact.⁽³⁰⁾

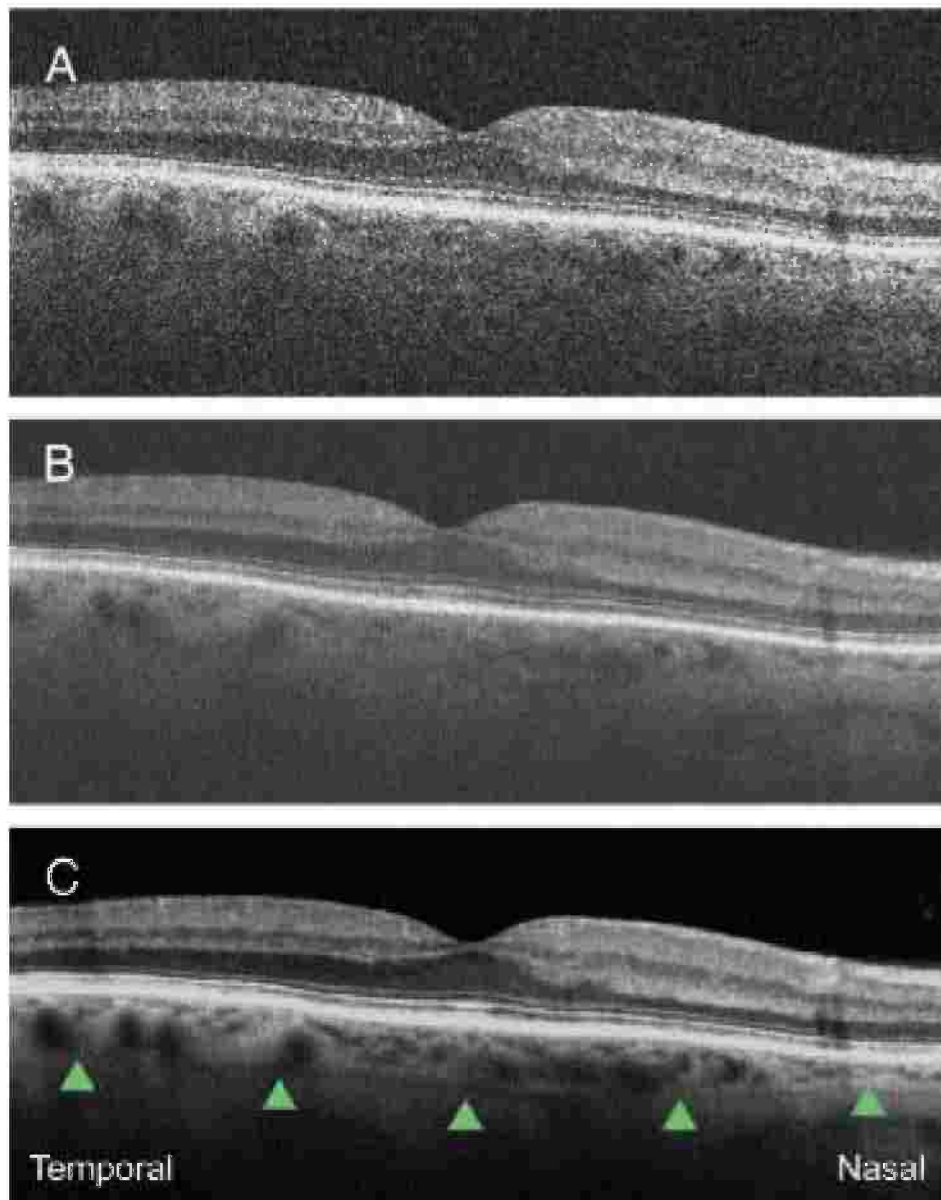


Fig. 1: Optical coherence tomography image demonstrating image averaging using Cirrus HD-OCT (Carl Zeiss Meditec, Inc). **A.** Image is a single B-scan. Note the low signal and indistinct border of the choroid and sclera. **B.** Image is 5 B-scans averaged together. Note the increase in signal from the choroid. **C.** Image is 20 B-scans averaged together. Note the further improvement in signal from image B and the distinct delineation of the choroid–sclera junction (green arrowheads).⁽²⁷⁾

Spectral domain detection cannot distinguish between positive and negative echo delays because the interference spectra are identical for an echo at a given positive or negative delay. Therefore, if a retinal structure crosses the zero delay line, it will appear reflected about the zero delay position, as a mirror-like image with reversed depth sensitivity.²⁷ This characteristic can be used to enhance choroidal imaging. If the scan operator advances the sample arm (patient interface) toward the patient so that the mirror image becomes fully apparent, this mirror image can be obtained as the main image. Because the mirror image is inverted about the zero delay line, the information from the choroid–sclera junction in the mirror image is maximized about the vitreous and inner retina. In other words, the decreased detection sensitivity in the choroid is minimized because the choroid–sclera interface is closer to the zero delay line in the mirror image (Figure 2). This technique was first reported by Spaide et al⁽²⁷⁾ as enhanced depth imaging (EDI). EDI is an acquisition software option on several commercial available devices. The EDI software automatically captures the cross-sectional image with the choroid close to the zero delay line to maximize the sensitivity on the outer limit of the choroid.⁽³¹⁾

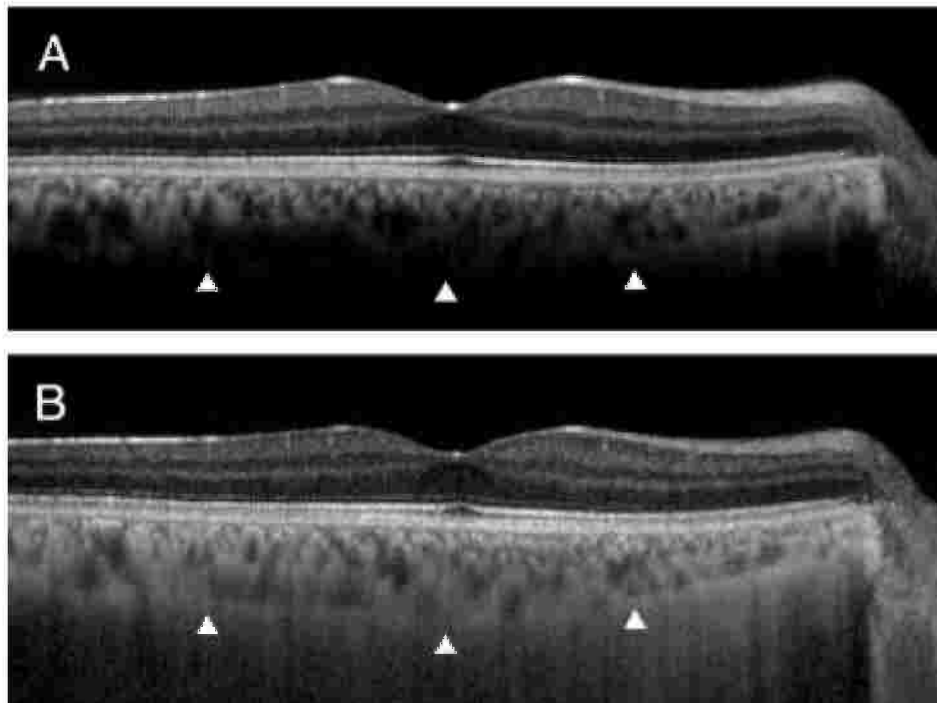


Fig. 2: Optical coherence tomography image demonstrating enhanced depth imaging on Spectralis (Heidelberg Engineering). **A.** Image was acquired with the inner retina adjacent to the zero delay. Note that the choroid–sclera junction is not distinct (white arrowheads) and there is low signal from the choroid but high signal from the inner retina. **B.** Image was acquired with the choroid adjacent to the zero delay by advancing the instrument toward the patient. Note that the choroid–sclera junction is distinct and that there is reduced signal from the inner retina but increased signal from the choroid.⁽²⁸⁾

Normal choroidal imaging

Normal choroidal thickness has been described by Margolis et al¹⁹ using the Spectralis (Heidelberg Engineering, Heidelberg, Germany) (Figure 3) and by Manjunath et al²² using the Cirrus HD-OCT (Carl Zeiss Meditec, Inc) (Figure 4). The choroidal thickness was measured manually perpendicularly from the outer edge of the hyperreflective RPE to the inner sclera (choroid–sclera junction) using the OCT software (Figure 4). Both groups report that the choroid is thickest subfoveally and thins nasally more than temporally. The reported subfoveal choroidal thickness was $287 \pm 76 \mu\text{m}$ (mean $\pm$ SD) on the Spectralis with a sample size of 30 patients (54 eyes) and $272 \pm 81 \mu\text{m}$ on the Cirrus device with a sample size of 34 subjects (34 eyes). Both groups also found a negative correlation between choroidal thicknesses and age.^(19,22)

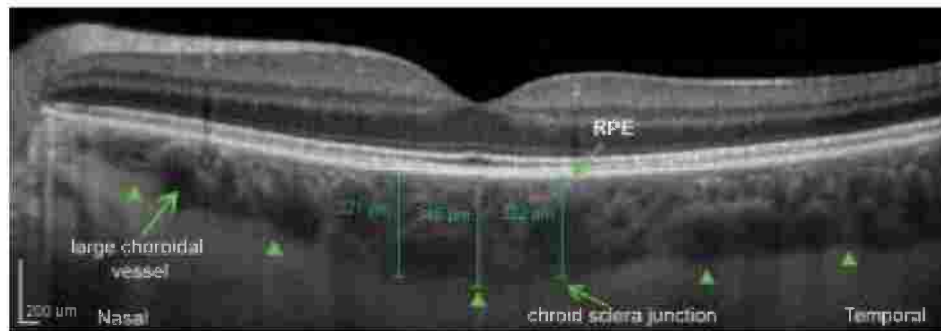


Fig. 3: Optical coherence tomography image of the normal choroid taken on Spectralis with EDI and over sampling. Choroidal thickness was measured in the fovea and at 500- μm intervals, nasal and temporal to the fovea with the measurements expressed in μm .⁽²²⁾

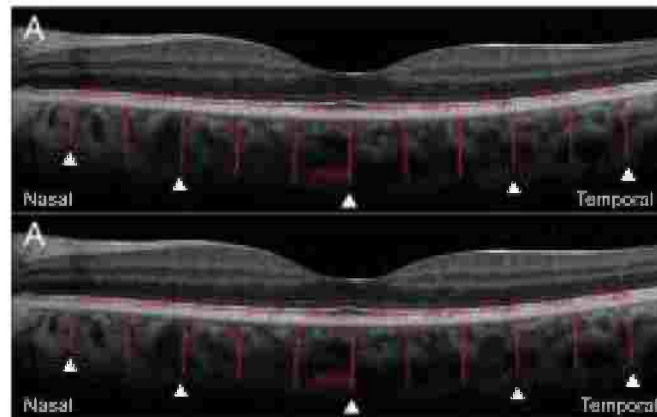


Fig. 4: Optical coherence tomography image of the normal choroid taken on Cirrus HD-OCT with 20 images averaged. **A.** Representative scan from a normal 24-year-old subject. **B.** Representative scan from a normal 76-year-old subject. Note the choroid–sclera.

A recent study demonstrated a high reliability and reproducibility for the choroidal thickness measurements in normal subjects. In this investigation, 6 independent examiners measured manually the subfoveal choroidal thickness on horizontal B-scan images. The reliability was evaluated by intraclass correlation coefficient, and the intervisit reproducibility was assessed by examining 10 volunteers, 4 months later. This study showed an interexaminer correlation of 0.970 (95% confidence interval [CI], 0.948–0.985) and an intervisit correlation of 0.893 (95% CI, 0.864–0.916).⁽²⁸⁾

The ability to reliably image the choroid with different SDOCT instruments makes this an emerging area of study. Consequently, it is important to validate the reproducibility of the choroidal measurements by different instruments to understand how precise and comparable the measurements are. A recent investigation analyzed the reproducibility between choroidal thickness measurements of images acquired with Cirrus HD-OCT (Carl Zeiss Meditec, Inc), Spectralis (Heidelberg Engineering), and Optovue RTVue (Optovue, Inc, Fremont, CA).⁽²⁸⁾ Choroidal thickness of normal eyes was manually measured in five areas. The measurements from any pair of three instruments (Cirrus vs. Spectralis, Cirrus vs. RTVue, Spectralis vs. RTVue) were strongly correlated. The intraclass correlation coefficient between all 2 system pairs of the 3 systems was >0.9 ($P < 0.0001$).

In addition, to evaluate the choroidal thickness, it is important to analyze the choroidal morphology, which may be altered in pathologic conditions. A recent study characterized the choroidal morphology in 42 eyes of 42 normal subjects. All subjects on this study had a bowl-shaped choroid–sclera junction; 98.8% of subjects had an even distribution of choroidal vessels in the nasal–temporal axis. The thickness of the large vessel layer at the fovea corresponded to 80% of the total subfoveal choroidal thickness.⁽²⁹⁾

Choroidal imaging and diseases

Table 1 summarizes the chorioretinal diseases that affect the choroidal thickness.

Table (1): Chorioretinal Diseases With Changes in the Choroidal Thickness.⁽¹⁸⁾

Diseases With Thicker Choroid	Diseases With Thinner Choroid
Central serous retinopathy	Neovascular AMD
Polypoidal choroidal vasculopathy	Dry AMD
Vogt–Koenig–Harada	Proliferative diabetic retinopathy
	Diabetic macular edema
	Mutifocal choroiditis
	Retinitis pigmentosa
	Glaucoma

Central serous chorioretinopathy

Central serous chorioretinopathy is a disease characterized by an exudative detachment of the neurosensory retina. Studies using ICG angiography revealed diffuse hyperpermeability of the choroidal vessels much larger than the active RPE leaks seen in the FA, suggesting generalized choroidal vascular disturbance.⁽³¹⁾ On OCT, CSC appears as an elevation of the full-thickness neurosensory retina from the highly reflective RPE–choriocapillaris complex separated by an optically empty zone. Retinal pigment epithelial detachments may also be detected usually within the areas corresponding to leakage in the FA. In addition, recent separated investigations using 3 different SDOCT devices,^(21,30,31) demonstrated significantly increased choroidal thickness in patients with acute CSC. Using Spectralis (Heidelberg Engineering), the mean ($\pm$ SD) choroidal thickness in 28 eyes of 19 patients with CSC was 505 ($\pm$ 124) μ m (range, 439 μ m–573 μ m), which was

214 μm (85%) greater than the mean choroidal thickness of age-matched normal eyes ($P \leq 0.001$) (Figure 5). Additionally, 2 studies evaluated both eyes of patients with unilateral CSC and observed that increased choroidal thickness was present in both eyes: the affected eyes ($445.58 \pm 100.25 \mu\text{m}$) and in the unaffected fellow eyes ($378.35 \pm 117.44 \mu\text{m}$) compared with normal eyes ($266.80 \pm 55.45 \mu\text{m}$).^(32,33) Although the pathophysiology of CSC remains elusive, these findings provide additional evidence that CSC may be caused by increased hydrostatic pressure in the choroid.^(21,31)

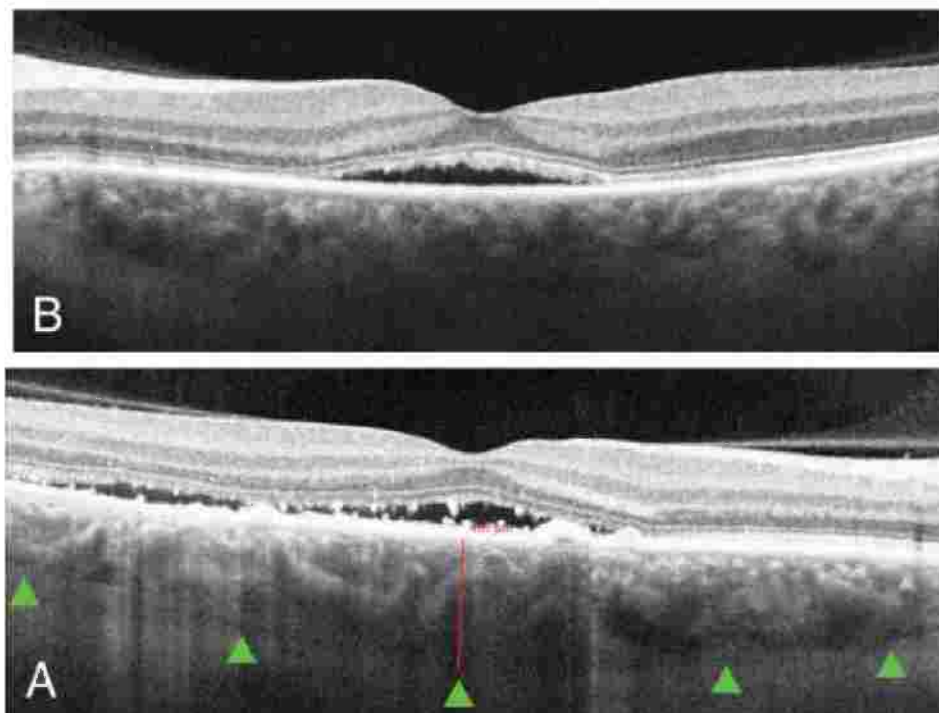


Fig. 5: Optical coherence tomography image of the choroid of a patient with central serous chorioretinopathy taken on Cirrus HD-OCT with 20 images averaged. **A.** Note that the choroid is thicker in the whole extension. The green arrowheads point the choroid–sclera junction. **B.** Note that it is not possible to visualize the choroid–sclera junction. The choroid may be so thickened as to lose signal penetration and intensity at the increasing depth because of signal roll-off distal to the zero delay line.⁽²¹⁾

In most eyes with CSC, the neurosensory retinal detachment resolves spontaneously within 3 months of duration. However, some patients present with persistent detachment and symptoms after 4 months, suggesting a chronic form of CSC. In these patients, treatment with laser photocoagulation or photodynamic therapy (PDT) is considered. Maruko et al⁽³⁴⁾ measured the subfoveal choroidal thickness before and after the treatment in 20 eyes of 20 patients with chronic CSC, using SDOCT. In this study, 8 eyes were treated with PDT and 12 eyes with laser photocoagulation. A significant decrease in the choroidal thickness after 4 weeks was found in all eyes treated with PDT ($389 \pm 106 \mu\text{m}$ at baseline vs. $330 \pm 103 \mu\text{m}$ after 4 weeks; $P < 0.001$). Patients treated with laser photocoagulation did not demonstrate a reduction in the choroidal thickness ($345 \pm 127 \mu\text{m}$ at baseline vs. $340 \pm 124 \mu\text{m}$ after 4 weeks; $P = 0.2$). A similar study examining choroidal thickness in 16 patients (16 eyes) with CSC before and after PDT showed a decrease in the subfoveal choroidal thickness after the treatment (from $421 \mu\text{m}$ [95% CI, 352–489 μm] to $346 \mu\text{m}$ [95% CI, 278–414]; $P = 0.0001$).³⁵ Another investigation also showed that half-

dose PDT resulted in thinner subfoveal choroidal thickness (80 μm , 20%) 1 month after treatment, decreased the choroidal vascular hyperpermeability, and maintained the remission for 1 year.⁽³⁶⁾ These findings may suggest that PDT reduces the choroidal vascular hyperpermeability observed in CSC.^(34,36)

The choroidal thickness may be used as an additional parameter to assist in the differentiation of CSC from other causes of serous retinal detachment. In addition, the choroidal thickness may indicate the activity of the disease on the follow-up after treatment with PDT.

Age-related macular degeneration

Age-related macular degeneration is the leading cause of blindness in both developed and developing nations in people older than 60 years.⁽³⁷⁾ Optical coherence tomography, together with clinical biomicroscopy, color fundus photography, FA, autofluorescence imaging, and ICG angiography, is used in the evaluation of patients with AMD. Macular thickness maps generated by the OCT have been shown to be useful in monitoring the progression and treatment response to disease states, such as neovascular AMD after antiangiogenic treatment.⁽³⁸⁾ Choroidal structure is of particular interest in AMD because abnormalities of the choroidal circulation have been hypothesized to contribute to the development of AMD. However, little is currently known about the choroid-related changes using SDOCT in this disease.

Spaide⁴ described a distinct entity, termed age-related choroidal atrophy, which has some overlap with dry AMD. He described 28 eyes of 17 patients with a mean age of 80.6 ± 7.3 years, with mean choroidal thickness less than 125 μm . The mean subfoveal choroidal thickness was 69.8 μm , and 35.7% (10 of 28) of the eyes presented with late-stage AMD (Figure 6). It suggests that choroidal thickness decreases with increasing age and the choroidal circulation might play a role in the pathophysiology of AMD.⁽⁴⁾

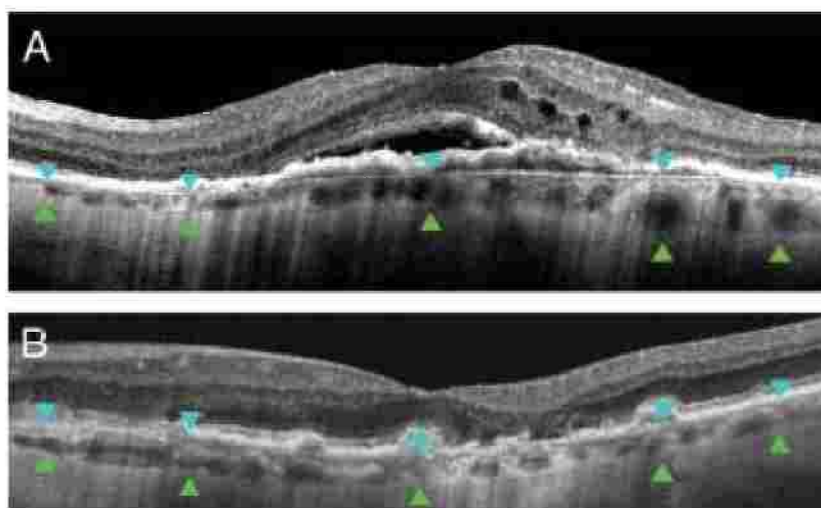


Fig. 6: **A.** Optical coherence tomography image of the choroid of a patient with exudative AMD taken on Cirrus HD-OCT with 20 images averaged. Note that the choroid is thin and that it can be visualized beneath choroidal neovascularization. The blue arrowheads point the outer edge of the hyperreflective RPE, and the green arrowheads point the choroid–sclera junction. **B.** Optical coherence tomography image of the choroid of a patient with dry AMD taken on Cirrus HD-OCT with 20 images averaged. Note that the choroid is thin, and in the area of RPE atrophy, it is possible to visualize a hyperreflective area on the choroid.⁽³⁷⁾

Another study compared subfoveal choroidal thickness in eyes with neovascular AMD (21 eyes of 21 patients) with that of eyes with polypoidal choroidal vasculopathy (PCV) (23 eyes of 23 patients). This study demonstrated that eyes with PCV have a thicker subfoveal choroid ($293 \pm 72.3 \mu\text{m}$) when compared with eyes featuring typical neovascular AMD ($245 \pm 73.1 \mu\text{m}$), $P = 0.032^{(39)}$ (Figure 6). Additionally, it was found that eyes with a subfoveal choroidal thickness of $\geq 300 \mu\text{m}$ are more than 5 times more likely to have PCV. Similar results were also observed in another study that compared the choroidal thickness among eyes with neovascular AMD, PCV, and CSC. It was demonstrated that the choroid was thicker in eyes with PCV and CSC than in normal subjects or in those with neovascular AMD.⁽³¹⁾ The thicker choroid could be partially attributed to the dilation of middle and large choroidal vessels or an increase in the choroidal vascular permeability that is observed by ICG.⁽³⁹⁾

Both PCV and neovascular AMD feature abnormal vascular lesions arising from the choroidal vessels, which lead to recurrent serous exudation and hemorrhages. Some studies suggest that PCV lesions respond better to focal treatment with laser or PDT as compared with lesions from neovascular AMD. Sometimes it is difficult to differentiate both diseases. Therefore, the choroidal thickness may play a role in the differentiation of these diseases because the choroid is thinner in eyes with neovascular AMD and the choroid is thicker in eyes with PCV.

Diabetic retinopathy

Clinical and experimental findings suggest that choroidal vasculopathy in diabetes may play a role in the pathogenesis of diabetic retinopathy. Various choroidal abnormalities, including obstruction of the choriocapillaris, vascular degeneration, choroidal aneurysms, and choroidal neovascularization, have been reported in histopathologic studies of diabetic eye.^(40,41) Esmacelpour et al⁽⁴²⁾ reported central choroid thinning in all Type 2 diabetic eyes regardless of disease stage. Sixty-three eyes were studied, and the choroidal thickness mapping of all diabetic patients demonstrated central and inferior thinning compared with healthy eyes (unpaired *t*-test; $P < 0.001$). Subfoveal choroidal thickness for healthy eyes was $372 \pm 74 \mu\text{m}$, which was found to be significantly thicker than all diabetic eyes: eyes with microaneurysm, $208 \pm 49 \mu\text{m}$; eyes with exudates, $205 \pm 54 \mu\text{m}$; and eyes with clinically significant macular edema $211 \pm 76 \mu\text{m}$ (ANOVA, $P < 0.001$; Tukey, $P < 0.001$).⁽⁴²⁾ It can be speculated that the thinner choroid indicates an overall reduction of choroidal blood flow in diabetic patients, as was previously demonstrated with laser Doppler flowmetry and ICG angiography.^(9,43,44) Therefore, it is likely that the decreased choroidal thickness may be related to retinal tissue hypoxia because the choroid is the major source of nutrition for the RPE and outer retinal layers.

High myopia

Myopia is the result of a mismatch between the optical power and the length of the eye, with the latter being too long. In high myopia, there is an increased risk of visual impairment and blindness due to ensuing pathologies, such as retinal detachments and choroidal neovascularization. The excessive elongation affects not only the retina but also the choroid and Bruch membrane. Histologic studies demonstrated choroidal thinning in eyes with high myopia due to significant thinning of the choriocapillaris and focal lack of vessels. Fujiwara et al⁽²⁰⁾ studied the subfoveal choroidal thickness in 55 eyes of 31 patients with high myopia (≥ 6 diopters) using SDOCT. The mean age was 59.7 ± 17.6 years and the mean refractive error was -11.9 ± 3.7 diopters. The authors demonstrated a

very thin choroid ($93.2 \pm 62.5 \mu\text{m}$) in highly myopic eyes (Figure 7), and the choroidal thickness was negative correlated with age ($P = 0.006$), refractive error ($P < 0.001$), and history of choroidal neovascularization ($P = 0.013$).⁽²⁰⁾ Additionally, this study suggested that subfoveal choroidal thickness decreases by $12.7 \mu\text{m}$ for each decade of life and by $8.7 \mu\text{m}$ for each diopter of increasing myopia. Furthermore, the choroidal thickness may be a predictive factor for visual acuity in highly myopic patients because the choroid is responsible for the oxygen and nutrient supply of the outer retina.

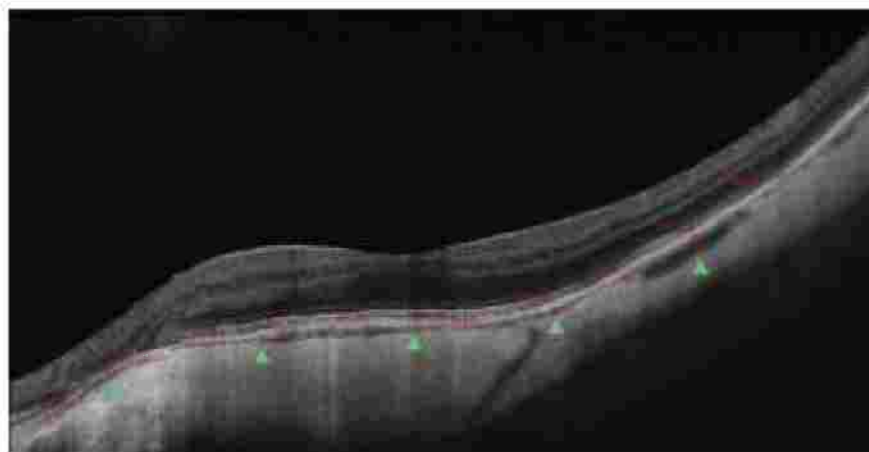


Fig. 7: Optical coherence tomography image of the choroid of a patient with high myopia taken on Cirrus HD-OCT (Carl Zeiss Meditec, Inc) with 20 images averaged. Note the choroid-sclera junction (green arrowhead). Choroidal thickness was measured in the fovea and at 500- μm intervals nasal and temporal to the fovea with the measurements expressed in micrometer (red lines). Note that the choroid is very thin.⁽²⁰⁾

Uveitis

The SDOCT is now proven to be an effective noninvasive investigation tool for detecting pathologic features in uveitis and is gaining more popularity as an ancillary examination in patients with posterior segment manifestations of uveitis.^(45,46)

Vogt-Koyanagi-Harada disease is a bilateral granulomatous panuveitis associated with autoimmunity against melanocytes. The acute uveitic stage of VKH is characterized by bilateral anterior and/or posterior segment involvement with exudative retinal detachment.⁽⁴⁷⁾ Indocyanine green shows patchy filling delays with hypofluorescent spots, and in addition, B-scan ultrasound shows anatomical alterations.⁽⁴⁸⁾ Maruko et al⁽⁶⁾ studied the choroidal thickness of 16 patients (32 eyes) with active VKH disease before and after corticosteroid treatment. All eyes had markedly thickened choroid ($805 \pm 173 \mu\text{m}$ vs. $287 \pm 76 \mu\text{m}$ of normal eyes) before the treatment, possibly related not only to inflammatory infiltration but also to increased exudation. Additionally, a reduction in the choroidal thickness was documented after 2 weeks of corticosteroid treatment (baseline: $805 \pm 173 \mu\text{m}$; 2 weeks of follow-up: $341 \pm 70 \mu\text{m}$, $P < 0.001$).⁶ Evaluating the choroidal thickness has potential importance for assessing treatment efficacy and recurrence in VKH disease. Another study investigated the choroid of six VKH patients in acute and convalescent stages. A loss of focal hyperreflectivity in the inner choroid. The presence of this feature in both acute and convalescent phases suggests that there is a permanent structural change to small choroidal vessels caused by VKH uveitis.⁴⁹ Choroidal imaging using SDOCT may change the treatment strategies used for VKH. In the past, treatment with high dose of corticosteroids was initiated

and then tapered off rapidly. If the corticosteroids were tapered too rapidly, the serous detachment returned. It is conceivable that gauging the proper corticosteroid taper can be more safely done by monitoring the choroidal thickness with SDOCT.

Choroidal changes are also described in multifocal choroiditis. Yasuno et al,⁽⁵⁰⁾ using an SDOCT with 1060 nm wavelength studied 1 patient with multifocal choroiditis and showed localized thinning of the choroid, occlusion of the choroidal vessels, and localized hyperreflectivity that may represent hyperpigmentation of the choroid. Another study investigated three cases with acute lesions of multifocal choroiditis using Spectralis (Heidelberg Engineering). This study reported the findings of acute lesions, which include presence of sub-RPE material, choroidal hyperreflectivity below these lesions, and overlying vitreous cells.⁽⁵¹⁾

Tumors

The SDOCT of the choroid has also been used to examine choroidal tumors. Torres et al⁵ found that features such as the delineation of the borders of the tumor both parallel and perpendicular to the visual axis along with evaluation of the inner choriocapillaris and outer large vessels within the tumor was possible: in around half of eyes examined (10 of 23), the axial thickness of the tumor was able to be identified using SDOCT as compared with ultrasound, which was able to detect the extent of all tumors examined. Another study evaluated 11 eyes with choroidal osteoma. The tumor was hyporeflective in two cases, isoreflective in seven cases, and hyperreflective in two cases. Additionally, it was observed that overlying choroid was compressed by the tumor in 72.7% and the retina exhibited degenerative changes in 45.5% of the cases.⁽⁵²⁾ These studies demonstrate that SDOCT allows characterization of the thickness and reflective quality of small (<3 mm thick) choroidal lesions including choroidal nevi and melanomas. Future improvements in image resolution and depth will allow better understanding of the mechanisms of visual loss, tumor growth, and tumor management.^(5,52,53)

Retinal dystrophies

The SDOCT has been used to investigate a number of structural abnormalities in the retina of subjects with inherited retinal dystrophies, including retinitis pigmentosa.⁵⁴ The choroid has been implicated in the pathophysiology of a number of these dystrophies as well.¹ Furthermore, using scanning Doppler flowmetry, choroidal blood flow has been demonstrated to be diminished in subjects with retinitis pigmentosa when compared with that of normal subjects and that this decrease correlates with cone function¹⁶ (Figures 8 and 9). Using EDI OCT, Yeoh et al⁵⁵ found changes in the structure of the choroid in a subset of a heterogeneous group of subjects with retinal dystrophies. The observed changes include both focal and diffuse choroidal thinning (Figure 8). These changes do not appear to be associated with visual acuity.⁽⁵⁵⁾

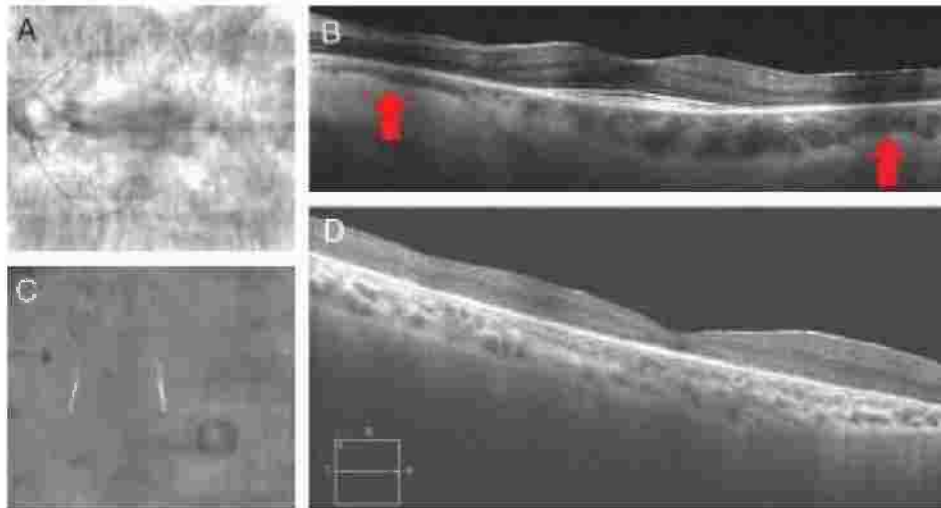


Fig. 8: Optical coherence tomography image of the choroid of 2 patients with retinitis pigmentosa taken on Cirrus HD-OCT (Carl Zeiss Meditec, Inc) with 20 images averaged. A. OCT fundus image of the subject whose B-scan appears in B. Green line indicates location.⁽¹⁶⁾

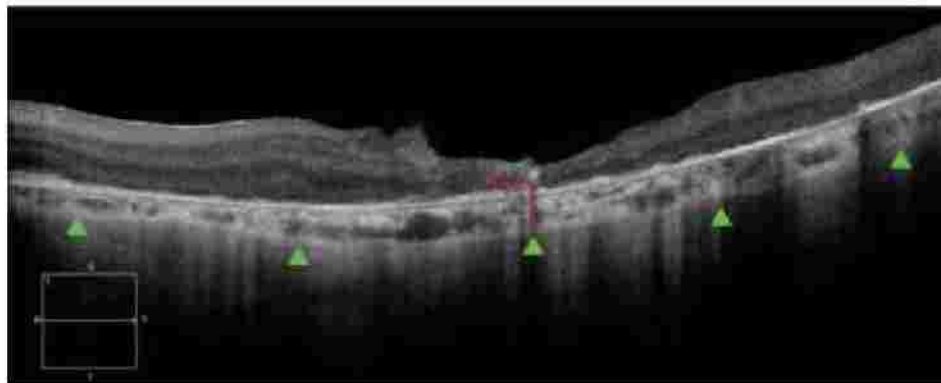


Fig. 9: Optical coherence tomography image of the choroid of a patient with late phase of retinitis pigmentosa taken on Cirrus HD-OCT (Carl Zeiss Meditec, Inc) with 20 images averaged. Note that the choroid is thin (choroid sclera junction - green arrowhead).⁽¹⁶⁾

Glaucoma

Choroidal imaging as a tool to investigate the vascular theory of glaucoma pathogenesis has evaluated both the macular choroid and the peripapillary choroid. Normal peripapillary choroidal thickness was described by Ho et al.⁽⁵⁶⁾ This investigation of 36 normal subjects imaged with both horizontal and vertical raster scans concluded that the inferior quadrant ($149.90 \pm 50.14 \mu\text{m}$) was significantly thinner than the superior ($229.41 \pm 50.96 \mu\text{m}$), nasal ($227.17 \pm 81.40 \mu\text{m}$), and temporal ($208.84 \pm 55.70 \mu\text{m}$) quadrants ($P < 0.001$).

Maul et al.⁽⁵⁷⁾ described a relationship between macular choroidal thickness as measured using EDI OCT and age, axial length, and nerve fiber layer loss. However, when compared with glaucoma suspects ($n = 37$), Maul et al.⁽⁵⁷⁾ did not find a significant difference in macular or peripapillary choroidal thickness in glaucomatous eyes ($n = 37$) ($14 \mu\text{m}$ [95% CI, -54 to 26]; $P = 0.5$).

Similarly, Mwanza et al⁽⁵⁸⁾ reported no significant difference in macular choroidal thickness between 38 normal subjects, 20 normal-tension glaucoma patients, and 56 primary open-angle glaucoma patients using EDI OCT. Ehrlich et al⁽⁵⁹⁾ confirmed these results in finding no difference in peripapillary choroidal thickness between glaucoma suspects ($n = 39$) and patients with primary open-angle glaucoma ($n = 31$) ($P \geq 0.13$). One consideration when interpreting these data is that both groups studied glaucoma suspects rather than healthy subjects as the control group.

In contrast, Usui et al⁽⁶⁰⁾ found significantly decreased choroidal thickness both in the macula and in the region of the ONH in highly myopic eyes with normal-tension glaucoma as compared with age- and refraction-matched healthy controls (fovea: 166 μm vs. 276 μm , $P < 0.001$; superior ONH: 172 μm vs. 241 μm , $P < 0.05$; superotemporal ONH: 161 μm vs. 244 μm , $P < 0.01$; temporal ONH: 110 μm vs. 161 μm , $P < 0.01$; and inferotemporal ONH: 115 μm vs. 159 μm , $P < 0.05$). Usui et al⁽⁶⁰⁾ used a prototypical swept-source OCT with a scan speed of 50,000 A-scans per second at 1060 nm.

Using EDI OCT, Park et al⁽⁶¹⁾ further investigated peripapillary imaging of glaucomatous eyes. They found in 139 eyes (73 patients) with glaucoma that the anterior surface of the lamina cribrosa was identifiable in 65% of eyes and that pores within the lamina cribrosa were identifiable in 76% of eyes. In 86% of eyes, the short posterior ciliary artery was identified. In 18% of eyes, the subarachnoid space was identified. Given that these structures have been implicated in the pathogenesis of glaucoma, in addition to choroidal imaging, SDOCT may be used in the future to investigate the relationship of these structures with glaucoma in vivo.

Future directions in choroidal imaging

Because of the posterior location of the choroid, OCT imaging is difficult because incident light is attenuated by the RPE and retinal structures. One method to increase the depth of choroidal penetration by OCT is to image with a longer wavelength of incident light to reduce attenuation from scattering.⁽⁶²⁾ One such approach has been to use an incident wavelength centered near 1050 nm, which is absorbed minimally by water in the vitreous. Prototype systems with 1050 nm wavelength and axial resolution of 7 μm have demonstrated increased choroidal visualization when compared with standard SDOCT using a wavelength centered ~ 800 nm.⁽⁶³⁾ Likewise, SDOCT systems using 1050 nm wavelength have demonstrated increased penetration through cataracts and media opacities when compared with systems using 800 nm wavelength.⁽⁶⁴⁾

Another method that may improve imaging the choroid is the swept source optical coherence tomography (SSOCT).^(65–67) The SSOCT uses a tunable frequency swept laser light source to increase signal quality deeper in tissue when compared with SDOCT. The light source of SSOCT is able to sequentially emit various frequencies in time. This means that the interference spectrum is measured by photodetectors rather than by a spectrometer and line scan camera, as is the case in SDOCT. Consequently, decreased sensitivity of a spectrometer to higher frequency modulation is eliminated. Furthermore, k-space is no longer unevenly sampled in SSOCT.^(67–69) This dramatically reduces sensitivity roll-off, thus making image quality deeper in tissues improved.

In SSOCT, the limiting factor in scan acquisition time is the speed at which the light source can sweep the needed frequencies. Because swept laser light sources can do this very rapidly, SSOCT is much faster than SDOCT with axial scan rates of up to 236,000 A-scans per second with 11- μm axial resolution in tissue.^(70,71) This allows for more data acquisition in a given time, making volumetric data sets feasible.⁽⁶⁷⁾

In addition to SS-OCT, Doppler OCT is a promising technology in the field of choroidal imaging.⁽⁷²⁻⁷⁴⁾ Doppler OCT can evaluate blood flow and volume of retinal vasculature.^(75,76) Also, it can highlight vessels where there is blood flow, without quantitatively measuring flow.⁽⁷⁷⁾ In contrast to FA or ICG angiography, which are 2 dimensional, Doppler OCT is depth resolved, meaning that cross-sectional images can be used to identify in which retinal layers vascular abnormalities occur. This method has been used to evaluate abnormalities in chorioretinal vasculature.⁽⁷⁸⁾ As such, this technology is promising to aid in monitoring chorioretinal diseases, especially neovascular AMD.

Along with modifications in hardware and image acquisition, improvements in software and processing of OCT data will probably be important in the future for evaluation and monitoring of choroidal changes. One such software advance is the use of en-face imaging. This technique allows the clinician to visualize three-dimensional data in a useful way as it shows a particular retinal layer at a given depth projected into an en-face view, which closely corresponds to a fundus image. It is hard to evaluate the health of the choroid using only a limited number of B-scans because the cross-sectional nature makes it such that vessels are transected at odd angles. En-face imaging makes subtle microstructural relationships much more apparent.⁽⁷⁹⁾ This promises to aid in the evaluation of chorioretinal diseases.