Detailed Visualization of the Anterior Segment Using Fourier-Domain Optical Coherence Tomography

Sanjay Asrani, MD; Marinko Sarunic, PhD; Cecilia Santiago, MD; Joseph Izatt, PhD

Objective: To study details of the anterior chamber drainage angle using Fourier-domain optical coherence tomography in healthy subjects and patients with angle abnormalities.

Methods: A high-speed anterior segment optical coherence tomography prototype was developed using a 1310-nm-wavelength swept light source. Six healthy subjects and 6 patients with glaucoma were imaged in an observational cross-sectional study.

Results: Schlemm’s canal and the trabecular meshwork were visualized in all of the patients. Fifteen-millimeter scans enabled entire anterior segment visualization providing configuration details of the iris with respect to the angle. Four-millimeter scans permitted detailed views of the angle configuration and its structures. Volumetric imaging was possible and Schlemm’s canal was visualized along part of its circumference.

Conclusion: Anterior segment Fourier-domain optical coherence tomography permits detailed noncontact imaging of the angle and its structures, providing a tool to improve our understanding of the pathogenesis of narrow-angle glaucoma.

Arch Ophthalmol. 2008;126(6):765-771

Visualization of the trabecular meshwork and Schlemm’s canal could be a valuable tool for glaucoma diagnosis and treatment, but it requires resolution in the range of tens of micrometers. Optical coherence tomography (OCT) is an emerging noncontact, noninvasive imaging modality for cross-sectional imaging of biological tissue. A time-domain (TD) OCT system operating at a wavelength of 1310 nm and a line rate of 2 kHz has been commercialized for ocular anterior segment imaging (Visante OCT; Carl Zeiss Meditec, Dublin, California). Narrow anterior chamber angles have been studied using a prototype TD-OCT imaging system. Optical coherence tomography was found to be a promising method for screening individuals at risk for narrow-angle glaucoma, providing higher resolution than ultrasound biomicroscopy with the additional advantages of a noncontact imaging modality compatible with slit-lamp biomicroscopy observation. However, Schlemm’s canal and the trabecular meshwork have not been readily visualized with the existing TD-OCT systems.

Optical coherence tomography has been revolutionized in recent years by the development of Fourier-domain (FD) techniques, which provide a substantial increase in sensitivity over traditional TD-OCT systems. This increase in sensitivity has been used to image at high scan rates without increasing the optical exposure or sacrificing image brightness. However, high-speed FD-OCT systems operating at a central wavelength of 1310 nm have been limited in useful imaging depths to less than 3 or 4 mm, hindering ocular anterior segment imaging. Images of the anterior chamber, the depth of which typically spans greater than 4 mm, are corrupted by a symmetric overlapping artifact, referred to in the literature as the complex conjugate artifact, that obstructs important physiological details. A doubling of the image depth can be achieved by resolving the complex conjugate artifact using various methods, thus permitting full-range imaging. A subset of these depth-doubling techniques has been previously applied for technology demonstrations of full-depth imaging of the anterior segment.

In this article, we describe a pilot study of high-speed, high-resolution imaging of the ocular anterior segment in vivo using a swept-source FD-OCT system. Full-range imaging is provided using a 3 × 3 fiber coupler topology and quadrature projection processing as described in detail.
Full-range FDOCT imaging of the entire anterior chamber width demonstrated the iris configuration with respect to the anterior lens surface. In 1 of the healthy subjects with myopia, a concave iris was detected on FDOCT and confirmed on clinical examination (Figure 1). No clinical evidence of pigment dispersion was detected. In the high-resolution 4-mm scans of the angle region, the configuration of Schlemm’s canal was visible along with the trabecular meshwork in eyes of healthy subjects (Figure 2A and B). The trabecular meshwork was seen as a triangular structure with the base attached at the scleral spur. The scleral spur was identified by its peaked outline and the slight projection into the anterior chamber (Figure 2C). Schlemm’s canal was seen as a curvilinear lucent area external to the trabecular meshwork. This lucent area extended from the scleral spur to the anterior tip of the trabecular meshwork located at the end of the Descemet membrane. This lucent area corresponding to Schlemm’s canal was distinct from the lucency of the corneal stroma. Although the ciliary body outline was visible in FDOCT images, details of the ciliary processes were not visible (Figure 2B). The gonioscopic appearance of the angle in the healthy subject shown in Figure 1 was open to the ciliary body band (grade IV by the Shaffer method; the angle between the iris and trabecular meshwork was 45°) and the 4-mm FDOCT scan demonstrated the cross-sectional view of the same in Figure 3A. In another healthy subject, gonioscopy revealed the angle to be open to the scleral spur (grade III by the Shaffer method; the angle between the iris and trabecular meshwork was < 45° but > 20°); its cross-sectional view is shown in Figure 3B.

Representative quantitative measurement was performed in a patient with gonioscopic grade II narrow-angle glaucoma. The angle opening distance at 500 µm (AOD500) is the length of a perpendicular from a corneal endothelial point 500 µm anterior to the scleral spur drawn to the iris. The angle recess area at 500 µm (ARA500) is the triangular area enclosed by the AOD500, the line along the corneal endothelium from the scleral spur to the AOD500, and the line following the contour of the iris from the scleral spur to the AOD500. The boundaries of the epithelium, endothelium, and iris were traced in raw coordinates manually. These lines were then converted to processed coordinates accounting for the refraction at each interface according to Snell’s law. The AOD500 (308 µm) and ARA500 (0.11 mm²) measurements were performed in processed coordinates from these boundary-delineating lines (Figure 4). The measurements have been described previously using TDOCT. Our measurements of AOD500 and ARA500 agree with the previously reported measurements and are within the range of values reported for narrow angles.23,24

Figure 1. Concave iris configuration seen in a patient with mild myopia. The full depth of the anterior chamber can be visualized.
Among the patients with narrow-angle glaucoma, 4 had primary narrow-angle glaucoma. In 1 patient, the right eye was examined before laser peripheral iridotomy for a potentially occludable narrow angle. Gonioscopy showed the angle to be narrow (grade I) with the anterior trabecular meshwork visible. The high-resolution FDOCT scan in the angle region demonstrated the extremely narrow space between the root of the iris and the trabecular meshwork (Figure 5A and B). In another patient, gonioscopy demonstrated the angle graded as 0. No angle structures were visible. The FDOCT scan shows the near touch between the trabecular meshwork and the root of the iris (Figure 5C and D). A patient with a clinical diagnosis of plateau iris who had episodes of high intraocular pressure and colored halos around lights despite a patent laser peripheral iridotomy was found to have a narrow angle (grade I) with the anterior trabecular meshwork visible on slitlamp gonioscopy. The FDOCT scan demonstrated a narrow space between the trabecular meshwork and the root of the iris when the room lights were on. This space was obliterated (resulting in a closed angle) when the room lights were switched off, and it returned to its narrow configuration once the room lights were turned on again (Figure 6). Although the ciliary body outline was visible in FDOCT images, details of the ciliary processes were not visible. In the fourth patient with narrow-angle glaucoma, scattered peripheral anterior synechiae were visible on gonioscopy. Areas of adhesion were observed between the root of the iris and the trabecular meshwork in the FDOCT cross-sectional image (Figure 7).

Volumetric imaging was performed by acquiring data sets consisting of $1024 \times 400 \times 60$ pixels (axial $\times$ lateral $\times$ elevation) in 4.5 seconds. The images were rendered using 3DView (RMR Systems, Suffolk, England), which limited the number of displayed pixels to 512. Cross-sectional planes of a volumetric image of a normal anterior segment are shown in Figure 8 and were acquired with the scan width reduced to 7 mm. Schlemm’s canal was visualized along part of the circumference by removing the overlying layers in the rendered 3-dimensional image. Additional information may be gleaned from the volumetric rendering by examining cross sections through the data set. A patient with secondary glaucoma due to a ciliary body tumor was also imaged. A volume scan was performed (Figure 9), and it showed the elevation of the peripheral iris in the inferior region of the eye causing a shallowing of the anterior chamber. However, the limited depth penetration of the FDOCT

Figure 2. Detailed angle structures. A, The trabecular meshwork is visible as a triangular body and Schlemm’s canal as a curvilinear lucent space external to the trabecular meshwork. B, The ciliary body outline is visible along with Schlemm’s canal and the trabecular meshwork in another eye of a healthy subject. C, Note the scleral spur as a structure projecting into the anterior chamber at the root of the trabecular meshwork.

Figure 3. Angle configurations. A, An eye clinically open to the ciliary body band on gonioscopy. B, An eye open to the scleral spur on gonioscopy.

Figure 4. Representative narrow angle measurement. The epithelium, endothelium, and iris were manually delineated in unprocessed coordinates. The angle opening distance at the 500-µm (AOD500) line is shown in blue, and the angle recess area at the 500-µm (ARA500) region is hatched with red lines in unprocessed coordinates. The values for the AOD500 and ARA500 were calculated in processed coordinates, and the ARA500 was obtained by integration of the equations describing the lines.
prevented imaging of the actual tumor located beneath the iris. Figure 10 shows the image from a patient with narrow-angle glaucoma secondary to a multiloculated iris cyst that is visible in the iris pigment epithelium.

COMMENT

In this study we have demonstrated, to our knowledge for the first time, high-resolution, noncontact, in vivo imaging of the anterior segment, trabecular meshwork, and Schlemm’s canal using a full-range FDOCT system. The axial resolution using the FDOCT system was less than 9 µm as compared with more than 11 µm with TDOCT.2,3 Resolution of the complex conjugate artifact permitted a scan depth in excess of 6.3 mm, allowing imaging of the entire depth of the anterior chamber while reducing artifacts. The reduced resolution observed in some of the images was introduced by imperfect sampling by the prototype wavelength swept interferometric system and was not an artifact of the complex conjugate–resolved imaging algorithm. High-resolution images (consisting of 1024 axial points by 800 lateral lines) provided a view of the trabecular meshwork and Schlemm’s canal. We have demonstrated the ability to view the structural relationship between the periphery of the iris and the trabecular meshwork as well as the visualization of peripheral anterior synechiae. Detailed imaging may help to improve understanding of the patho-

Figure 5. High-resolution Fourier-domain optical coherence tomographic scans of narrow angles with gonioscopic grade I and grade 0. A, A very narrow space is visible between the root of the iris and the trabecular meshwork in the right eye of a patient with narrow-angle glaucoma (gonioscopic grade I). B, The left eye of the same patient (A) shows similar findings. C, Near touch between the trabecular meshwork and the root of the iris is visible in the right eye of another patient with narrow-angle glaucoma (gonioscopic grade 0). D, The left eye of the same patient (C) shows a similar angle configuration.

Figure 6. Images with the room lights turned off (A) and back on (B). A, A closed angle with the iris root almost in approximation with the trabecular meshwork when the room lights are off. B, On turning the room lights on, the pupil constricts with ensuing stretching of the iris and widening of the angle.
genesis of pupillary block syndrome,24 plateau iris syndrome, and possibly malignant glaucoma.

Visualization of the angle nasally and temporally along a line that horizontally crosses the pupil provides data about the angle across only 2 points along the circumference. This typically may represent the overall nature of the angle as seen gonioscopically in most cases. However, there are eyes with uneven angle configuration as they progress from having open to narrow angles. To visualize the angle and Schlemm’s canal along the entire circumference, volumetric scans may be useful. Volumetric scanning was possible with the FDOCT system at 13.4 frames per second (thus minimizing motion artifact), which is useful for quantitative measurements of the anterior chamber in 3 dimensions. Variations in the depth of the anterior chamber such as in the case of the eye with a ciliary body tumor can be easily visualized with the help of volumetric scanning. However, it is important to note that FDOCT was unable to image the ciliary processes or structures beneath the pigmented layer of the iris.

Proper gonioscopy to observe the configuration of the angle of the anterior chamber requires a learned skill that involves recognition of the landmarks of the angle with and without compression by the contact gonioscopic lens. Recognition of landmarks can be challenging during gonioscopy on eyes with minimal pigmentation in their trabecular meshwork. However, even with proper technique, gonioscopy requires the use of slitlamp illumination, which typically causes pupillary constriction. Constriction of the pupil causes the iris to be drawn away from the angle and may thus artificially open up a very narrow angle. Compression caused by inadvertent pressure by the examiner holding the lens during gonioscopy can also falsely open the angle. This may lead to an erroneous diagnosis of an open angle with the potential for major clinical consequences, resulting in an acute angle-closure glaucoma or chronic angle closure. The incidence of both of these types of glaucoma may be reduced if narrow angles are correctly identified and prophylactic laser iridotomy is performed before any synechial formation occurs.

A noncontact OCT system that operates in the infrared spectrum permits viewing of the angle in its natural state without pressure from a contact lens or the effects of illumination.25 Because details of the trabecular meshwork are visible with the FDOCT system, the triangular space between the root of the iris and the trabecular meshwork can be evaluated for the relative position of the root of the iris with regard to the corneoscleral bands of the trabecular meshwork. Widespread application of anterior segment FDOCT to glaucoma may permit accurate diagnosis even by a general eye care practitioner without training in gonioscopy. This may aid in the diagnosis of angle closure, in both acute and chronic states, which constitutes one-half of the total glaucoma cases in the world.26

Numerous studies have investigated the structures of the trabecular meshwork and Schlemm’s canal, resulting in better understanding of their microscopic and molecular composition. These structures have been studied using either histopathological examination or perfusion studies of postmortem specimens.27-29 Despite these studies on the outflow system, our knowledge has been limited owing to the lack of a noninvasive imaging system that permits visualization of these structures in vivo. Imaging by FDOCT may permit studies to detect changes of the trabecular meshwork and Schlemm’s canal with accommodation, with different pharmacological agents, and following surgery. Detailed imaging of
Schlemm’s canal may also help in understanding the pathogenesis and evolution of open-angle glaucoma.

Nonpenetrating glaucoma surgical procedures have not yet been widely adopted secondary to the high skill level required to identify Schlemm’s canal and thus long surgical times. Each of these surgical procedures is heavily dependent on precise localization of Schlemm’s canal. Adapting the FDOCT system to the operative microscope may allow intraoperative visualization of Schlemm’s canal, permitting easier, more accurate, and more efficient surgery on the canal. Volumetric scanning can also aid in the study of patients who have undergone surgical procedures such as viscocanalostomy and deep sclerectomy by cross-sectional image evaluation in different planes. This may permit study of multiple parameters, for example, the extent of the spread of the viscoelastic along the circumference of Schlemm’s canal or the state of the Descemet window.

The limitations of TD systems include image degradation due to motion artifact secondary to patient’s microsaccades. The fast scan rate of 13.4 frames per second for FDOCT allows real-time cross-sectional imaging at high resolution and thus may permit image-guided interventions such as injections into Schlemm’s canal. Intraoperative FDOCT imaging may permit development of precise and targeted glaucoma surgery such as removal of the trabecular meshwork using the trabectome. Imaging may also aid in the placement of transcanalicular shunts (microtube shunt; Glaukos Corp, Laguna Hills, California), which enable aqueous humor to bypass the trabecular meshwork. Various pharmacotherapeutic treatments such as gene transfer to the trabecular meshwork could potentially be injected directly into Schlemm’s canal using FDOCT imaging to guide the needle.

In summary, anterior segment FDOCT may assist in understanding the pathogenesis of different types of glaucoma, aid in accurate diagnosis, and provide new avenues of treatment.

Submitted for Publication: October 15, 2006; final revision received November 13, 2007; accepted November 13, 2007.

Correspondence: Sanjay Asrani, MD, Duke University Eye Center, Box 3802, Durham, NC, 27710 (asran002@mc.duke.edu).

Financial Disclosure: Drs Sarunic and Izatt are inventors of intellectual property, assigned to Duke University, related to the subject of this article. Dr Izatt is a founder and part-time employee of Bioptigen, Inc.

Funding/Support: This work was supported in part by an unrestricted grant from Research to Prevent Blindness, Inc, and by grant EY013516 from the National Eye Institute. Dr Asrani is a recipient of the Career Development Award from Research to Prevent Blindness, Inc.

Previous Presentation: This study was presented at the annual meeting of the American Glaucoma Society; March 4, 2006; Charleston, South Carolina.

Additional Contributions: Kevin Hsu, PhD, Micron Optics, Inc, loaned the high-speed swept laser source, and Bioptigen, Inc, supplied the base software package for acquisition and display.

REFERENCES

3. Radhakrishnan S, Huang D, Smith SD. Optical coherence tomography imaging...


**Call for Papers**

*The Archives* will be publishing a special theme issue on translational research. The issue will include (1) reviews of successes and failures in translating laboratory research into clinical applications; (2) descriptions of new experimental models as test beds for novel therapies; (3) discussion of end points for clinical trials, including surrogate measures; (4) examples of new preclinical research with a high likelihood of translation into clinical studies. The issue will include a combination of invited papers and papers presenting original research. Leonard A. Levin, MD, PhD, will be editor for this special issue. We invite and encourage all investigators to submit manuscripts describing original research and reviews by September 1, 2008. All manuscripts submitted for this issue are subject to an expedited peer review. Early receipt ensures the best chance for acceptance for this special issue. Accepted manuscripts not included in this issue will be published in other issues of the *Archives*. Please note in the cover letter that the submission is for the “Translational Research” theme issue. The expected publication date for this issue is April 2009.