High-resolution retinal imaging in young children using a handheld scanner and Fourier-domain optical coherence tomography

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PURPOSE
To test the feasibility and applicability of a handheld probe for Fourier-domain optical coherence tomography (Fd-OCT) retinal imaging in infants and children.

METHODS
Thirty children ages 7 months to 9.9 years, with (10 of 30) or without (20 of 30) retinal pathology, were imaged with Fd-OCT. Imaging was performed under sedation in 10 of 30 children ages 7 months to 3.7 years. A high-resolution Fd-OCT system (axial resolution: 4.5 μm; acquisition speeds: 1000 A-scans/frame, 9 frames/second), constructed at the UC Davis Medical Center, in conjunction with a handheld scanner, was used for retinal imaging.

RESULTS
Useful images were obtained from all selected patients. Image acquisition was possible in a conscious state in children as young as 3 years of age. All children tolerated the tests well. The most challenging situation for young children was the lack of an internal fixation target and the moving scanning line, which usually distracted them from a steady fixation. Despite these problems, image quality was comparable with scans previously obtained from an adult population.

CONCLUSIONS
The flexible handheld scanner in association with high acquisition speed and high-resolution Fd-OCT allows retinal imaging in infants and children. This technology provides high-resolution documentation of retinal structure in a pediatric population for the first time. (J AAPOS 2009;13:72-74)

The purpose of this study was to test the feasibility and applicability of a handheld probe for Fourier-domain optical coherence tomography (Fd-OCT) retinal imaging in infants and children. Few reports pertaining to retinal imaging in children using commercially available time-domain OCT systems have been published, with patient ages in these studies as follows: 4 to 17 years, in which 4- and 5-year-old patients were uncooperative; 9 years (1 case); 7 and 9 years (2 cases); 9 years (1 case); and 7 years (1 case); detailed ages not provided. To the best of our knowledge, the use of high-speed Fd-OCT systems in infants or children has not been previously reported. We performed such retinal imaging successfully in sedated as well as conscious infants and children. The handheld scanner accommodated the child’s ability to either hold still on a chin rest or be tested in a reclined position.

Methods
Subjects
Thirty children, ages 7 months to 9.9 years, with (10) or without (20) retinal pathology were imaged with retinal Fd-OCT. (One subject had retinal disease in one eye and a normal retina in the other.) Clinical diagnoses were maculopathy (6), retinal dystrophy (4), and posttraumatic choroidal neovascularization (1). Examination under chloral hydrate sedation (10 patients aged 7 months to 3.7 years) was used only for standard-of-care diagnostic procedures and not primarily for this research study. One patient with a retinal dystrophy (Leber’s congenital amaurosis) had horizontal pendular nystagmus. Pupils were dilated in 27 of 30 subjects: 3 children did not want eye drops but were cooperative enough for successful retinal imaging.

Written informed consent and/or assent were obtained from all participants and/or guardians. The project was approved by the Research Ethics Board at The Hospital for Sick Children in Toronto and the UC Davis Medical Center Institutional Review Board, and conducted in accordance with the Tenets of Helsinki.
younger than 6 years of age, was the lack of an internal fixation target for the nontested eye. The most external fixation target and the moving scanning line, which usually distracted them from a steady fixation. External fixation targets were successful in only a few cases. The localization of the macular area was easier in conscious than in sedated children because of lack of simultaneous fundus viewing. Retinal layer identification was possible in all subjects. Images obtained using the handheld mode contained movement artifacts caused by the examiner and/or the child. It still allowed retinal layer identification but reduced the image quality for further possibility of 3D reconstruction.

Macular Scans Obtained from Controls
Scans from control subjects of different ages were obtained under sedation or conscious conditions. The resolution of the images was similar in all tested subjects (Figure 2). Macular scans shown in Figure 2 demonstrate well-formed macular layering. Retinal layers are labeled as previously published.8 The small sample size in this pilot study did not allow us to address potential age-related changes of different retinal layers. Macular scans of patients with retinal diseases that illustrate the usefulness of high-resolution imaging in children are available in e-Supplement 1 (available at jaapos.org).

Discussion
The application of high-resolution retinal imaging in infants and children without and with sedation using a flexible setup is described here for the first time. The high-speed acquisition allowed imaging in young children with a low attention span or intermittent fixation as well as in one patient with nystagmus and reduced vision. In addition, very cooperative children younger than 6 years of age could be imaged without pupil dilation. These cases highlight the benefit of having high-speed, high-resolution retinal imaging modalities for pediatric assessments. Microstructural retinal abnormalities can now be detected and correlated with visual function in this population. The ability to recognize small intraretinal abnormalities is an important objective measure in cases where nonorganic vision loss is suspected. Identification and quantification of retinal layer changes and localization of intraretinal deposits is an important investigative tool in children with early stages of retinal dystrophy (e-Supplement 1).

The handheld retinal imaging system with the flexible probe has overcome the current technical limitations of using OCT systems in a supine position. The probe can be mounted to a slit lamp or articulating arm or held, depending on the child’s cooperativeness and ability. Further improvements of this technology, such as simultaneous fundus viewing and a fixation target, will advance the applicability in a routine clinical setting. The use of such a device will contribute to our further understanding of normal retinal morphology, develop-
ment, and aging, and prove useful in early detection of retinal diseases in children.

**Literature Search**

PubMed and MEDLINE were searched for the period 1990 to the present for the following terms: OCT, optical coherence tomography, infant, pediatric, and children.

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**References**

e-Supplement 1. Macular scans of patients with retinal diseases illustrate the usefulness of high-resolution imaging in children.

Composite of horizontal Fd-OCT scans through the fovea and fundus photographs from 3 patients with different retinopathies are discussed.

Fd-OCT was valuable in the investigation of a 9.1-year-old neurologically challenged boy with sudden vision loss (0.7 log MAR in both eyes), minimal macular RPE changes on fundus exam, and normal electroretinogram (ERG) responses. Macular scans identified loss within the foveal outer and inner segment layer (OSL/ISL; denoted by stars), which could explain the vision loss and excluded a nonorganic cause.

![Macular scans of a patient with sudden vision loss, minimal macular RPE changes, and normal ERG responses.](image)

Macular scans of an 8.9-year-old girl with nyctalopia and vision loss to 0.4 logMAR in both eyes showed inner retinal splitting in the right macula (denoted by an error) as well as OSL and ISL disruption (marked by stars). These findings complemented the diagnosis of the ERG abnormalities in support of Enhanced S-cone syndrome.

![Macular scans of a patient with nyctalopia and vision loss, showing inner retinal splitting and OSL/ISL disruption.](image)

A 7.3-year-old with nystagmus and reduced visual acuity (RE, 0.5 logMAR; LE, 1.6 logMAR) was diagnosed with Leber's congenital amaurosis caused by a homozygous RPE65 mutation. Fd-OCT images demonstrated well-preserved inner retinal layers. Photoreceptor layers were thinned with nondistinguishable OSL or ISL (marked by stars). This information will be useful in selecting an optimal potential therapeutic approach.

![Macular scans of a patient with Leber's congenital amaurosis, showing well-preserved inner retinal layers.](image)