Semiautomated Assessment of Anterior Segment Structures in Pediatric Glaucoma Using Quantitative Ultrasound Biomicroscopy

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Abstract: The study was a prospective observational study comparing semiautomated to manual quantitative ultrasound biomicroscopy image analysis among 82 images from 41 eyes of 32 subjects (21 controls and 11 glaucoma) enrolled in the Pediatric Anterior Segment Imaging Innovation Study. Intraclass correlation coefficients and correlation coefficients were >0.8 for all parameters, and comparison of respective analysis speed was 7 times faster for the semiautomated method compared with manual image quantification.

Key Words: ultrasound biomicroscopy, high frequency ultrasound, image analysis, cornea, anterior segment, pediatric, congenital glaucoma

Primary and secondary causes of early childhood glaucoma can cause physical changes to individual structures of the anterior and posterior segment. Glaucomatous structural changes can be clinically subtle or dramatic, depending on the severity, and include increased corneal diameter and thickness, horizontal striations, and increased axial length. Structural changes are thought to be because of the unique state of collagen in the pediatric eye featuring decreased stromal collagen fibril cross-linking, increased hydration, and increased collagen fiber crimp structure. The specialized collagen architecture of youth results in more elastic biomechanical corneal properties, and allows for enlargement and distortion of the collagenous corneal and scleral stroma. The ability to precisely quantify and monitor corneal axial change is the mainstay of congenital glaucoma monitoring, in addition to frequent measurement of intraocular pressure (IOP).

Ultrasound biomicroscopy (UBM) allows for a high-resolution view of the anterior segment and offers a unique opportunity for visualization of the anatomy impacted by pediatric glaucoma. Because of high-frequency sound waves in the range of 40 to 50 MHz, UBM offers a span of depth of ~5 mm, and thus can be used to image the cornea, lens, iris, and ciliary body. UBM has demonstrated several clinically useful characterizations for structures involved in pediatric glaucoma and, unlike all other available imaging modalities, UBM is not hindered by clouding of the cornea. Anterior segment optical coherence tomography has similar clinical utility to UBM, with better resolution, but is inferior to UBM in the setting of corneal clouding or for imaging structures posterior to the iris. Established standardized UBM image measurement protocols with manual image analysis using ImageJ have previously demonstrated good intraobserver repeatability and interobserver agreement.

Some of the major drawbacks to use of UBM in a clinical setting are lack of standardized normative values, time-intensive image analysis, and the requirement for high level knowledge and extensive training. Because of these challenges, UBM use is typically limited to cases where the corneal view is profoundly compromised, or when clinical concern relates to structures posterior to the iris. While these applications are worthy of a dedicated imaging tool, the potential use of the UBM is much greater than these few niche applications. A rapid quantitative assessment tool may allow clinicians additional objective data that may influence diagnosis or management. The development of rapid quantitative analysis for UBM images introduces the emerging field of Quantitative UBM or Q-UBM.

The purpose of our study was to develop a novel open access Q-UBM image analysis tool and evaluate its use in subjects with and without history of congenital glaucoma. Our research team utilized Python open source programming language for semiautomated analysis of UBM images. This program, EyeMark, is able to measure anterior segment structures, including those that may be altered in patients with glaucoma, improving the utility of UBM in a clinical setting and addressing the problems of manual UBM image interpretation.
TABLE 1. Demographics and Measurement Results in Glaucoma and Control Groups

|                      | Glaucoma | Controls |
|----------------------|----------|----------|
| Number of subjects (eyes) | 11 (20)  | 21 (21)  |
| Sex (% female)        | 52       | 36       |
| Mean age (mo)         | 9 ± 14   | 9 ± 12   |
| Median age (mo)       | 3        | 4        |
| Age range (mo)        | 0-564    | 0-600    |
| Race/ethnicity (%)    |          |          |
| African American      | 73       | 52       |
| Caucasian             | 27       | 29       |
| Hispanic              | 0        | 14       |
| Multiple              | 0        | 5        |
| AA (mm)               | 12.7 ± 1.7 | 12.7 ± 1.7 |
| CCT (mm)              | 0.642 ± 0.141 | 0.634 ± 0.130 |
| ACRC (radius) [diopters] | 8.23 [41.00] ± 1.43 | 7.56 [44.75] ± 1.61 |
| PCRC (radius) [diopters] | 7.42 [45.50] ± 1.03 | 7.42 [45.50] ± 1.03 |
| ACD (mm)              | 3.59 ± 0.76 | 3.62 ± 0.75 |

AA indicates angle-to-angle distance; ACD, anterior chamber depth; ACRC, anterior corneal radius of curvature; CCT, central corneal thickness; PCRC, posterior corneal radius of curvature.

MATERIALS AND METHODS

UBM images were obtained from 41 eyes of 32 subjects (21 healthy age-matched subjects and 11 subjects with history of infantile glaucoma) enrolled in the Pediatric Anterior Segment Imaging Innovation Study (PASIIS) (Baltimore, MD). PASIIS is a collaborative program between the University of Maryland and Children’s National Medical Center designed to apply advances in technology and image analysis specifically to clinical evaluation and management of pediatric anterior segment disease. Demographics are displayed in the Table 1.

Participants were enrolled at the time of ocular examination. Consent for participation in the UBM imaging and analysis was obtained from the parent(s) or participant. Inclusion criteria for infantile glaucoma participants required onset of glaucoma before age 2 years. Glaucoma was defined as IOP > 21 mm Hg, with 1 or more of the following: corneal enlargement, progressive myopic shift coupled with increasing corneal diameter and/or axial length, increased optic nerve cup-to-disc ratio ≥ 0.2, or a surgical procedure for IOP control. Exclusion criteria for control participants included eyes with abnormal visual function for age, history of traumatic injury, past or present glaucoma or glaucoma suspect status, abnormal gonioscopy, refractive error > 4 diopters of hyperopia or myopia, or > 2 diopters of astigmatism.

This study adhered to the ethical principles outlined in the Declaration of Helsinki as amended in 2013. The Institutional Review Board has approved the above referenced protocol and the associated consent forms. Collection and evaluation of protected health information was compliant with the Health Insurance Portability and Accountability Act of 1996.

A total of 82 UBM images (2 from each eye) were obtained using the Aviso Ultrasound Platform A/B UBM with 50 MHz linear transducer (Quantel Medical, Clermont-Ferrand, France) or the Accutome UBM Plus Platform with 48 MHz linear transducer (Keeler Accutome Inc., Malvern, PA). Studies have shown no significant difference in measurements of anterior segment structures in children and adults between different anterior segment imaging platforms, including UBM. The PASIIS standardized protocol was developed to extract quantitative data from both UBM and anterior segment optical coherence tomography images. Imaging techniques have been previously described. Good repeatability and reliability of landmark identification in axial UBM images for structural measurements has been previously established. Our imaging protocol included 2 UBM images, 1 horizontal axial, and 1 vertical axial image per eye, in focus at the pupil center. Images were selected based on overall image quality, centration of the pupil, symmetry of the 2 angles, with resolution sufficient to localize the scleral spur on each side of the image, and identification of landmarks including the corneal center and angles. All images were de-identified and measured by masked observer.

Parameters measured included angle-to-angle distance (AA), central corneal thickness (CCT), anterior corneal radius of curvature (ACRC), posterior corneal radius of curvature (PCRC), and anterior chamber depth (ACD). Images were measured in 2 ways: manual measurements using ImageJ and semiautomated measurements using Python EyeMark. ImageJ 1.52a (National Institute of Health, Bethesda, MD), is a Java-based open access image processing program (https://imagej.net/ImageJ). The ImageJ measurements were performed according to the PASIIS protocol. Reliability and repeatability analysis of the ImageJ analysis protocol has been previously published. The semiautomated UBM image analysis protocol was developed using Python 3.7.4 (Python Software Foundation, Frederick, VA; http://www.python.org). EyeMark relies on the user to select 8 points (2 points on the caliper reference line, 2 angle apices, central corneal endothelium, central corneal epithelium, and 2 points on the peripheral corneal epithelium), to calculate the various metrics of the eye as seen in Figure 1. Using these 8 points, EyeMark is able to calculate AA, CCT, ACRC, PCRC, and ACD (Fig. 1, Table 1).

Statistical analysis was performed with Microsoft Excel (2019 Edition; Microsoft). Means for each parameter were determined (Table 1). Corresponding values were compared using analysis of variance (ANOVA) and the interclass correlation coefficient was calculated across each set of 82 data points, with an intraclass correlation coefficients (ICC) of > 0.8 indicating very good reliability and an ICC of > 0.9 indicating excellent reliability. Conditional correlation was determined by computing the correlation coefficient (R value) for each parameter along with the P-value. An R value > 0.8 was accepted to indicate a strong positive correlation. Bland-Altman plots were constructed to evaluate agreement between manual (ImageJ) and semiautomated (Python EyeMark) methods.

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The time to measure each image with ImageJ was recorded for each of 10 images. The EyeMark program was run 3 times for all 82 images and the overall time was divided by the number of images to determine the average amount of time required to measure 1 image with EyeMark.

RESULTS

ICC and correlation coefficients (R values) for each parameter are shown in Table 2. Bland Altman plots and correlation curves are shown in Figure 2A. When participants are separated by glaucoma status, the correlation between ImageJ and EyeMark remains unchanged across all parameters regardless of whether the participant has a diagnosis of congenital glaucoma (Fig. 2B). The R value for participants with glaucoma is 0.99 (P < 0.001) and the R value for participants without glaucoma is similar at 0.99 (P < 0.001). The average evaluation time per image using ImageJ is 6 minutes 57 seconds with a SD of 2 minutes and 38 seconds. The average evaluation time per image using EyeMark is 54 seconds per image (Table 2, Fig. 2).

DISCUSSION

The purpose of this project was to determine if the clinical utility of UBM can be improved by semi-automation. A useful image analysis process must yield results that are: (1) Consistent with known values; (2) Repeatable and reliable compared with gold standard; (3) Applicable in both disease and nondisease states; and (4) Faster than current standard image analysis. Based upon our results, the EyeMark automated analysis tool fit each of these criteria when compared with ImageJ manual analysis. Previous studies established cross-platform reproducibility, longitudinal reproducibility, and interoperator reproducibility,7–11 while the current study focused exclusively on the comparison of semiautomated and manual techniques. Semiautomation represents a key translational step in advancing the clinical relevance of Q-UBM imaging, to expand on the use of UBM for predominantly qualitative purposes.

ImageJ and EyeMark both extracted data consistent with expected values (Table 1), and well-established patterns were observed when comparing between the glaucoma and control groups. Greater cross-sectional chamber distance (AA), thicker CCT, flatter curvature (ACRC and PCRC), and deeper chamber (ACD) were measured among participants with congenital glaucoma. In comparing the repeatability and reliability of semiautomated compared with manual image analysis, the ICC for all parameters was 0.81 or greater, indicating good reliability with the use of EyeMark. The ICC for AA and PCRC in particular showed excellent reliability. The R values for all parameters was 0.86 or greater, implying strong correlation between measurements taken by ImageJ and EyeMark. Bland Altman

![FIGURE 1. A–E, Sample measurements of angle-to-angle distance, central corneal thickness, anterior corneal radius of curvature, posterior corneal radius of curvature, anterior chamber depth, respectively. F, Manual ImageJ user interface with sample measurements. G, Automated EyeMark user interface with sample user input.]

### TABLE 2. Intraclass correlation coefficients (ICC) and correlation coefficient (R) for automated compared with manual image analysis for 5 parameters.

| Parameter      | ICC  | R    | P    |
|----------------|------|------|------|
| AA             | 0.98 | 0.98 | < 0.001 |
| CCT            | 0.86 | 0.86 | < 0.001 |
| ACRC           | 0.81 | 0.89 | < 0.001 |
| PCRC           | 0.93 | 0.93 | < 0.001 |
| ACD            | 0.93 | 0.93 | < 0.001 |

AA indicates angle-to-angle distance; ACD, anterior chamber depth; ACRC, anterior corneal radius of curvature; CCT, central corneal thickness; PCRC, posterior corneal radius of curvature.
plots suggested no systematic proportional bias. The difference between methods did not systematically tend to grow larger or smaller as the mean increases. Variability was consistent without increase in scatter as the mean increases (or decreases), with an exception in CCT for values >800 µm. This feature of the measurement highlights the importance of using the same measurement tool consistently for longitudinal assessment of corneal thickness. Data among glaucoma and nonglaucoma age-matched control eyes show similar correlation, demonstrating that our semiautomated Q-UBM EyeMark program produced accurate measurements regardless of glaucoma status (Fig. 2). Lastly, the time-to-analyze was 7-fold less in the automated analysis, confirming the efficiency advantage of semiautomation.

A relatively small sample size was evaluated in this study. While appropriate to answer the question of reliability in this specific use case, absolute values may not apply at the population level. Subjects included eyes with and without history of congenital glaucoma according to well-defined diagnostic criteria, and a broad range of patient ages for both glaucoma and control subjects. Results may generalize to images of similar quality and features, but may not generalize to image analysis involving more profound anterior segment pathology or disorganization, or lower image quality. Measurement of parameters with more complex geometry than the 5 parameters in this study may likewise have lower repeatability and reliability. The accuracy and precision of EyeMark is dependent on proper identification of anatomical features on ultrasound images. The Python EyeMark program makes assumptions about the geometry of the eye. The epithelium and endothelium of the cornea are assumed to fit into the curvature of a circle, a well-established assumption which governs the use of diopters to describe the cornea. Some disease processes violate this assumption, including trauma and keratoconus. More complex geometric patterns must be taken into account when applying this tool to various disease categories. Unique ocular pathology should be considered at the time of user input. The time-to-evaluate analysis does not take into account training time of the analyst before use. We do not intend to disregard that early steps in image analysis training are relevant and sometimes challenging hurdles to overcome.

Python EyeMark semiautomated UBM image analysis provides a more time efficient quantitative UBM analysis without sacrificing data accuracy when compared with ImageJ manual image analysis among subjects with and without history of congenital glaucoma. The clinical implications of using EyeMark for UBM of the anterior segment are broad. Longitudinal studies will be essential to fully understand the utility of this program in clinical context. As with all novel ophthalmic imaging tools and programs, potential applications are discovered and optimized through use. Please visit github.com/karunkannan/EyeMark to access the program.

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