Optical Biometry Versus Ultrasound Biometry

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ABSTRACT

Background: Implantation of an intraocular lens (IOL) is the gold standard in modern-day cataract surgery. Over the last decade, IOL power calculations have become a focal point of cataract surgery. Over the last decade, IOL power calculations have become a focal point of cataract surgery.

Objective: To compare the sensitivity and specification of optical biometry and applanation ultrasound biometry in the measurement of intraocular lens.

Patient and methods: This prospective cohort study was done on 100 eyes from one hundred patients undergoing phacoemulsification with primary intraocular lens implantation. Fifty eyes of patients were measured by ultrasound measurement (by A-Scan, Group 1) and the other 50 eyes by optical biometry (by IOL Master, Group 2).

Result: in Group 1, there were 22 eyes of 22 males (44%), and 28 eyes of 28 females (56%) with a mean age was 58.4 ±0.13 years. While, in Group 2, there were 16 eyes of 16 males (32%), and 34 eyes of thirty-four females (68%) and the mean age of the patients in this group was 61.2±9.25 years. The mean IOL of the patient in Group I was (18.86±2.57) with minimum IOL power (16.29) and maximum IOL power (21.43). Whereas, the mean IOL potent ion of the patient in group II was (20.5±1.98) with minimum IOL power (18.52) and maximum IOL power (+22.48).

Conclusion: We can conclude that IOL measurements performed by using the Zeiss IOL Master, using partial coherence interferometry, resulted that a significantly better IOL power forecast and therefore the refractive result in cataract surgery than Applanation US biometry.

Keywords: Optical Biometry, Ultrasound Biometry, Intraocular lens.

INTRODUCTION

Intraocular lens (IOL) calculation is an essential step in obtaining the exact target that includes the refractive result and is a critical goal in modern cataract surgery. Several devices and formulas are currently available, allowing the precise IOL precision needed to reach target refraction (1,2).

To achieve targeted refraction, the axial length (AL), anterior chamber depth (ACD), and corneal radius (K1 and K2) must be completely accurate (3). Axial length measurement (AL) uses the principle of signal reflection to calculate the difference between the numerous ocular structures and the overall length of the eye. The reflection time of the signal from an interface is measured, divided into two, and multiplied by the speed of the signal in the corresponding medium (4). Distance is measured using the rules: distance = speed * time / 2 the transmitting signal can be ultrasonic. Ultrasound biometrics have been the gold standard for decades. A specific crystal embedded in the probe oscillates to create a high-frequency sound wave to the eye. There are two types of ultrasound biometrics

Biometrics of contact type made by applying an ultrasound probe to the cornea; It carries the risk of infection. Immersion-type biometrics require immersion of the probe in a saline-filled shell. A decade ago, optical biometry was established in clinical practice, and over time optical biometry replaced ultrasound biometry as the standard calculation for measuring the axial length of the eye because it is a fast, easy to use, and contact-free method. Optical biometrics use a 780nm laser diode infrared light to transmit the signal (5). Interface phenomena between the reflected signal and the reference signal and used to determine distances between interfaces.

Previous comparisons between ultrasound and optical biometrics showed equal or better results with ophthalmometric biometrics (6). A recent study concluded that ultrasound biometrics and optical biometrics can be applied interchangeably for IOL calculations (4).

AIM OF THE STUDY

This study aimed to compare the sensitivity and specification of optical biometry and applanation ultrasound biometry in the measurement of intraocular lens.

PATIENT AND METHODS

Study Design: This prospective cohort study was done on patients scheduled for cataract surgery in the Ophthalmology Department of Menoufia University hospitals and El-Mahalla Ophthalmology Hospital. The study was conducted on 100 eyes of one hundred patients undergoing phacoemulsification with primary IOL implantation after the purpose of the study was explained to them. 50 eyes of patients will be measured by ultrasound measurement (by A-Scan, Group 1) and the other 50 eyes by optical biometry (by IOL Master, Group 2).
Ethical consideration: The study was approved by the ethics committee of the department of ophthalmology, faculty of medicine, Menoufia University. Written informed consent was obtained from all patients.

Inclusion criteria:
Patients have a visually considerable cataract suitable for phacoemulsification and primary implantation of rearward chamber intraocular lens in one or both eyes. Also, Preoperatively, Snellen visual acuity was evaluated and all patients go through a cycloplegics refraction, IOP gauging, slit lamp examination for studying the morphology of cataract and under examination by indirect ophthalmoscopy, as well, spherical equivalent < -6 D and or Axial Length < 25.0 mm as measured by Zeiss IOL Master or A-scan biometry, and the patient is willing and able to comply with scheduled visits and other study procedures.

Exclusion criteria:
Patients history of trauma, Patients have an ophthalmic condition other than cataract that could affect have vision or axial length measurements such as optic neuropathy, age regarding macular retrogression, macular edema, retinal detachment, proliferative diabetic retinopathy, ocular inflammation, retinitis pigmentosa, or glaucoma. Also, corneal opacities or irregularities: previous scarring, dystrophy, and ectasia.

Surgical exclusion criteria: Patients were excluded from the study when the following complications were encountered through surgery: failure to achieve secure 'in-the-bag' placement of the IOL. Also, the use of corneal sutures and multiple operative procedures at the time of IOL implantation. Also, Post-implantation exclusion criteria.

Methods:
Patient Preparation:
Patients were submitted to thorough history taking, clinical, examination, and investigations. All the patients underwent: History taking included: age, sex, residence, special habits of the patients or their relatives, main complaint (painless gradual diminution of vision, analysis of the complaint: onset, course, and duration. Also, previous operations, past history of previous ocular surgery, ocular trauma, drug intake, and family history of any ocular disease. General examination and vital signs and ocular examination included first: anterior segment examination: an examination of the external eye. Second: posterior segment examination: fundus examination patient underwent dilated fundus examination with an indirect ophthalmoscope to detect any abnormalities included: vitreous humor, retina, and optic disc.

The preoperative evaluation: All patients were undergone phacoemulsification and posterior chamber in-the-bag foldable IOL implantation, preoperatively, Snellen visual acuity was appreciating and all patients underwent non-cycloplegic autorefraction and fundus examination, preoperatively, patients underwent autorefraction, keratometry measurement, and axial length measurement, and USG B scan was done in patients where media opacity was dense obscuring fundus visualization.

Third, Assessment from the Axial Length of the eye included: Group I (Ultrasound group, 50 eyes) had axial length measurements by A-scan ultrasound and K measurements with manual keratometer. A-scan-guided biometry is done by an ophthalmic ultrasound scanner. The first step in the prediction of IOL power is an accurate keratometric reading. Next, the axial length was measured using the contact A-scan biometry. The patients were prepared by instillation of one drop of tropicamide (mydriacil) 1% then one drop of surface anesthesia in the form of benoxinate hydrochloride 0.4% is dropped into the patient's eye. The data including two keratometric readings and the average axial length plus a constant of IOL were introduced in the calculating program of the Ophthalmic ultrasound scanner.

This procedure of introduction was repeated with the addition of anterior chamber deepness ACD in the form of the Haigis formula. The intraocular lens power was based on the Haigis formula and all patients underwent uncomplicated cataract surgery by phacoemulsification within the bag IOL implantation through an interim clear corneal incision. Group II (IOL Master Group) (50 eyes) had AL and K measurements with the IOL Master. An axial length (AL) data, keratometric values (Ks), and anterior chamber depth (ACD) were acquired with the IOL Master by an experienced doctor.

The surgery:
Phacoemulsification surgery was performed through a 2.3 mm. superior limbal clear corneal incision. All patients underwent in-the-bag implantation of the same IOL power which is soft hydrophobic acrylic foldable IOL obtained by each scan. A standard postoperative topical antibiotic and the anti-inflammatory regime was administered. Patients were studied at the following intervals: 1 day after surgery, 1 week after surgery, 6 weeks after surgery, and 3 months. The primary outcome measure of the study was post-operative spherical equivalent. The actual postoperative spherical equivalence (SE) was recorded 3 months following the surgery by an auto-refractor (Topcon AR, Tokyo, Japan).

The IOL Master permitted calculation using Haigis formula (to calculate the strength of the implanted IOL and predicted postoperative spherical equivalent (SE). The mean absolute error (MAE) was calculated for the Haigis formula which represents the difference between actual postoperative SE and the predicted postoperative SE. This is an absolute rate of the numeric error. Postoperative refraction to compare the results.
Statistical Methodology

The data collected were tabulated & analyzed by SPSS version 15 on IBM compatible computer. Quantitative data were expressed as mean & standard deviation (X+SD) and analyzed by applying student t-test. Qualitative data were expressed as number and percentage (No & %), Fisher exact test, Chi-square test, Student t-test, and Spearman’s correlation were used. p<0.05 was considered a statistically significant level.

RESULTS

Group 1 (A-scan ultrasound biometry group) Included 50 eyes subjected to biometry with A-Scan ultrasound biometry. There were 22 eyes of 22 males (44%), and 28 eyes of 28 females (56%). And the mean age was 58.4 ± 10.13 years. While Group 2 (Optical biometry (IOL Master group)) Included 50 eyes subjected to biometry with IOL Master Optical biometry. There were 16 eyes of 16 males (32%), and 34 eyes of thirty-fourth females (68%), and the mean age of the patient in this group was 61.2 ± 9.25 years.

Also, a total of 100 eyes of one hundred patients were enrolled in our study (eyes in 62 females (62%) and eyes in 38 males (38%). Group I: 22 eyes of 22 males (44%), and 28 eyes of 28 females (56%), and Group II: 16 eyes of 16 males (32%), and 34 eyes of thirty-fourth females (72%), as shown in (table 1).

<table>
<thead>
<tr>
<th>Item</th>
<th>Group 1 (n = 50)</th>
<th>Group 2 (n = 50)</th>
<th>t</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>%</td>
<td>N</td>
<td>%</td>
</tr>
<tr>
<td>Age group</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30-40y</td>
<td>4</td>
<td>8.0%</td>
<td>2</td>
<td>4.0%</td>
</tr>
<tr>
<td>40-50y</td>
<td>6</td>
<td>12.0%</td>
<td>0</td>
<td>0.0%</td>
</tr>
<tr>
<td>50-60y</td>
<td>20</td>
<td>40.0%</td>
<td>22</td>
<td>44.0%</td>
</tr>
<tr>
<td>&gt;60y</td>
<td>20</td>
<td>40.0%</td>
<td>26</td>
<td>52.0%</td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>58.4 ± 10.13</td>
<td>61.2 ± 9.25</td>
<td>1.444</td>
<td>0.152</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Male</td>
<td>22</td>
<td>44.0%</td>
<td>16</td>
<td>32.0%</td>
</tr>
<tr>
<td>Female</td>
<td>28</td>
<td>56.0%</td>
<td>34</td>
<td>68.0%</td>
</tr>
</tbody>
</table>
The mean axial length of the patient in Group I was (23.95 ± 0.89 mm) with minimum axial length 23.02 mm and maximum axial length 24.84 mm, while, in group II was (23.17 ± 0.64 mm) with minimum axial length 22.53 mm and maximum axial length 23.81 mm. Also, the mean k1 reading of the patient in Group I was (44.5 ± 1.78) Dioptr. While in group II was (45.12 ± 2.01) Dioptr. Additionally, group I had mean k2 reading of the patients as (44.85 ± 1.89) Dioptr, and group II: had a mean of (44.86 ± 1.46) Dioptr. Group I, had mean k reading of the patient of (44.6 ± 1.81) Dioptr with minimum average k 42.01D and maximum average k 46.3D, whereas, group II had mean of (45 ± 1.48) Dioptr with minimum average k 42.7 D and maximum average k 46.7 D, as shown in (table 2).

**Table (2):** Mean Axial Length and K reading distribution among both groups.

<table>
<thead>
<tr>
<th>Item</th>
<th>Group 1 (n = 50)</th>
<th>Group 2 (n = 50)</th>
<th>t</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Axial length</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>23.95 ± 0.89</td>
<td>23.17 ± 0.64</td>
<td>5.012</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>k1</td>
<td>44.5 ± 1.78</td>
<td>45.12 ± 2.01</td>
<td>1.635</td>
<td>0.105</td>
</tr>
<tr>
<td>k2</td>
<td>44.85 ± 1.89</td>
<td>44.86 ± 1.46</td>
<td>0.032</td>
<td>0.975</td>
</tr>
<tr>
<td>Average k</td>
<td>44.6 ± 1.81</td>
<td>45 ± 1.48</td>
<td>1.218</td>
<td>0.226</td>
</tr>
<tr>
<td><strong>ACD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- Range</td>
<td>2.84: 3.56</td>
<td>2.81: 3.57</td>
<td>0.145</td>
<td>0.885</td>
</tr>
<tr>
<td>- Mean ± SD</td>
<td>3.2 ± 0.36</td>
<td>3.19 ± 0.38</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ACD: Anterior Chamber Depth

The mean ACD of the patient in Group I was (3.2±0.36) with minimum ACD 2.84 mm and maximum ACD 3.56 mm, while, in group II was (3.19±0.38) with minimum ACD 2.81mm and maximum ACD 3.57mm. Also, the mean IOL of the patient in Group I was (18.86 ± 2.57) with minimum IOL power (16.29) and maximum IOL power (21.43). Whereas, the mean IOL strength of the patient in group II was (20.5 ± 1.98) with minimum IOL power (18.52) and maximum IOL power (21.43). Also, the mean lens thickness in group I was (4.02 ± 0.75) with a minimum of (3.27) and maximum of (4.77). While in group II was (4.17±0.44) with a minimum of (3.73) and a maximum of (4.61). Mean postoperative refraction spherical equivalent of the patient in group I was (-0.65±0.49) in group I: with minimum postoperative refraction spherical equivalent (-0.16) and maximum postoperative refraction spherical equivalent (-1.14). And in group II was (-0.36±0.27) with minimum postoperative refraction spherical equivalent (-0.9) and maximum postoperative refraction spherical equivalent (-0.63), as shown in (table 3).
Ultrasonic Biometry In measuring IOL.

In our study which included 100 eyes of one hundred patients (eyes in 62 females) (62%) and (eyes in 38 males) (38%) with axial lengths less than 25.0 mm with cataract as the only ocular pathology. Postoperative Visual and Refractive outcome out of 100 patients, 50 were implanted IOL calculated by IOLM and 50 by ultrasound. For IOLM patients 48 (96.0%) had postoperative spherical refraction in the range of 0.50 to 0.50 and 2 (4.0%) were outside this range. For ultrasonic patients, 40 (80.0%) had postoperative spherical refraction in the range of 0.50 to 0.50 and 10 (20.0%) were outside this range.

In our study, we used the Haggis formula (a fourth-generation formula) to IOL power calculation in the A-scan guided biometry device and the IOL Master device for biometry, which give the best refractive outcomes due to its inclusion of the IOL Master-measured anterior chamber depth (ACD)\(^{(10)}\). Third-generation formulae such as the Hoffer Q and SRK/T are 2-variable formulae that rely on AL and central corneal power to predict the postoperative IOL position. These formulae do not use actual measurements of the ACD; they assume that short eyes will have shallower ACDs and long eyes will have deeper ACD\(^{(11)}\). In our study, the mean absolute prediction error of optimized IOL Master biometry was significantly smaller (P<0.0001) than that of optimized ultrasound. In our study, the improvement in the refractive outcome of 16% was noticed. In Shah et al.\(^{(12)}\) the mean absolute prediction error of optimized IOL Master biometry was significantly smaller (P<0.0001) than that of optimized ultrasound and an improvement in the refractive results of 23% was noticed. Drexler et al.\(^{(13)}\) and Hitzenberger et al.\(^{(14)}\) reported an improvement regarding 30% when the SRK II formula was used and Rajan et al.\(^{(15)}\) reported a 16% improvement on retrospective IOL power calculations using the IOL Master. Contrary to our study, Gantenbein et al.\(^{(16)}\) Drexler et al.\(^{(17)}\) set up high precision and reproducibility with both methods postoperatively compared to the preoperative aim (P<0.001). There was no statistical difference in the mean absolute error between the two groups. Nevertheless, despite the improvement of the refractive outcome, outliers still exist. This may be due to various cataract characteristics, as the IOL Master utilizes the same group refractive index for all cataract grades. IOL Master Biometry was found to be more accurate in the measurement of the ocular axial length than planplanation.

CONCLUSION

We can conclude that IOL measurements performed with the Zeiss IOL Master, using partial coherence interferometry, yielded significantly best IOL power prediction and therefore refractive results in cataract surgery than Applanation US biometry.

REFERENCE