Article

Repeatability of the Novel Intraocular Pressure Measurement From Corvis ST

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Purpose: To assess the repeatability of intraocular pressure (IOP) measured with the Corvis ST (CST) and the Ocular Response Analyzer (ORA).

Methods: A total of 141 eyes from 141 subjects were studied, including 35 healthy eyes and 106 glaucomatous eyes. All subjects underwent IOP evaluations with Goldmann applanation tonometer, CST, and ORA. With CST, biomechanical corrected IOP (bIOP) was calculated; bIOP is purported to be less dependent on biomechanical properties. For ORA, corneal-compensated intraocular pressure (IOPcc) and Goldmann-correlated IOP (IOPg) were derived. The repeatability of the various IOP values was assessed using the coefficient of variance (CV) and the intraclass correlation coefficient (ICC).

Results: The CV with bIOP (5.5 ± 3.1: mean ± standard deviation) was significantly smaller than the CVs measured with IOPg (7.3 ± 4.3) and IOPcc (7.2 ± 4.4). ICC values were 0.90, 0.80, and 0.86 with IOPg, IOPcc, and bIOP, respectively.

Conclusions: The bIOP showed a better prevision and repeatability for IOP measurement.

Translational Relevance: The bIOP measurement from CST had a better reproducible than IOPcc measurement from ORA.

Introduction

Glaucoma can severely damage a patient’s visual function, including the visual field and visual acuity; it remains the second leading cause of blindness worldwide, affecting 60 million people.1 It is widely acknowledged that the irreversible damage to visual fields, caused by progressive retinal ganglion cell loss, is intensified by elevated intraocular pressure (IOP), which is an established risk factor of glaucoma.2–10

The Goldmann applanation tonometer (GAT) is the most common method for measuring IOP, especially in the management of glaucoma. Importantly, the accuracy of GAT can be affected by many factors. Previous studies have shown that GAT IOP may be overestimated when central cornea thickness (CCT) is large, and underestimated when CCT is small.11–13 However variations in CCT only account for up to 12% of the measured variation of GAT-IOP,14 and hence correction nomograms that adjust GAT IOP based solely on CCT are neither valid nor useful in individual patients.16 A recent study with a simulated cornea biomechanical model revealed that corneal biomechanics across individuals may have greater impact on IOP measurement errors than CCT.17 For instance, it has been reported that corneal hysteresis (CH) measured with the Ocular Response Analyzer (ORA; Reichert Ophthalmic Instruments, Buffalo, NY)18 influences the GAT IOP measurement,19 and reflecting this, ORA generates the following two IOP measurements: a corneal compensated IOP (IOPcc), which is corrected for CH, as well as the IOP Goldmann (IOPg).

In addition, there has recently been renewed interest in measuring IOP more accurately;
particular, the Corneal Visualization Scheimpflug Technology instrument (Corvis ST tonometry: CST; Oculus, Wetzlar, Germany) is a new noncontact tonometer designed to measure IOP while correcting for the biomechanical properties of the cornea. Similar to ORA, a rapid air-puff is used in CST, but unlike ORA, an ultra-high speed Scheimpflug camera is used to directly visualize the associated corneal movement. CST generates a biomechanical corrected IOP (bIOP) measurement, which is corrected for CCT and other properties of the cornea. A previous study suggested that bIOP is not dependent on CCT in a normative population. Thus, bIOP may be useful in the management of glaucoma, however, the reproducibility and repeatability of the measurement have not been investigated. Thus, the purpose of the current study was to investigate the repeatability of these IOP values.

Materials and Methods

The study was approved by the Research Ethics Committee of the Graduate School of Medicine and Faculty of Medicine at The University of Tokyo. Written informed consent was given by patients for their information to be stored in the hospital database and used for research. This study was performed according to the tenets of the Declaration of Helsinki.

Subjects

The current study investigated 141 eyes of 141 subjects (35 healthy eyes and 106 glaucoma eyes). Inclusion criteria for glaucomatous patients were as follows: no abnormal eye-related findings except for primary open-angle glaucoma (POAG) or normal-tension glaucoma (NTG) on biomicroscopy, gonioscopy, and funduscopy. Eyes with corneal pathologic features, such as Fuch’s endothelial dystrophy or keratoconus, were excluded. Eyes that had undergone any intraocular surgery, including cataract surgery, were also excluded. Only subjects aged more than 20 years were included. IOP was not used as an exclusion criterion so that a wide range of IOPs were considered in the analysis. As a result, ocular hypertensive eyes were included. Normative subjects were defined as having no abnormal findings except for clinically insignificant senile cataract on biomicroscopy, gonioscopy, and funduscopy, and no history of any ocular disease or surgery, including cataract surgery. Both glaucoma patients and normative subjects were recruited at the glaucoma clinic in the University of Tokyo hospital.

Corvis ST Tonometry Measurements

Measurements with CST were carried out three times per participant. An interval of approximately 1 minute was given between each measurement during which data storage and processing operations were carried out by the CST instrument. All CST measurements were considered reliable according to the “OK” quality index displayed on the device monitor. The principles of the CST have been described in detail elsewhere. Briefly, a high-speed Scheimpflug camera records over 4000 frames/sec to monitor the corneal response to an air-puff pulse that forces the cornea inward until it reaches a concavity phase. A number of CST parameters are produced including bIOP, which is an IOP value corrected for CCT. The formula to calculate bIOP was initially suggested as:

\[
\text{bIOP} = \left( C_{\text{CCT1}} \times C_{\text{CST-IOP}} + C_{\text{CCT2}} \right) \times C_{\text{age}} + C,
\]

where

\[
C_{\text{CCT1}} = 4.67 \times 10^{-7} \times \text{CCT}^2 - 7.8 \times 10^{-4} \times \text{CCT} + 0.63
\]

\[
C_{\text{CCT2}} = -1.73 \times 10^{-5} \times \text{CCT}^2 + 2.02 \times 10^{-3} \times \text{CCT} - 0.97
\]

\[
C_{\text{CST-IOP}} = 10 + \left( \text{CST-IOP} + 1.16 \right)/0.389
\]

\[
C_{\text{age}} = -2.01 \times 10^{-5} \times \text{age}^2 + 1.3 \times 10^{-3} \times \text{age} + 1.00
\]

\[
C = 1.5 \text{ mmHg}
\]

However, an updated formula is used in the latest version of the software (personal communication with Oculus):

\[
\text{bIOP} = C_{\text{CCT1}} \times C_{\text{AP1}} \times C_{\text{age1}} + C_{\text{CCT2}} \times C_{\text{age2}} + C_{\text{DCR}} + \text{a19}
\]

where

\[
C_{\text{CCT1}} = (a_1 \times \text{CCT}^3 + a_2 \times \text{CCT}^2 + a_3 \times \text{CCT} + a_4)
\]

\[
C_{\text{AP1}} = (a_5 \times \text{AP1} + a_6)
\]

\[
C_{\text{age1}} = (a_7 \times \text{Ln(Beta)}^2 + a_8 \times \text{Ln(Beta)} + a_9)
\]

\[
C_{\text{CCT2}} = (a_{10} \times \text{CCT}^3 + a_{11} \times \text{CCT}^2 + a_{12} \times \text{CCT} + a_{13})
\]
\[ C_{age2} = \left( a_{14} \times [ \ln(\text{Beta})]^2 + a_{15} \times [\ln(\text{Beta})] + a_{16} \right) \]

\[ \text{Beta} = 0.5852 \times \exp(0.0111 \times \text{Age[year]}) \]

\[ C_{DCR} = a_{17} \times \text{highest concavity radius} + a_{18} \]

where the highest concavity radius is a CST parameter which represents the curvature radius when the cornea is at the point of highest concavity.

**Ocular Response Analyzer Measurements**

The ORA measurement was carried out three times on the same day with the GAT measurement. An interval of approximately 1 minute was given between each repeated measurement. ORA records two applanation pressure measurements, during the inward and outward corneal movements, following application of a rapid air-puff. Due to its viscoelastic property, the cornea resists the air-puff, resulting in delays in the corneal movement, which cause a measurable difference in the applanation pressure values at the inward and outward corneal movements. The average of the inward and outward applanation pressure values defines the IOPg measurement, and the difference in the two values describes CH, which is primarily an indication of viscous damping in the corneal tissue. The corneal resistance factor (CRF) is also calculated from the two applanation pressure values, but places greater emphasis on the first applanation pressure because this gives more information about the elastic properties of the cornea.

IORpc is another IOP measurement generated by ORA that incorporates the CH metrics in its derivation. Only reliable data, as indicated by a waveform score more than 7.0, were used in the analyses.

The order of ORA and CST measurements was decided randomly.

**Other Ocular Measurements**

GAT measurements were carried out after the CST and ORA measurements, and after instillation of topical 0.5% tetracaine. The tonometer was set at 10 mm Hg before each reading. AL was measured three times using the IOL master (Carl Zeiss Meditec, Dublin, CA) and CCT was measured three times using the CST. Average values were used in analyses.

**Statistical Analysis**

The relationship between the different IOP measurements was analyzed using linear regression and the paired Wilcoxon test. The agreement between devices, (1) IOPpc and GAT IOP, and (2) bIOP and GAT IOP, measured using Bland-Altman plot. The coefficient of variation (CV) and the intraclass correlation coefficient (ICC) were calculated for bIOP, IOPg, and IOPpc. CV was calculated as follows:

\[ CV = \frac{\text{standard deviation of three IOP readings}}{\text{average of three IOP readings}} \]

CV values were compared using the paired Wilcoxon test.

All statistical analyses were performed using the statistical programming language R (R version 3.2.3; the Foundation for Statistical Computing, Vienna, Austria).

**Results**

Characteristics of the study subjects are summarized in Table 1. The mean ± standard deviation (SD) (range) age of patients was 55.2 ± 16.5 (24–86). Seventy-one participants were male and seventy participants were female. Mean ± SD GAT IOP was 13.1 ± 2.7 (7.8–22) mm Hg. CCT was 537.7 ± 34.6 (458.7–644.0) μm. Corneal curvature was 7.7 ± 0.3 (7.2–8.3) mm. CH was 9.5 ± 1.2 (6.4–12.4) mm Hg. CRF was 8.8 ± 1.6 (4.2–15.2) mm Hg.

The ICC and CV values of IOPg, IOPcc, and bIOP are summarized in Table 2. The CV value of IOPg was 7.3 ± 4.3 (0.37–21.4)%; IOPcc was 7.2 ± 4.4 (0.41–24.2)%, and bIOP was 5.5 ± 3.1 (0.59–18.1)%. The CV value of bIOP was significantly smaller than those with IOPg and IOPcc (P < 0.001, Wilcoxon test). ICC of IOPg, IOPcc, and bIOP were 0.90 (95% confidence interval [CI]: from 0.82–0.93), 0.80 (95% CI: from 0.75–0.85), and 0.86 (95% CI: from 0.82–0.89), respectively.

Mean ± SD values of IOPg, IOPcc, bIOP, and GAT IOP are shown in Table 3. IOPcc was significantly higher than GAT IOP (P < 0.001, paired Wilcoxon test), whereas bIOP was not significantly different from GAT IOP (P = 0.34, paired Wilcoxon test). Figure 1 shows the relationships between IOPpc and GAT IOP, and between bIOP and GAT IOP. There was a significant relationship between these values (R² = 0.39 and 0.38, respectively, P < 0.001, linear regression).

Figure 2 shows the agreement between IOPpc and GAT IOP, and between bIOP and GAT IOP, using the Bland-Altman plot. Mean difference between
IOPcc and GAT IOP was 1.4 mm Hg, whereas it was 
\(-0.2\) mm Hg between bIOP and GAT IOP. There 
was no significant relationship between the differ-
ence between IOPcc and GAT IOP and the average 
of these values; however, the difference between 
bIOP and GAT IOP was significantly larger when 
the average of these values was small (\(P < 0.001\)); the 
bIOP value was smaller than the GAT IOP 
measurement when the average of these values was 
small.

**Discussion**

In the current study, IOP measurements were 
carried out using GAT, CST, and ORA in glauco-
matous patients and normative subjects. Among the IOP 
measurements, bIOP (CST) had a lower CV and a 
higher ICC compared with IOPg (ORA) and IOPcc 
(ORA), suggesting a high repeatability of bIOP. All 
IOP measurements were significantly related to one 
another as follows: IOPg tended to be lower than 
GAT IOP while IOPcc tended to be higher than GAT 
IOP; there was not a significant difference between 
bIOP and GAT IOP.

In a previous report, we investigated the repeat-
ability of ORA IOP measurements in eyes with 
POAG. The CV and ICC values of IOPg were 6.5 
\(\pm\) 4.0\% and 0.91, respectively.\(^{26}\) The CV and ICC 
values of IOPcc were 6.7 \(\pm\) 4.0\% and 0.84, 
respectively.\(^{26}\) Many others have also investigated 
the reproducibility of these measurements. Wang et 
al.\(^{27}\) reported that the CV and ICC of IOPg is 7.0\% 
and 0.79, respectively, while these same values are 
9.8\% and 0.57 for IOPcc. Kopito et al.\(^{28}\) reported that 
the CV values of IOPg and IOPcc were 7.7 and 10.1\%. 
Moreno-Montañez et al.\(^{29}\) reported that the ICC 
values of IOPg and IOPcc were 0.93 and 0.78. In the 
current study, the CV values of IOPg and IOPcc were 
7.3 and 7.2, and the ICC values of IOPg and IOPcc 
were 0.90 and 0.80; these values align well with those 
published in previous reports. The results in the 
current study suggested lower CV (5.5 \(\pm\) 3.1\% with 
bIOP and 7.2 \(\pm\) 4.4\% with IOPcc) and higher ICC 
values (0.86 with bIOP and 0.80 with IOPcc) with 
bIOP compared with IOPcc, suggesting better repeat-
ability of bIOP compared with IOPcc. The entire 
reason of this finding is not clear, but it may be 
because of the different calculations between these 
two IOP values. bIOP is calculated using age, CCT, 
and highest concavity radius. Age does not change 
between the CST measurements. We previously 
reported CST measured CCT was highly repeatable 
(CV = 0.9 \(\pm\) 0.9\% and ICC = 0.99) although highest 
concavity radius (formally named as highest concav-
ity curvature) had a moderate repeatability (CV = 8.1 
\(\pm\) 8.7\% and ICC = 0.68) in a previous study.\(^{30}\) The 
high repeatability of bIOP would be beneficial when 
used at the clinical settings.

Many earlier reports have suggested a difference 
between IOPcc and GAT IOP. Hager et al.\(^{31}\) 
compared IOPcc and GAT IOP in eyes with 
glaucoma, and reported that IOPcc was significantly 
higher than GAT IOP by 3.6 mm Hg (17.9 \(\pm\) 5.9: 

| Variable | Value |
|----------|-------|
| Age, mean \(\pm SD\) (range), \(y\) | 55.2 \(\pm\) 16.5 (24–86) |
| Male/female | 71/70 |
| Right/left | 113/28 |
| AL, mean \(\pm SD\) (range), mm | 25.5 \(\pm\) 1.6 (22.3–29.2) |
| GAT IOP, mean \(\pm SD\) (range), mm Hg | 13.1 \(\pm\) 2.7 (7.8–22.0) |
| bIOP, mean \(\pm SD\) (range), mm Hg | 12.9 \(\pm\) 2.2 (8.6–20.3) |
| Corneal curvature, mean \(\pm SD\) (range), \(\mu m\) | 7.7 \(\pm\) 0.3 (7.2–8.3) |
| CCT, mean \(\pm SD\) (range), \(\mu m\) | 537.7 \(\pm\) 34.6 (458.7–644.0) |
| CH, mean \(\pm SD\) (range), mm Hg | 9.5 \(\pm\) 1.2 (6.4–12.4) |
| CRF, mean \(\pm SD\) (range), mm Hg | 8.8 \(\pm\) 1.6 (4.2–15.2) |

AL, axial length.
mean ± SD and 14.3 ± 4.3 mm Hg). Martinez de la Casa et al. compared IOPcc and GAT IOP in POAG eyes and reported that IOPcc was significantly higher than GAT IOP by 8.3 mm Hg (25.1 ± 5.4 and 16.8 ± 3.4 mm Hg). Ehrlich et al. have reported that IOPcc was significantly higher than GAT IOP by 5.4 mm Hg (19.8 ± 3.4 and 14.4 ± 3.4 mm Hg) in NTG eyes. Oncel et al. reported that IOPcc was higher than GAT IOP by 1.0 mm Hg in healthy volunteers (15.8 ± 2.9 and 14.8 ± 3.1 mm Hg). Pepose et al. reported that IOPcc was higher than GAT IOP by 1.6 mm Hg (15.4 ± 3.2 and 13.8 ± 3.3 mm Hg) among eyes with myopia. In the current study, IOPcc was significantly higher than GAT IOP (P < 0.001); however, the difference was much smaller (by 1.4 mm Hg: 14.5 ± 2.6 and 13.1 ± 2.7 mm Hg) compared with these previous reports, but similar to that observed in our previous report where IOPcc was significantly higher than GAT IOP by 1.6 mm Hg. Thus, the reason for the small difference between IOPcc and GAT IOP observed in the current study is not clear, but may be attributable to the racial difference in study populations. Indeed, Morita et al. also compared IOPcc and GAT IOP in a Japanese population and likewise reported a much smaller difference between IOPcc and GAT IOP (by 2.1 mm Hg: 15.2 ± 2.0 and 13.1 ± 1.3 mm Hg) in NTG eyes, and no significant difference in healthy eyes (IOPcc: 13.6 ± 2.0 mm Hg and GAT IOP: 13.2 ± 1.4 mm Hg). In contrast to IOPcc, there was not a significant difference between biOP and GAT IOP in the current study. It would be of further interest to investigate whether similar results are obtained in other ethnicities.

A recent report from the United Kingdom Glaucoma Treatment Study suggested that, among IOPg, IOPcc, GAT IOP, and IOP with dynamic contour tonometry, IOPcc from ORA had the highest probability of being the best predictor of glaucoma progression (Lascaratos, et al. IOVS. 2014;55:ARVO E-Abstract 128). Further, Hong et al. reported that rapid visual field (VF) progression was more likely to occur in patients with high IOPcc, low CH, and a large recorded difference between IOPcc and GAT IOP. CST is a relatively new noncontact tonometry, and we have shown glaucomatous VF progression and also severity can be even better analyzed using CST-derived parameters (biOP was not analyzed, because of the older software used). Albeit with the high repeatability suggested in the current study, biOP may be useful to assess the progression of glaucomatous visual field (VF) progression. Another aspect to be considered is the relationship between IOP reading with each device and corneal biomechanical properties, because IOP measurements, such as GAT can be affected by CCT, and also progression of glaucoma is associated with various corneal properties such as CCT.

### Table 3. Summary of IOPg, IOPcc and biOP Measurements and Their Correlation With GAT IOP

| Measure          | Mean ± SD (range) | Correlation to GAT IOP (P value) |
|------------------|-------------------|---------------------------------|
| IOPg, mm Hg      | 12.6 ± 3.2 (6.2–26.3) | <0.001                          |
| IOPcc, mm Hg     | 14.5 ± 2.6 (8.7–23.2) | <0.001                          |
| biOP, mm Hg      | 12.9 ± 2.2 (8.6–20.3) | <0.001                          |
| GAT IOP, mm Hg   | 13.1 ± 2.7 (7.8–22.0) |                                  |
and also CST measured corneal biomechanical characteristics. In the current study, there was a significant difference between IOPcc and GAT IOP, whereas this was not the case between bIOP and GAT IOP. A future study would be of interest to investigate whether bIOP is a better predictor of glaucoma progression, preparing longitudinal data.

A limitation of the current study is that IOP data were obtained from a hospital clinic, hence true IOP could not be collected and compared against the various IOP readings, as in a previous study. Further, the effects of antiglaucomatous eye drops on corneal biomechanical properties was not considered. In addition, GAT IOP measurement was conducted either once, twice, or three times, and not in a masked fashion (i.e., the GAT dial was not set to a random number and then the final reading was recorded), as in a previous study, because the repeatability of GAT IOP has already been reported and it was not the purpose of the current study.

In conclusion, the CST-derived bIOP measurement has a good repeatability.

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References

1. Quigley HA. Glaucoma. Lancet. 2011;377:1367–1377.
2. Heijl A, Leske MC, Bengtsson B, et al. Reduction of intraocular pressure and glaucoma progression: results from the Early Manifest Glaucoma Trial. Arch Ophthalmol. 2002;120:1268–1279.
3. Garway-Heath DF, Crabbe DP, Bunce C, et al. Latanoprost for open-angle glaucoma (UKGTS): a randomised, multicentre, placebo-controlled trial. Lancet. 2015;385:1295–304.
4. Leske MC, Heijl A, Hyman L, Bengtsson B, Dong L, Yang Z. Predictors of long-term progression in the early manifest glaucoma trial. Ophthalmology. 2007;114:1965–1972.
5. Holmin C, Thorburn W, Krakau CE. Treatment versus no treatment in chronic open angle glaucoma. Acta Ophthalmol (Copenh). 1988;66:170–173.
6. Pajic B, Pajic-Eggspuehler B, Hafliger IO. Comparison of the effects of dorzolamide/timolol and latanoprost/timolol fixed combinations upon intraocular pressure and progression of visual field damage in primary open-angle glaucoma. Curr Med Res Opin. 2010;26:2213–2219.
7. Migdal C, Gregory W, Hitchings R. Long-term functional outcome after early surgery compared with laser and medicine in open-angle glaucoma. Ophthalmology. 1994;101:1651–1656; discussion 1657.
8. Jay JL, Murray SB. Early trabeculectomy versus conventional management in primary open angle glaucoma. Br J Ophthalmol. 1988;72:881–889.
9. Musch DC, Gillespie BW, Lichter PR, Niziol LM, Janz NK. Visual field progression in the Collaborative Initial Glaucoma Treatment Study the impact of treatment and other baseline factors. Ophthalmology. 2009;116:200–207.
10. The Advanced Glaucoma Intervention Study (AGIS): 7. The relationship between control of intraocular pressure and visual field deterioration. The AGIS Investigators. Am J Ophthalmol. 2000;130:429–440.
11. Ehlers N, Bramsen T, Sperling S. Applanation tonometry and central corneal thickness. Acta Ophthalmol (Copenh). 1975;53:34–43.
12. Whitacre MM, Stein RA, Hassanein K. The effect of central corneal thickness on intraocular pressure measured by pneumotonometry, non-contact tonometry, the Tono-Pen XL, and Goldmann applanation tonometry. Br J Ophthalmol. 1993;115:592–596.
13. Doughty MJ, Zaman ML. Human corneal thickness and its impact on intraocular pressure measures: a review and meta-analysis approach. Surv Ophthalmol. 2000;44:367–408.
14. Tonnu PA, Ho T, Newson T, et al. The influence of central corneal thickness and age on intraocular pressure measured by pneumotonometry, non-contact tonometry, the Tono-Pen XL, and Goldmann applanation tonometry. Br J Ophthalmol. 2005;89:851–854.
15. Ku JY, Danesh-Meyer HV, Craig JP, Gamble GD, McGhee CN. Comparison of intraocular pressure measured by Pascal dynamic contour tonometry and Goldmann applanation tonometry. Eye (Lond). 2006;20:191–198.
16. Weinreb RN, Brandt JD, Garway-Heath D, Medeiros F. Intraocular pressure. Wayne: Kugler Publications; 2007.

17. Liu J, Roberts CJ. Influence of corneal biomechanical properties on intraocular pressure measurement: quantitative analysis. *J Cataract Refract Surg*. 2005;31:146–155.

18. Terai N, Raiskup F, Haustein M, Pillunat LE, Spoerl E. Identification of biomechanical properties of the cornea: the ocular response analyzer. *Curr Eye Res*. 2012;37:553–562.

19. Broman AT, Congdon NG, Bandeen-Roche K, Quigley HA. Influence of corneal structure, corneal responsiveness, and other ocular parameters on tonometric measurement of intraocular pressure. *J Glaucoma*. 2007;16:581–588.

20. Joda AA, Shervin MM, Kook D, Elsheikh A. Development and validation of a correction equation for Corvis tonometry. *Comput Methods Biomech Biomed Engin*. 2016;19:943–953.

21. Vinciguerra R, Elsheikh A, Roberts CJ, et al. Influence of pachymetry and intraocular pressure on dynamic corneal response parameters in healthy patients. *J Refract Surg*. 2016;32:550–561.

22. Koprowski R. Automatic method of analysis and measurement of additional parameters of corneal deformation in the Corvis tonometer. *Biomed Eng Online*. 2014;13.

23. Luce DA. Determining in vivo biomechanical properties of the cornea with an ocular response analyzer. *J Cataract Refract Surg*. 2005;31:156–162.

24. Roberts CJ, Liu J. Corneal Biomechanics: From Theory to Practice. Wayne: Kugler Publications; 2017.

25. Ayala M, Chen E. Measuring corneal hysteresis: threshold estimation of the waveform score from the Ocular Response Analyzer. *Graefes Arch Clin Exp Ophthalmol*. 2012;250:1803–1806.

26. Matsuura M, Hirasawa K, Murata H, et al. The Relationship between Corvis ST tonometry and ocular response analyzer measurements in eyes with glaucoma. *PLoS One*. 2016;11:e0161742.

27. Carbonaro F, Andrew T, Mackey DA, Spector TD, Hammond CJ. The heritability of corneal hysteresis and ocular pulse amplitude: a twin study. *Ophthalmology*. 2008;115:1545–1549.

28. Kopito R, Gaujoux T, Montard R, et al. Reproducibility of viscoelastic property and intraocular pressure measurements obtained with the Ocular Response Analyzer. *Acta Ophthalmol.* 2011;89:e225–230.

29. Moreno-Montanes J, Maldonado MJ, Garcia N, Mendiluce L, Garcia-Gomez PJ, Segui-Gomez M. Reproducibility and clinical relevance of the ocular response analyzer in nonoperated eyes: corneal biomechanical and tonometric implications. *Invest Ophthalmol Vis Sci*. 2008;49:968–974.

30. Asaoka R, Nakakura S, Tabuchi H, et al. The relationship between Corvis ST tonometry measured corneal parameters and intraocular pressure, corneal thickness and corneal curvature. *PLoS One*. 2015;10:e0140385.

31. Hager A, Loge K, Schroeder B, Fullhas MO, Wiegand W. Effect of central corneal thickness and corneal hysteresis on tonometry as measured by dynamic contour tonometry, ocular response analyzer, and Goldmann tonometry in glaucomatous eyes. *J Glaucoma*. 2008;17:361–365.

32. Martinez-de-la-Casa JM, Garcia-Feijoo J, Fernandez-Vidal A, Mendez-Hernandez C, Garcia-Sanchez J. Ocular response analyzer versus Goldmann applanation tonometry for intraocular pressure measurements. *Invest Ophthalmol Vis Sci*. 2006;47:4410–4414.

33. Ehrlich JR, Radcliffe NM, Shimmyo M. Goldmann applanation tonometry compared with corneal-compensated intraocular pressure in the evaluation of primary open-angle Glaucoma. *BMC Ophthalmol*. 2012;12.

34. Oncel B, Dinc U, Orge F, Yalvac B. Comparison of IOP measurement by ocular response analyzer, dynamic contour, Goldmann applanation, and noncontact tonometry. *Eur J Ophthalmol*. 2009;19:936–941.

35. Pepose JS, Feigenbaum SK, Qazi MA, Sanderson JP, Roberts CJ. Changes in corneal biomechanics and intraocular pressure following LASIK using static, dynamic, and noncontact tonometry. *Am J Ophthalmol*. 2007;143:39–47.

36. Morita T, Shoji N, Kamiya K, Hagishima M, Fujimura F, Shimizu K. Intraocular pressure measured by dynamic contour tonometer and ocular response analyzer in normal tension glaucoma. *Graefes Arch Clin Exp Ophthalmol*. 2010;248:73–77.

37. Hong Y, Shoji N, Morita T, et al. Comparison of corneal biomechanical properties in normal tension glaucoma patients with different visual field progression speed. *Int J Ophthalmol*. 2016;9:973–978.
39. Hirasawa K, Matsuura M, Murata H, et al. Association between corneal biomechanical properties with ocular response analyzer and also CorvisST tonometry, and glaucomatous visual field severity. *Transl Vis Sci Technol*. 2017;6(3):18.

40. Whitacre MM, Stein R. Sources of error with use of Goldmann-type tonometers. *Surv Ophthalmol*. 1993;38:1–30.

41. Kotecha A, Elsheikh A, Roberts CR, Zhu H, Garway-Heath DF. Corneal thickness- and age-related biomechanical properties of the cornea measured with the ocular response analyzer. *Invest Ophthalmol Vis Sci*. 2006;47:5337–5347.

42. Feltgen N, Leifert D, Funk J. Correlation between central corneal thickness, applanation tonometry, and direct intracameral IOP readings. *Br J Ophthalmol*. 2001;85:85–87.

43. Bhan A, Browning AC, Shah S, Hamilton R, Dave D, Dua HS. Effect of corneal thickness on intraocular pressure measurements with the pneumotonometer, Goldmann applanation tonometer, and Tono-Pen. *Invest Ophthalmol Vis Sci*. 2002;43:1389–1392.

44. Foster PJ, Baasanhu J, Alsbirik PH, Munkhbayar D, Uranchimeg D, Johnson GJ. Central corneal thickness and intraocular pressure in a Mongolian population. *Ophthalmology*. 1998;105:969–973.

45. Gunvant P, Baskaran M, Vijaya L, et al. Effect of corneal parameters on measurements using the pulsatile ocular blood flow tonograph and Goldmann applanation tonometry. *Br J Ophthalmol*. 2004;88:518–522.

46. Shah S, Chatterjee A, Mathai M, et al. Relationship between corneal thickness and measured intraocular pressure in a general ophthalmology clinic. *Ophthalmology*. 1999;106:2154–2160.

47. Shimmyo M, Ross AJ, Moy A, Mostafavi R. Intraocular pressure, Goldmann applanation tension, corneal thickness, and corneal curvature in Caucasians, Asians, Hispanics, and African Americans. *Am J Ophthalmol*. 2003;136:603–613.

48. Stodtmeister R. Applanation tonometry and correction according to corneal thickness. *Acta Ophthalmol Scand*. 1998;76:319–324.

49. Wolfs RC, Klaiver CC, Vingerling JR, Grobbee DE, Hofman A, de Jong PT. Distribution of central corneal thickness and its association with intraocular pressure: the Rotterdam Study. *Am J Ophthalmol*. 1997;123:767–772.

50. Jonas JB, Holbach L. Central corneal thickness and thickness of the lamina cribrosa in human eyes. *Invest Ophthalmol Vis Sci*. 2005;46:1275–1279.

51. Wells AP, Garway-Heath DF, Poostchi A, Wong T, Chan KC, Sachdev N. Corneal hysteresis but not corneal thickness correlates with optic nerve surface compliance in glaucoma patients. *Invest Ophthalmol Vis Sci*. 2008;49:3262–3268.

52. Zhong Y, Shen X, Yu J, Tan H, Cheng Y. The comparison of the effects of latanoprost, travoprost, and bimatoprost on central corneal thickness. *Cornea*. 2011;30:861–864.

53. Nielsen CB, Nielsen PJ. Effect of alpha- and beta-receptor active drugs on corneal thickness. *Acta Ophthalmol (Copenh)*. 1985;63:351–354.

54. Inoue K, Okugawa K, Oshika T, Amano S. Influence of dorzolamide on corneal endothelium. *Jpn J Ophthalmol*. 2003;47:129–133.

55. Kaminski S, Hommer A, Koyuncu D, Biowski R, Barisani T, Baumgartner I. Influence of dorzolamide on corneal thickness, endothelial cell count and corneal sensitivity. *Acta Ophthalmol Scand*. 1998;76:78–79.

56. Kotecha A, White E, Schlottmann PG, Garway-Heath DF. Intraocular pressure measurement precision with the Goldmann applanation, dynamic contour, and ocular response analyzer tonometers. *Ophthalmology*. 2010;117:730–737.

57. Pandav SS, Sharma A, Gupta A, Sharma SK, Gupta A, Patnaik B. Reliability of proton and Goldmann applanation tonometers in normal and postkeratoplasty eyes. *Ophthalmology*. 2002;109:979–984.

58. Dielemans I, Vingerling JR, Hofman A, Grobbee DE, de Jong PT. Reliability of intraocular pressure measurement with the Goldmann applanation, dynamic contour, and ocular response analyzer tonometers. *Ophthalmology*. 2010;117:730–737.

59. Drielemans I, Vingerling JR, Hofman A, Grobbee DE, de Jong PT. Reliability of intraocular pressure measurement with the Goldmann applanation tonometer in epidemiological studies. *Graefes Arch Clin Exp Ophthalmol*. 1994;232:141–144.

60. Tonnu PA, Ho T, Sharma K, White E, Bunce C, Garway-Heath D. A comparison of four methods of tonometry: method agreement and interobserver variability. *Br J Ophthalmol*. 2005;89:847–850.