Clinical comparison of the ProTon and Tono-Pen tonometers with the Goldmann applanation tonometer

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Abstract
A clinical evaluation of a new electronic ProTon tonometer was performed comparing the values of intraocular pressure (IOP) measured using this instrument with those determined by a similar instrument, Tono-Pen XL, and by Goldmann applanation tonometry. The mean IOP measured in 106 eyes with the ProTon tonometer was not significantly different from that determined with Goldmann applanation, while the IOP values measured with Tono-Pen XL were significantly lower. The 95% limits of agreement between applanation tonometry and ProTon tonometry were between −4 mm Hg and 5 mm Hg and between applanation tonometry and Tono-Pen XL tonometry between −3 mm Hg and 8 mm Hg. The ProTon tonometer appears to have a higher level of accuracy than the Tono-Pen XL tonometer in clinical practice.

In recent years, an automatic microprocessor controlled tonometer, Tono-Pen, has been developed for the measurement of intraocular pressure (IOP). Recently, a new, similar instrument, the ProTon tonometer has been introduced in Norway. The advantage of these small, hand held instruments is that tonometry can be performed more easily and conveniently for both patient and examiner than with classic instruments, such as the Schiitz tonometer, based on mechanical measurement principles. General practitioners and opticians have begun to use these new tonometers, while for ophthalmologists Goldmann applanation tonometry is still a standard method for determining IOP. However, for some patients — for example, those who do not cooperate with applanation tonometry or who have corneal epithelial irregularities, the new tonometers which come into contact with a very small area of the cornea, may be considered as alternative instruments.

Before these new tonometers can be accepted as adequate alternatives to standard applanation tonometry, their accuracy should be assessed and they should be tested in a routine clinical examination procedure. Until now, several studies have been performed with the Tono-Pen tonometer, but its reliability has not been confirmed. Some investigators have suggested that Tono-Pen measurements of IOP were accurate when compared with Goldmann applanation tonometry, while others have reported significant differences between these two methods or poor reproducibility of measurements. Several studies have shown that the Tono-Pen has a tendency to overestimate pressures at low IOP levels and underestimate pressures at high IOP levels, or it consistently overestimated the actual IOP in an unpredictable manner. It was thus found to be difficult to accept this instrument for clinical use.

The purpose of this study was to perform a clinical examination of the new ProTon tonometer using Goldmann applanation tonometry as a standard method. Furthermore, a comparison of the results obtained with the ProTon and Tono-Pen XL tonometers was also performed in order to determine their relative accuracy.

Materials and methods
The two electronic tonometers ProTon (Tomey Technology, Inc, Cambridge, MA, USA) and Tono-Pen XL (Bio-Rad Inc, Santa Ana, CA, USA) share similar measurement principles, developed on the basis of the Mackay-Marg tonometer. However, their design is different. In both instruments, IOP is converted to an electronic signal which is stored and analysed by a microprocessor. The means of several measurements are given on a digital display, which also shows the range of the single measurements as a measure of reliability. The digital readout minimises user bias. In the Proton tonometer, the probe and the display unit are separated, while these are located together in the Tono-Pen XL. The Tono-Pen XL stores and averages four accepted measurements, while the ProTon tonometer usually averages three to five. Both instruments emit an audible beep, making the observer aware of the accepted readings. In this study, only the readings with a reliability of 5% were accepted for Tono-Pen XL and with the highest accuracy (SEM <0.5 mm Hg) for the ProTon. The IOP was measured according to the instruction manual and both instruments were calibrated daily.

Goldmann applanation tonometry was performed with the applanation tonometer on a Rodenstock slit-lamp. An average systolic diastolic measurement was obtained for each eye. Two measurements were performed on each eye. However, if the difference between the two measurements exceeded 2 mm Hg, three readings were averaged.

Examination of the patients was performed in a masked fashion. A total of 53 consecutive patients admitted for a routine clinical examination at our department gave their permission to be included in the study. Patients with corneal irregularities were excluded. The refractive error of all of the patients was within −8.5 D to +4.0 D in both eyes with the exception of three
patients with aphakia in the left or right eye. The refractive error of these three eyes was 11.5-13.0 D. The astigmatism of one of the aphakic eye was −4.0 D, while astigmatism of all of the other eyes did not exceed −2.0 D.

IOP was measured in the right and left eyes of the patients by three investigators (including the author (AW) and two ophthalmic assistants). Three different instruments were used: the Goldmann applanation tonometer, the Tono-Pen XL tonometer, and the ProTon tonometer. All investigators were trained to use these instruments before the start of the study.

All patients were examined in the same sequence starting with applanation tonometry, followed by ProTon tonometry and Tono-Pen tonometry. The set of readings with these three instruments was obtained within intervals not exceeding 15 minutes.

The results were read into a computer and subjected to statistical analysis. An initial study included all 106 eyes. Separate analyses were then performed for the right and left eyes. Generally, the test was found appropriate to test the significance of differences between measurements obtained with any two of the instruments. A p value of less than 0.05 was considered statistically significant. A regression analysis was performed to test for irregularities such as heteroscedasticity and autocorrelation. As a final check, the Walsh test was used to ascertain whether the readings came from symmetrical populations. The 95% confidence intervals (CI) and 95% limits of agreement (mean difference (SD 1.96)) were also calculated according to Bland and Altman.+

### Results

The results for IOP measured with the Goldmann applanation tonometer, ProTon tonometer, and Tono-Pen XL tonometer in both eyes of 53 patients are shown in Table 1. The pressure range of the 106 eyes was 8-38 mm Hg with applanation (94 eyes were between ≥10 mm Hg and ≤20 mm Hg). Statistical analysis revealed a slight deviation from the normal distribution for tonometry data from the right eye with small variations between the three instruments. The left eye readings were very close to a normal distribution for all tonometers. However, there was one outlier in the left eye (IOP 38 mm Hg with applanation). As shown by regression analysis, this outlier did not have any noticeable effect on the slope of the regression line and could have safely been included.

Comparing the mean values, the IOP measured with the ProTon tonometer was not significantly different from that determined by applanation tonometry (p=0.166 in the right eye and p=0.204 in the left eye). With the Tono-Pen XL, significantly lower values of IOP were measured in comparison with the Goldmann applanation tonometry in both eyes (p<0.001).

As shown in Table 2, the 95% limits of agreement between the ProTon tonometer and the applanation tonometer were within −4 mm Hg and +5 mm Hg. With Tono-Pen XL the same limits were within −3 mm Hg and +8 mm.
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Figure 3  The IOP measured in the left eye of each patient with Goldmann applanation and with ProTon tonometry in relation to the IOP level.

Figure 4  The IOP measured in the left eye of each patient with Goldmann applanation and with Tono-Pen XL tonometry in relation to the IOP level.

The ProTon readings were distributed more symmetrically about the applanation values, indicating a closer correlation between these two instruments.

Discussion

After the introduction of the new microprocessor controlled tonometers as alternative instruments for measurement of IOP in many countries, it can be expected that these will gradually replace the classic older instruments such as the Schiötz tonometer. The greatest advantage of the new instruments is the simplicity of their use. Before they can be accepted as equivalent devices to standard tonometers, it is necessary to perform extensive clinical testing and evaluation. Discrepancies in IOP measurements could lead to incorrect clinical decisions – for example, in the detection and treatment of glaucoma.

This is the first clinical study to evaluate the accuracy of the new tonometer ProTon. The results of this study showed a good agreement between the ProTon tonometer and the Goldmann applanation tonometer. The overall accuracy of this instrument was higher than that of the Tono-Pen XL tonometer. In contrast with the Tono-Pen XL, the ProTon tonometer did not show a tendency to underestimate significantly the IOP over the tested IOP range. The limits of agreement between the values obtained with the ProTon and with the Goldmann applanation tonometer in the present study are less wide than those obtained with Tono-Pen XL. For all eyes, 89% of the ProTon readings and 63% of the Tono-Pen XL values were within 3 mm Hg of the Goldmann applanation. In the clinical studies, a mean error of less than 3 mm Hg has been suggested as tolerable in screening situations.4

With the Tono-Pen XL, the results obtained in the present study confirmed a statistically significant difference between this tonometer and the Goldmann applanation tonometer reported previously.5,11 In this study, a tendency to underestimate the IOP using this instrument in the pressure range 8–38 mm Hg with applanation was observed. A possible explanation for that could be a decline of the IOP caused by stress from repeated measurements of IOP using three instruments in each eye. With the applanation tonometer, a decline of the IOP of 2–4.6 mm Hg was reported to occur after several repeated readings (for a review see Whiteacre and Stein5). Because applanation tonometry was performed first, the possible decline of the IOP as a result of repeated measurements with this instrument will influence all subsequent measurements with the other tonometers in the same manner. Among the other sources of error with the use of the Goldmann applanation tonometer, the influence of ocular rigidity, corneal irregularity, or astigmatism is of little importance because of the selection of the patients in this study.

How measurement with the ProTon tonometer influences the IOP level is unknown. With the Tono-Pen XL, a possible decline of the IOP after tonometry with this instrument has been reported.1 This was explained as a result of
repeated indentations of the cornea during the measurement. On the other hand, despite using the Tono-Pen uniformly after applanation, a consistent overestimation of the IOP was also observed to occur in both normal and post-keratoplasty eyes.

The results of the present study show that 95% of the differences between ProTon and Goldmann application tonometers were within plus or minus $5\text{ mm Hg}$. These limits are relatively wide. Some investigators have suggested that an average error of more than plus or minus $3\text{ mm Hg}$ cannot be tolerated in the diagnosis and treatment of vision threatening diseases when more accurate techniques are available. On the other hand, in complicated cases such as uncooperative patients, where the error rate with any instrument can be expected to be higher than normal, the new tonometers can be considered to be convenient and accurate means of IOP screening. Our findings suggest that the ProTon tonometer represents a better choice than the Tono-Pen XL tonometer, although further investigations with these instruments are still needed to confirm this.

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