Applanation tonometry: a comparison of the Perkins handheld and Goldmann slit lamp-mounted methods

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Purpose: To compare intraocular pressure (IOP) measurements, taken using Perkins applanation tonometry (PAT) and Goldmann applanation tonometry (GAT).

Methods: 100 eyes of 100 patients underwent Perkins and Goldmann applanation tonometry, with a randomized order of modality, performed by a masked observer. The right eye was measured, for all subjects, and the data used in statistical analysis. The comparability of results given by the two instruments was evaluated using the Bland–Altman method.

Results: IOP measurements for 100 eyes were obtained (range: 10–44 mmHg). The mean GAT reading was 21.63 mmHg, with standard deviation (SD) 5.69 mmHg. The mean PAT reading was 21.40 mmHg, with SD 5.67 mmHg. The mean difference between readings from Goldmann versus Perkins tonometry was 0.22 mmHg (SD: 0.44 mmHg). The limits of agreement were calculated to be −0.64–+1.08 mmHg (1.96 SD either side of the bias).

Conclusion: The Perkins applanation tonometer yields IOP measurements that are closely comparable with GAT. Therefore, PAT may be used in routine clinical practice, as part of the implementation of national guidelines, or preferred practice patterns, for glaucoma and ocular hypertension.

Keywords: glaucoma, intraocular pressure, ocular tonometry, Perkins applanation tonometry, Goldmann applanation tonometry

Introduction

Intraocular pressure (IOP) measurement is an integral part of any comprehensive ophthalmic examination, and is especially important for individuals who are at risk with respect to glaucoma and ocular hypertension, as IOP is the most important risk factor for development of glaucomatous optic neuropathy, and is the best evidenced target for management.1–6 Slit lamp-mounted Goldmann applanation tonometry (GAT) is regarded as the optimum modality for the measurement of IOP;7 as such, it is recommended as part of the UK National Institute for Health and Clinical Excellence (NICE) guidelines for chronic open angle glaucoma and ocular hypertension.8 However, these guidelines do not similarly endorse Perkins handheld applanation tonometry (PAT),9 and state: “Hand-held methods of tonometry such as Perkins may be useful in a case finding/screening scenario where a person may have difficulty being examined on a slit lamp (for example with curvature of the spine). However there is no evidence to suggest that these methods are equivalent to slit lamp mounted GAT.” This is due to a lack of comparable published evidence in Europeans, using what would be considered appropriate statistical methods (for modern standards). As PAT is in widespread use in primary care optometric practice (the source of the majority of glaucoma referrals...
to the Hospital Eye Service in the UK), it is timely to assess comparability of the two methods. Therefore, the purpose of our study was to compare PAT with GAT across a wide range of IOP, and to present and analyze the results, applying suitable statistical techniques.

**Patients and methods**

One hundred subjects were enrolled into the study, by invitation, during their attendance at routine HES clinics, at which they were due to undergo IOP measurement. Adult subjects with capacity were chosen, for ease of explanation and the gaining of informed consent. The subjects were generally fit, with mild systemic disease (typical of American Society of Anaesthesiologists grade 2; they were attending the hospital for ambulatory day case cataract surgery). None was morbidly obese. The ability to tolerate corneal applanation under topical anaesthesia was included, for the sake of completeness. During the study, no potential patient failed this inclusion criterion. The International Standardization Organization (ISO) draft standard for tonometers subdivides IOP measurements into four defined ranges, to better describe and confirm fitness for purpose of tonometers, at all levels of IOP likely to be met in clinical situations. Therefore, we recruited subjects into groups, according to these IOP ranges, as shown in Table 1.

We chose to exclude patients with significant corneal astigmatism (>3 D), so as to eliminate the requirement for alteration of the orientation of the tonometer prism from the convention of the horizontal meridian. Refractive cylinder has been used as a surrogate for corneal astigmatism. Astigmatism of 3D cylinder would be associated with a potential for “error” of 1–2 mmHg, thus, it may be taken as a limit of the same order of magnitude as the level of accuracy used in recording measurements of IOP.

Other exclusion criteria were related to physical difficulty in performing applanation (infirmity preventing suitable positioning at a slit lamp, or inability to cooperate with eye opening), pathology of the cornea that could affect biomechanics and, thus, the size and appearance of the tear meniscus semicircles, like keratoconus, corneal scarring, previous corneal surgery, corneal infection, microphthamas, buphthalmos, nystagmus, and blepharospasm. Ocular pathologic changes, other than glaucoma, mild nuclear sclerosis, and rare drusen, were also excluded.

Consecutive patients were considered against the inclusion and exclusion criteria until sufficient subjects had been allocated to each of the following intraocular pressure range groups: <18 mmHg, 18–24 mmHg, and ≥24 mmHg, in accordance with the ISO guidelines for ocular tonometry.

The local research ethics committee was consulted regarding our study design, and confirmation was given that formal ethics approval was not required. As the patients were having GAT as part of their general ophthalmic assessment; the taking of additional IOP measurements by PAT was not considered as requiring formal institutional ethics review. Nevertheless, we are able to certify that all applicable institutional regulations concerning the ethical use of human volunteers were followed during this study, consistent with the tenets of the Declaration of Helsinki. Informed patient consent was gained for the additional IOP readings.

The same Goldmann tonometer (Haag-Streit, Koeniz, Switzerland) and Perkins Mk2 tonometer (Haag-Streit) were used for all IOP measurements. The calibration of each instrument was checked at the beginning of each session, according to the manufacturers’ instructions. The patients were randomized into two groups: Group A (where GAT was used first, followed by PAT) and Group B (where PAT was used first, then GAT). A standard technique for IOP measurement was followed. Topical anesthesia was applied (proxymetacaine with fluorescein) (Bausch + Lomb Pharmaceuticals, Tampa, FL, USA), and the eyes were dabbed, to remove excess fluid from the ocular surface. Applanation was performed by the same masked investigator. Prior to each applanation, the tonometer was set to an IOP value of 10 mmHg. Then, the knob was turned until the applanation visual endpoint was reached. IOP values were recorded by a second observer (a member of the nursing or technical staff, who had received specific training in applanation methods), who then reset the tonometer knob, unobserved by the applanator.

All measurements were taken with the patient in a sitting position. For GAT, the face was in a vertical plane. For PAT, the head was inclined slightly backward, through a degree of neck extension, typically in the order of 30 degrees.

Reusable tonometer prisms were used. These were cleaned and disinfected between uses, in accordance with the manufacturer’s recommendations.

IOP may vary significantly, as a consequence of pulsatile ocular blood flow during the cardiac cycle. Therefore, during applanation, the tear meniscus semicircles may appear to

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**Table 1** Numbers of eyes having IOP within each sub-range of ISO draft standard on tonometers

<table>
<thead>
<tr>
<th>IOP range</th>
<th>Number of eyes</th>
</tr>
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<tbody>
<tr>
<td>&lt;18 mmHg</td>
<td>36</td>
</tr>
<tr>
<td>18–24 mmHg</td>
<td>30</td>
</tr>
<tr>
<td>≥24 mmHg</td>
<td>34</td>
</tr>
</tbody>
</table>

Abbreviations: IOP, intraocular pressure; ISO, International Organization for Standardization.
change size to such an extent as to make it difficult to choose a value for the IOP reading. To eliminate this potential for variation not arising from any systemic bias associated with the tonometer type, we elected to include only those eyes with an IOP pulse variation of ≤2 mmHg (one graduated interval on the tonometer measuring wheel).

IOP in the right eye was measured first, for all subjects, and these values were used in statistical analysis. A set of three measurements for each applanation technique was obtained, for each study eye, so that a mean value could be used in the analysis of results. All of the measurements for each patient were taken within a single period of examination, and were completed within 15 minutes. A Bland–Altman plot was constructed, to show the differences between IOP-G and IOP-P, against their means, for each pair of values. The mean of the differences between pairs of values was calculated (ie, the bias), as also were the limits of agreement, defined as the bias ±1.96 times the standard deviation (SD) of the differences between pairs of values.

Results

From 141 consecutive potential subjects, 100 satisfied the inclusion and exclusion criteria; 31 subjects were excluded for inter-reading variation of ≥2 mmHg, and 10 subjects were excluded for frequent blinking.

Thus, 100 eyes of 100 subjects were included. IOP values (GAT taken as giving the definitive value) for each eye ranged between 10–44 mmHg, with an even distribution (minimum of 20 eyes). The numbers of eyes having IOP values within the three specified ranges are shown in Table 1.

A possible source of overstated agreement might have come from our resetting the tonometer to a value of 10 mmHg, rather than to a random level, between applanations. However, we feel that this is not likely to have been of clinical significance— it was of less importance than the simplicity of the method we chose, given the use of non-clinical assistants for resetting.

Only one eye of each subject was included for analysis. As a standard, we used the first eye to be tested, which was the right eye for each subject. The mean of the GAT readings was 21.63 mmHg, with SD of 5.69 mm Hg. The mean of the PAT readings was 21.40 mmHg, with SD of 5.67 mmHg. The mean difference between readings, by Goldmann or Perkins tonometry, was 0.22 mmHg (SD: 0.44 mmHg).

In the <18 mmHg IOP group, the mean difference was 0.27 mmHg; in the 18–24 mmHg group, the mean difference was 0.08 mmHg; and in the >24 mmHg group, the mean difference was 0.28 mmHg.

The patients were randomized into two groups: Group A (where GAT was used first, followed by PAT) and Group B (where PAT was used first, then GAT). In Group A, the mean Goldmann reading was 21.58 mmHg (SD: 5.71 mmHg), and the mean Perkins reading was 21.77 mmHg (SD: 5.95 mmHg). In Group B (where PAT was used first), the mean Goldmann reading was 21.35 mmHg (SD: 5.72 mmHg), and the mean Perkins reading was 21.49 mmHg (SD: 5.92 mmHg). The mean difference between readings in Group A was 0.21 mmHg (SD: 0.29 mmHg), and in Group B, was 0.32 mmHg (SD: 0.35 mmHg).

The data were entered into a Microsoft Excel 2009 spreadsheet (Microsoft Corp, Redmond, WA, USA), and Bland–Altman graphs were generated, according to the authors’ formula, as shown on the chart.

The Bland–Altman plot (Figure 1) shows the difference between pairs of (averaged) values for GAT and PAT, plotted against their mean. The mean difference (bias) between Goldmann and Perkins IOP readings was 0.22 mmHg (SD: 0.44 mmHg). The limits of agreement were calculated as –0.64 to +1.08 mmHg (1.96 SD either side of the bias).

A clear correlation between GAT and PAT was found (r=0.85; P<0.001).

Discussion

Precision in the measurement of IOP is a prerequisite for any glaucoma care pathway. The “landmark” glaucoma studies have emphasized the importance of IOP in clinical decision making and management.2–6 GAT is still the method most commonly used by the majority of ophthalmologists. This is because it has proved to be accurate, precise, easy to incorporate into routine slit lamp eye examination, and shows low intra- and interobserver variability.14 Corneal factors, like astigmatism, corneal curvature, and central corneal thickness, affect the accuracy of applanation tonometers. Studies have shown that there was a positive correlation between corneal curvature and tonometer readings. For each 3-D increase in corneal power in this sample, the average IOP increased 1 mmHg.16 The thickness of the cornea also affects most noninvasive methods, by varying resistance to the tonometer probe. A thick cornea gives rise to a greater probability of an IOP being overestimated, and a thin cornea can lead to IOP being underestimated.15–17

Given that there is a degree of variability in the values obtained for each eye, by each method in our study, we recommend the practice of using more than one IOP value in a patient’s clinical record (eg, two or three).
The Perkins tonometer was devised as a portable handheld applanation tonometer, for use in children, patients unable to cooperate for slit lamp examination, and anesthetized and bedridden patients. PAT is currently in widespread use in the primary care setting. One reason may be historical, from a time when optometrists did not routinely use slit lamp microscopes, relying on handheld direct ophthalmoscopy for ocular examinations.

The accuracy and compatibility of PAT in animals is supported by several studies. Andrade et al reported a strong correlation between the IOP values obtained by direct ocular manometry and PAT in horses and cattle. However, limited data in humans is available. Having the same physical principles and similar basic instrument construction, PAT should be comparable with GAT. Previous comparison reports have either used small numbers of subjects (thus, providing relatively weak evidence) or used only correlation coefficients in their statistical analyses. Such an approach does not infer or confirm comparability, nor provide sufficient evidence to allow for interchangeability between techniques. Therefore, there is a need for a more robust evidence base for using PAT in routine human ophthalmic practice. The Bland–Altman Plot has become the universally-adopted statistical method technique for comparison of measurement methods. Limits of agreement are defined as two standard deviations either side of the mean difference between values given by each method.

Ours is a simple study to answer a simple question, addressing a small but significant gap in the tonometry literature, namely: “Is Perkins an acceptable alternative to Goldmann?” Our results show a suitably compact grouping of difference values, not just within the defined Bland–Altman limits of agreement, but also well within the tolerance limits given in the ISO standard for a tonometer. Our report shows that the Perkins applanation tonometer measures IOP to a much closer level of comparability than other tonometer types. This is to be expected, given the physical properties of the applanating prisms, visual endpoints, and internal workings of the instruments. Our low value of 1.5 mmHg, for the spread of 95% of difference values between the techniques, could arise (in part) both as a result of our use of the mean of three IOP readings for each tonometer, for each eye (suggesting limited usefulness for common clinical practice), and due to our exclusion of any eye with a wide amplitude of pulse-induced intraocular pulse variation (>2 mmHg) (Table 1). We consider the former cause unlikely, as similar results were achieved when using just one IOP value per eye, and the latter issue of limited relevance, considering the demonstrable lack of any systematic bias in the tonometric modalities.

Perkins handheld tonometry would appear to be comparable with the “gold standard” of slit lamp-mounted Goldmann applanation tonometry, for eyes satisfying the inclusion and exclusion criteria. Therefore, we consider it to be acceptable for routine clinical practice, not merely to be considered as a “second best”, for when slit lamp-mounted GAT is not possible. We would suggest that PAT may be permissible for IOP measurement, as part of care pathways...
for open angle glaucoma and ocular hypertension. In the UK, this should include modification of the 2009 NICE guidelines. But corroborative evidence from others would be required to confirm this.

**Summary**

**What was known before?**
- Goldmann applanation tonometry (slit lamp-mounted) is the accepted “gold standard” method for IOP measurement.
- Perkins applanation tonometry (handheld) is in widespread use, particularly in the primary care setting, but evidence of comparability with Goldmann applanation tonometry is not substantial.

**What this study adds**
- Measurements of IOP by Perkins applanation tonometry are sufficiently comparable with Goldmann tonometry to be acceptable for routine clinical practice.

**Acknowledgment**

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**Disclosure**

The authors declare they have no conflict of interest in this work.

**References**


