Clinical comparison of the Keeler Pulsair 3000 with Goldmann applanation tonometry

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Abstract

Aim—To confirm the accuracy of the Pulsair 3000 before introducing the instrument into clinical practice.

Method—A masked study by two experienced tonometrists comparing the mean of the Goldmann intraocular pressure (IOP) readings against the Pulsair 3000 reading (average of four puffs). Results of 150 eyes were compared with an IOP range of 10 mm Hg–44 mm Hg.

Results—Correlation between the two Goldmann tonometry results was 0.9830 with a standard deviation of 1.1085 mm Hg. Correlation between the mean of the two Goldmann readings against the Pulsair 3000 reading was 0.982 with a standard deviation of 1.1179 mm Hg. Bland–Altman analysis confirms a satisfactory outcome.

Conclusion—The Pulsair 3000 provides an accurate and objective method of measuring IOP with many advantages over traditional Goldmann tonometry.

Methods

Two experienced Goldmann tonometrists performed a masked study of 150 eyes. The mean pressure of the 150 eyes tested by tonometrist no 1 was 21.06 mm Hg and by tonometrist no 2 was 21.17 mm Hg. In order to cover a comprehensive range of IOP we included no less than 40 eyes in each of the following pressure groups: 7–18 mm Hg, >18–24 mm Hg, and >24 mm Hg. The total range was 10–44 mm Hg. These ranges were chosen according to the International Standards Organisation draft standard on tonometers.

Subjects were selected from glaucoma clinics according to the following exclusion criteria:

1. High corneal astigmatism—that is, those eyes displaying an oval contact image with the Goldmann tonometer
2. Corneal scarring or corneal surgery including corneal laser surgery
3. Microphthalmos
4. Buphthalmos
5. Blepharospasm
6. Manifest nystagmus
7. Keratoconus
8. Known corneal or conjunctival infection.

Both Goldmann tonometers were calibrated at the beginning of each session and the results of tonometrist no 1 were unknown to tonometrist no 2 until the end of each session. The Pulsair 3000 test was performed by tonometrist no 1 after the first Goldmann reading. The Pulsair provides a digital readout of the IOP, therefore prior knowledge of the Goldmann result could not influence the result. The Goldmann reading only was taken by each tonometrist no 1 after the first Goldmann reading. One second Goldmann reading was taken by tonometrist no 2 with as little delay as possible between the reading (average 20 minutes). One Goldmann reading only was taken by each tonometrist to avoid the tendency for IOP to decrease on multiple testing.

Results

Correlation between the first and second Goldmann readings was 0.9830 with a standard deviation of 1.1085 mm Hg.

The mean of the two Goldmann readings was used in comparison with the Pulsair 3000 system that accurately controls the amount of air delivered to the eye enabling a precise, repeatable puff producing precise and repeatable readings with less discomfort to the patient. Software improvements have made the instrument easier to use with prompts and error messages ensuring accurate results.

Clinical proof of the accuracy of the Pulsair 3000 was required before introducing the instrument into our clinics.
result (average of four puffs). The correlation was 0.982 with a standard deviation of 1.1179 mm Hg.

Clinically, it is important to prove that the Pulsair could replace applanation tonometry or the two methods could be interchangeable. Therefore, a further and more appropriate statistical analysis would be to use the Bland-Altman method. Rather than simply plotting the results of one method against those of another, a plot of the difference between the methods against their mean would be more informative; 95% of the differences should be less than two standard deviations from the mean difference in order for the instrument to be acceptable.

The numbers on the figure show the mean difference (−0.48) and the mean difference plus or minus 2 standard deviations of the differences (−2.72 and 1.75).

This shows that 95% of Pulsair 3000 results fall between 1.75 and −2.72 mm Hg with a mean value of 0.48 mm Hg compared with the Goldmann average.

If the same method is used to analyse the two Goldmann readings it shows that 95% of the results fall between 2.10 and −2.33 mm Hg with a mean value of −0.11 mm Hg.

**Discussion**

The results of this study show that the Pulsair 3000 has been accurately calibrated to within an acceptable margin of error. There is little difference between the correlation of the two Goldmann readings compared with the correlation of Pulsair against the Goldmann mean.

The results using the Bland-Altman method also show good agreement and that the Pulsair shows acceptable readings on comparison with Goldmann tonometry.

The changes to the Pulsair have produced a user friendlier instrument for the operator and less discomfort to the patient. Advantages include removing the need for local anaesthetic and fluorescein drops and no risk of corneal abrasion or cross infection.

With practice, the Pulsair can provide an accurate measurement of IOP within approximately 20 seconds per eye and eradicates most operator influences or error.

The Pulsair 3000 is hand held and portable and is very useful clinically when testing wheelchair bound or immobile patients. It has also been used on many occasions in our clinic to test the IOP of babies and children avoiding the need for general anaesthesia.

In conclusion, the Pulsair 3000 is as accurate as Goldmann tonometry in the population studied and has distinct advantages.

The authors have no financial interest in the Pulsair 3000.