No differences in central corneal thickness between open-angle and pseudoexfoliation glaucoma patients

Marcelo Ayala1
Johanna Karlsson2

1Eye Department, Skaraborg Hospital, Skövde, Sahlgrenska Academy, Gothenburg University and Karolinska Institute, 2Faculty of Medicine, Örebro University, Örebro, Sweden

Introduction: Measurements of intraocular pressure (IOP) with Goldmann applanation tonometry are affected by central corneal thickness (CCT), as thinner corneas underestimate and thicker corneas overestimate the true IOP value. The literature is controversial regarding CCT values in patients with primary open-angle glaucoma (POAG) and exfoliation glaucoma (XFG). The aim of this study was to evaluate CCT in patients with XFG and POAG.

Methods: CCT was evaluated with optical coherence tomography (OCT). All participants who were previously diagnosed with either POAG or XFG underwent ophthalmological examinations. Contact lens users and patients with corneal diseases were excluded.

Results: Totally, 145 patients were enrolled in this study. The mean CCT was 535±30.4 µm in patients with POAG and was 536±33.7 µm in patients with XFG. The result was not statistically significant (P=0.98). The mean age for all participants was 73.8±7.7 years. The study included totally 61 women and 84 men. The two groups were similar in their demographic data, and mean deviation was the only parameter that differed statistically when comparing POAG with XFG (P=0.02).

Conclusion: Our data indicate that patients with XFG do not have thinner corneas than those with POAG, and therefore, CCT cannot explain why they progress differently.

Keywords: primary open-angle glaucoma, exfoliative glaucoma, central corneal thickness, optical coherence tomography

Correspondence: Marcelo Ayala
Eye Department, Skaraborg Hospital, 54185 Skövde, Sweden
Tel +46 500 43 1000
Fax +46 500 43 2179
Email marcelo.ayala@vgregion.se

Introduction: Glaucoma is the second leading cause of blindness worldwide. It consists of a heterogeneous group of diseases, all characterized by optic-nerve damage causing irreversible visual field loss and vision impairment, which slowly can progress to blindness if not treated. Open-angle glaucoma (OAG) is the most common form of glaucoma in the Western world.

The etiology of OAG is still unknown, but genetic and environmental risk factors are thought to be involved. The intraocular pressure (IOP) is considered to be the most important risk factor for developing glaucoma. IOP is an important factor for monitoring glaucoma treatment and determining glaucoma severity, and it is also the only treatable risk factor. Other risk factors are, for example, older age, thin central cornea, ethnicity, family history and pseudoexfoliation.

There are different types of OAG, of which the most common type is primary open-angle glaucoma (POAG). Exfoliative glaucoma (XFG) is a secondary form of OAG with faster progression and worse prognosis than POAG. XFG is characterized by accumulation of abnormal fibers in the anterior segment of the eye, causing...
disrupted drainage of aqueous humor and elevation of IOP. The progression of XFG is correlated with the level of IOP, and patients with XFG more often require not only medical treatment but also laser and surgery to lower the IOP. By lowering IOP in patients with OAG, further progression of this condition can be prevented or delayed.

The “gold standard” method to measure IOP is Goldmann applanation tonometry (GAT). It is well known that measurements of IOP with GAT are affected by central corneal thickness (CCT), as thinner corneas underestimate and thicker corneas overestimate the true IOP value. Moreover, thinner CCT has been shown to be a predictor for the development of POAG in patients with ocular hypertension. Patients with thinner corneas are more likely to have visual field progression and a worse visual field defect.

The CCT differs in different populations and in different types of glaucoma. Genetic factors have been shown to be of major importance in CCT. It has been shown that Afro-Americans have a thinner CCT than other populations. However, the literature is controversial regarding the differences in corneal thickness in patients with POAG and XFG. Some studies have shown eyes with XFG to have thinner CCT compared to POAG and/or normal eyes, while other studies have found thicker CCT in XFG compared to POAG and/or normal eyes.

The aim of this study was to evaluate central corneal thickness in patients with XFG and POAG.

Methods

Study protocol

A comprehensive medical and ocular history was obtained from patients. Ophthalmological examination was performed before including these patients in the study. Visual acuity, IOP measurements, optic-nerve status, gonioscopy, Humphrey visual fields (HFA, 24-2) and presence or absence of exfoliation were recorded. Visual acuity was recorded using Snellen’s chart. IOP was measured using a Goldmann applanation tonometer. Three measurements were done and the average value was calculated. Then, the pupils were dilated and exfoliation was checked and recorded as present or absent. Afterward, the optic-nerve status was evaluated using a 90-D lens, and hence, stereo photographs were taken. Previous eye surgery was also registered.

Glaucoma was defined, following the European guidelines for glaucoma, as the presence of at least two repeatable Humphrey visual fields showing glaucoma damage in patients, using the software 24-2 and the optic nerve showing typical glaucoma damage. Demographic data, such as gender, age, ocular history, visual field and number of antiglaucomatous medical methods, were also recorded. Inclusion criteria were patients who were previously diagnosed with POAG or XFG according to their medical records. Diagnosis was based on optic disc appearance and visual field damage. Subjects were excluded if they were contact lens users or had any corneal disease.

POAG was defined as the condition, as above, without the presence of exfoliation in dilated pupil. XFG was defined as open angle, concomitant with the presence of exfoliation material, a grayish-white material, observed at the anterior lens capsule and/or at the pupillary border with dilated pupil. Patients who were operated for cataract were diagnosed with exfoliation before cataract operation.

CCT was measured by optical coherence tomography (OCT, 3D OCT-2000 and 3D OCT-1000; Topcon Corporation, Tokyo, Japan) using the anterior segment imaging protocol. Only images of good quality were recorded (>60 signal strength). CCT was measured by two examiners, a senior consultant ophthalmologist (M.A.) and a medical student (J.K.). The measurements were done during the daytime, from 8 am to 5 pm. Both eyes were measured in all patients once. If patients had bilateral glaucoma, one eye was later randomly selected to be included in the statistical analysis.

Statistical methods

Statistical analysis was performed with SPSS (SPSS, Chicago, IL, USA). First, the data were tested for normality using the Kolmogrov-Smirnov test. Then, the data were tested for equality of variance using Levene’s test. Student’s $t$-test was used to compare CCT values and demographic data between patients with POAG and XFG. For comparison of gender between the two groups, a chi-squared test was used. Averages ± standard deviations (SDs) were used to report data. A $P$-value of 0.05 or less was considered to be statistically significant. Sample size was estimated to be 63 individuals in each group with a significance level of 0.05 and power of 90%.

Ethical considerations

Ethical approval was received from the Institutional Review Board (Ethical approval number: 717-13), Gothenburg University, Gothenburg, Sweden. The study followed the tenets of the Declaration of Helsinki. A written informed consent was obtained from all patients who participated in this study, and their medical records were controlled, before they were included into this study.
Results
Totally, 145 patients were included in this study between December 2013 and December 2016. Of them, 66 patients were diagnosed with POAG and 79 with XFG. All patients were born in Scandinavia. The mean age for all participants was 73.8±7.7 years, and the age range was 52–93 years. There were totally 61 women and 84 men included in this report. The demographic properties of two glaucoma groups are presented in Table 1.

The two groups were similar in their demographic data, and mean deviation (MD) was the only parameter that differed statistically when comparing POAG with XFG (P=0.02).

Regarding CCT, the data are found to be normally distributed (Kolmogrov–Smirnov; P=0.20), and the variances between the groups are shown to be equal (Levene’s test; P=0.36). The mean CCT was 535±30.4 µm in patients with POAG and was 536±33.7 µm in patients with XFG. The result was not statistically significant (t-test; P=0.98). The mean was adjusted for confounding factors. The adjusted mean was 469.40 µm in POAG patients and was 480.04 µm in XFG patients.

Data from four patients were excluded due to difficulties in recording of the measurements (n=3) and failure of the equipment (n=1).

Discussion
Central corneal thickness is a well-known factor affecting IOP readings with GAT. It was earlier shown by Goldmann and Schmid that IOP readings with GAT could give falsely elevated or decreased IOP readings depending on the corneal thickness. Ehlers et al reported that accurate measurements only were given at a central corneal thickness of 520 µm. They found that for each 10 µm it varies from 520 µm, a correction of 0.7 mm Hg is needed. Other reports have later revealed average correction to be lower, namely 0.18–0.23 mmHg, 0.19 mmHg and 0.5 mmHg per 10 µm.

CCT values among normal individuals without glaucoma vary and are normally found to be 540±30 µm. The literature is controversial regarding CCT in patients with POAG and XFG and how they correlate with each other and with normal individuals without glaucoma. In our report, no statistical significant differences in CCT could be detected between these glaucoma types. These results are in concordance with other studies. Reports by Tolesa and Gessesse, Ventura et al, Shah et al and Ozkok et al did not find any significant difference in corneal thickness between XFG and POAG. Tolesa and Gessesse found, in their multiethnic report, thicker CCT in patients with POAG (520±38.95 µm) compared to XFG (507±35), but these results were not significant. In the report by Ventura et al, the mean CCT was 515±35 µm in eyes with POAG and was 507±25 µm in XFG. Shah et al reported mean CCT to be 530.7 µm in XFG and 550.1 µm in POAG, and Ozkok et al reported mean CCT to be 546±34.9 µm vs 550±25 µm. In this study, we found the mean CCT to be 536±33.7 µm in XFG eyes and 535±30.4 µm in POAG eyes. Previous reports cited in this article have reported mean CCT in patients with XFG to be between 493–546 µm and 507–556 µm. Our results accord with previous studies.

Reports by Kitsos et al, Bechmann et al, Gorezis et al and Kniesledt found CCT to be significantly lower in XFG compared to POAG. They had included only 32, 24, 50 and 12 XFG patients, respectively. No evidence was found in the literature about published studies comparing XFG to POAG with the same number of XFG patients than the ones included in this report. Furthermore, according to our estimation of sample size, 63 individuals in each group with significance of 0.05 and power of 90% were needed. Often, estimation of sample size has not been included in previous studies, which may contribute to explain why some studies found a difference and some not in CCT between POAG and XFG. Another possible explanation about the small number of XFG included in previous studies may be that XFG is quite common in Scandinavian countries. It might be difficult to recruit enough number of participants in studies performed in other countries, and therefore, leading to too small sample size of exfoliation patients.

Disagreement between studies could be explained by different techniques used to measure CCT. Currently, CCT can be measured by ultrasound pachymetry (USP), optical low-coherence reflectometry (OLCR), anterior segment

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<th>Table 1 Demographic data of the study groups</th>
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<td>Characteristics</td>
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<tr>
<td>Age (SD), years</td>
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<td>Sex (male/female)</td>
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<td>IOP (SD), mmHg</td>
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<td>Trabeculectomy</td>
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Note: *Two-tailed t-test; **Chi-square test.

Abbreviations: POAG, primary open-angle glaucoma; XFG, exfoliation glaucoma; SD, standard deviation; IOP, intraocular pressure; VFI, visual field index; MD, mean deviation; PSD, pattern standard deviation.
optical coherence tomography and Scheimpflug camera. Comparison in CCT between XFG and POAG has mostly been done using USP.26,30,32,33 Bechmann et al38 used OCT, Ventura et al31 used OLCR and Gorezis et al32 used a peculiar microscope. USP is currently the gold standard for measuring corneal thickness, but it has some limitations. The examiner must place the probe at the exact same position at the center of the cornea, otherwise a different CCT value will be obtained.44 It also requires topical anesthesia before measurement can be accomplished, which can affect cornea and the results.45 OCT has the advantages of being a noninvasive, noncontact technique, and therefore, no topical anesthesia is needed to perform the measurement. It is also less affected by the examiner’s knowledge and experience, and has high degree of repeatability and reproducibility.46 Additionally, incorrect placement of a probe cannot bias OCT results.47

Previous reports have shown that OCT gives lower CCT readings compared to ultrasound.47-49 The opposite, overestimation with OCT compared to USP has also been reported.50,51 Wells et al48 compared optical pachymetry, OCT and ultrasound, and found that these devices are not interchangeable in clinical practice. Garcia-Medina et al49 compared Fourier-Domain OCT with USP in 80 patients with POAG and found a significant difference between the mean CCT. However, they indicated that this difference should not be clinically significant in IOP estimation. There is also a report by Adibelli et al,31 showing a significant difference between OCT and USP in 26 POAG patients, where mean CCT was higher with OCT compared to USP. Comparison by Ayala and Strandås50 between OCT and USP did not show any difference between these two devices. However, this might be a reason for different outcomes when measuring CCT worldwide. As reported by Wells et al,48 CCT can vary up to 30 µm when using different instruments.

There is a lack of information in the literature about how interchangeable the devices are concerning XFG patients. Maybe, the exfoliation material at the endothelium disturbs the measurement with OCT, creating a false thicker cornea and, therefore, we did not find any difference between these two glaucoma types. Further studies are needed to investigate the issue. However, in our study patients suffering from POAG and XFG were measured with the same technique (OCT) minimizing the risk of bias. Regarding demographic data, our two groups were very similar. Only mean deviation differed statistically. This is in concordance with The Early Manifest Glaucoma Trial, showing a faster progression rates in MD in patients with XFG than in POAG.52

The study has several limitations. The examiners were not blinded to condition the patient had, and therefore there are some risks for bias. The fact that two different examiners performed the measurements can induce certain bias to the study. However, a recent study by the same author has shown that there are no differences between the examiners measuring with OCT.50 There is also risk of selection bias as the recruitment of the patients was performed at an eye clinic. Variation in measurements due to lack of patients’ cooperation may also be a limitation. During measurements with OCT, the patient should be able to focus her or his gaze on something during the measurement. This was difficult only for a few patients because of the inability to understand instructions owing to very poor hearing or dementia. However, the same difficulties would be present in the both glaucoma types included. Another possible bias in the study was that patients were recruited directly from the clinic, and patients were in different stages of their illnesses as same as using different amount of eye drops.

Conclusion
Our data indicate that patients with XFG do not have thinner corneas than those with POAG, and therefore CCT can not explain why they progress differently.

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Disclosure
The authors report no conflicts of interest in this work.

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