Comparison of Corneal Biomechanical Properties of Physiological Macrodiscs and Glaucomatous Macrodiscs

Funda Ebru Onmez, Ayse Cigdem Altan, Banu Satana, Berna Basarir, Isil Pasaoglu, Muhittin Taskapili
University of Health Sciences Beyoglu Eye Training and Research Hospital, Istanbul, Turkey

Abstract

Objectives: This study was designed to compare optic nerve head (ONH) parameters and corneal biomechanical properties of healthy and glaucomatous macrodiscs.

Methods: This cross-sectional study included total of 234 eyes, of which 92 served as normal controls, 92 had healthy macrodiscs, and 50 had glaucomatous macrodiscs. Measurement of ONH parameters in all patients was performed using optic coherence tomography (OCT). Corneal hysteresis (CH) was measured in each patient using Ocular Response Analyzer (ORA; Reichert, Inc., Depew, NY, USA). Central corneal thickness (CCT) was determined by ultrasonic pachymetry. All OCT and ORA parameters of healthy and glaucomatous macrodiscs were compared.

Results: Optic disc area of healthy macrodiscs and glaucomatous macrodiscs was similar, and larger than observed in control group (p=0.70, p=0.0001, respectively). Rim area was significantly thinner in glaucoma group than healthy macrodisc group or control group. (p=0.0001, p=0.022, respectively) Mean cup area of glaucomatous discs was larger than healthy macrodiscs (p=0.001). Cup/disc (C/D) area ratio, and horizontal and vertical C/D ratios were higher in glaucoma group than healthy macrodisc group (p=0.002, p=0.002, p=0.018, respectively) ORA analyses revealed that CH of glaucoma patients was lower than that of healthy macrodisc or control group (p=0.048, p=0.035, respectively), whereas, mean CH of macrodisc group was similar to that of control group (p=0.988). Mean CCT of macrodisc group was higher than that observed in control or glaucoma patients (p=0.015, p=0.045).

Conclusion: It was concluded that OCT analyses can help differentiate healthy macrodiscs from glaucomatous discs. CH and corneal thickness measurements of healthy macrodiscs are greater than those of glaucomatous optic discs.

Keywords: Glaucoma, large optic disc, macrodisc, ocular response analyzer, optic nerve head analysis.

Introduction

Glaucoma is chronic optic neuropathy that leads to progressive injury to the optic nerve and retinal nerve fiber layer (RNFL). Elevated intraocular pressure (IOP) is the best-known risk factor for glaucoma. Recent studies revealed that corneal biomechanical factors affect IOP measurements and ocular effects of IOP (1). The Ocular Response Analyzer (ORA) (Reichert Inc., Depew, NY, USA) evaluates biomechanical properties of the cornea in situ. Corneal hysteresis (CH) measurement obtained using ORA demonstrates cornea's ability to dampen and buffer fluctuations in IOP (2).

It has been suggested that eyes with greater CH tend to have more capacity to buffer increases in IOP, providing a protective property (1, 3). Low CH might increase risk for developing glaucomatous optic neuropathy, possibly due to reduced capacity of the eyewall to buffer IOP spikes (3, 4). In addition, it has been proposed that lower CH might influence both glaucoma progression and severity (5–7).

Clinical estimation of vertical cup/disc ratio (C/D) remains the most frequently performed assessment of the optic disc in diagnosis and follow-up of glaucoma. C/D ratio is
physiologically related to optic disc size (8–10). As large optic nerve heads (ONHs) have large cups, they can be misdiagnosed and treated as glaucoma. Therefore, differentiation of ONH parameters between normal-sized discs and macrodiscs is extremely important for differential diagnosis of healthy macrodisc from glaucomatous optic neuropathy. Our previous study revealed that macrodiscs have larger cup area, horizontal and vertical C/D ratio, and C/D ratio compared with normal-sized discs using optical coherence tomography (OCT). Moreover, RNFL thicknesses, rim area, and visual field (VF) indices (mean deviation [MD], pattern standard deviation [PSD]) of macrodiscs were similar to those of normal eyes (11).

The aim of this study was to assess whether biomechanical properties of corneal tissue of the eye in healthy patients with macrodiscs are different from those of glaucoma patients with macrodiscs.

**Methods**

Cross-sectional, comparative study was conducted. Before being included in the study, each individual was informed of its purpose and provided written consent to participate. All participants ranged in age from 40 to 70 years. All tested eyes had best corrected visual acuity above 20/25, spherical refractive error within ±2.0 diopters, cylindrical error within ±2.0 diopters, and intraocular pressure ≤21 mmHg.

In this study, we defined macrodiscs as larger than 2.80 mm². Normal disc size was defined as disc area between 1.4 and 2.80 mm² (11). According to this definition, 3 groups were formed: normal control patients, patients with healthy macrodiscs, and patients with glaucomatous macrodiscs.

Normal control patients and those with healthy macrodiscs were patients with no remarkable medical or ocular history who came to ophthalmology clinic for regular eye examination. They underwent a complete ophthalmic evaluation, including past medical history, IOP measurement using Goldmann applanation tonometry, gonioscopy, VF testing, central corneal thickness (CCT) measurement with corneal ultrasound pachymeter, undilated and dilated biomicroscopy, and dilated fundus examination. Ophthalmic evaluation, ORA measurement, VF testing, and OCT imaging were all performed within 1 month.

Glaucomatous participants with macrodiscs had already been diagnosed and presented for antiglaucoma treatment; they underwent some ophthalmological examinations as healthy individuals to confirm diagnosis. Patients with angle-closure, normal-tension, pigmentary, inflammatory, or aphakic glaucoma, and patients who had undergone glaucoma surgery, were excluded from the study.

ORA was used to measure CH and obtain 3 other output variables: corneal resistance factor (CRF), corneal-compensated IOP (IOPcc), and Goldmann-correlated IOP (IOPg). For this study, 3 measurements were performed for each eye, and average was accepted as final value used for statistical analysis. Readings from instrument required consistent and clean raw signal morphology (well-defined raw signal peaks with repeatable characteristics for multiple measurements).

All optic disc and RNFL measurements were performed using fast optic disc scanning protocol and automated ONH analysis using Stratus OCT device and version 3.0 software (Carl Zeiss Meditec, Inc., Dublin, CA, USA). Maximum of 2 ONH scans of each patient were obtained and better of 2 scans was chosen for interpretation. Fast RNFL algorithm was used to measure RNFL thickness with Stratus OCT. Three scan images were obtained from each participant, with each image consisting of 256 A-scans along a 3.4-mm-diameter ring around the optic disc. These values were averaged to yield 12 clock-hour thicknesses, 4 quadrant thicknesses, and global average RNFL thickness measurement.

Default axial length and refraction for optic disc measurement in every OCT scan were set to 24.46 mm and 0 D, respectively. Therefore, exact disc size in an eye with axial length other than 24.46 mm and/or refraction <0 D is different from printout values and manual correction of optic disc measurements is necessary. In order to correct axial length-related ocular magnification, Littmann formula \( t = p \cdot q \cdot s \), as modified by Bennett and later adopted by Leung et al. and Kang et al. was applied (12–15). In this formula, \( t \) is actual fundus dimension, \( s \) is measurement on OCT, \( p \) is magnification factor related to imaging system, and \( q \) is magnification factor related to the eye. Factor \( p \) is instrument-dependent and remains a constant in telecentric imaging system; for Stratus OCT, this figure is 3.382. Ocular magnification factor \( q \) of the eye can be determined with formula \( q = 0.01306 \cdot \text{axial length} - 1.82 \) (14). Therefore, given a value, \( s \), obtained with OCT, the real size of RNFL peripapillary scan circle can be determined by means of the formula \( t = 3.382 \cdot 0.01306 \cdot \text{axial length} - 1.82 \). Because Littmann formula refers to linear magnification, for this study, equation was modified to \( t^2 = p^2 \cdot q^2 \cdot s^2 \) for area magnification, according to suggestion of Leung et al. (16).

Patients with significant ocular disorder, history of intraocular surgery, or systemic disease with possible ocular involvement, such as diabetes mellitus, were excluded from the study. OCT scans with signal strength of less than 6 were also excluded. All participants had reliable (fixation loss, false-positive, and false-negative error less than 10%), and normal [absence of all 3 of Anderson and Patella’s criteria (15)] Humphrey 30–2 Swedish Interactive Threshold Algorithm-standard testing. One eye of each participant was enrolled. If both eyes met inclusion criteria, randomization was performed.
Statistical Analysis
All statistical analyses were performed with SPSS software (SPSS for Windows, Version 15.0; SPSS Inc., Chicago, IL, USA). Categorical variables were compared using χ² analysis, and continuous variables with normal distributions were compared with one-way analysis of variance test. Kolmogorov–Smirnov nonparametric test was used to evaluate normal distribution of numerical data. Results of measurements were expressed as mean ±SD. Pearson's correlation coefficient analyses were performed to evaluate relationship between CH and other parameters. P<0.05 was considered statistically significant.

Results
Table 1 is a summary of baseline characteristics of normal eyes, healthy macrodiscs, and glaucomatous macrodiscs. Study included total of 234 eyes of 234 patients, of whom 92 participants served as normal controls, 92 had glaucoma, and 50 had healthy macrodiscs.

Optic disc area of healthy macrodiscs and glaucomatous eyes was similar, and larger than that of control group (p=0.70, p=0.0001, respectively). Rim area was significantly thinner in glaucoma than healthy macrodisc group and control group (p=0.0001, p=0.022, respectively). Cup area was determined to be significantly smaller in control group than other groups, and mean cup area of glaucomatous discs was larger than seen in healthy macrodiscs (p=0.0001, p=0.001, respectively). C/D area ratio, and horizontal and vertical C/D ratios were higher in glaucoma group than in healthy macrodisc group (p=0.002, p=0.002, p=0.018, respectively) (Table 2).

ORA analyses revealed that mean IOPg values were similar between study groups (p=0.19), but mean IOPcc of glaucoma group was significantly higher than seen in healthy macrodisc or control group (p=0.048, p=0.035, respectively), whereas, mean CH of macrodisc group was similar to that of control group (p=0.988). Mean CCT of healthy macrodisc group was found to be higher than that of control group or glaucoma group (p=0.015, p=0.045) (Table 3).

There was no statistically significant correlation between age and CH in control or healthy macrodisc group (p=0.229); however, significant negative correlation was determined in glaucoma group (p=0.004).

MD was similar between the 3 groups. There was significant positive correlation between MD and CH in glaucoma group. PSD was significantly higher in glaucoma group (p=0.0001). Correlation analysis revealed that PSD was not related to CH in control patients or glaucoma group; however, there was positive correlation between CH and PSD in healthy macrodisc group (p=0.026).

Discussion
Present study was designed with primary objective of evaluating and comparing CH and other ORA parameters of healthy macrodiscs, glaucomatous macrodiscs, and normal-sized discs. To the best of our knowledge, this is first study to investigate corneal biomechanical properties of macrodiscs.

We found that CH was significantly lower in glaucomatous macrodiscs and that IOPcc was significantly higher in this group. We also determined that healthy macrodiscs had thicker CCT than glaucomatous macrodiscs or control group.

As large ONH have large cups, they can be misdiagnosed and treated as glaucomatous. Therefore, differentiation of ONH parameters of normal and macrodiscs is extremely important for differential diagnosis of healthy macrodisc from glaucomatous optic neuropathy. Our study revealed that glaucomatous eyes had lower rim area value than healthy macrodiscs, as well as higher C/D ratio and horizontal and
vertical C/D ratio than seen in healthy macrodiscs.

CH is biomechanical property that can provide insight into normal development and pathological changes of the cornea. As Luce et al. reported, CH is not correlated with corneal curvature, astigmatism, VA, or axial length (17).

CH represents a dynamic resistance component of the cornea (18). Distensible ocular structures may be associated with progression of glaucomatous lesions and biomechanical properties of the cornea may reveal weakness in the lamina cribrosa. As previously reported, lower CH has been associated with progressive worsening of VF in patients with glaucoma. Bochmann et al. compared CH measurements in glaucoma patients and patients with acquired pit of the optic nerve head (APON). The latter condition mainly occurs in normotensive glaucoma and is associated with higher risk of progressive optic disc damage (19). They found that CH

Table 2. Comparison of optical coherence tomography parameters (magnification corrected) between healthy macrodiscs, glaucomatous macrodiscs, and normal-sized discs

<table>
<thead>
<tr>
<th></th>
<th>Healthy macrodiscs (n=92)</th>
<th>Glaucomatous macrodiscs (n=50)</th>
<th>Control group (n=92)</th>
<th>F</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disc area (mm²)</td>
<td>3.12±0.23</td>
<td>3.18±0.28</td>
<td>2.40±0.28</td>
<td>209.14</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Rim area (mm²)</td>
<td>1.77±0.54</td>
<td>1.38±0.54</td>
<td>1.59±0.51</td>
<td>7.55</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Cup area (mm²)</td>
<td>1.34±0.61</td>
<td>1.75±0.62</td>
<td>0.80±0.56</td>
<td>48.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Horizontal C/D ratio</td>
<td>0.67±0.16</td>
<td>0.78±0.14</td>
<td>0.57±0.19</td>
<td>28.25</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Vertical C/D ratio</td>
<td>0.59±0.13</td>
<td>0.67±0.13</td>
<td>0.51±0.17</td>
<td>23.07</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>C/D area ratio</td>
<td>0.42±0.17</td>
<td>0.54±0.17</td>
<td>0.32±0.21</td>
<td>24.55</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Superior RNFL thickness (µm)</td>
<td>126.7±17.5</td>
<td>116.7±17.7</td>
<td>122.4±17.0</td>
<td>3.05</td>
<td>0.004</td>
</tr>
<tr>
<td>Inferior RNFL thickness (µm)</td>
<td>131.2±17.9</td>
<td>121.9±19.2</td>
<td>126.6±19.8</td>
<td>1.48</td>
<td>0.01</td>
</tr>
<tr>
<td>Average RNFL thickness (µm)</td>
<td>102.4±10.7</td>
<td>97.3±12.0</td>
<td>100.3±9.65</td>
<td>2.32</td>
<td>0.02</td>
</tr>
</tbody>
</table>

SD: standard deviation; C/D: cup/disc; RNFL: retinal nerve fiber layer.

Tukey’s multiple comparison test

<table>
<thead>
<tr>
<th></th>
<th>Disc area</th>
<th>Rim area</th>
<th>Cup area</th>
<th>Horizontal</th>
<th>Vertical</th>
<th>Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control/macrodiscs</td>
<td>0.0001</td>
<td>0.317</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>Control/glaucomatous</td>
<td>0.0001</td>
<td>0.022</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>Macrodiscs/glaucoma</td>
<td>0.705</td>
<td>0.0001</td>
<td>0.001</td>
<td>0.002</td>
<td>0.018</td>
<td>0.002</td>
</tr>
</tbody>
</table>

Table 3. Comparison of Ocular Response Analyzer measurements of healthy macrodiscs, glaucomatous macrodiscs, and control group

<table>
<thead>
<tr>
<th></th>
<th>Healthy macrodiscs</th>
<th>Glaucomatous macrodiscs</th>
<th>Control</th>
<th>F</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>IOPg</td>
<td>15.63±3.92</td>
<td>16.38±4.23</td>
<td>15.04±4.51</td>
<td>1.66</td>
<td>0.193</td>
</tr>
<tr>
<td>IOPcc</td>
<td>15.73±3.49</td>
<td>17.56±3.89</td>
<td>15.27±4.34</td>
<td>5.71</td>
<td>0.004</td>
</tr>
<tr>
<td>CH</td>
<td>10.6±1.61</td>
<td>9.76±1.98</td>
<td>10.56±2.12</td>
<td>3.62</td>
<td>0.028</td>
</tr>
<tr>
<td>CRF</td>
<td>10.6±1.94</td>
<td>10.1±2.31</td>
<td>10.4±2.13</td>
<td>0.83</td>
<td>0.436</td>
</tr>
<tr>
<td>CCT</td>
<td>564.8±34.2</td>
<td>550.0±35.8</td>
<td>553.0±33.2</td>
<td>4.84</td>
<td>0.009</td>
</tr>
</tbody>
</table>

Tukey’s multiple comparison test

<table>
<thead>
<tr>
<th></th>
<th>IOPcc</th>
<th>CH</th>
<th>CCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control /healthy macrodiscs</td>
<td>0.703</td>
<td>0.988</td>
<td>0.015</td>
</tr>
<tr>
<td>Control/glaucomatous macrodiscs</td>
<td>0.003</td>
<td>0.048</td>
<td>0.99</td>
</tr>
<tr>
<td>Healthy macrodiscs /glaucomatous macrodiscs</td>
<td>0.024</td>
<td>0.035</td>
<td>0.045</td>
</tr>
</tbody>
</table>

CCT: central corneal thickness; CH: corneal hysteresis; CRF: corneal resistance factor; IOP: intraocular pressure; IOPcc: corneal compensated IOP; IOPg, Goldmann-correlated IOP; SD: standard deviation.
was significantly lower in APON patients than in glaucoma patients (20). This finding also shows that lower CH could be marker for possible susceptibility of the optic nerve to glaucomatous damage, independent of CCT. As such, our study showed that CH of these healthy macrodiscs was similar to that of control group. Glaucomatous eyes with large disc area had lower CH than healthy macrodiscs.

Large discs have large total lamina cribrosa area and more lamina pores than small discs. These pores allow more space for nerve fibers to travel through, and therefore, they reduce risk of compression to the optic nerve axons (21). On the other hand, pressure differential across the lamina cribrosa can produce an increased deformation and displacement of the central tissue in macrodiscs, leading to greater glaucoma susceptibility in these eyes (22, 23). Burgoyne et al. suggested that mechanical failure of the connective tissue of the lamina cribrosa underlies glaucomatous cupping. Therefore, large discs may be more susceptible to pressure damage, as per Laplace’s law (24). Burk et al. concluded that statistically normal IOP readings should not be considered protection against future glaucomatous damage, especially in ONH with increased cupping and large disc areas (25). Hence, we can suggest that eyes with large optic disc area may be susceptible to glaucomatous damage, but high CH and corneal thickness can provide protection.

Weizer et al. demonstrated that CCT diminished after mean follow-up of 8 years, and this reduction was more pronounced in glaucomatous patients than in healthy subjects. However, exact mechanism of this reduction in glaucoma patients has not been explained. In this study, we found negative correlation between age and CCT only in glaucoma group (26).

Limitation of this study was that our glaucoma patients had been taking glaucoma medications, including prostaglandins, for significant length of time. There is chance that this may have affected ORA measurements. Prospective studies should be carried out to investigate this hypothesis.

In conclusion, macrodisc may have macrocup and should not be misdiagnosed as glaucoma. Present study has shown that healthy macrodiscs can be differentiated from glaucomatous discs using OCT parameters of rim area, cup area, C/D area ratio, and vertical and horizontal C/D ratios. We suggest that macrodiscs may be susceptible to glaucomatous damage due to mechanical failure of lamina cribrosa, but high level of CH and corneal thickness can provide protection against this damage.

Disclosures

Peer-review: Externally peer-reviewed.

Conflict of Interest: None declared.

Authorship Contributions: Involved in design and conduct of the study (FEO, BS, CA, BB, IBP, MT); preparation and review of the study (FEO, BS, CA, MT); data collection (FEO, CA, BB, IP); and statistical analysis (FEO, IP, BB).

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