Differentiating untreated and cross-linked porcine corneas of the same measured stiffness with optical coherence elastography

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Abstract. Structurally degenerative diseases, such as keratoconus, can significantly alter the stiffness of the cornea, directly affecting the quality of vision. Ultraviolet-induced collagen cross-linking (CXL) is an emerging treatment for keratoconus, which increases corneal stiffness. In addition to structural changes within the corneal tissue caused by CXL, intraocular pressure (IOP) also has an effect on the measured stiffness of the cornea. Therefore, there is a possibility that a cornea may be structurally weakened by keratoconus, yet have a “normal” measured stiffness due to an elevated IOP. Current techniques are not able to measure the true IOP in vivo without consideration of the effect of corneal biomechanical properties.

Distinguishing corneas that have the same measured stiffness but are at different IOPs is still a challenge. Optical coherence elastography (OCE) is an emerging non-invasive technique that can map the local biomechanical properties of tissues. Similar to ultrasound elastography (USE) and magnetic resonance elastography (MRE), OCE is usually composed of an external loading component that produces displacements within the tissue. In OCE, imaging this tissue displacement is performed with optical coherence tomography (OCT), which has superior spatial resolution compared with USE and MRE. From the velocity of an induced elastic wave (EW), or stress-strain curve measured by OCE, tissue elasticity can be quantitatively estimated.

In this work, we present a method utilizing OCE which is capable of distinguishing corneas of the same measured stiffness but at different IOPs. Validation experiments were performed on agar and gelatin phantoms of the same stiffness. This method was then applied to untreated (UT) and UV-induced collagen CXL porcine corneas. Artificial IOP control was used to induce the same measured corneal stiffness in the UT and CXL eyes.

2 Methods and Results

A home-built phase-stabilized swept-source OCT (PhS-SSOCT) system consisted of a focused air-pulse delivery system and a phase-stabilized swept-source OCT (PhS-SSOCT) system. Details of the system can be found in our previous work. Briefly, a short duration-focused air-pulse was expelled through an electronic solenoid-controlled air gate and induced an EW in the sample. A pressure gauge provided air source pressure control and measurement. The localized air-pulse excitation was positioned with a three-dimensional linear micrometer stage. The PhS-SSOCT system was composed of a broadband swept laser source (HSL2000, Santec, Inc., Torrance, California) with a central wavelength of 1310 nm, bandwidth of ~150 nm, scan rate of 30 kHz, and output power of ~29 mW. A-scan acquisition was triggered by a fiber Bragg grating. The axial resolution of the system was ~11 μm in air. The experimentally measured phase stability of the system was ~16 mrad, which corresponded to ~3.3-nm displacement in air. By synchronizing the focused air-pulse with consecutive M-mode images, the EW velocity and a two-dimensional depth-resolved elasticity were calculated. Previous studies have demonstrated that OCE is feasible for quantitatively assessing the elasticity of a sample. A validation study was initially conducted on 14.0% gelatin (w/w) and 1.1% agar (w/w) phantom samples (n = 5 for each type) with the same cylindrical dimensions of diameter D = 33 mm and height H = 11 mm. As shown in Fig. 1.

1 Introduction

Detecting changes in the biomechanical properties of ocular tissues can aid in the diagnosis of structurally degenerative diseases. For example, keratoconus can pathologically decrease the stiffness of the cornea, leading to a loss in the quality of vision. Ultraviolet (UV)-induced collagen cross-linking (CXL) is an emerging treatment for keratoconus, which increases corneal stiffness. In addition to structural changes within the corneal tissue caused by CXL, intraocular pressure (IOP) also has an effect on the measured stiffness of the cornea. Therefore, there is a possibility that a cornea may be structurally weakened by keratoconus, yet have a “normal” measured stiffness due to an elevated IOP. Current techniques are not able to measure the true IOP in vivo without consideration of the effect of corneal biomechanical properties.

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the EW velocity, \( c \), measured by PhS-SSOCE in the gelatin samples was 3.76 ± 0.2 m/s, which was very similar to the EW velocity in the agar samples of 3.64 ± 0.3 m/s. The acoustic surface wave equation [Eq. (1)] was used to estimate the Young’s moduli of the samples, where the density, \( \rho = 1.02 \text{ kg/m}^3 \) and Poisson ratio, \( v = 0.49 \). As shown in Fig. 1(b), the Young’s modulus for the 14.0% gelatin and 1.1% agar phantoms obtained by the analytical model were 48.7±9.2 and 46.6±8.2 kPa, respectively. Uniaxial mechanical compression tests (Model 5943, Instron Corp., Massachusetts) were conducted on the phantoms for elasticity validation. The measured Young’s modulus was 47.6±5.3 kPa for the 14% gelatin and 44.9±6.6 kPa for the 1.1% agar phantoms as shown in Fig. 1(b). These results demonstrated that the 14.0% gelatin sample and 1.1% agar phantoms were of similar stiffness, as confirmed by both analytical model and uniaxial compression tests.

\[
E = \frac{2\rho(1 + v)^3c^2}{(0.87 + 1.12v)^2} \quad (1)
\]

To compare the damping characteristics between any two normalized displacement amplitude attenuation curves of the EWs, a customized ratio, \( r \),

\[
r_{ND_i/ND_2} = \text{mean}(r_i) + \text{std}(r_i) \quad \text{with} \quad r_i = \frac{ND_i}{ND_2} \quad (2)
\]

was used, where \( ND_i \) and \( ND_2 \) were the normalized displacement of the induced EW at the \( i \)th measurement position for samples 1 and 2, respectively. Displacement amplitudes were normalized by dividing the EW displacement amplitude at each measurement position by the displacement amplitude at the excitation position. If \( r \) was significantly greater than 1, the displacement in sample 2 damped faster than in sample 1. If \( r \) was significantly less than 1, sample 1 damped faster than sample 2. If \( r \) was close to 1, the damping was similar in both samples.

This ratio was first calculated for the same 14.0% gelatin phantom to examine the effects of different initial position displacements by changing the focused air-pulse pressure on the sample to 11 and 22 Pa. The normalized displacement attenuation curves are shown in Fig. 2(a) with the ratio \( r_{22/11} = 0.95 \pm 0.12 \), which was very close to 1. As anticipated, this indicated that the initial displacement amplitude did not affect the damping speed of the EW.

This ratio was then calculated to compare the gelatin and agar phantoms. As shown in Fig. 2(b), the normalized displacement in the agar phantoms was higher than in the gelatin phantoms at the same scan position. By using Eq. (1), \( r_{\text{agar/gelatin}} = 1.56 \pm 0.47 \), which demonstrated that the 14% gelatin damped faster than the 1.1% agar. This result was in agreement with previous findings that gelatin has a higher viscosity than agar, which corresponds to faster damping. Therefore, these comparisons showed that this method could be successfully utilized to distinguish two materials of similar stiffness.

To induce a similar measured corneal stiffness in the UT and CXL porcine corneas, the IOP of the whole eye was controlled by a custom-built controller comprising a pressure transducer and microinfusion pump connected in a feedback loop. The EW was measured in a porcine cornea by the PhS-SSOCE system before and after UVA-Riboflavin-induced CXL. EW measurements were taken at IOPs from 15 to 35 mm Hg with 5-mm Hg increments. The EW velocities of the EW in the UT and CXL corneas at the various IOPs are presented in Table 1.

<table>
<thead>
<tr>
<th>IOP (mm Hg)</th>
<th>UT EW velocity (m/s)</th>
<th>CXL EW velocity (m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>1.5 ± 0.1</td>
<td>2.7 ± 0.1</td>
</tr>
<tr>
<td>20</td>
<td>2.3 ± 0.1</td>
<td>3.6 ± 0.1</td>
</tr>
<tr>
<td>25</td>
<td>3.0 ± 0.3</td>
<td>4.0 ± 0.1</td>
</tr>
<tr>
<td>30</td>
<td>3.6 ± 0.4</td>
<td>4.2 ± 0.2</td>
</tr>
<tr>
<td>35</td>
<td>3.7 ± 0.4</td>
<td>4.7 ± 0.5</td>
</tr>
</tbody>
</table>

Note: The bold values indicate that exactly same values were obtained under different tissue manipulations.
after the CXL treatment. In addition, the normalized displacement attenuation curves were fitted by \( y = ae^{bx} \) in which the parameter \( b \) was treated as the damping speed. According to the fitted results, the damping speed of the UT cornea (\( b_{UT} = -0.031 \text{ mm}^{-1} \)) was almost twice the damping speed of the CXL cornea (\( b_{CXL} = -0.017 \text{ mm}^{-1} \)), which confirmed that the damping speed decreased after CXL treatment. One possible reason for this result is that the CXL treatment is a procedure which displaces water from the cornea tissue. The UT cornea contains more water which is responsible for a higher viscosity. Therefore, the EW damps faster in the UT cornea than the CXL cornea.

Kotecha et al. measured biomechanical parameters of UT eyes at different IOPs with the ocular response analyzer (ORA, Reichert Inc., Depew, New York) and discussed how the viscosity was negatively correlated with measured corneal stiffness, indicating that the CXL cornea has a lower viscosity than the normal one, which corroborates with our results. However, the ORA only provides the index of corneal hysteresis to reflect the corneal damping ability, but is unable to provide the information about the cornea stiffness. Furthermore, the induced displacement in ORA is in the order of mm, which is hundreds of times larger than in the present method.

3 Conclusion

We have demonstrated a method using PhS-SSOCE to distinguish UT and CXL corneas of the same measured stiffness but at different IOPs. This noninvasive method has potential to evaluate the biomechanical properties of the cornea in vivo for detecting the onset and progression of corneal degenerative diseases such as keratoconus. Future work would entail extracting and separating the elasticity and viscosity of the cornea.

Acknowledgments

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References