Attenuation coefficient as a quantitative parameter for analyzing cataracts with optical coherence tomography

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Abstract. Crystalline lenses of mice were imaged in vivo with a custom-made swept-source optical coherence tomography system. The use of the attenuation coefficient as a quantitative parameter for investigating the lens opacities magnitude is proposed, demonstrating a significant difference between the values retrieved from cataractous and normal mouse lenses.

1 Introduction

Cataracts are the major cause of blindness worldwide, they are characterized by a progressive loss of transparency of the ocular lens which ultimately leads to poor vision [1]. Optical coherence tomography (OCT) has been proposed as a non-invasive imaging technique to visualize and quantify the morphology of these opacifications in human patients and animal models [2,3]. In biomedical applications, the attenuation coefficient of the tissue recovered by optical methods has been used as a parameter for classification and diagnosis [4]. Previously we presented OCT as a tool to visualize different opacifications in vivo in the crystalline lens of mice [3]. Here we want to extend our previous results with the use of the optical attenuation coefficient as a further measure to analyze and characterize cataract formations.

2 Methods

In this paper we used the custom-made OCT system illustrated in Fig. 1(A), which operates at a central wavelength of 1310 nm and provides an axial resolution of ~6.5 μm in air with a sensitivity of 100 dB. The OCT device was described in detail elsewhere along with the experimental procedure of the data used for this work [3]. Data sets with 400 × 400 × 768 pixels (3 × 3 × 5 mm) were acquired in the mouse anterior segment, repeating 4 scans per position to increase the signal-to-noise ratio. A post-processing pipeline was implemented in MATLAB (MATLAB, R2015b, MathWorks). Every acquired spectrum was aligned with a unique spectrum using cross-correlation to reduce shifts induced by A-line trigger jitter. Background removal and spectral shaping were then performed before averaging the resulting repeated scans after Fourier transformation. The lens surface was segmented and the equation

\[ y = A \cdot \exp(-\mu z) \]  

(1)

was fitted to the intensity profiles in all A-scans for a range spanning 0.5 mm in depth underneath the surface. In the fit, the scaling factor \(A\) and the attenuation coefficient \(\mu\) were the free running parameters to be optimized by the algorithm. En-face attenuation maps were then calculated. A similar region of interest consisting of 0.16 mm² was analyzed for cataract and non-ctaract eyes and compared as shown in Figure 1(B). A Student’s t-test was used to analyze the difference between the average attenuation coefficients of the two groups.

Fig. 1. A) Home-built OCT device used for mouse lens imaging. PBS: beam splitter, BD: balanced detector, L: lens, M: mirror, MEMS: microelectromechanical scanner, PC: polarization controller, QWP: quarter wave plate, RR: retroreflector. B) Average attenuation coefficient compared between normal and cataract eyes.

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3 Results

An attenuation map of a mouse eye without cataract is shown in Figure 1(A). In the eyes where no opacification was observed, only a very subtle scattering signal was detected in the lens due to its transparency, which led to low attenuation coefficients as shown in the Figure 1(B) and 2(ACD). An intensity profile of a lens without opacification is shown in Figure 2(D). When an opacity was present in the mouse eye, the attenuation within the lens was stronger as shown in Figure 1(E). This was due to a cloudy hyperscattering signal caused by the cataract formed in the lens cortex as shown in Figure 2(F). In the lateral attenuation profile of a B-scan with opacification, higher values can be observed in addition to the ones corresponding to the iris on the left as shown in Figure 2(G). This is also visible in the axial intensity profile shown in Figure 2(H) where higher values, rising as well as a steeper slope of the fitted exponential can be observed. When comparing similar regions between cataracous and non cataracous eyes, a higher attenuation ($5.5 \pm 0.7 \text{ mm}^{-1}$) was found in cataracous areas compared to the non-ctaracts regions ($1.8 \pm 0.5 \text{ mm}^{-1}$), showing a significant difference.

4 Conclusion

A custom-made OCT system was used for volumetric in vivo visualization of the crystalline lens in mice. A post-processing pipeline was implemented to obtain the attenuation coefficient, which revealed a significant difference between cataracts and normal eyes.

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