Single-camera polarization-sensitive full-field optical coherence tomography with polarization switch

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Abstract. We present a single-channel detection-based polarization-sensitive full-field optical coherence tomography (PS-FF-OCT) for simultaneous acquisition of high-resolution OCT and linear retardance images. A linearly polarized sub-10-fs laser was used as a broadband light source for the OCT system. A bi-stable polarization switching device, composed of a ferroelectric liquid crystal cell and an analyzer, was employed to get the horizontal and the vertical polarization components of a full-field interference signal. The time-switched two perpendicularly polarized interference signals were sequentially recorded by a single charge-coupled device camera, then processed to extract en-face functional images of a biological sample. The rat tail tendon was imaged ex vivo to confirm the imaging feasibility of the proposed system, which showed axial and transverse resolutions of 2 and 1.3 μm, respectively. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.18.10.100504]

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Polarization-sensitive optical coherence tomography (PS-OCT) is a functional extension of OCT and able to produce images of the structural and polarization properties of biological tissue with a micrometer resolution.1 Phase retardance, being a key feature of PS-OCT and previously providing supplementary tissue information, now offers new insight into the evaluation of tissue inhomogeneity, which has contributed to a broad range of OCT applications including ophthalmology,2 dermatology,3 gynecology,4 and oncology.5 Recent PS-OCT systems have been based on frequency-domain OCT (FD-OCT) because of the advantages of the high-speed imaging rate and high sensitivity. As an alternative to the FD-based PS-OCT, the PS full-field OCT (PS-FF-OCT) has been proposed.6 Despite its rather low sensitivity and imaging speed compared with the FD-OCT, the PS-FF-OCT is attractive for its ability to provide images of the functional structure at a subcellular level.

Previous studies on PS-FF-OCT, however, used two identical charge-coupled device (CCD) cameras as detectors to get two perpendicularly polarized interference images. In such a dual-channel detection scheme, precise positioning between the two CCD cameras is imperative so to prevent pixel mismatching between the two CCD images, which causes image deformation. Further, the use of an additional CCD camera also requires added cost and extra space in system implementation. To address these issues, we propose a single-channel detection scheme of PS-FF-OCT. Most of the previously reported single-channel PS-OCTs were based on fiber-scanning OCT techniques,9,10 in which a one-dimensional line CCD camera, not a two-dimensional area CCD camera, was used.

To get the full-field interference images in both polarization states with a single CCD camera, a bi-stable polarization switching (BSPS) device was implemented, which consists of a ferroelectric liquid crystal (FLC) cell and a linear polarizer or a polarization optics.11 Typically, the standard FLC cell is fabricated as a half-wave retarder,12 which causes chromatic or phase errors in broadband applications. However, in our experiment, an achromatic FLC waveplate (Displaytech Inc., Carlsbad, California) was used, which was designed and assembled so that the retardance was a half-wave in the broadband spectral range (~300 nm). The achromatic FLC cell works at a center wavelength of 820 nm, which can switch its fast axis in 80 μs and be modulated up to 20 kHz.

A schematic of the experimental setup is shown in Fig. 1. As a broadband light source, a Ti-Sapphire sub-10 femto-pulsed laser (Rainbow, Femtolasers Inc., Viena, Austria) was used in experiments. Spectral bandwidth of the source was 130 nm at full-width half-maximum (FWHM) with an 800-nm center wavelength, and the average output power was approximately 320 mW. An extremely broad bandwidth of the laser was partially cut off from 720 nm, due to the limited spectral sensitivity of the achromatic quarter wave plates (AQWPs), linear polarizers (LPs), and the FLC cell. We applied a spectral filter to minimize optical aberration, allowing the transmitted spectral bandwidth with 127 nm at FWHM. Accordingly, the axial resolution of the system was calculated to be ~1.7 μm in water, but it was measured as 2.0 μm experimentally, which was affected by residual dispersion or the non-Gaussian spectral shape of the source spectrum. The output beam from the laser was delivered through a 22-m long multimode fiber (MMF) with a core diameter of 400 μm to reduce spatial coherence within the FF-OCT light source.13

The beam from the MMF passed through the field stop, aperture stop, and achromatic doublet lens, designed for the Köhler illumination. After that, the beam was split by a beam splitter (BS) into reference and sample arms. Both arms were equipped with identical water immersion microscope objectives and the AQWPs. Also, a neutral density filter for attenuation of light intensity and a glass plate to match the dispersion with the reference arm were positioned in reference and sample arms, respectively. For the sample arm, the AQWP was set to be oriented at 45 deg so that the linearly polarized input beam changes into a...
circularly polarized one. For the reference arm, the AQWP was set to be oriented at 22.5 deg; the polarization direction of the round trip beam was oriented at 45 deg, and vertical and horizontal polarization components of the round trip beam thus had identical amounts. The beams reflected and backscattered from the reference and the sample arms recombined at the BS. The combined beams passed through the BSPS device and then were detected by the single CCD camera (CCD1020, 10-bits, 512 × 512 pixels, 20 frames/s, VDS).

As depicted in the inset of Fig. 1, the piezoelectric transducer (PZT), the CCD camera, and the FLC cell were synchronized to each other and operated at 5, 20, and 2.5 Hz, respectively. A set of four CCD frames \( \Gamma \kappa ; E \kappa \), \( \Gamma \kappa \), \( \Gamma \kappa \kappa \), \( E \kappa \) were captured during one period of PZT under sinusoidal phase modulation to capture the vertical component of the interfered beam. The consecutive set of four CCD frames \( E \kappa, E \kappa, E \kappa, E \kappa \) were taken during the next period of the PZT modulation to provide the horizontal component of the interfered beam. Finally, intensity and phase retardation images were calculated from measured vertical and horizontal polarization components as explained in Ref. 8. Those can be expressed as 

\[
\Gamma \kappa = (E \kappa - E \kappa)^2 + (E \kappa - E \kappa)^2 \\
\Gamma \kappa = (E \kappa - E \kappa)^2 + (E \kappa - E \kappa)^2
\]

The retardation image could be constructed by taking the ratio of \( \Gamma \kappa \) and \( \Gamma \kappa \). The intensity image was also made by the summation of \( \Gamma \kappa \) and \( \Gamma \kappa \) at the same time.

To validate the system performance, we measured the transmission spectrum after passing through the BSPS and measured the retardation using the proposed FF-PS-OCT system with a sample of known retardation state. Figure 2(a) shows the wavelength dependency of the BSPS with the switching signals. The vertically polarized light was passing through the BSPS device, while the transmission axis of the linear polarizer was installed in the vertical direction. The transmission was suppressed around −11 dB at −5 V switching signal.

The Babinet compensator was used to generate a certain birefringence-induced phase retardation to the system and placed between the BS and the microscope objective (MO) in the sample arm. To match the dispersion with the sample arm, a glass plate of appropriate thickness corresponding to the compensator was positioned in the reference arm. While rotating the gradations on the compensator by the amount giving 15-deg phase retardation, phase retardation values were measured. Every measurement was carried out by averaging 10 frames. The measured retardation was coplotted with given retardation, as shown in Fig. 2(b). The polarization-induced phase retardation was linearly increased with the actual retardation, but it showed relatively high errors around the zero retardation. This discrepancy between expected and measured retardations can be caused by the remaining transmission light from the low-extinction ratio of the FLC cell.

To confirm the performance of the system in the given biological tissue, a rat tail tendon was prepared and imaged with water immersion micro-objectives (Olympus, 10×, NA 0.3, MO), which gives a theoretical spatial resolution of ~1.3 μm. The skin of the rat tail was peeled by a stripper so to view and image the tendon directly. During the experiment, the prepared rat tail tendon was immersed in phosphate-buffered saline to maintain a constant pH in the biological tissue immersion medium. To prevent CCD camera image saturation, the launched power was heavily attenuated by a linear polarizer and an attenuator. The illumination power on the sample was about 25 μW. The acquisition time for one PS-FF-OCT image took approximately 0.4 s, and 10 images were averaged for one depth-resolved image.

Figure 3 shows PS-FF-OCT images of the rat’s tail tendon with its polarization-induced phase retardation images. Several strands of tendon with crimp patterns adjoin each other, as shown in Fig. 3(a). Secondary fiber bundles (fascicles) and even subfascicles (digits 1 to 4) of the tendon, which are invisible with conventional PS-OCT in general, are clearly visible in the XY and YZ images, respectively. Figure 3(b) shows the linear retardance image taken at the same depth of the intensity image of Fig. 3(a). The phase retardation from 0 to 90 deg...
Ex vivo images of the rat tail tendon by PS-FF-OCT: (a) En-face intensity images at a depth of 43 μm; XZ and YZ cross-sectional images.
(b) En-face retardation image at the same depth with (a); XZ and YZ cross-sectional retardation images.

The cross-sectional image shows the variation of phase retardation along and across the tendons.

In summary, we have constructed a single-channel PS-FF-OCT using a BSPS device, which consists of a FLC cell and an LP. The BSPS device was positioned in front of the CCD camera to rotate the polarization state of the scattered light, providing a geometric separation of two perpendicular components of interfered light. As a result, the temporal split of two polarization components enabled a single CCD camera to collect two polarization components. Thanks to the high-resolution nature of PS-FF-OCT, the fascicle structure of the tendon was apparent with its retardation image. This method provides a simple process and relatively easy alignment of the PS-FF-OCT system.

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