

Endoscopic Functional Fourier Domain Common-Path Optical Coherence Tomography for Microsurgery

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(Invited Paper)

Abstract—A single-arm interferometer-based optical coherence tomography (OCT) system known as common-path OCT (CP-OCT) is rapidly progressing towards practical application. Due in part to the simplicity and robustness of its design, Fourier domain CP-OCT (FD-CP-OCT) offers advantages in many endoscopic sensing and imaging applications. FD-CP-OCT uses simple, interchangeable fiber optic probes that are easily integrated into small and delicate surgical tools. The system is capable of providing not only high-resolution imaging but also optical sensing. Here, we report progress towards practical application of FD-CP-OCT in the setting of delicate microsurgical procedures such as intraocular retinal surgery. To meet the challenges presented by the microsurgical requirements of these procedures, we have developed and initiated the validation of applicable fiber optic probes. By integrating these probes into our developing imaging system, we have obtained high-resolution OCT images and have also completed a demonstration of their potential sensing capabilities. Specifically, we utilize multiple superluminescent diodes to demonstrate sub-3- μm axial resolution in water; we propose a technique to quantitatively evaluate the spatial distribution of oxygen saturation levels in tissue; and we present evidence supportive of the technology's surface sensing and tool guidance potential by demonstrating topological and motion compensation capabilities.

Index Terms—Biomedical optical imaging, optical fibers, optical interferometry, optical tomography.

I. INTRODUCTION

OPTICAL coherence tomography (OCT) has recently emerged as a valuable 3-D imaging technology for medical applications [1]–[3]. One of the earliest uses of optical interferometric techniques in ophthalmology was to accurately measure the thickness of cornea [4], [5]. With additional technological and commercial refinements, OCT is extensively used in the assessment of ocular pathology with highest clinical utility

Manuscript received July 30, 2009; revised August 26, 2009; accepted August 26, 2009. Date of publication October 23, 2009; date of current version August 6, 2010. This work was supported in part by the National Institutes of Health under Grant BRP 1R01 EB 007969-01 and Grant R21 1R21NS063131-01A1, and by the National Science Foundation under Grant NSF101485.

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Digital Object Identifier 10.1109/JSTQE.2009.2031597

in the area of diagnostic assessment of macular disease in the retina. The earlier interferometric technique used by Fujimoto *et al.* [4] was based on femtosecond Michelson interferometry, whereas Fercher *et al.* [5] used a simpler common-path interferometric method. Since then the Michelson interferometer-based OCT has emerged as the technique of choice. A key factor in the emergence of the Michelson interferometer was its use of a high-peak power femtosecond laser that allowed for increased resolution and a higher signal-to-noise ratio (SNR). Since its first application to clinical retina in the field of ophthalmology, OCT scanning has emerged as one of the most utilized diagnostic applications in this field [6]. Ophthalmological application has proven to be a particularly applicable field for OCT applications due in part to the relative transparency of relevant eye tissues such as the cornea and retina [7]. In the last decade, numerous technological breakthroughs have been made and OCT now offers unparalleled resolution in real time [8]–[27]. One relatively unexplored yet promising application of OCT is in the area of endoscopic 3-D imaging and optical sensing. Endoscopic optical coherence tomography (EOCT) is, however, associated with significant technical challenge that leads to a choice between reduced imaging sensitivity or a significant increase in the level of technical complexity of the required system [28]. Ultrahigh resolution EOCT is even more difficult to implement because of the issues of polarization and dispersion mismatch (over a wide range of wavelengths) between the two arms of the Michelson interferometer [29]. However, each of these difficulties has been largely addressed via a strategic choice of components and ingenious system design.

Despite great progress in the development of Michelson interferometer-based technologies there remains significant interest in further development of common-path interferometer-based OCT technology. The common-path approach utilizes a simple interferometer design where the sample and reference arms share a common path. This allows a much simpler system design, lower associated costs, and the ability to use interchangeable probes as well as the freedom to use any arbitrary probe arm length. A common-path interferometer-based frequency-domain OCT (FDOCT) has been reported [30] and other common-path interferometer-based approaches for time-domain OCT, using a free space interferometer, have also been demonstrated with limited success [31], [32]. In the last several years, there has been rapid development in the fiber optic-based

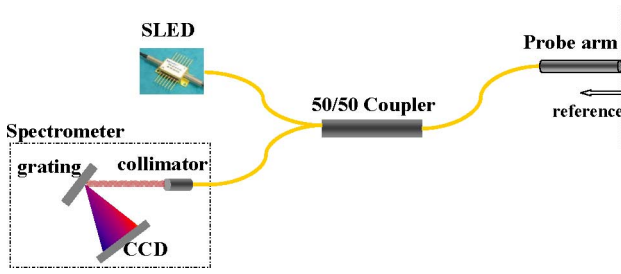


Fig. 1. Schematic diagram of a FD-CP-OCT system.

technologies that is applicable to both frequency-domain and time-domain common-path OCT systems [33]–[39].

The fiber optic common-path OCT (CP-OCT) is compatible with existing endoscopic medical instrumentation. In recent years we have been developing functional Fourier domain CP-OCT (FD-CP-OCT) for its potential application in the field of microsurgery with intraocular surgical applications relevant to retinal surgery as the model application [40], [41]. In this paper, we will describe in detail the results of our recent progress towards practical FD-CP-OCT microsurgical application. Section II details an overview of FD-CP-OCT theory as well as the general experimental setup. In Section III, we briefly discuss a CP-OCT fiber optic probe design with the integration of a simple surgical tool, in this case, a surgical needle. Section IV details our experimental results following intraocular 3-D, *in vivo* imaging of ocular tissue in a fish eye model. Section V details experimental results and a processing method to obtain ultrahigh axial resolution. Section VI presents an analysis method and experimental results from oxygen saturation (SO_2) measurement experiments. Finally, Section VII presents a method and demonstration of FD-CP-OCT-based topological and motion compensation.

II. FD-CP-OCT

A. System Configuration

The schematic representation of a simple and cost effective FD-CP-OCT system is shown in Fig. 1. The interrogation of a target starts from a broadband source that couples to one of the arms of a fiber coupler and exits out to the probe arm, which acts as both the sample and the reference arm. The reference signal is obtained from a partial reflector near the distal end of the probe arm. The reference and the signal couple back to the fiber and exit out of the third arm of the fiber coupler to the optical spectrometer and the resultant spectrum is analyzed to obtain a longitudinal image. Commercial superluminescent diodes (SLED) from Exalos were used as broadband sources. The spectrometer was in-house built using a 2048 pixel Aviva line scan camera from e2v Technologies. The spectrometer is capable of up to 28,000 line scan per second.

B. SNR Study

The FD-OCT signal amplitude can be written as [42]

$$S_{\text{peak}} = \frac{S\eta\tau}{Nh\nu_0} \sqrt{P_r P_s} = \frac{1}{N} \sqrt{N_{P_r} N_{P_s}} \quad (1)$$

where S_{peak} is a dimensionless quantity. S is spectrometer efficiency, η is detector quantum efficiency, τ is charge-coupled device (CCD) exposure time, N is the number of CCD pixels. h is the Planck constant, ν_0 is the center frequency of the light source spectrum. P_s is sample power and P_r is reference power. In the system, optical energy incident onto each pixel is detected and digitized by CCD system. The summation of counts at all the pixels is proportional to the optical energy incident on the spectrometer. N_{P_r} and N_{P_s} are the summations of counts corresponding to the reference power and the sample power, respectively. We can readily obtain N_{P_r} and N_{P_s} using (1).

However, (1) assumes reference and sample light are perfectly coherent, which is not practically valid. First, the experimental pulse spread function (PSF) is broader than the theoretical PSF, which means that the signal amplitude has to be diminished to conserve energy. Second, the interferometer is not an ideally sinusoidal signal modulator. For the above reasons, the measured peak amplitude $\widehat{S}_{\text{peak}}$ of A-scan obtained by Fourier transforming the interferogram is much smaller. Taking the ratio $r = \widehat{S}_{\text{peak}}/S_{\text{peak}}$ into consideration, the SNR can be expressed as

$$\text{SNR} = 10 \log_{10} \left(r^2 \frac{S_{\text{peak}}^2}{\sigma_{\text{noise}}^2} \right) \quad (2)$$

where σ_{noise}^2 is the noise. The main noise sources in FD-OCT systems are receiver noise, shot noise and excess noise.

The excess noise of the CP-FD-OCT is given by

$$\sigma_{\text{excess}}^2 = \frac{1+V^2}{2\delta v'} \left(\frac{S\eta}{h\nu_0} \frac{P_r + P_s}{N} \right)^2 \tau. \quad (3)$$

Here, V is the degree of polarization and $\delta v' = \Delta v_{\text{eff}}/N$. Δv_{eff} is the effective spectral line width [12].

Shot noise is given by

$$\sigma_{\text{shot}}^2 = \frac{S\eta\tau}{h\nu_0} \frac{P_r + P_s}{N}. \quad (4)$$

If we assume the noise originated from each CCD pixel is additive, independent and white, than they combine incoherently after FFT. In other words, they add in intensity instead of amplitude. The total noise after FFT is then given by

$$\sigma_{\text{noise}}^2 = (\sigma_{\text{shot}}^2 + \sigma_{\text{excess}}^2 + \sigma_{\text{dark}}^2 + \sigma_{\text{read}}^2) \frac{1}{N}. \quad (5)$$

A mirror is used as a sample for analyzing SNR of the system. Only does a small fraction of light reflected by the mirror couple back to the probe. We changed the source power, while keeping the mirror and fiber probe fixed. The curve in Fig. 2 shows SNR as a function of reference power calculated by (5), and the open circles are the measured value.

It is noted that the SNR calculated by (2) and experimental SNR match well in Fig. 2. This result shows our noise model is reasonable, but (1) overestimates the signal amplitude.

III. CP ENDOSCOPIC FIBER OPTIC PROBE

Endoscopic fiber optic probes for FD-CP-OCT, in order to be integrated into existing surgical catheters and needles for

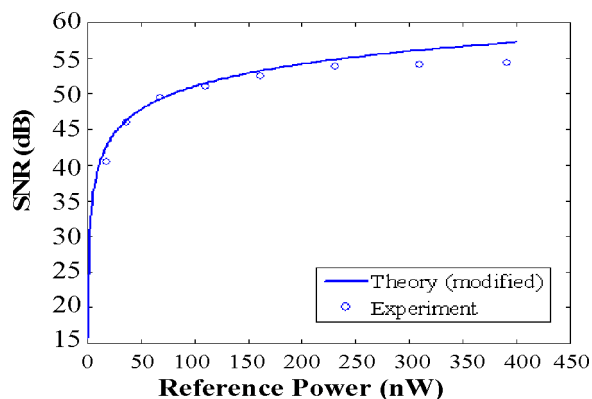


Fig. 2. Theoretical and experimental SNR as function of reference power.

assisting microsurgeries, needs to be simple and small. In order to achieve these design objectives, we have devised fiber optic probes that use Fresnel reflection at the distal end of the fiber as a reference. For operating in air, a simple end-cleaved fiber probe works well. However, for the cases where the optical probe needs to work in an intravitreal and/or intraocular environment, Fresnel reflection dramatically decreases from 3.59% to 0.24% due to the reduced refractive index difference between optical fiber and aqueous medium. For relatively high-incident laser powers >0.2 mW, this in fact increases the SNR of the system and better OCT images can be obtained. However, for low power levels <0.2 mW, a higher reference level is needed in order to maintain an appropriate level of SNR. To achieve this, we have coated the distal end of fiber with thin gold (Au) film. Thus, the reference amplitude does not depend on the reflectivity of the interface media even when the probe is submerged in the liquid or in contact with the tissue. Gold was chosen because it is chemically stable and biomedically safe for medical applications. Here, no focusing lens was implemented with the Au-coated fiber probe (Au-FP) in order to limit the probe size to the current fiber diameter of $125\ \mu\text{m}$, as well as not to introduce any unwanted reflections from between the components. We have experimentally measured the lateral resolution and the working distance of the bare fiber probe and these were found to be $\sim 28\ \mu\text{m}$ at 1 mm from the fiber probe tip and 3 mm, respectively. Au-FP was fabricated using a standard sputtering process. Au-FPs were fabricated from standard single-mode optical fibers. The fiber was cleaved using an ultrasonic cleaver (FK11) and a thin-film gold-palladium (Au-Pd) alloy was deposited on the end-face using a standard sputtering technique (Denton Vacuum DESK II). The palladium (Pd) alloy helps the gold layer deposit evenly in a (few tens of nanometer thin) uniform-coated layer without beading as could otherwise result from the surface tension of the pure gold.

We performed Au-deposition for 0 s (0 s represents for normal bare fiber without coating), 5, 10, 15, and 25 s, and the applied incident power of 1 mW for testing. For less than or equal to 15 s of Au-thin film deposition, we were able to achieve a stable level of reflectance in the water. As expected, the reflectance without any coating (0 s) becomes $\sim 1/15$ compared to the value in the air from 3.59% to 0.24%. However, with more than 15 s of depo-

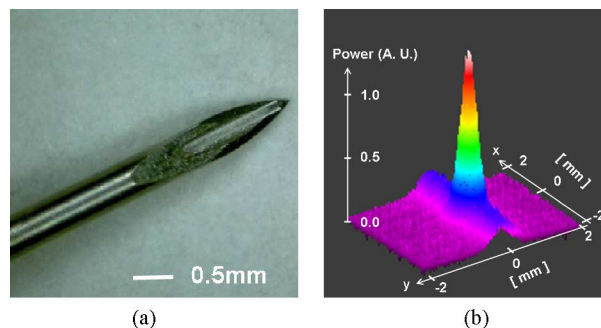


Fig. 3. OCT probe. (a) OCT fiber probe integrated surgical needle (prototype). (b) Output beam far field profile.

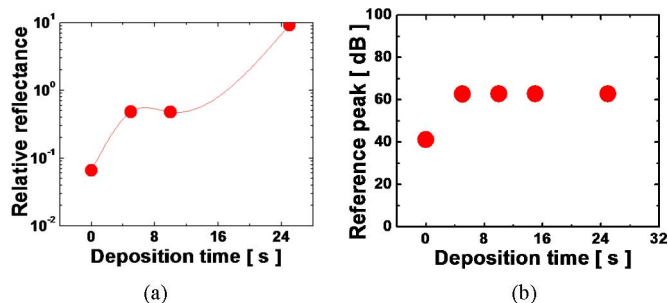


Fig. 4. Characteristics of reference magnitude in aqueous medium. (a) Ratio to the reflectance in the air. (b) Reference amplitude level at 1 mW source power.

sition, the layer thickness became too thick and the metal layer acted as a highly reflective mirror. The resulting reflectance was too large and did not permit any light transmission. An ideal deposition time was found to be around 15 s, which provided an ideal thickness of the coated layer that resulted in approximately 1.7% reflection in water, which is more than seven times greater than that without the coating. Afterwards, Au-FP was then integrated into a metallic hypodermic needle. B-mode lateral scanning was achieved simply by sweeping the whole probe using with a stepper motor.

Fig. 3(a) is a magnified microscope image of a fiber optic probe integrated into a 25-gauge surgical needle where the diameters of the needle and fiber (cladding) are ~ 500 and $125\ \mu\text{m}$, respectively. A sample 2-D far-field beam profile of the fiber probe was measured by an infrared (IR) CCD camera as shown in Fig. 3(b) in which the correlation between the beam profile and the perfect Gaussian shape is greater than 0.9 and the ellipticity, which is the ratio between the smaller and larger beam widths, was larger than 0.95. The reflectance of the Au-FP in water is shown in Fig. 4(a), where it is normalized to the reflectance in the air (3.59%) for comparison. The result shows that one can obtain the same amount of reference as a glass-air interface from the gold-coated probe in water when the deposition time is greater than 15 s. The deposition time of gold linearly relates to the total film thickness. The corresponding reference level in the OCT system is shown in Fig. 4(b). The reference peak saturates for the deposition time greater than ~ 7 s to ~ 62 dB above the noise level. Furthermore, the Au-coated probe showed no Fresnel reflection dependence on the interfacing medium. Uncoated fibers showed the same ~ 62 dB

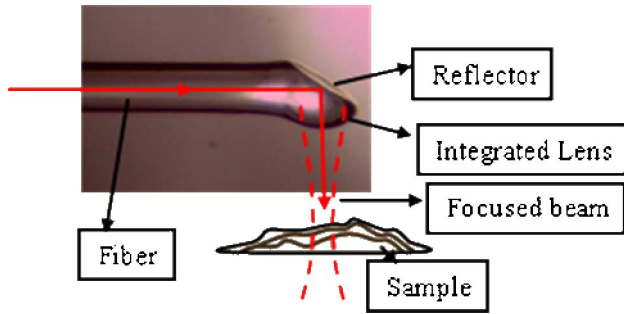


Fig. 5. Photograph of a side viewing fiber probe with integrated lens.

peak in the air but it was dropped by more than 20 dB in the water.

We have also demonstrated integrated side viewing probes as shown in Fig. 5. This design can also easily integrate into different microsurgical tools. The details of such probe design will be reported elsewhere. In addition, we are developing fiber bundle-based probe system that would eliminate the need for the physical sweeping of the probe for the B-mode imaging.

IV. OCT IMAGING

A. Intraocular Tissue Imaging

The CP-FD-OCT system setup for imaging is the same as that shown in Fig. 1. The source is a SLED light source centered near $0.8 \mu\text{m}$ (40 nm width), a fiber optic directional 2×1 coupler, a fiber probe, and a high-resolution spectrometer. The integrated probe consists of a bare single mode fiber (the numerical aperture of the fiber is 0.14) having a core size of $\sim 5.7 \mu\text{m}$ and outer polymer buffer diameter of $\sim 240 \mu\text{m}$ which is inserted and bonded to a 25-gauge hypodermic surgical needle having inner diameter of 0.24 mm, outer diameter of 0.5 mm, and $1\frac{1}{2}$ in. in length. For imaging the cornea, the subject eye was placed vertically to simulate a potential operating position. In order to reduce the strong surface reflection from the cornea and keep the sample from drying out, we submerged the sample in a solution of isotonic (0.9%) NaCl saline (index ~ 1.33). The specimen used were eyes from *Spirinchus lanceolatus* (or simply Shishamo) fish and *Engraulis japonicus* (or simply anchovy) fish.

For the lateral scans (B-mode and C-mode), the probe was fixed to the 2-D linear translational stage with stepper motors in the x and y axes, and with a high accuracy (step size between 1 and $5 \mu\text{m}$). The measured axial resolution ($\sim 8 \mu\text{m}$ in air) in the A-mode scan matched well with the theoretical resolution determined by the source coherence length. With an eye of *Spirinchus lanceolatus*, we performed lateral B-scan of the cornea with $2 \mu\text{m}$ step size. The anterior chamber (index ~ 1.336), lens and iris edge can be observed in Fig. 6(a) (image size: $3 \text{ mm} \times 0.86 \text{ mm}$). We can also clearly observe detailed features inside of the ocular tissue (e.g., cystic spaces seen as dark colored spots). We then performed a 3-D scan on an eye from *Engraulis japonicus*, with both B-scan and C-scan steps of $5 \mu\text{m}$. Fig. 6(b) displays the 3-D sliced volume of $2000 \mu\text{m} \times 2000 \mu\text{m} \times 860 \mu\text{m}$, and Fig. 6(c) is an en-face image recon-

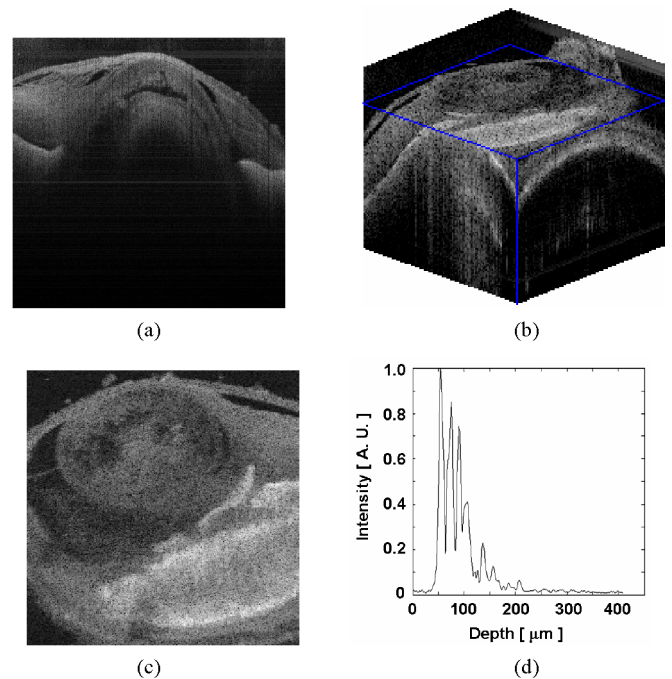


Fig. 6. Scanned OCT Images of intraocular tissues. (a) Cornea (2-D, *Spirinchus lanceolatus*). (b) Cornea (3-D, *Engraulis japonicus*). (c) Cornea en-face image reconstructed from (b) at depth = $150 \mu\text{m}$. (d) Retina (1-D, rat).

structed from the volume data at a depth of $\sim 150 \mu\text{m}$ from the anterior cornea. The lens structure and the iris are clearly visible. The retinal structures are evidenced by numerous peaks in Fig. 6(d). Retinal OCT is frequently used to assess retinal thickness and to evaluate pathology presenting as an altered pattern of reflective interfaces in clinical medicine.

B. Polarization Dependent Imaging of Birefringent Beef Sample

It is known that many tissues exhibit polarization dependent scattering and reflection [43]. This can lead to enhanced imaging for OCT systems that are capable of discriminating between different polarizations [44]. To test whether this principal is applicable to CP-OCT, we built a simple setup to conduct preliminary testing. To control the polarization, fiber-type circular retardation plates have been introduced and inserted between the bare fiber probe and the directional fiber coupler for manipulating the birefringence of the sample. The rest of the system is identical to the one presented in the previous section. The test tissue was a piece of fresh beef in this preliminary evaluation.

The tomograms of a fresh beef sample were compared in Fig. 7 for different polarizations and positions in the specimen. If the polarization was not discriminated, the OCT image would appear simply as a depth-resolved intensity profile. By controlling the polarization, we can obtain images that have different features depending on the collagen tissue characteristics. This is shown in Fig. 7(b) with dotted lined layers where clearer layered configuration for than that of Fig. 7(a) can be seen. In case of fat and muscle, they have different scattering features so that a larger signal contrast between them occurs in

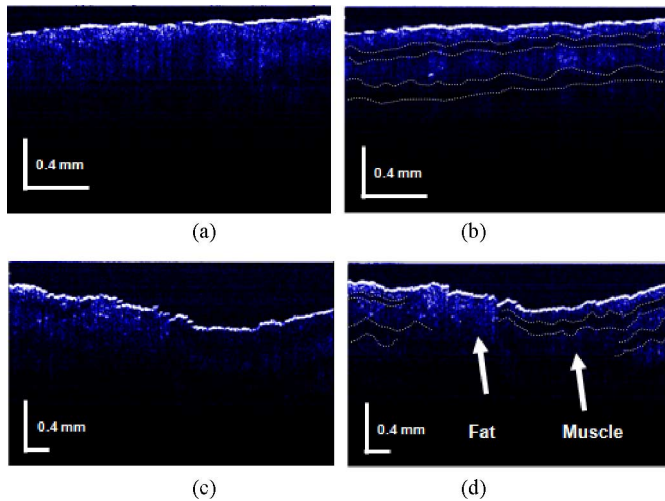


Fig. 7. Polarization-dependent FD-CP-OCT images of birefringent beef sample: (a) without polarization control (muscle only); (b) with polarization control (muscle only); (c) without polarization control (fat and muscle interface); and (d) with polarization control (fat and muscle interface).

birefringence-matched case. This is shown in Fig. 7(d), where the clear interface between fat and muscle tissues can be seen compared to that in Fig. 7(c).

V. SIMPLE HIGH-RESOLUTION OCT USING MULTIPLE SLEDs

A. High-Resolution CP-OCT and Signal Processing

The main challenge of constructing an ultrahigh resolution OCT system is the dispersion compensation between reference and sample arm of the interferometer. This is because an extremely broadband source with a bandwidth larger than 100 nm is required for a system and there is a large amount of dispersion between the two arms as a result. Both hardware and software compensations have been applied to make the axial resolution approach the theoretical value [45]. The hardware compensation requires extra optical components and precise alignment that together, complicate the system. When OCT is used to image retina, the dispersion induced by the eye varies between the subjects, hence the system requires the ability to adjust compensation for each subject. In terms of software compensation [46] using an iterative algorithm is usually time consuming and hence is not applicable for real time processing.

The shared sample and reference arm in CP-OCT helps to circumvent the problem of dispersion mismatch, which makes it appropriate for high-resolution OCT imaging. Nevertheless, the CP-OCT does not compensate for the dispersion arises from the sample and the extent to which this affect CP-OCT is same as that in the Michelson interferometer-based OCT systems. We built an ultrahigh resolution FD-CP-OCT using three combined SLEDs [47] as the broadband source. There have been a number of reports that have used multiple SLEDs with different central wavelengths to increase the effective bandwidth and hence axial resolution. However, due to the phase and amplitude dissimilarities between the SLEDs, optimum imaging results have not been obtained when used in conventional OCT systems. To solve problems caused by the asymmetrical and non-Gaussian-shaped

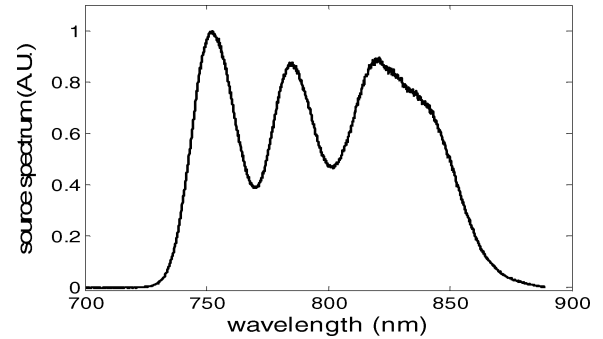


Fig. 8. Source spectrum by combining three SLEDs.

source spectrum, Wiener deconvolution [48], [49], and spectral shaping [50] were used to process the data. The resolution obtained in our three-source FD-CP-OCT without any dispersion compensation effort was $3.6 \mu\text{m}$ (in air).

The single SLED source FD-CP-OCT described in Section II was replaced with a broadband source made up of three multiplexed SLEDs to obtain higher axial resolution. By adjusting the ratio of output power of each source, we are able to optimize the overall spectral shape of the source. Fig. 8 shows the spectrum of the combined source, with full width half maximum (FWHM) of 106 nm and central wavelength of 800 nm. The broad bandwidth of the combined source results in a very narrow PSF, however, the ripples within the band would cause sidelobes in PSF that would blur the image.

In order to compare the differences in resolution, SNR and sidelobe suppression of the three signal-processing methods considered, we applied three signal-processing algorithms to extract the OCT signal from the interfered spectra. The methods compared include the following.

- 1) Standard signal-processing procedure for FD-OCT: the data is resampled to wave number space after reference power is subtracted from the interference data and fast Fourier transformation (FFT) is performed.
- 2) Spectral shaping: the spectra were shaped to have a Gaussian envelope $g(\lambda)$, which has the same central wavelength and the same e^{-2} bandwidth as source spectrum, $g(\lambda)$ centers at 800 nm and has FWHM bandwidth of 70 nm. This is followed by the processing described in 1) to obtain OCT image.
- 3) Wiener deconvolution: Wiener filter $W(k)$, designed to minimize the error between the restored and the original signal in the mean squared sense, is used to filter the interfered spectrum. $W(k) = S_0^*(k)/[|S_0(k)|^2 + \text{SNR}^{-1}]$, where $S_0(k)$ is the source spectrum, k is wave number $k = 2\pi/\lambda$. The resulting filtered data was analyzed by the procedure described in 1).

To characterize the resolution of our system and the different methods, we theoretically calculated the PSF by Fourier transforming the source spectrum $S_0(k)$, and experimentally measured the PSF by using a mirror as a sample and processed the data with the standard FD-OCT signal-processing procedure. The theoretical PSF has a $2.7 \mu\text{m}$ FWHM and the experimental PSF has a $3.6 \mu\text{m}$ FWHM. The spectra are also processed

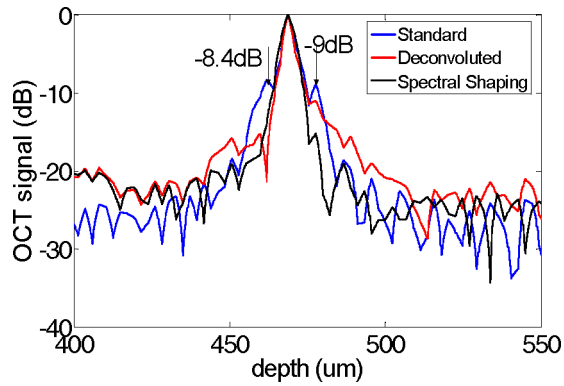


Fig. 9. Coherence function obtained from a mirror in air processed with different algorithms.

by the other two above-mentioned methods to obtain OCT A-scans. The results are plotted in Fig. 9, which clearly show that the sidelobes can be effectively suppressed by the deconvolution and spectral shaping methods used.

B. Results

Fig. 10 shows the images of onion obtained by different signal-processing methods. In addition, we also compared the A-scans obtained of onion by different methods in Fig. 10(d).

We also imaged a blood vessel in the chicken embryo chorioallantoic membrane (CAM), and the results are shown in Fig. 11. CAM is commonly used as a model tissue for surgical retinal research [51]. As a reference, Fig. 11(d) shows the histological cross section of a blood vessel at CAM [51].

Notably, Fig. 10(a)–(c) were obtained by using the same raw spectral data to form images, and so, were Fig. 11(a)–(c). These results demonstrated that Wiener deconvolution and spectral shaping can effectively suppress sidelobes in scattering samples other than the specular surface. However, as shown in Fig. 11, Wiener deconvolution and spectral shaping did not improve the quality of vessel images to the extent that the onion images were improved. Onion possesses clear imaging boundaries that are highly reflective, while the vessel in CAM does not have these optically favorable boundaries.

C. Discussion

In this paper, two signal-processing algorithms are applied to deal with the non-Gaussian shape of the source spectrum, in addition to the standard FD-OCT processing. Both the methods effectively suppress the blur caused by source spectrum modulation. The drawback of deconvolution is an increased noise level. The spectral-shaping method seems to provide a good balance between sidelobe suppression and SNR, but this comes at the expense of a small loss of resolution. The improvement in image quality also depends on the property of samples. A sample, such as onion, with high-reflectivity specular boundaries will have ghost signals if the data were analyzed directly by FFT. Wiener deconvolution and spectral shaping can improve this artifact and hence the image quality significantly. For samples lacking large changes in the reflectivity, such as the CAM vessel

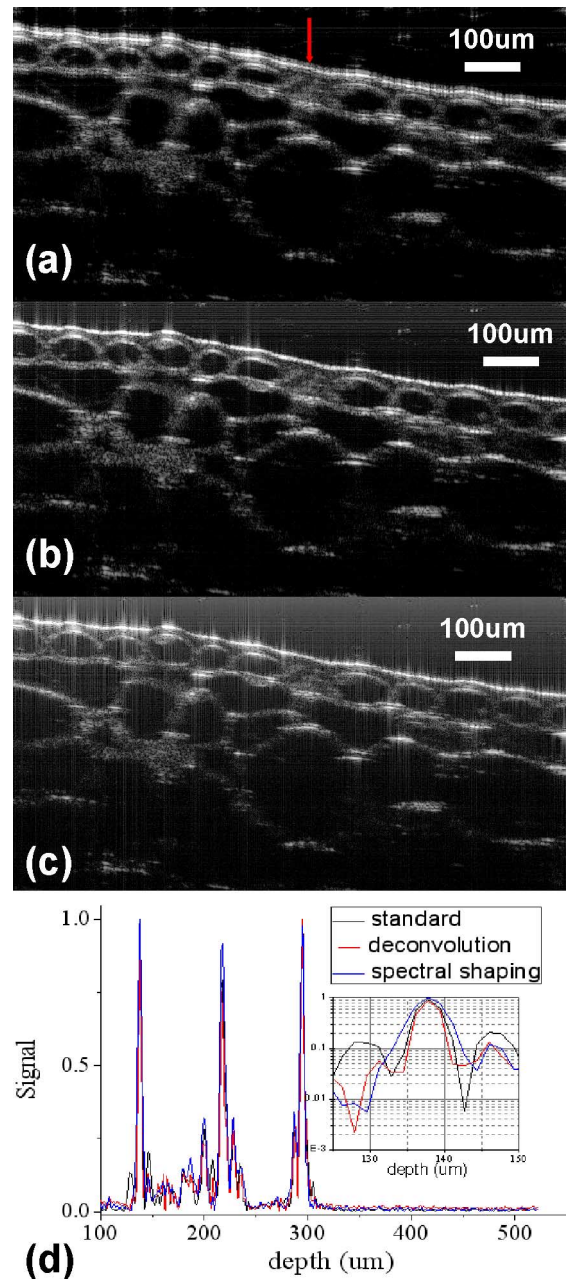


Fig. 10. OCT images of an onion. (a) Image obtained by standard signal-processing procedure for FD-OCT. (b) Image obtained by spectral shaping. (c) Image obtained by deconvolution. (d) Comparison of A-scan signal at lateral position is indicated by the arrow in (a).

that was imaged, standard processing procedures are sufficient to obtain high-resolution OCT images.

Whether or not the deconvolution is effective in improving the quality of OCT image is questionable. Nevertheless, our goal of using the deconvolution algorithm is to suppress the sidelobes caused by the ripples in the source spectrum, in order to better identify the highly reflective boundaries. This objective has been achieved as shown in Fig. 10(c). However, there are several problems that do exist. The prerequisite of using Wiener deconvolution is that the noise in the system is additive and the spectrum is known. However, as in any other coherent

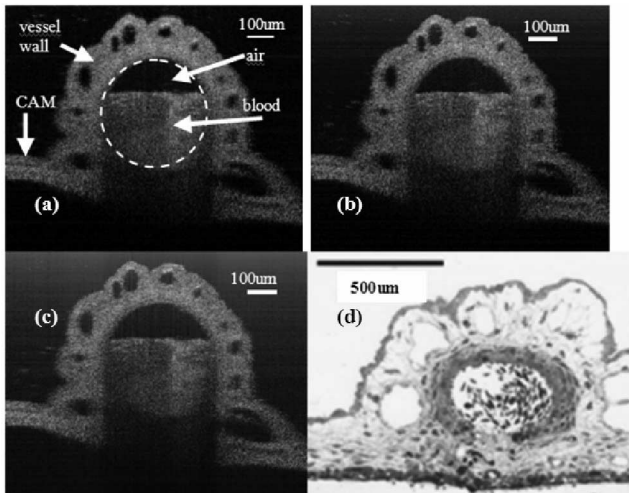


Fig. 11. OCT images of a vessel at CAM. (a) Image obtained by standard signal-processing procedure for FD-OCT. (b) Image obtained by spectral shaping. (c) Image obtained by deconvolution. (d) Histological cross section of a blood vessel at CAM.

imaging modalities, there exists speckle noise in OCT, which is not additive. Therefore, Fig. 10(c) still contains speckled noise. A speckle reduction algorithm [52] might further improve the image quality. Moreover, the frequency dependency of SNR is not clear, because the noise spectrum also depends on the sample structure, which is unknown in prior. For simplicity, we use the same estimated signal to noise ratio for the whole spectrum. In other words, we assumed the white noise, which is not completely accurate as discussed before. This is why we have a broadened pedestal in our deconvoluted image, shown as vertical lines in at the vicinity of bright spots.

There are two factors that allow our system to remain almost free of chromatic dispersion. First, the reference and sample arms share the same optical path in our FD-CP-OCT, therefore, there is no system-induced dispersion mismatch, which is a common problem in OCTs with separated reference and sample arms. The other reason is that the probe located in close proximity to the site imaged. This decreases the mismatch caused by the media present between the probe tip and the sample to a negligible level. Based on part in these two factors, our FD-CP-OCT system was able to approach the theoretical resolution limit in the absence of a physical compensation scheme. The residual difference between theoretical and experimental resolution is in part, most likely due to remaining calibration error in the spectrometer.

VI. FUNCTIONAL OCT SENSING

A. Introduction

Hemoglobin (Hb) oxygen saturation (SO_2) level, the ratio of oxyhemoglobin (HbO_2) to total Hb, is potentially a relevant indicator of tissue perfusion. However, traditional blood oxygen monitoring devices such as pulse oxymeters or blood gas analyzers are usually nonlocal or invasive. OCT as a high resolution, high-speed imaging modality, has the potential to measure lo-

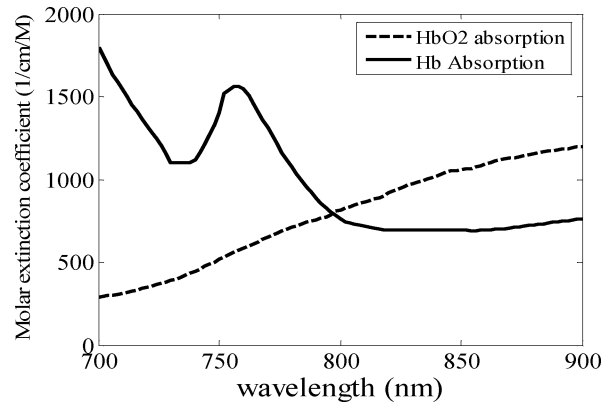


Fig. 12. Molar extinction coefficient spectra of Hb and HbO_2 .

calized blood oxygen saturation level via spectroscopic analysis of the sample's spectral features [53].

In [54], Faber *et al.* used a spectroscopic OCT technique to measure the absorption of nonscattering Hb and HbO_2 , by analyzing the spectra at the glass-liquid boundaries from A-scans of a time domain OCT, illuminated by a broadband Ti-sapphire laser source. An extension of this paper is reported in [55], the differential attenuation coefficients of whole-blood samples as a function of SO_2 were determined by the same method used in [54]. However, in *in vivo* measurements, there is often not a clear boundary between blood and other tissues. Lu *et al* reported using a spectroscopic spectral domain OCT to measure SO_2 [56]. To achieve this A-mode profiles were fit with exponential functions to derive the decay constants, at long (>800 nm) and short (<800 nm) wavelengths, respectively. The difference in decay constants was then used to determine the SO_2 level. *In vivo* measurements of the wavelength-dependent optical density of veins and arteries in human retina have been performed, with ultrahigh resolution OCT [57].

B. Theory and Data analysis

The underlying principle of oxygen saturation measurement with OCT is based on the differing light absorption properties of HbO_2 and Hb. As shown in Fig. 12, the absorption spectra of HbO_2 and Hb are different. HbO_2 absorbs more light at longer wavelength (>800 nm) and Hb absorbs more light at shorter wavelength (<800 nm). In order to derive the localized SO_2 , we need to know how much spectral distortion blood causes in the long and short wavelength range, for each specific optical path length. The detected OCT interference signal in spectral domain OCT is

$$I(k) = \eta S_0(k) \int r_s(k, l) \cos(j2kl) dl \quad (6)$$

where l is the delay between reference and sample light; k is wave vector ($k = 2\pi/\lambda$); η is a constant; $S_0(k)$ is the source spectrum; $r_s(k, l)$ is the wavelength-dependent path length resolved reflectance, which is used to extract SO_2 level.

Morlet wavelet Transformation [53] can be used to get $r_s(k, l)$.

First, an inverse Fourier transform was performed on $I(k)$ to get $i_0(l)$, which is the A-scan. Afterwards, Morlet wavelet transformation is applied to $i_0(l)$. And then the result is normalized by source spectrum $S_0(k)$, as described in (7)

$$\hat{r}_s(k, l) = \frac{\left| \mathbb{F} \left[i_0(l+x) \exp \left(-4 \ln 2 \frac{x^2}{L^2} \right) \right] \right|^2}{S_0^2(k)} \quad (7)$$

in which $\mathbb{F}[\]$ indicates to take Fourier transform. The Gaussian window size L determines the spatial resolution of $\hat{r}_s(k, l)$, which is $14 \mu\text{m}$. Accordingly, the spectral resolution of this analysis is about 20 nm , inversely proportional to $L \cdot \hat{r}_s(k, l)$ shows the sample reflectivity as a function of both wavelength and optical path length l .

Generally for a given k , $\hat{r}_s(k, l)$ decays as increasing l exponentially, with an attenuation coefficient, which depends on the scattering coefficient $\alpha_s(k)$, absorption coefficient $\alpha_a(k)$, as well as the measurement apparatus. $\hat{r}_s(k, l)$ can be expressed as

$$\hat{r}_s(k, l) = r_{s0} \exp \left\{ -2 \int_0^l [\alpha_s(k, l') + \alpha_a(k, l')] dl' \right\}. \quad (8)$$

For simplicity, we assume α_s does not depend on wavelength.

By averaging $\hat{r}_s(k, l)$ in long (808 and 848 nm) and short (743–766 nm) wavelength range, we have $\hat{r}_{s_long}(l)$ and $\hat{r}_{s_short}(l)$. They both depend on oxygen saturation level, because the blood absorption coefficient $\alpha_a(k, l)$ is a function of SO_2

$$\alpha_a(k, l) = [\text{SO}_2(l)] \alpha_{\text{HbO}_2}(k) + [1 - \text{SO}_2(l)] \alpha_{\text{Hb}}(k) \quad (9)$$

where α_{HbO_2} is the wavelength dependent absorption coefficient of HbO_2 and α_{Hb} is the absorption coefficient of Hb.

To get the difference between $\hat{r}_{s_short}(l)$ and $\hat{r}_{s_long}(l)$, we defined $R(l)$ as following:

$$R(l) = \frac{\hat{r}_{s_short}(l) - \hat{r}_{s_long}(l)}{\hat{r}_{s_short}(l) + \hat{r}_{s_long}(l)}. \quad (10)$$

The difference between $\hat{r}_{s_short}(l)$ and $\hat{r}_{s_long}(l)$ is assumed to be much smaller than $\hat{r}_{s_short}(l)$ and $\hat{r}_{s_long}(l)$. In (10), factors independent of wavelength cancels out, hence $R(l)$ reveals the absorption difference between long and short wavelength light, which could be considered as depth resolved relative SO_2 . Notably, the spectral distortion in (8) is cumulative. In other words, $\hat{r}_s(k, l)$ is not only determined by light absorption in the vicinity of l , but also determined by absorption from the end of the probe to l . This makes it very difficult to explain the straightforward physical meaning of $R(l)$.

The setup of our experiment has been described in Section V. Data measured from a vessel in the CAM of a chicken embryo is used to extract SO_2 information. The relative SO_2 image is formed by laterally scanning the fiber probe above the surface of the sample to get $R(l)$ at different lateral locations.

C. Result

Fig. 13(a) shows the spatial variation of SO_2 . The value of $R(l_x, l_z)$, is converted into the range of $[0 \ 1]$ and is assigned to every pixel as gray level. Pixels with larger brightness correspond to larger R -value. In Fig. 13(b), we fuse SO_2 image and

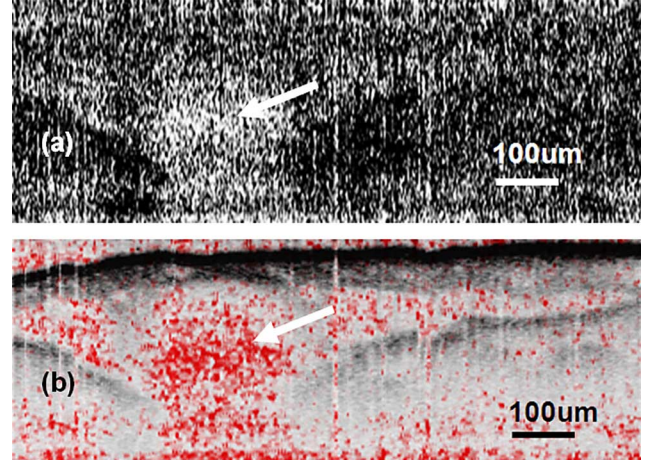


Fig. 13. (a) SO_2 image (b) overlapped SO_2 image and OCT image.

OCT image together as described in Fig. 13(a). Hue, saturation, value (HSV) color space instead of red green blue (RGB) space was used. H (hue) component is set as constant 0, which is red. S (saturation) is assigned with values of R of that pixel. When R is small, the colors are unsaturated and are simply shades of gray. When R is large, it becomes fully saturated red. V (value or brightness) is assigned by OCT signal amplitude. In Fig. 13(b), area enclosed by the vessel which is marked with an arrow has larger R -value compared with other part of the image. There is distinct difference in R for blood and other tissue. This verifies that our spectroscopic analysis correctly reveals oxygen saturation variation in living tissue samples.

D. Discussion

2-D relative SO_2 imaging is presented in this paper. As in (3), the depth–wavelength resolved reflectance is a very complicated function. Therefore, calibration is a challenging task. First, an exponential decay assumption does not necessarily hold because of the random nature of scattering events in turbid tissue. Second, as previously noted the spectral distortion at depth l is determined by light absorption from the end of the probe to l , which is cumulative. Nevertheless, we have measured highly localized relative hemoglobin oxygen saturation levels in the chicken embryo, based on the SO_2 dependent absorption spectrum for blood around 800 nm. Here, we have presented a 2-D SO_2 image obtained from OCT measurements for the first time. In doing so, we have verified the ability of FD-CP-OCT to distinguish between saturated and unsaturated hemoglobin.

VII. OCT-GUIDED MOTION CONTROL AND COMPENSATION

A. Motivation

Traditional intraocular microsurgery such as vitreoretinal surgery is guided by binocular surgical microscopes that provide 3-D view, however, the surgeon's operating view of intraocular tissues such as retinal is largely limited to the en-face level and retinal surface detail. OCT imaging has the potential to extend the imaging field to the in-depth level and is, thus, a

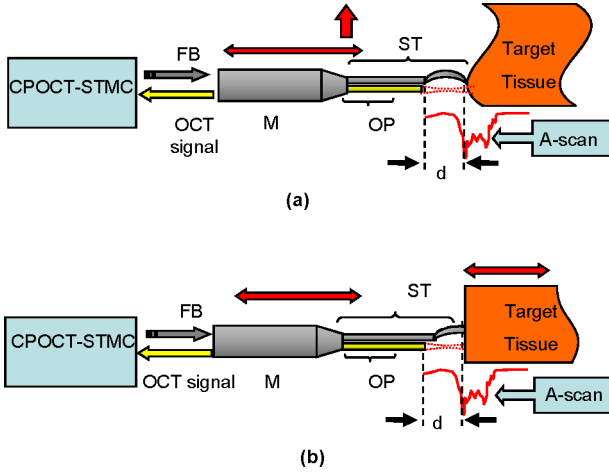


Fig. 14. Schematic representation of CPOCT-STMC system. (a) Topology compensation. (b) Motion compensation (here, we mainly consider the axial motion). ST: surgical tool; OP: CP-OCT probe; M: mechanical module; FB: feedback control signal.

promising new modality of intraoperative microsurgical guidance that may lead to enhanced surgical abilities, less iatrogenic injury and better patient outcomes [16]. An advantage of OCT as compared to other potential intraocular imaging modalities is that it is more compact and has a much higher resolution. These are potential advantages for integration into microsurgical tools.

Conventional OCT systems have only 1–3 mm coherence-gate-limited imaging depth that restricts its practical applications when the target tissue surface has a large topological variance [2]. Moreover, OCT imaging artifacts resulting from unavoidable motion inherent to the surgical tissue (e.g., vascular pulsation or respiratory induced motion) could misguide surgical efforts if these artifacts are not accounted for [58], therefore, at the present time an experienced and highly dexterous surgeon is required in order to avoid related complications. A simple and efficient solution for these issues is an adaptive ranging technique based on depth-tracking, which first locates the subject tissue surface by OCT imaging and then analyzes the result for depth information in order to adjust the reference arm [59], [60]. By these methods the topological variance and motion effect can be compensated for and the surgical procedure can proceed on a virtual “plain and static” surface. The schematic representation of the CP-OCT surface topology and motion compensation (STMC) system is presented in Fig. 14. A CPOCT fiber-optic probe is integrated into a surgical tool like needle or microtweezers, which is driven by mechanical module such as a computer-controlled actuator or robotic arm. Every A-scan measurement is input into the CPOCT-STMC system and analyzed for the surface position. The system then feeds back the control signal to correct the tool’s position.

CP-OCT is a nearly ideal approach for these types of compensation because the reference and sample signals share the same path, thus, the reference offset can be changed by adjusting the probe position without synchronizing the reference arm.

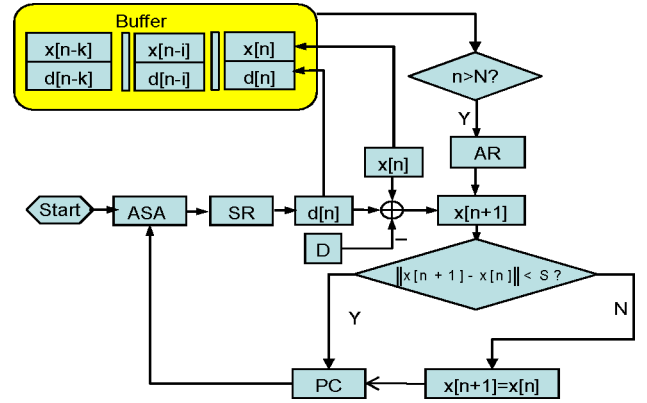


Fig. 15. CPOCT-STMC flowchart. ASA: A-scan acquisition; SR: surface recognition; AR: autoregressive predictor; PC: probe control.

B. Schematic of CP-OCT STMC

Fig. 15 shows the CPOCT-STMC flowchart while the experimental setup is mainly based on Fig. 14. The CP-OCT probe is driven by high-precision motorized actuators (Newport 850G). The whole system is controlled and coordinated by LabView program through general-purpose interface bus (GPIB) control. The probe–sample distance is set to a fixed value of D . Each A-scan is processed by a depth-tracking algorithm to find the actual probe–sample distance, denoted as $d[n]$. In this setup, there is a potential error caused by random scattering particles suspended between probe and sample, so rogue jumps can happen. To eliminate these unwanted events, a temporal comparison critic is applied prior to starting the actuator. This is done by defining a safe distance S , which is pre-estimated, according to the observed properties of the specific target’s topology and motion as well as the temporal response of the mechanical module. The k -order autoregressive (AR) predictor is applied to actively track the rhythmic motion of the tissue target [61]

$$x[n+1] = \sum_{i=0}^{k-1} \alpha_i (y[n-i]) - D$$

$$= \sum_{i=0}^{k-1} \alpha_i (d[n-i] + x[n-i]) - D \quad (11)$$

where $y[n-i] = d[n-i] + x[n-i]$ is the target position of the tissue. Before starting the predictive tracking, k values of $y[n]$ is acquired and registered in a memory buffer, which is then used to determine the model coefficients α_i by the least-square regression. AR tracking starts at a preset step number N ($N > k$). The B-scan image is reconstructed, according to the probe position for each A-scan.

C. Topological Compensation

To evaluate the topological compensation property, we made a phantom sample possessing highly curved surfaces consisting of eight-layers semitransparent tape. Fig. 16(a) is obtained by conventional fixed-reference OCT while Fig. 16(b) is obtained by CPOCT-STMC system. The motion trace of the

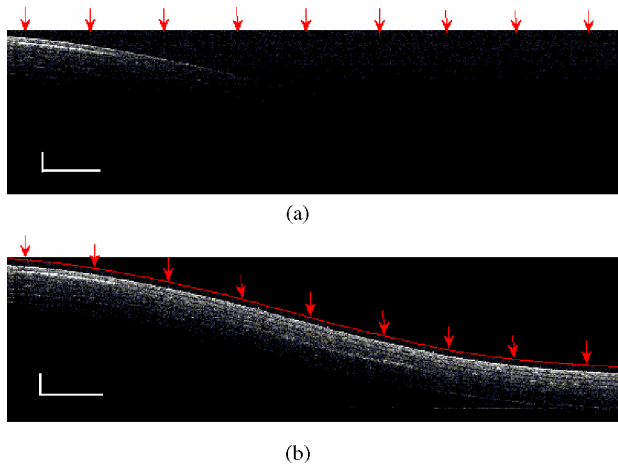


Fig. 16. Images of a phantom sample by CPOCT-STMC system. (a) Traditional OCT with limited imaging depth. (b) Topological compensation with extended imaging depth. The bar indicates $500 \mu\text{m}$.

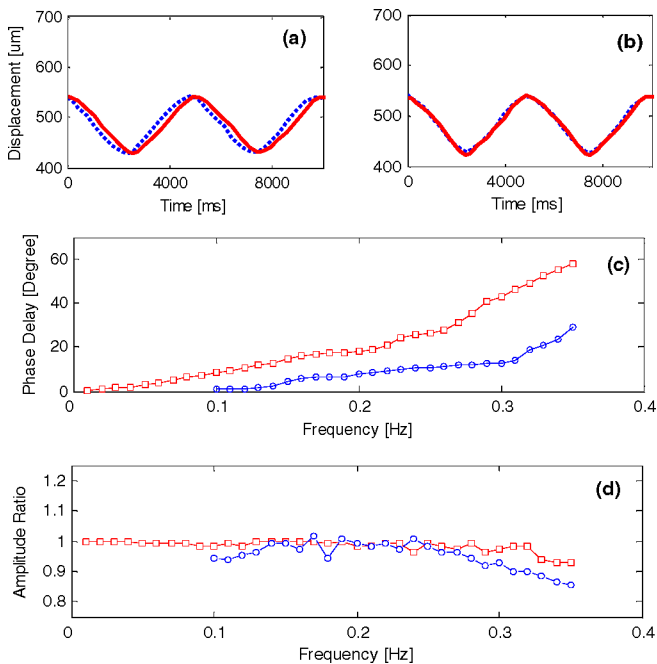


Fig. 17. CPOCT-STMC Motion compensation properties. (a) and (b) Response to a target motion without/with AR predictor. Dotted line: target motion; solid line: system response. (c) and (d) Phase difference and amplitude response to rhythmic motion. Square mark: without predictor; circle mark: with predictor.

scanning probe is indicated by the arrows. The conventional OCT has a working depth of $\sim 1 \text{ mm}$, therefore, the image fades with the imaging depth. In the STMC mode, the scanning probe can penetrate the surface geometry while acquiring useful A-scan information at significantly greater depths. The solid line in Fig. 15(b) shows the moving trace of the probe, which correlates with the surface geometry very well.

D. Motion Compensation

Fig. 17 presents the testing of SMTC temporal response to periodic motion where a phantom sample is periodically moving

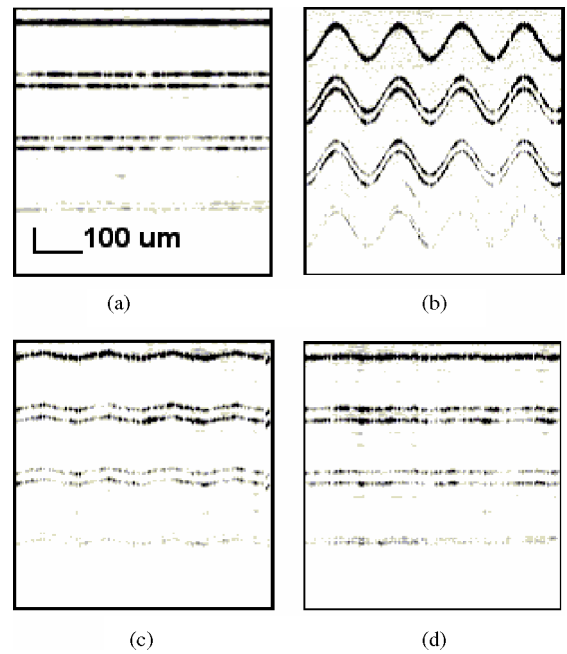


Fig. 18. Motion compensation of the CPOCT-STMC system. (a) and (b) Traditional B-scan image of static/periodical moving phantom. (c) and (d) STMC B-scan image of periodically moving phantom without/with AR predictor.

back and forth in the axial direction at an amplitude of $110 \mu\text{m}$. The period of target motion is set to be 5 s due to the limited actuator speed ($500 \mu\text{m/s}$). Apparent phase delay of approximately 20° between the response and the motion can be seen in Fig. 17(a). When AR predictor is applied, as in Fig. 17(b), this phase delay is significantly decreased to about 5° . The advantages of AR predictor is further investigated in Fig. 17(c) by slowing down the phase delay increase with frequency. A faster mechanical module can improve the STMC response speed to capture real tissue motion in real time. Fig. 18 illustrates the phase difference effect on the image. A phantom sample is made of three semitransparent layers and scanned by a conventional OCT probe while static and while moving at a 10-s period. From Fig. 18(b), we can see a fluctuating surface that is apparently artifact caused by sample motion. This is a critical problem for microsurgery as stated earlier. Fig. 18(c) and (d) show the results of re-imaging by the STMC system with diminished artifact signal.

VIII. CONCLUSION

We have developed a simple all-fiber endoscopic FD-CP-OCT for biological imaging and sensing, which is eventually targeted for the use in the setting of delicate microsurgical procedures such as intraocular retinal surgery. We have demonstrated that this system can achieve extremely high axial resolution $< 3 \mu\text{m}$ without any external dispersion compensation. We have achieved this simply by combining multiple SLEDs using cascaded fiber optic couplers. We have demonstrated that its probe can be easily integrated into surgical tools that provide high resolution *in vivo* imaging and can be used in aqueous media

similar to those of vitreous by optimizing a semitransparent, gold thin film layer with 1.7% reference reflectance. We have also developed a spectroscopic signal-processing method that allows measurement of relative spatial varying oxygen saturation level in tissue. The high spatial resolution of OCT ensures that this spectroscopic analysis is highly localized. Finally, we have successfully demonstrated the feasibility of this system for the assessment of motion and topological compensation.

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