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Abstract. Quantitative phase microscopy is applied to image temporal changes in the refractive index (RI) distributions of solutions created by microbicidal films undergoing hydration. We present a novel method of using an engineered polydimethylsiloxane structure as a static phase reference to facilitate calibration of the absolute RI across the entire field. We present a study of dynamic structural changes in microbicidal films during hydration and subsequent dissolution. With assumptions about the smoothness of the phase changes induced by these films, we calculate absolute changes in the percentage of film in regions across the field of view. © 2011 Society of Photo-Optical Instrumentation Engineers (SPIE). [DOI: 10.1117/1.3665439]

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Microbicides are topical products that are designed to reduce the risk of HIV/AIDS transmission and can be deployed in vivo using various delivery forms, such as films, gels, tablets, ring devices, or injections. 1 Although each of these dosage forms has advantages and drawbacks, they are all designed to provide discreet protection from HIV that young women, who are 3–5× more likely to be infected than men, can use independently and proactively.1 Recently, thin polymeric films have been evaluated as delivery vehicles for microbicidal active pharmaceutical ingredients (APIs).2 Microbicidal films are homogeneous sheets ranging from tens to hundreds of microns in thickness and typically contain APIs, water soluble polymers, and plasticizers (e.g., cholesterol, ethanol, carbopol, etc.).2 Depending on their composition, these films can exhibit highly variable hydration profiles and dissolution dynamics. Because these films are designed to prevent aqueous degradation of the API by hydrating only after topical application, the parameters that characterize API release include inherent moisture content, disintegration time, dynamic hydration rates, and drug-content uniformity. Current methods of film assessment include bulk physical, chemical, and mechanical testing on a macroscopic scale; however, microscopic investigation of film dissolution has not been previously conducted and could be used to inform the compositions of clinical microbicidal films.

Quantitative phase microscopy (QPM) is well adapted to imaging microbicidal film solutions. The method is based on recording the complex amplitude and phase of light that is transmitted through a sample with weak refractive index (RI) variations, which gives rise to relative optical path length changes. We have previously developed QPM as a tool for imaging various cell types and studying dynamic biological phenomena.3, 4 Other studies of imaging cell samples have sought to decouple optical thickness from RI in phase measurements using various approaches,5–8 however these methods and their assumptions are not necessarily applicable to studying microbicidal films that dissolve over time. Therefore, to enable effective studies of film dissolution dynamics, an alternative method of determining absolute RI is needed with a single measurement in time preferred.

In this letter, we present the novel method of using a sloped physical calibration structure in QPM imaging to facilitate accurate calculation of the absolute RI from measured optical path lengths. We use a set chamber thickness and a calibration standard, a polydimethylsiloxane (PDMS) ramp fabricated by soft lithography, in the field of view (FOV) to determine the absolute RI of the sample at every two-dimensional spatial location using only a single temporal acquisition. On the basis of the assumptions of smoothness in the phase variations over time and the final spatial distribution, the PDMS ramp allows straightforward calculation of the absolute RI without sensitivity to temporal phase drift. To demonstrate the utility of this technique, we use this novel time-resolved imaging refractometer to image temporal changes in the RI distributions of solutions created by microbicidal films undergoing hydration.

Quantitative phase maps are acquired using an interferometric microscopy system similar to that previously described in Ref. 9, but with slight modifications. In the present experiments, we use a 532-nm laser (LaserGlow Technologies, Toronto, Canada) for illumination. The sample consists of a flow chamber (FCS3 chamber, Bioptechs, Butler, Pennsylvania) with No. 1 coverslips as upper and lower surfaces at a fixed physical thickness determined by an intermediate spacer (250 μm nominal thickness), and inlet/outlet ports to allow fluid flow. Transmitted light through the sample and reference arms of the interferometer is collected with matched 10x objectives (“A-Plan” 0.25NA, Carl Zeiss Microscopy, Thornwood, New York) and recombined prior to being imaged by an f = 150 mm achromatic tube lens onto a CCD (CoolSNAP cf 1392 × 1040 pixels, Photometrics, Tuscon, Arizona). The use of identical objectives allows us to minimize curvature mismatch of the optical wavefronts; these are arranged to produce a linear off-axis interference pattern,
which may be optimized by axially and laterally adjusting the reference arm objective and its incident beam.

This QPM system records off-axis interferograms, which are processed to produce phase maps by spatially filtering to remove the complex conjugate and autocorrelation terms, and then recentering in Fourier space. Inverse Fourier transformation of the result yields the complex wavefront at the image plane. After taking the angle of the complex data, the resulting phase information is unwrapped using a combined temporal and spatial method to produce three-dimensional data sets free of 2π ambiguities. Under assumptions of temporal and spatial smoothness (i.e., the phase between two adjacent data points is not more than π), the following phase unwrapping algorithm is employed: (i) use a one-dimensional unwrapping algorithm to temporally unwrap the phase of each lateral pixel; (ii) use a two-dimensional quality map-guided algorithm to unwrap the phase of the last frame; and (iii) add the unwrapped phase from the last frame to each of the previous frames that have been temporally unwrapped in a pointwise fashion. This phase unwrapping method yields unambiguous phase images over an extended time-course experiment, but requires further analysis to determine the absolute RI.

Relative optical path delays, \( D_\phi(r) \), across a FOV as well as the average RI, \( \bar{n} \), are related to the measured relative phase shifts, \( \varphi_\phi \), as \( D_\phi(r) = \varphi_\phi(r) / \lambda = \bar{n}(r)d \), where \( d \) is the total thickness of the chamber. From this expression, the relative RI can be calculated. However, the measured phase only accounts for relative path differences across the FOV. One approach to obtaining the absolute RI is to introduce some known landmark into the FOV that can be used to aid unwrapping within the area of interest. As a reference, we used a molded polydimethylsiloxane (PDMS) ramp structure [Fig. 1(b)] with a maximum height that was matched to the sample chamber spacer thickness and a ramp length that almost spans the width of the interferometric microscope’s FOV. After being UV-epoxied to the lower coverslip of the chamber, the ramp makes contact with the up-
average water content of 7.33%; it also had an average refractive index of 1.4699, which was determined by decomposing the measured RI, \( n(\lambda) \), as a linear combination of the bulk film RI, \( n_f \), and the RI of water, \( n_{\text{H}_2\text{O}} \), which had been measured to be 1.4699 and 1.3351, respectively. Figure 2 shows the volume fraction percentages (% v/v) of film at three different 100-\( \mu \text{m}^2 \) regions of interest (ROIs), as indicated by the corresponding colored squares in Fig. 1(c)–1(e), and also for the entire FOV. Relatively fast changes occurred in the first 10 s of dissolution, and the system gradually progressed toward a more uniform distribution of film and water over the full 106 s time course. The total amount of film in the FOV also increased gradually during this process as shown by the solid black curve in Fig. 2. As seen from the changes in ROI 3, which is farthest from the film at initial hydration, the percent of film increased from 0 to 1.240%. Without the use of the PDMS calibration ramp, it would be impossible to decouple this change from the temporal phase drift of the optical system and both the average integral RI measurements and percent film calculations would be inaccurate. As defined by the standard deviations of the PDMS RI measurements described above, the absolute film percentages shown in Fig. 2 have an accuracy of 0.3 percentage points (pp) and a precision of 0.004 pp.

In summary, we have presented a novel method of time-resolved imaging refractometry based on QPM. Both two-dimensional spatial phase unwrapping and pointwise temporal phase unwrapping were combined to remove phase ambiguities from the complete data set. By using a set chamber thickness and a physical calibration ramp structure in the FOV, the system is immune to temporal drift and we are able to calculate not only relative but also absolute RI. These measurements are accurate to \( \sigma = 4.1 \times 10^{-4} \), and the relative changes in RI are bounded by a standard deviation of \( \sigma = 5.5 \times 10^{-6} \). This technique is a highly accurate measurement tool that can be used to image RI distributions as a function of time for a variety of transparent samples. It should be noted that while a frame rate of 10 fps was sufficient to investigate the dynamics of film dissolution in this study, it is also possible to use this method at higher speeds and it is only limited by the camera acquisition rate and illuminating power level. The spatial and temporal film dissolution profiles obtained with this method can be used to model the transport of APIs into tissue. We anticipate that this measurement approach will be useful for evaluating the diffusion characteristics of various film compositions and optimizing their overall effectiveness.

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