High-contrast multilayer imaging of biological organisms through dark-field digital refocusing

Ahmad Faridian
Giancarlo Pedrini
Wolfgang Osten
High-contrast multilayer imaging of biological organisms through dark-field digital refocusing

Ahmad Faridian, Giancarlo Pedrini, and Wolfgang Osten
Universität Stuttgart, Institut für Technische Optik, Pfaffenwaldring 9, D-70569 Stuttgart, Germany

Abstract. We have developed an imaging system to extract high contrast images from different layers of biological organisms. Utilizing a digital holographic approach, the system works without scanning through layers of the specimen. In dark-field illumination, scattered light has the main contribution in image formation, but in the case of coherent illumination, this creates a strong speckle noise that reduces the image quality. To remove this restriction, the specimen has been illuminated with various speckle-fields and a hologram has been recorded for each speckle-field. Each hologram has been analyzed separately and the corresponding intensity image has been reconstructed. The final image has been derived by averaging over the reconstructed images. A correlation approach has been utilized to determine the number of speckle-fields required to achieve a desired contrast and image quality. The reconstructed intensity images in different object layers are shown for different sea urchin larvae. Two multimedia files are attached to illustrate the process of digital focusing, © 2013 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.18.8.086009]

Keywords: digital holographic microscopy; dark-field microscopy; digital focusing; image reconstruction; three-dimensional imaging; living organism.

1 Introduction

Imaging through various layers of a biological specimen is a demanding technology in the field of biology and biomedicine. Many improvements have been done using scanning fluorescence techniques for three-dimensional (3-D) imaging of micro- and sub-millimeter organisms. Confocal fluorescence microscopy has been shown to be promising in creating a 3-D image of a specimen using point by point scanning through different stacks of the specimen.1 Other scanning techniques like 4pi microscopy,2 stimulated emission depletion microscopy3 and structured illumination microscopy4 have been developed to improve the lateral and/or axial resolution in cellular fluorescence microscopy. Different imaging approaches, like selective plane illumination microscopy5 and double-sided plane illumination6 were aimed at imaging larger organisms. Although these fluorescence methods have enhanced the resolution, they have some limitations, like being able to image only stained particles inside the organism. To investigate biological samples for which no marker or dye is introduced, or staining is not of interest for the given investigation, some label-free 3-D microscopy methods have been developed based on auto-fluorescence or scattering properties, like optical projection tomography7 and spontaneous Raman scattering.8 However, either the weak auto-fluorescence for different materials or the presence of pump background reduces the contrast in these methods. In coherent anti-Stokes Raman scattering microscopy,9 the signal yield is a few orders of magnitude higher than what can be obtained through the spontaneous Raman scattering process and it is also possible to remove the background using stimulated Raman scattering.10 Later on, the technique was improved by Ozeki et al.11 and made it possible to record high contrast 3-D images of cells. Dark-field microscopy was also shown to be an effective approach in contrast and resolution enhancement.12 Lee et al. took advantage of this approach and combined it with optical sectioning microscopy to achieve a high contrast 3-D image of a biological specimen.13 The dark-field approach was also shown to be promising in the contrast enhancement in optical coherence microscopy by suppressing low spatial frequencies and increasing the signal for high spatial frequencies.14 Nevertheless, like the fluorescence techniques, the mentioned 3-D methods need to scan over the sample, which makes them impractical in imaging many dynamic processes and moving organisms. Being a scan-less and also a label-free technique, digital holographic microscopy (DHM) can be used to extract the 3-D information of the organism using a single image (digital hologram).15–17 In this technique, the quantitative phase and amplitude information of the object wave-front can be digitally obtained along the depth of the object from the recorded hologram, which makes it possible to digitally focus on different layers of the specimen any time later and reconstruct a 3-D profile of the optical thickness of the organism using the quantitative phase value.18–20 A short-coherence light source21–23 and the spatial coherence effect24 have been utilized to reduce the coherent noise and improve the image contrast. Park et al. showed that the contrast can be enhanced in DHM using speckle-field illumination.25 Dubois et al.26 and Verpillat et al.27 both implemented dark-field approach in a digital holographic system to image and track nanoparticles. Because of having no background in dark-field imaging, the noise coming from the coherent light in the background is suppressed, leading to a significant contrast enhancement.

1086009-1 August 2013 • Vol. 18(8)
Unlike imaging nanoparticles, due to the speckle noise coming from the scattered light, performing dark-field DHM for imaging larger organisms seems to be challenging. In this paper, we present an imaging system utilizing various speckle-fields to overcome the speckle noise and enhance the contrast. Employing the presented system, a multilayer imaging of the organism has been performed without scanning the specimen using digital refocusing. Adult sea urchins of the species *Arbacia lixula* have been selected as a sample organism. To avoid movement of the larvae for the current investigation, they were fixed first with formaldehyde and then immersed in water.

2 Method

2.1 Dark-Field Microscopy

To achieve a high contrast image of the specimen, a dark-field approach has been utilized. Generally, in dark-field microscopy, the specimen is illuminated with a hollow cone of light and therefore, the zero diffraction order, which falls out of the view of the imaging objective, is absent in the image, leading to a dark background. Hence, the highly diffracted orders, being scattered from the fine structures of the specimen, make the main contribution in the image formation (Fig. 1). Collecting only the high diffraction orders, the system delivers a higher resolution. On the other hand, due to the absence of the bright background, the small structures are visible with an enhanced contrast.

2.2 Speckle-Field Illumination

In spite of transferring high spatial frequency, the scattered light creates a strong speckle noise in the case of using a coherent source for illumination. The created speckle pattern reduces the image quality and the high frequency information will be lost in the image. Therefore, implementing the dark-field approach in a coherent microscopic system seems to be challenging. We took advantage of the speckle noise itself to improve the image contrast. In this method, instead of using a uniform spot for illumination, the specimen is illuminated with a speckle-field (the experimental setup is illustrated in detail in the following subsection). Figure 2 represents the images of a given sea urchin larva recorded with different illumination conditions. A bright-field image of the larva is shown in Fig. 2(a), illuminated by a short coherence source (55 nm) with a wavelength of 405 nm. Figure 2(b) was obtained by illuminating the larva with the same light source as Fig. 2(a), but with 200 different speckle-fields and by averaging over the fields. Although a short coherence source has been utilized, Fig. 2(a) still suffers from the presence of coherence noise, which can be improved using a speckle-field illumination approach [Fig. 2(b)]. Figure 2(c) represents a dark-field image of the same larva illuminated by a 660 nm laser diode with a coherence length of >1 m. Due to the long coherence length, the speckle noise is dominant in the image, which is apparent in the enlarged section of Fig. 2(c). For the image shown in Fig. 2(d), the larva was illuminated in the dark-field mode with 200 speckle-fields and the final image has been derived by averaging over the fields. In the enlarged section of Figs. 2(c) and 2(d), one can clearly realize the contrast enhancement after illuminating with and averaging over various speckle-fields.

2.3 Dark-Field Digital Holographic Setup

Figure 3 represents a schematic of the dark-field digital holographic microscope. For dark-field illumination, a Nikon bright/dark-field objective with the magnification of 20× and the numerical aperture of 0.45 has been utilized. The objective has the ability to operate both in dark-field and bright-field modes. An off-axis holographic setup has been arranged for
the experiment. The blue colored (in the online version) beam path shows the capability of bright-field holographic microscopy. For simplicity, the reference beam path is only shown for the dark-field mode in the scheme. The illumination beam is redirected toward the dark-field objective using a ring-shaped mirror. The portion of the beam which passes through the center of the ring mirror is used as the reference beam. A pinhole is placed in the reference path to create a uniform reference wavefront. A laser diode with the wavelength of 660 nm has been selected for the dark-field illumination, as the biological tissue has a relatively higher scattering rate in this region of the visible spectrum. A short coherence (55 μm) laser diode operating at 405 nm was used for the bright-field mode. A SVCam-ECO charged coupled devices camera (SVS-VISTEK GmbH, Seefeld, Germany) with the pixel size of $3.45 \times 3.45 \mu m^2$ ($2448 \times 2050$ pixels) and the maximum frame rate of 10 fps captured the holograms. To create a speckle-field, a diffuser was placed in the dark-field illumination path. The diffuser was being rotated by a stepper motor to produce different fields for each capturing shot. A hole in the middle of the diffuser allows the passage of the central portion of the beam that goes to the reference path, without being altered by the diffuser.

3 Results and Discussion

3.1 Image Reconstruction

A dark-field hologram of a larva is shown in Fig. 4(a); the interference fringes can be seen in the magnified inset image. This hologram was taken using a single speckle-field illumination. An angular spectrum algorithm has been used to perform image reconstruction and digital refocusing. Figure 4(b) shows the Fourier transform (FT) of the given hologram. One of the lobes of the first-order FT was selected (highlighted by a square) to reconstruct the intensity image [Fig. 4(c)]. Because of using the diffuser, the specimen has a random phase distribution and due to the dark background, the rest of the image shows a noisy phase distribution [Fig. 4(d)]. In addition, because of the nature of dark-field microscopy (i.e., having no light without any specimen), it is not possible to take a reference hologram and compensate for the phase contribution of the optical elements. As a result, the derived phase does not provide us with valuable quantitative information and is only used for wave-front propagation and digital focusing. To extract the 3-D profile (optical thickness) of the specimen from the phase distribution, the bright-field mode can be used. The bright-field hologram of the larva and the reconstructed phase distribution are shown in Figs. 4(e) and 4(f), respectively.

3.2 Optimal Number of Speckle Fields for Real-Time Imaging

As previously mentioned, several holograms should be captured using different speckle fields to reconstruct a high contrast intensity image. To be able to apply the technique for the imaging of living organisms, the optimal number of speckle-fields that is required to achieve a desired contrast has been evaluated. For this purpose, 100 holograms have been recorded for a given larva and the correlation between images, obtained by averaging over the different number of speckle-fields, have been plotted in two different ways. In the first case, the correlation between the image obtained from 100 speckle-fields, and the image obtained
from $N$ number of fields, has been plotted [Fig. 5(a)]. From the inset figure of the plot, it can be seen that images obtained using more than 60 speckle fields have a correlation factor of more than 0.99 with the image derived from 100 fields. For the second case, the correlation between the image obtained from $N$ and $(N - 1)$ speckle-fields has been plotted [Fig. 5(b)]. After the 30th averaging, the correlation factor seems to follow a flat trend and, as shown in the inset figure, for 60 speckle-fields it reaches a value of 0.9996. It should be noted that to obtain a more accurate value of the correlation factor, the dark background of the images in which the specimen has no contribution, has been eliminated in the calculation. Otherwise, unrealistically high correlation factors would be obtained.

To better demonstrate the resolution and contrast enhancement, a negative United States Air Force (USAF) test target has been utilized and the corresponding images have been presented in Fig. 6(a). The finest structure of the target, which is highlighted and magnified in the subimages, has a resolution of 228 line-pairs/mm. The magnified subimages, placed below the main image, were obtained by averaging over a different number of speckle-fields, indicated below each subimage (SF, speckle-fields); including the image taken with normal illumination and introducing no speckle-field (first sub-image from bottom-left). (b) The plot shows the improvement of the signal-to-noise ratio, $S/N$, (right axis) and the reduction of speckle contrast, $C$, (left axis) by increasing the number of averaging speckle fields, $N$, which has been calculated for the given USAF target.

Summing over a given number of speckle-fields, the speckle contrast, $C$, and the signal-to-noise ratio, $S/N$, have been expressed by Goodman \(^{31}\) as,

$$C = \frac{\sigma}{\bar{I}}$$

$$\frac{S}{N} = \frac{1}{C}$$

in which $\sigma^2$ is the variance and $\bar{I}$ is the mean value of the total intensity. The plot, shown in Fig. 6(b), represents $C$ and $S/N$ versus the number of speckle-fields, $N$, which has been calculated for the image shown in Fig. 6(a).

To perform real-time imaging, the number of required holograms should be adapted to the movement speed of the specimen and the desired signal-to-noise ratio. Based on correlation factor and $S/N$ calculations, an optimal number of 60 speckle-fields or more is suggested to achieve a desirably high contrast. However, referring to Fig. 6, a fewer number of speckle-fields may be selected if the corresponding resolution and $S/N$ satisfies the requirement for a specific purpose of study. Selecting the given number of speckle-fields, $N$, the minimum required time to capture these holograms is limited to the frame rate of the camera, fps, by $N$/fps.

### 3.3 Digital Focusing

Due to the variant phase distribution of the diffuser for each speckle-field, the fringe pattern is different in each hologram. Therefore, it is not possible to combine all holograms at first and then perform the image reconstruction using a coherently superimposed hologram. Thus, each hologram should be analyzed separately and the corresponding intensity image should be reconstructed. The final image will be obtained by incoherent averaging over the reconstructed images. Figure 7 shows the superimposed reconstructed intensity image of the larva for
different focus planes, illuminated using 60 speckle-fields. Video 1 represents the digital refocusing process for this larva. In the video, the reconstructed images were obtained from 60 sequential holograms. Sequential frames have been reconstructed by propagating the wavefront for 1 mm for each frame toward infinity. The parameter $Z$ in the video represents the axial position of the object plane relative to the initially focused plane for which the hologram has been recorded. A thin lens equation has been applied to calculate the axial position. The separation of the layers cannot be distinguished. Figure 8 and Video 2 show another two neighboring larvae in the same medium. Like Fig. 7, this figure also represents different focus planes for the given larvae. It can be seen that in each subfigure, different parts of the larva skeleton, along with the internal structure of the specimen, is in focus that results in more accurate 3-D information of the skeleton.

4 Conclusion
In summary, we introduced a scan-less multilayer imaging of a biological organism. The high contrast images have been obtained by implementing a dark-field approach in a digital holographic microscope. The speckle noise has been removed...
by acquiring sequential holograms with various speckle-fields and incoherently superimposing the final reconstructed images. The presented speckle-field illumination technique has been shown to be practical in regaining the contrast, which is achievable in conventional dark-field microscopy. The 3-D information of the organism was extracted without moving the specimen or performing any other mechanical scanning, by only capturing sequential holograms and performing digital refocusing. The imaging process can be performed in a short time, utilizing a high speed camera that provides researchers with the opportunity to image all layers of a live specimen without being required to fix it, making it possible to investigate internal structural interactions and to track particles inside the organism with a high contrast. Unlike fluorescence or optical sectioning techniques, the out-of-focus planes of the specimen introduce a background and we are continuing our investigation by designing an opposed-view dark-field DHM system to derive more accurate 3-D information of the specimen and developing an algorithm to remove the background imposed by the out-of-focus planes.

**Acknowledgments**

The authors would like to acknowledge the support of the DFG-Deutsche Forschungsgemeinschaft (German Research Foundation) under grant No. OS 111/33-1. The authors would like to gratefully acknowledge Ms. Julia Maxi Kanold and Prof. Franz Brümmer at the Department of Zoology, Biological Institute, University of Stuttgart, to provide the biological samples.

**References**