Digital holographic microscopy for the three-dimensional dynamic analysis of in vitro cancer cell migration

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In oncology, combating the dispersal of tumor cells constitutes a clinical need that currently remains unsatisfied. Novel types of compounds with antimigratory actions are thus necessary, and their identification requires efficient in vitro tests. Many developments in the fields of videomicroscopy and image analysis have enabled considerable advances in cell tracking in the 2-D environments in which living cells can be easily observed. Unfortunately, the biological characteristics of cells migrating in a 2-D environment do not fit in with those of cells migrating in a 3-D environment, which corresponds to reality in the case of cancer patients. The quantitative analyses of cell locomotion in a 3-D environment can be carried out by means of wide-field (standard) fluorescence, with the subsequent deconvolution of the data series, or phase-contrast microscopy. Techniques for sample analysis in three dimensions also include confocal microscopy and the related two-photon and multiphoton techniques, optical projection tomography, and selective plane illumination microscopy (SPIM). Techniques also exist that make use of interferometry, such as optical coherence tomography and time-domain digital holography in reflection. The principle consists of using an optical source with a low temporal coherence effect to record the interferometric information between the beam reflected by the sample slice under investigation and the reference. Due to the reduced temporal coherence effect, the interference occurs when there is an accurate equalization between the object and the reference beams. The post-processing consists of keeping the interfering slice contribution while eliminating the noninterfering background. As these systems work in a reflection mode, the sectioning effect is high and reduces the temporal coherence. The interference capability makes imaging through turbid media that has been also investigated with a multiple-quantum-well photorefractive holographic camera possible.

These previously cited techniques use optical sectioning to eliminate out-of-focus disturbances. However, to monitor 3-D dynamic cell processes, optical sectioning requires the storage of a large number of microscope images acquired over time with accurate position changes along the optical axis. By overcoming these limitations, transmission digital holography microscopy (TDHM) is able to record the full 3-D sample information without scanning, thanks to a computer based in depth refocusing ability. A sample hologram recorded with a CCD camera is numerically reconstructed to provide a stack of slice images refocused at incremental depths. The main advantages are a reduction in the amount of data to be stored.

Abstract. Cancer cell motility and invasion are critical targets for anticancer therapeutics. Whereas in vitro models could be designed for rapid screening with a view to investigate these targets, careful consideration must be given to the construction of appropriate model systems. Most investigations focus on two-dimensional (2-D) assays despite the fact that increasing evidence suggests that migration across rigid and planar substrates fails to recapitulate in vivo behavior. In contrast, few systems enable three-dimensional (3-D) cell migration to be quantitatively analyzed. We previously developed a digital holographic microscope (DHM) working in transmission with a partially spatial coherence source. This configuration avoids the noise artifacts of laser illumination and makes possible the direct recording of information on the 3-D structure of samples consisting of multiple objects embedded in scattering media, such as cell cultures in matrix gels. The software driving our DHM system is equipped with a time-lapse ability that enables the 3-D trajectories of living cells to be reconstituted and quantitatively analyzed. © 2006 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.2357174]

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and the time required to record the full 3-D information. With the phase-stepping technique, a small number (typically four) of successive video frames must be recorded.\textsuperscript{17} Since holographic recording requires optical coherence between the beams, the usual approach involves the use of highly coherent laser sources.\textsuperscript{17–21} However, this type of illumination suffers from an important noise artifact that significantly restricts the TDHM performances. This complicates the analysis of the actual 3-D samples, which consist of multiple objects embedded in scattering media, such as living cells in a 3-D matrix gel. To the best of our knowledge, the use of digital holography for the analysis of living cells is limited to low cell density, cells cultured on planar and solid substrates, or possibly, in very thin layers of gel.\textsuperscript{19–21} In a transmission mode one way to reduce the coherent noise consists of using a partial spatial coherent source that must be distinguished from the partial temporal coherence that is useful in reflection.\textsuperscript{22} Only a few implementations of partially spatial coherent sources in transmission digital holographic microscopes were reported. Based on a work by Leith et al.,\textsuperscript{23} and Indebetouw and Klysubun report that the spatiotemporal digital holography approach benefits from the partial spatial coherent source for imaging through a turbid media.\textsuperscript{24} Although the transmission microscope mode is mentioned, the experimental demonstration is shown in a reflection case. The same authors implemented a transmission-type system where the beam is split after the sample and the reference beam filtered by a pinhole.\textsuperscript{25} However, the recording of the hologram requires that the sample be defocused to match the partial spatial coherence function on the detector.

The microscope used for this work benefits from a partially spatial coherent illumination and is originated from a previous implementation.\textsuperscript{26} The sample is placed inside a MachZehnder interferometer (Fig. 1) to remove the constraints on the sample location and to maximize the contrast ratio of the interferometric patterns. With respect to our previous configuration,\textsuperscript{26} optical condensers are placed inside the interferometer to provide more flexibility in adjusting the ob-

Fig. 1 Description of the device: (A): Light-emitting diode (LED), optical fiber (OF), lenses (L1 through L6), beamsplitters (BS1 and BS2), mirrors (M1 through M5), microscope lenses 10X (ML1 and ML2), optical flat (Sr), culture dish (So), and Hamamatsu Orca camera (CCD). The relative orientation of the two subsystem assemblies (SE1) and (SE2) (each holds a beamsplitter and a mirror) enable the optical paths in the two arms of the interferometer to be accurately equalized. M4 is mounted on a piezoelectric transducer to implement the four-frame phase-stepping technique. (B) A computer drives all the microscope functionalities. (C) The culture dish is placed in the thermoregulated chamber on the microscope focusing stage. The microscope is inverted and the specimen is observed through the bottom of the culture dish. (D) The scattering features of a collagen gel (shown with contrast-phase microscopy).
ject and the reference beams. The image degradation caused by the noise of the coherent laser source is eliminated (Figs. 2 and 3), and the quality of the image is much improved. A partial spatial coherent illumination also provides a low temporal coherence-like effect. In scattering media such as collagenous gels [Fig. 1(d)], this approach further reduces the noisy coherent contribution by selecting the ballistic photons for the interferometric information. Moreover, as the optical source is an LED of low temporal coherence, the selection of the ballistic photons is further improved. The system provides the optical phase and the intensity images on one plane of a sample, with a typical acquisition time of 1/4 s [Figs. 2(a) and 2(b)]. The resulting complex optical field is used to refocus the optical intensity and phase fields on parallel planes without any mechanical scanning or loss of time [Figs. 2(a), 2(c), and 3]. As the information needed to perform the refocusing is recorded on the single imaged plane by the CCD, the point spread function of the optical system is constant over the experimental volume, so that the digital holographic refocusing does not introduce a loss of information. The magnification corresponding to that of a standard optical microscope is about 100X, the lateral resolution computed on the optical field transmitted by the sample is re-

Fig. 2 Information provided by the digital holography microscopy (DHM) system. Images (343×343 μm, corresponding to 1/4 of the complete microscope field actually observed) obtained by means of our digital holographic microscope that show unstained HT-1080 fibrosarcoma cells cultured in a 3-D collagen gel. The image shown in (A) is refocused over a distance of 85 μm by means of digital holography reconstruction to achieve the focused cell image shown in (C). (B) The optical phase map corresponding to (A) and (D) the differential interference contrast (DIC) mode emulation obtained from the holographic data corresponding to (C).

The resolution of the optical thickness computed on the phase map is about 2 nm. This value is established by assessing the noise that occurs on the phase maps. As the full information on the optical field transmitted by the sample is recorded, it is possible to emulate standard optical microscopy modes, such as the DIC [see Fig. 2(d)]. This latter mode is successfully implemented thanks to the low noise level and is particularly useful in the observation of living cells by providing the scientists with a common tool of visualization.

To test our systems and to illustrate their performances (see Figs. 2 to 4) we carried out experiments with unstained HT-1080 fibrosarcoma cell suspensions added to a solution of 1.6 mg/ml collagen type 1 and a cell culture medium supplemented with fetal calf serum (10%). The cell-collagen mixture was transferred to a Starna® glass dish (Anadis Instruments, Malden, The Netherlands) [cf. Fig. 1(c)], onto which a very fine coat of cell-free collagen had previously been applied. After collagen polymerization, the cells were embedded in a 3-D collagen gel approximately 1500 μm thick. The dish, which was provided with a two-neck cover (Fig. 1(c)), was then filled with culture medium. After having closed the cover, we added culture medium via the cover outlets to pre-

Fig. 3 Refocusing ability of the DHM system. Examples of refocusing by the 4-D interactive software interface using the mouse wheel are shown in holograms (A1 to A4) and (B1 to B4) (171×171 μm).
vent the propagation of any free-surface vibration, which could disturb the accuracy of the interferometric measurements. The culture dish was then maintained at 37 deg in a thermoregulated chamber placed on the microscope stage. The software driving the DHM has a time-lapse capability and recorded the holographic images every 4 min for 48 h. A semiautomated software package enables an operator to perform a 4-D analysis of the cancer cell dynamics on the time-lapse images on the basis of three visualization modes, i.e., the bright field, the phase image, and the DIC modes. This C++ custom software enables us to determine a region of interest (ROI) and to displace it in the hologram. Inside this ROI, the operator can refocus, with the mouse wheel, the different cells embedded in the gel (Fig. 3), and the recording of the \((r, X, Y, Z)\) locations is carried out with a mouse click on a central point inside the cell body, a usual way to record cell locations in computer-assisted cell tracking experiments.\(^1\) Stacks of 4-D images allow for the observation of the cell displacement during the experiment.

Figure 4(a) illustrates 22 different cell trajectories established by this method over a period of 13 h. Figure 4(b) illustrates the quantitative characterization of these trajectories by means of two features. The first is the individual cell average speed (i.e., the total length of the trajectory divided by the period of observation), and the second is the largest linear distance covered by each cell from its initial location (at time \(t = 0\)) to the farthest point reached by the cell in its trajectory.\(^2\) In contrast to the cell speed, this latter feature is able to distinguish between cell trajectories consisting of many small movements (around the original cell location) and...
those presenting a few large displacements. This latter feature was previously demonstrated as a particularly robust measurement of cell displacements and was used to rank the different trajectories analyzed, as illustrated in Fig. 4(b). The 22 trajectories illustrated in Fig. 4(a) are thus numbered according to their rank reported in Fig. 4(b). The largest covered distances observed in this set range from 73 to 359 μm and the cell speeds range from 14 to 52 μm/h. A relatively small level of positive correlation was observed between these two features (Spearman ρ = 0.47, p = 0.03), meaning that the speedy cells have a slight tendency to cover larger distances than the slow ones. These data illustrate the level of heterogeneity encountered in the migratory behavior of an a priori homogeneous cell population in a 3-D matrix gel, which, in fact, mixes migrating and less migrating cells, in terms of speeds as well as covered distances.

In conclusion, efficient noninvasive methods are necessary to validly analyze cancer cell migration in a 3-D environment and the possible changes due to drug action. For this purpose, the TDHM system that we developed records, without any significant time distortion or loss of information, the 3-D structure of a sample that is a posteriori analyzed with appropriate software. The partially spatial coherence nature of the optical source provides high quality images, even in the case of light-scattering samples (e.g., cell cultures in thick gels). Accurate, local optical thickness measurements and standard phase-contrast imaging modes are available. TDHM is of particular use in studying marker-free living biological (transparent) specimens. Consequently this approach does not require the insertion of fluorescent probes into cells, which sensitize them to photodamage and thus facilitates the long-term time-lapse imaging of living cells. Because of these advantages, DHM makes possible large-scale cell culture analyses in 3-D gels in the experimental conditions commonly encountered in cell biology laboratories. Since the magnification can be modified by using different microscope lenses, detailed cell morphology and its dynamical changes can also be monitored.

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