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# 3D localization and tracking of gold particles in biological environment using digital holography.

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**Abstract:** By using the dark field holographic microscopy technique described in Opt. Express, 19 pp 26044-26055 (2011) we have track gold particles in brownian motion in living cells. First results are presented. The particles remain confined to a region a few microns in size. We interpret this as a cage effect related to the cell structure.

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## 1. Introduction

The study of cellular processes at the single-molecule level is a flourishing field of research in Biology. Individual molecules labeled with sub-micron markers can now be tracked in a cellular environment, and quantitative information about their dynamics can be obtained by reconstructing their trajectory. One of the most used techniques for this purpose is single-molecule fluorescence microscopy (SMFM), but standard SMFM provides no information on the axial position of the marker, limiting this technique to 2D tracking. Digital Holographic Microscopy (DHM) [1] of gold nano particles circumvents this drawback. Moreover gold nano particles are not toxic for cells, and can be used as markers in biology [2]. As the scattering cross section of a particle scales as the sixth power of its radius [3], how easily and accurately a particle can be detected strongly depends on its size. Micron-sized colloids or particles can be easily detected by DHM in transmission geometry [4] or by In Line Holography [5], which is able to localize and track in 3D micron-sized latex particles with nanometer accuracy [6].

Since the signal is more than three orders of magnitude lower, Nano particles detection requires DHM with dark field illumination scheme. Atlan et al. [7] and Warnasooriya et al. [8] uses for example Total Internal Reflection (TIR) on a prism to detect and localize  $d = 50$  nm and  $d = 40$  nm particles, which are respectively spin coated on glass slide or attached to a cell membrane. The Atlan and Warnasooriya TIR configuration does not allow to track moving particles, since it yields a standing wave: when a moving particle crosses a node, the illumination (and thus the signal) goes down to zero, and the tracked particle is lost. More recently, Verpillat et al., which use a standing wave free dark field illumination scheme, localize and track  $d = 100$  nm particles in brownian motion in water [9].

By using the Verpillat method, we have track gold particles within HeLa cells. We present here our first results. The particle diameter is  $d = 50$  nm, and the random motion of the particle remains confined to a region a few microns in size. We interpret this as a cage effect related to the cell cytoskeleton.

## 2. Digital holography setup

The DHM setup is shown on Fig.1. The motion of  $d = 50$  nm gold particles in HeLa cells is investigated. The particles are brought into the cell by Pinocytosis [10, 11] with initial particles concentration  $10^{-9}$  m/l. The laser (Crystal Laser:

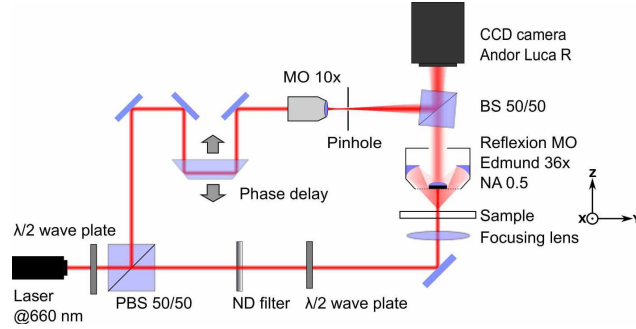


Fig. 1. Experimental setup. The sample is located in the X,Y plane. Z is the optical axis of the microscope objective.

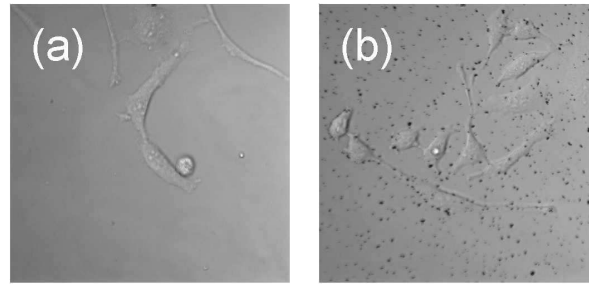


Fig. 2. White light images of HeLa cells without (a) and with (b) particles.

Diode Pump Solid State laser, 660 nm, 0.6 mm coherence length) is split into two beams by a Polarizing Beam Splitter (PBS), a half-wave plate PBS setting the beams energy ratio. The reference beam passes through a dove prism fixed on a micrometer translation stage to adjust the length of the optical path. It is spatially filtered through a  $35\ \mu\text{m}$  diameter pinhole and expanded as to uniformly cover the CCD chip of the camera (Andor Luca R:  $512 \times 512$  pixels, 22.5 Hz). The illumination beam is focused on the sample (waist diameter  $200\ \mu\text{m}$ ,  $250\ \text{W}/\text{cm}^2$ ). The light scattered by the particles is collected in transmission with a dark-field reflecting objective (Edmund Optics; ReflX series:  $\text{NA}=0.5$ ,  $36\times$ ). A small mask on the input of the objective limits the collection of light between  $\text{NA}=0.2$  and  $\text{NA}=0.5$ , so the illumination beam is totally blocked after passing through the sample. This dark-field configuration prevents the saturation of the CCD chip. Since the illumination is along the optical axis  $z$ , standing waves are minimized. A non-polarizing 50/50 beam splitter (BS) behind the objective combines the scattered light with the reference beam. BS is tilted by few degrees to be in off-axis holographic configuration.

### 3. Results and conclusion

Figure 2 shows the results we have obtained. The details of the holographic reconstruction, localization and tracking calculations are given in [9]. Figure 2 (a,b) show white light images of HeLa cells without and with gold particles. The particles are labeled with streptavidin, and many particles aggregates. Most of the black points of Fig.2 (b) correspond thus to aggregates, which are attached to the sample coverslip. Figure 3 (a) shows an holographic reconstructed images of the sample displayed in arbitrary log scale for the optical field intensity. Most of the bright point corresponds to particles or aggregates attached to the coverslip. They do not move. Some of the bright point like the one marked by the white arrow move slightly. They correspond particles in motion within the cell. We have track the particle marked by the arrow, and we have plot its 3D motion on Fig.3 (b). Since there is no standing waves, the particles can be tracked

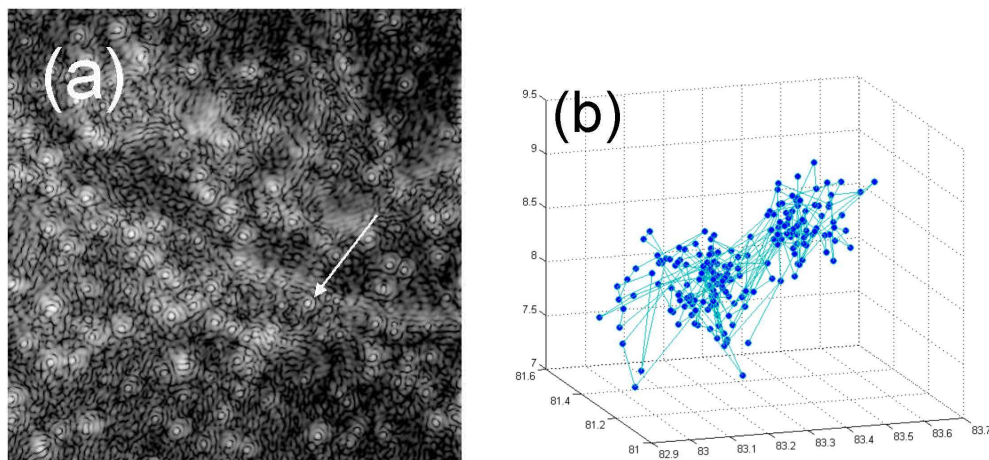


Fig. 3. Reconstructed holographic images of HeLa cells with  $d = 50$  nm gold particles (a). Plot of the 3D motion of the particle marked by the white arrow (b). x,y and z coordinates are in  $\mu\text{m}$  Units.

for long sequences of images (400 images or steps). For clarity, we have displayed only 200 steps of the motion on Fig.3 (b).

As it can be seen, the particle in motion remains located in confined region of the 3D space. We interpret this as a cage effect related to the cell cytoskeleton. It could be interesting to analyze in detail this effect, which should depend on the particle diameter. This preliminary work shows that it is possible to track  $d = 50$  nm gold nanoparticles in brownian diffusion motion within a biological sample, and to get by the way biological information.

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