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We report a multi-perspective scanning microscope based on the Talbot effect of a periodic focal spot array. Talbot illumination decouples the lateral scanning and the focal spots tuning. Large field of view fluorescence Talbot Microscope has been demonstrated by globally changing the incident wavefront gradient. Here, we explore the design freedom of adjusting the wavefront locally within each period and thus engineer the point spread function of the focal spots. We demonstrate an imaging system capable of reconstructing multi-perspective microscopic images in both bright field and fluorescence mode. With the multi-perspective imaging capability, we envision a more robust microscopic imaging system for large field of view fluorescence microscopy applications. This method is also suitable for compact imaging systems for multi-layer microfluidic systems.

Since its first discovery\textsuperscript{1} and explanation by Lord Rayleigh in diffraction theory,\textsuperscript{2} self-imaging phenomena of periodic structures, known as the Talbot effect, have been extensively investigated under various circumstances\textsuperscript{3,4} and in different wavelength regimes. Applications of the Talbot effect have been explored in precision metrology,\textsuperscript{5,6} spectroscopy,\textsuperscript{7} illumination design,\textsuperscript{8,9} and x-ray phase imaging.\textsuperscript{10} Recently, the self-imaging effect has been demonstrated in large field of view fluorescence microscopy, termed Fluorescence Talbot Microscope.\textsuperscript{11} Different from other scanning microscope setup where the sample is directly scanned by the focal spots of a MicroLens Array (MLA),\textsuperscript{12} Fluorescence Talbot Microscope scans the sample at Talbot distance away from the original focal spots, resulting in a longer working distance and an improved intensity uniformity of the focal spots. More importantly, the phase sensitivity of the Talbot effect decouples the scanning of the focal spot array and the modification of the intensity profile of individual focal spot. By generating a slight gradient to the global incident wavefront in the Fluorescence Talbot Microscope setup, we can scan the spot array for a significant distance without introducing off-axis aberrations, as illustrated in Fig. 1(a).

Different from changing the global wavefront gradient, changing the wavefront locally within each period provides another degree of freedom of design to engineer the point spread function of the focal spots, which can improve the performance of the scanning microscope. Hulskin et al. optimized the local wavefront to compensate for the higher order propagation phase term and achieved a high resolution Talbot focal spots array.\textsuperscript{13} The application is also focused on large field of view microscopic imaging. In fact, for large field of view microscopy applications, calibration and alignment are major challenges: a small tilt of the sample plane or microlens array can lead to a substantial defocus of the image. Here, we demonstrate a method to generate Talbot images in multiple perspectives by changing the local wavefront. With multi-perspective views of the sample, we could numerically compensate the defocusing through post-processing, which is similar to the concept of plenoptic microscope,\textsuperscript{14} and thus leads to a more robust large field of view system.

Talbot effect, in essence, is a two-dimensional periodic wavefront repeating itself periodically along the wave propagation direction. This period in the propagation direction is called the Talbot distance, $D_T$

$$D_T = \frac{2d^2}{\lambda}, \quad (1)$$

where $d$ is the period of the wavefront, and $\lambda$ is the wavelength of the light. In the setups where the period is $\sim$100 $\mu$m and wavelength is $\sim$0.5 $\mu$m, the working distance is $\sim$40 mm, which is greater than the working distance of a scanning microscope with an objective with similar Numerical Aperture (NA). With a conventional microscope objective, changing the wavefront gradient at the back focal plane shifts the imaging perspective and the field of view simultaneously. As mentioned earlier, in Talbot illumination set-up, we can adjust the global and local wavefront separately to scan the focal spots and change the imaging perspective, respectively. As shown in Fig. 1(b), the local wavefront can be adjusted by selectively illuminating the different regions of individual microlens. Shifting the illumination in the Fourier plane generates a “tilt” of the focal spots, which is equivalent to a shift in the scanning image perspective.

Our setup is based on the Talbot microscope system described previously\textsuperscript{15} and is shown in Fig. 1(c). The microlens array (SUSS MicroOptics) has a pitch of 100 $\mu$m, and each microlens has a numerical aperture of 0.17. The local wavefront is changed by shifting a binary transmission aperture mask (HTA photomask) mounted on a motorized translation stage (LHA-HS, Newport). The Talbot image of the aperture mask is optically relayed to the microlens array. The diameter of individual aperture on the mask is 50 $\mu$m, and the period of the aperture array is 100 $\mu$m, matching that of the microlens array. The aperture size of the mask can be selected according to applications: smaller aperture diameter increases the focal depth and provides higher angular resolution, but with compromise in spatial resolution.
This trade-off is similar to that of the light field microscopy.\textsuperscript{14} We use a 488 nm laser (OBIS, Coherent) as the source. The sample is placed at half Talbot distance, 20.5 mm, from the focal plane of the microlens array. To scan the sample, the global wavefront is adjusted by a piezo-driven mirror (AG-M100L, Newport). The detection system consists of an objective (4X, NA = 0.13, Nikon), a tube lens (f = 150 mm, Newport), and a camera (Manta G-145B, AVT). It is worth noting that the spatial resolution of the system is determined by the illumination from the microlens array, and thus the objective in the detection system does not necessarily limit the field of view. Here, the field of view is 1.69 mm by 1.43 mm. A field of view as large as 15 mm has been demonstrated previously.\textsuperscript{15} The sample is a two-layer patterned photoresist. SU-8 (MicroChem) is patterned on both side of a microscope cover slip. The height of the pattern is 15 \mu m, and the thickness of the cover slip are 155 \mu m. The pattern of the sample is an array of cylinders, each having 20 \mu m diameter, and the period of the pattern is 40 \mu m. The bright field image of the sample is shown in Fig. 2(a) by a conventional microscope objective (10X, NA = 0.3, Nikon).

Talbot Microscope images the sample from 5 successive perspectives by translating the aperture mask. The translation step size is 12 \mu m. The reconstructed images of these five perspectives of the two-layer sample are shown in Figs. 2(c1)–2(c5). The dark boundaries of the cylinders can be clearly observed due to the scattering at the edges. The total shift of the relative location of patterns on two layers is 11.1 \mu m.

We further quantify the spatial resolution and the focal depth of the multi-perspective Talbot Microscope. The spatial resolution is measured by imaging a USAF resolution target, the reconstructed image is shown in Fig. 3(a1). The discernable line width is 2.2 \mu m, as shown in Fig. 3(a2), and the profile of the pattern is shown in Fig. 3(a3), with a visibility of 0.24. We also measure the focal depth of the grid by scanning the focal spots along the axial direction by a microscope objective (20X, NA = 0.45, Nikon). The scanning step size is 25 \mu m. The intensity profile of the grid spot, shown as red line in Fig. 3(b), is compared with the simulation result in blue. The full width at half maximum (FWHM) of the grid is 4.4 \mu m.
The experimental profile is 133 μm, and the simulated FWHM is 118 μm. The point spread functions with the aperture placed at the −24 μm, 0, and 24 μm from the center of the microlens are also simulated, as shown in Figs. 3(c1), 3(c2), and 3(c3), respectively. The angular perspective shift is 4.9°. For the two layers that are 155 μm apart, the relative shift is calculated to be 16.5 μm, which is comparable with the experimental measurement of 11.4 μm. The difference could be caused by the inaccuracy of the focal length of the microlens and the thickness of the coverslip.

We also demonstrate the capability of the system operating in the fluorescence mode. The fluorescent sample is a two-layer sample of air dried fluorescent beads (d = 15 μm, FS07F, Bangs Laboratories) on both side of a coverslip. The scanning step size is 0.63 μm. The dwell time of each step is 0.138 s. We have inserted a fluorescence filter (D535/40, Chroma) in the detection path. The full field of view image from the perspective of 4.9° is shown in Fig. 4(a). Figs. 4(b) and 4(c) show a magnified region viewed from 4.9° and −4.9° angular perspectives. The area highlighted by the red box shows two beads from different layers, and the relative distance of the beads changes in different perspectives. We also checked the sample in conventional microscope and verified that the two beads were located on different layers. The change of the distance between two beads is 11.25 μm. Since the point spread functions are modified on the illumination side, the fluorescence collection efficiency is not affected by changing the perspective. In complex microfluidic systems, multiple layers can be distinguished from multi-perspective views with long imaging depth and could thus eliminate the adjustment of focus. Here, we perform a two-perspective fluorescence imaging of a two-layer microfluidic channels. The vertical separation of the two layers is
170 \mu m, and the width of the microfluidic channels is 28 \mu m. The quantum dots solutions (753769, Aldrich) are dried on the bottom and side walls of each channel, giving the fluorescence light. The reconstructed images are shown in Figs. 4(d)–4(f). The lateral shift of two channels is 11.3 \mu m from different perspectives. Here, we would like to mention the difference between our system and integral imaging\textsuperscript{14,16} which also generates multi-perspective images using microlens array. The depth resolution of the reconstructed 3D image is limited by the angular range from different perspectives. The angular range of our method is limited by the NA of individual microlens, while the angular range of integral imaging is limited by the extent of the MLA or the objective lens. Given the NA = 0.17 of the microlens, our setup could achieve a depth resolution of 12.7 \mu m.

In summary, we have demonstrated a scalable field of view multi-perspective microscope that can operate in both bright field and fluorescence mode. Different from a conventional scanning microscope, the Talbot Microscope decouples the lateral scanning Talbot spot and the focal spots tuning. The multi-perspective imaging can be achieved by adjusting the local wavefront. We expect further development based on the concept of engineering the point spread function of the Talbot illumination. With the long focal depth and multi-perspective imaging capability, we envision a more robust large field of view fluorescence imaging system. The demonstrated method is also suitable for a compact imaging system for multi-layer microfluidic system.

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