ABSTRACT: Determining the axial position of an emitter with nanoscale precision is critical to a fundamental imaging methodology. While there are many advanced optical techniques being applied to high-resolution imaging, high-axial-resolution topography imaging of living cells is particularly challenging. Here, we present an application of metamaterial-assisted photobleaching microscopy (MAPM) with high-axial resolution to characterize morphological properties of living cells. Quantitative imaging of changes in the morphology of live cells is obtained by topographic and statistical analysis. The time-lapse topography image using the metamaterial-induced photostability provides information about growth factor induced changes in the cell morphology with high-axial resolution.

KEYWORDS: Metamaterials, Hyperbolic metamaterials, Superaxial-resolution microscopy, Topographic imaging, Morphology changes, Photobleaching

In the last two decades, fluorescence microscopy has been widely used to optically image and analyze biological specimens. Although imaging techniques such as 4Pi-reversible fluorescent saturable optical transition (RESOLFT), interferometric photoactivation and localization microscopy (iPALM), and stochastic optical reconstruction microscopy (STORM) provide three-dimensional super-resolution imaging of the biological cell structure, these optical imaging methods meet technical difficulties in achieving axial resolutions below 10 nm. In addition, complicated optical systems are also required in order to obtain a topographic imaging of cell morphology. The monitoring of dynamic changes in the cell morphology is certainly useful for a better understanding of its dynamic processes. We believe that in order to obtain accurate statistical analysis of the cell morphology, it is necessary to acquire information on the local distance associated with cell movement through the measurement of time-lapse topography images.

A combination of the super-resolution imaging techniques and plasmonics can fundamentally increase the capabilities of fluorescence microscopy. Recently, we have developed an imaging technique, metamaterial assisted photobleaching microscopy (MAPM), to determine the axial position of fluorophores above a substrate surface with nanometer accuracy (~2.4 nm). The photobleaching lifetime of fluorophores is lengthened when brought into the vicinity of the metamaterial surface. The photobleaching lifetime modification of dipole emitters is well described by the distance-dependent Purcell effect. Therefore, metamaterial-induced photobleaching lifetime modification can be used to axially localize the position of the fluorophore above the metamaterial with nanometer accuracy. MAPM using the photobleaching lifetime measurement does not require any additional optical modules. Since every pixel acquires an emission signal at the same time with a reasonable imaging speed, it is adaptable for the analysis of cell morphology changes with nanometer accuracy.

In this study, we present MAPM imaging applications for analyzing the epidermal growth factor (EGF)-induced cell morphology changes in live cells (Figure 1a,b). The extended MAPM technique to analyze image stacks from time-lapse sequences allows functionalyzed imaging of dynamic morphology changes of live cells. Figure 1c,d shows the simplified representation of the fluorescent protein as a dipole emitter within a cell membrane situated close to a planar hyperbolic metamaterial (HMM) substrate with separation distance $d$. The correlation between the photobleaching lifetime and distance above the HMM surface is given by the distance-dependent Purcell effect (Figure S1). The planar HMM provides an enhanced Purcell effect over a broadband spectral range.
Figure 1. Metamaterial assisted photobleaching microscopy (MAPM) imaging method of EGF-stimulated changes in cancer cell adhesion. (a, b) Treatment of cells with EGF causes cell morphology changes. (c, d) Schematics showing the metamaterial assisted photobleaching microscopy (MAPM) configuration. The photobleaching lifetime of fluorophores near the metamaterial is lengthened due to the strong near-field coupling. The bleaching of the fluorescent signal over time provides positions of fluorescent molecules above an HMM surface.

Figure 2. Conventional fluorescence images and MAPM reconstructed super-resolution images of Cos-7 cells. (a) Averaged fluorescence image of the fixed Cos-7 cell. (b) Stack of fluorescence microscopy images (500 frames) is converted into a photobleaching-lifetime mapping image. (c) MAPM reconstructed super axial-resolution image of the Cos-7 cell situated on the HMM substrate. Scale bar: 20 μm. (d) Zoom of the boxed region in (c).
Figure 3. MAPM imaging of the EGF-induced morphology changes of HeLa cells. Averaged intensity images of fluorescently labeled HeLa cell membrane (a) before and (b) 15 min after EGF treatment. (c) Merged image. Photobleaching lifetime image and MAPM topography image of the Hela cell which shows the cell morphology changes with nanometer accuracy (d, e) before and (f, g) after EGF treatment, respectively. Scale bar is 20 μm.

range,12,13 which is essential for multicolor MAPM imaging technology. The time-lapse topography image using the metamaterial-induced photostability provides information about cell morphology changes with superaxial resolution.

Cos-7 cells were grown on top of the HMM substrate and then transiently transfected (see Methods section Cell Transfection for details). Cell samples with fluorescently labeled actin (Lifeact-Venus) situated on the HMM substrate were excited by a CW laser of wavelength 488 nm at the laser intensity of about 23 W/cm². The fluorescence intensity images were collected by an objective lens (40×/0.6 NA) and a magnification tube lens (2×) and then detected by a sCMOS camera with an emission filter (520/40 nm) (see Methods and Figure S1 for details). In the experiment, a stack of fluorescence microscopy images consisting of 500 frames was collected in 200 s (2.5 frames per second; 2.5 fps). An averaged fluorescence image of a Cos-7 cell is shown in Figure 2a. Figure 2b shows the photobleaching-lifetime image with a strong spatial dependence. The distance from the HMM surface to the fluorescent protein can be obtained for each pixel according to the precalculated and premeasured axial distance dependence of the photobleaching lifetime, and the resulting MAPM images are shown in Figure 2c,d. The distance image of the fluorescently labeled actin shows the spatial variation of the distance in the range of 2−160 nm. Additional morphological images of Cos-7 cells are shown in Figure S2. The reproducibility of our MAPM technique was tested by repeating the MAPM imaging procedures with over 20 different Cos-7 cells. The spatial distribution of the distance between the cell and the HMM surface could result from the spatial difference in cellular structures or the heterogeneity of the cell.3,14

Having discussed the basic principle, we now test the capability of MAPM to observe the influence of the drug treatment on cell morphology. It is well-known that the epidermal growth factor (EGF) stimulates differentiation, migration, cell growth, and cell adhesion changes15,16 by binding to its receptor. HeLa cells were grown on top of the HMM substrate and then transiently transfected with a fluorescent protein targeted to the plasma membrane using the C-terminal CAAX sequence of KRas (Venus-CAAX) (see Methods section Cell Transfection for details). The fluorescence intensity image sequences of 500 frames of HeLa cells were collected in 200 s (2.5 frames per second; 2.5 fps), while the fluorophores are excited with a constant laser intensity of about 23 W/cm² for the wide field of view. An averaged fluorescence image of the HeLa cells is shown in Figure 3a. After 15 min of incubation with epidermal growth factor (EGF, 10 ng/mL), another 500 frames of images of the growth-factor-treated cells were collected. An averaged fluorescence intensity image of the EGF-treated HeLa cells is shown in Figure 3b with the merged image (Figure 3c). The same exposure time, pH of buffer solutions, and temperature of the chamber were used during the data collection period for each measurement data set. Cells showed negligible movement during a short photobleaching measurement cycle (500 frames, 200 s).

In the postreconstruction processing of the collected data, the photobleaching signals were fitted for each pixel, and the photobleaching lifetimes were extracted (Figure 3d,f). In addition to information on the modification of the size and shape of growth-factor-treated HeLa cells in the intensity images, the photobleaching-lifetime images show additional information about spatial heterogeneity. By measuring photobleaching lifetime, positions of the plasma membrane localized proteins are obtained with nanometer axial precision. From the MAPM reconstructed images (Figure 3e,g) showing the distances between the fluorescence proteins to the HMM surface with a superaxial resolution, one can easily observe that the cells spread out after the drug treatment. Treatment with EGF significantly changes the cell morphology. Arrows denote cell presettled/ floating/spreading over the HMM substrate. Additional representative images and analysis of the changes in the cell morphology are provided in Figure 4.
Imaging of the time-lapse series was performed to observe EGF stimulated cell morphology changes over a longer time span after drug treatment. A total of 500 frames were collected every 400 ms over a period of 200 s, in which sufficient distance-dependent photobleaching signals were acquired for each pixel. Cell movement during each imaging cycle was minimal. Averaged fluorescence intensity images of the HeLa cell stimulated with EGF after the indicated time are shown in Figure 5a,b with the merged image (Figure 5c). Interestingly, MAPM reconstructed images (Figure 5d−f) show significant information about cell morphology that does not appear in the intensity images. Direct quantification is presented in the MAPM reconstructed images, showing the cell protrusion and retraction. The observed cell morphology changes in the three snapshots describe the formation of individual cell−substrate contacts, i.e., domains which are located in the vicinity of the top surface of the substrate, with nanometer accuracy. Additional representative images are shown in Figure S3.

In conclusion, we demonstrated MAPM imaging applications for analyzing the epidermal growth factor (EGF)-induced morphology changes in live cells with superaxial resolution. We experimentally developed an extended MAPM imaging technique for analyzing image stacks from time-lapse sequences for the combined time-lapse and topographic imaging, and in particular of cellular systems, based on the use of MAPM. For the purpose of obtaining precise distance information with axial resolution below 10 nm between the cell and the substrate, MAPM could be one of the simplest and most user-friendly imaging methods. We believe MAPM is a useful imaging method for live cell morphology studies, and future studies will combine this method with strategies for lateral resolution improvement.

■ METHODS

Experimental Setup. A custom-modified fluorescence microscope, Olympus IX83 with an upright configuration, was utilized. A 488 nm laser coupled through a multimode fiber was used to excite fluorescently labeled biological samples, and 500 frames of the fluorescence signal were collected by an objective lens (0.6 NA) and an sCMOS camera with proper color filters. A 488 dichroic mirror and a 520/40 nm band-pass filter were adopted. The reduction in emission intensity over time of the acquired fluorescence signal was mathematically fitted and analyzed for each pixel using MATLAB. We use MATLAB to adjust the data acquisition card (DAQ) voltage output module to synchronize laser beam shutters and acquisition modules properly. All cells were kept in a humidified incubator at 37 °C with a 5% CO2 atmosphere during time-lapse MAPM imaging.

Cell Transfection. Cells were cultured in Dulbecco’s modified Eagle medium (DMEM; Gibco) containing 4.5 g/L glucose and supplemented with 10% (v/v) fetal bovine serum (FBS, Sigma) and 1% (v/v) penicillin-streptomycin (Pen-Strep, Sigma-Aldrich). All cells were maintained in a humidified incubator at 37 °C with a 5% CO2 atmosphere. At 24 h prior to transfection, cells were seeded onto the HMM substrate and grown to 50−70% confluence.
Cos-7 Cell Transfection. Cos7 cells were then transfected with 100 ng of pcDNA3-Lifeact-Venus using Lipofectamine 2000 (Invitrogen) and grown an additional 24 h before fixation. Cells were washed with phosphate-buffered saline (PBS) before fixation with 4% paraformaldehyde and 0.2% glutaraldehyde PBS for 10 min at room temperature. Cells were quickly rinsed in PBS after fixation and quenched with freshly made 0.1% NaBH₄ ice-cold PBS. After quenching, cells were washed three times for 5 min each with PBS on a shaker. Cells were imaged at room temperature.

HeLa Cell Transfection. HeLa cells were then transfected with 100 ng of pcDNA3-Venus-CAAX (87612, Addgene) for plasma membrane labeling using Lipofectamine 2000 (Invitrogen) and grown an additional 24 h. Cells were imaged at room temperature.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.0c04529.

Metamaterial-assisted photobleaching microscopy, superaxial-resolution image of the Cos-7 cells, and time-lapse MAPM imaging of a HeLa cell after EGF treatment (PDF)

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Notes
The authors declare no competing financial interest.

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