Organic Hyperbolic Material Assisted Illumination Nanoscopy

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Resolution capability of the linear structured illumination microscopy (SIM) plays a key role in its applications in physics, medicine, biology, and life science. Many advanced methodologies have been developed to extend the resolution of structured illumination by using subdiffraction-limited optical excitation patterns. However, obtaining SIM images with a resolution beyond 40 nm at visible frequency remains as an insurmountable obstacle due to the intrinsic limitation of spatial frequency bandwidth of the involved materials and the complexity of the illumination system. Here, a low-loss natural organic hyperbolic material (OHM) that can support record high spatial-frequency modes beyond 50k0, i.e., effective refractive index larger than 50, at visible frequencies is reported. OHM-based speckle structured illumination microscopy demonstrates imaging resolution at 30 nm scales with enhanced fluorophore photostability, biocompatibility, easy to use and low cost. This study will open up a new route in super-resolution microscopy by utilizing OHM films for various applications including bioimaging and sensing.

1. Introduction
Super-resolution fluorescence microscopy has enabled a plethora of studies of deep subwavelength structures such as biological molecules, proteins and subcellular structures that cannot be achievable with traditional microscopes. Among various super-resolution microscopy techniques, structured illumination microscopy (SIM) has recently attracted much attention due to its advantages in high spatial-temporal resolution and low photo-toxicity[1,2] compared to other super-resolution techniques such as STED, PALM, and STORM.[3–5] Typically, SIM can enhance the resolution about twice beyond the diffraction limit by using diffraction-limited harmonic illumination patterns.[6,7] Yet one requirement of standard SIM is the precise knowledge of the illumination patterns to reconstruct a super-resolution image and distortion in illumination will usually cause reconstruction artifacts.[8] To alleviate such a limitation, speckle-structured illumination microscopy (speckle-SIM) has recently been developed[9] with the consideration of a statistical prior that the averaged speckle illuminations are homogeneous.[8] However, resolution improvement of speckle-SIM is still about twofold as the speckle patterns are typically also diffraction limited.
To further improve the resolution capability of SIM, a material system that supports higher spatial bandwidth needs to be sought. For example, plasmonic-structured illumination microscopy (plasmonic-SIM)[10–12] utilizing plasmonic waves as illumination sources has further extended the resolution beyond the conventional SIM. Metamaterial is another platform people used to achieve super resolution.[13–15] One of the most widely used metamaterials is metal and dielectric multilayer hyperbolic metamaterial (HMM)[16–18] (Figure 1a, left), implemented as the first hyperlens by curving the multilayers and converting the evanescent waves from an object directly into propagating waves.[19–23] In another aspect, HMMs can be also implemented to provide illumination patterns far beyond the diffraction limit.[19–23] Therefore, with such high-k illuminations, the metamaterial assisted illumination nanoscopy (MAIN) currently represents a very unique SIM approach with greatly extended resolving power.[19–23] Challenge still remains to obtain images with a resolution beyond 40 nm at visible frequency.

In this Letter, we demonstrate a new OHM-based SIM method with 30 nm resolution capability. With periodicity at 1.6 nm, hyperbolic dispersion of the developed OHM film supports remarkably high-k optical modes in the visible frequency. Comparing to inorganic counterparts, the OHMs not only provide better deep subwavelength-scale illuminating field confinement in visible spectral range, but also offer many advantages such as extremely enhanced photostability of fluorophores near the OHM,[27] easy fabrication process, and excellent biocompatibility[28,29] for bioimaging applications.

Figure 1b shows the calculated optical dispersion of an Ag-based HMM using the effective medium theory (EMT) where its isofrequency surface, with the open hyperbola topology[30,31] can be clearly seen in k-space. Note that the maximum available wavevector and photonic local density of states (LDOS)[32,33] are limited because the EMT description of the HMM eventually fails when taking into account the limited unit cell size as shown in Figure 1b,c (see Supporting Information S1 for details). Besides, optical losses and nonlocal response[34,35] limit the maximum available wavevector of propagating optical modes as shown in Figure 1d. Natural hyperbolic materials[36–38] could circumvent some of the limitations on the HMMs, e.g., the finite structural periodicity and nonlocal effect. For example, Bi2Te3,[39] and GaTe[40] have been experimentally demonstrated as the natural hyperbolic materials. Unfortunately, these materials are very lossy leading to hyperbolic modes with a very small wavevector (Figure 1d, green star)—much worse than that provided by the Ag-based HMMs (Figure 1d, red star). Compared to the traditional HMMs and other natural hyperbolic materials, the OHMs used in this work (Figure 1a, right) feature a low optical loss[41] and thus support record high-k modes (Figure 1d, cyan star). Therefore, the OHM presents an excellent material platform achieving the highest possible resolution improvement for super-resolution microscopy.

2. Results and Discussion

To produce the OHM, 100 mg of 98% regioregular poly(3-hexylthiophene-2,5-diyl) (rr-P3HT) was dissolved in 1 mL chlorobenzene (CB) solution (P3HT:CB = 100 mg mL⁻¹). From this solution, thin films were spin-coated on top of a 0.17 mm thick plasma-cleaned glass coverslip as described in reference[41]
(see the Experimental Section for details of the film fabrication). The fabricated 182 nm thick rr-P3HT films exhibit a hyperbolic dispersion in the visible spectral range 400–560 nm.[41] The capability of super-resolution imaging using the OHM film is estimated by the measured permittivity (see details in Supporting Information S2, Figure S2) and resultant optical transfer function (OTF) based on transmission coefficient calculation (see details in Supporting Information S1, Equation S5). The OTF of the OHM has an exceptionally high cutoff spatial-frequency \( k_{\text{cut-off}} = 54k_0 \) when OTF \( (k_{\text{cut-off}}) = 0.01 \) at \( \lambda = 465 \) nm. The wavelength of the propagating waves and thus the resolution supported by this OHM can reach below 10 nm. A comprehensive comparison of HMM multilayers in various combinations with the same cutoff spatial-frequency (see a detailed comparison in Supporting Information S3) reveals that the metal film thickness has to be around 1 nm scales introducing significant fabrication challenges. In addition, the optical property of such thin metallic film has to be modified by nonlocal or quantum size effect, which makes the achievement of such OTF highly questionable. Momentum-resolved electron energy loss spectroscopy \( (k\text{-EELS}) \) measurement in a transmission electron microscope (TEM) further confirms the high-\( k \) mode excitation in the OHM. Figure 2b shows the measured energy-momentum (scattering angle) dispersion of the OHM at 200 keV incident energy of electron (see the Experimental Section for details of the \( k\text{-EELS} \) measurement). The \( k\text{-EELS} \) spectra integrated over the scattering angles (Figure 2c and Figure S4, Supporting Information) show additional evidence of excitation of the high-\( k \) hyperbolic polariton modes at the hyperbolic dispersion wavelength range 400–560 nm (2.2–3.1 eV).

The high-\( k \) illumination brings super-resolution to a traditional microscopy.[44] We tested the imaging performance of the OHM assisted illumination nanoscopy with 20 nm diameter fluorescent beads as imaging objects. The fluorescent beads (see details in the Experimental Section) were drop-casted on the OHM-coated coverslip substrate. It has been shown previously that the high-\( k \) speckle patterns on the top surface of hyperbolic materials can be generated and controlled by tuning the incident non-uniform illumination patterns.[14,24] In the experimental setup (see details in the Experimental Section and Supporting Information S5), the high-\( k \) speckle illumination on the OHM were varied by adding mechanical forces on the multi-mode fiber with a stepping motor. At the sample plane, the high-contrast and high-\( k \) speckles excited the fluorophores in a specimen, and after passing through an emission filter (520/40 nm band-pass), the fluorescence signal was collected by an sCMOS camera. By varying the speckle illumination \( \times \text{N-times} \), \( N \)-different fluorescence image frames were obtained via an 80×0.6 NA objective lens. A set of \( N = 300 \) frames (5 frames per second) were captured in the whole sequence, which were used to reconstruct the final super-resolution image by the Blind-SIM reconstruction algorithm[8] (see the Experimental Section for details of the reconstruction).

Figure 3 shows the diffraction-limited image (left panel) and super-resolution image (middle panel) of fluorescence beads. As can be seen, the super-resolution image reveals much greater details that cannot be observed by any individual diffraction-limited image. The cross-section (white solid line) curves are plotted for two adjacent beads with the center-to-center distance of 610 nm (Figure 3a,b), 60 nm (Figure 3c,d), 43 nm (Figure 3e,f), and 31 nm (Figure 3g,h), respectively. After fluorescence imaging, the imaged locations were fixed for SEM by using a reference marker on the sample. The distributions of fluorescent beads were identified by SEM images (insets of the middle panel of Figure 3). To confirm the actual resolution achieved, a Fourier ring correlation (FRC)[45] analysis was subsequently performed. The FRC is a resolution criterion that is widely accepted for super-resolution microscopy. Two reconstructed images based on two independent subsets of data were used. The standard 1/7 FRC resolution criterion illustrates that 25 nm resolution has been achieved (see Figure 3j), representing more than 17 times spatial resolution improvement beyond the diffraction limit.
To test the biocompatibility, we present reconstruction results of fluorescently labeled plasma membrane and actin filaments in Cos-7 cells. Cells were grown on top of the OHM substrate and then transiently transfected with fluorescently labeled F-actin (Lifeact-Venus) or a fluorescent protein targeted to the plasma membrane using the C-terminal CAAX sequence of KRas (Venus-CAAX) (see the Experimental Section for Cos-7 cell transfection). In the imaging experiments, 200 frames were acquired with 10 W cm⁻² illumination intensity. Figure 4a–d shows the diffraction-limited images of Venus targeted to the plasma membrane in Cos-7 cells. The corresponding reconstructed images are shown in Figure 4e–h.

Figure 5 shows diffraction-limited images (Figure 5a) and super-resolution images (Figure 5b) of Venus labeled actin in Cos-7 cells via our super-resolution imaging technique. The FRC analysis and resolution of the reconstructed images (outlined by a white box in Figure 5b) as a function of the number of subframes, N, used to generate a super-resolution image are shown in Figure 5c–e. The resolution of ≈40 nm scales is achieved by only 120 frames (Figure 5d) for a 5 fps acquisition rate. The nature of a stochastic blinking of fluorophores is one of the main factors in limiting the temporal resolution of probe-based localization microscopy. On the other hand, predetermined speckles in OHM assisted illumination microscopy can represent much more prior information enabling improved image reconstruction with less measurements. This eventually can lead to higher imaging speed. Note that during the image acquisitions, almost negligible photobleaching was observed because of the

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**Figure 3.** Images of fluorescent beads with the averaged diameter of 20 nm. The diffraction-limited images (left panel), the super-resolution images and scanning electron microscope (SEM) images (middle panel) of fluorescent beads, and the cross-section (white solid line) curves (right panel) are plotted for two adjacent beads with the center-to-center distance of a,b) 610 nm, c,d) 60 nm, e,f) 43 nm, and g,h) 31 nm, respectively. i) Diffraction limited image and reconstructed image of two fluorescence beads (center-to-center distance of 23 nm). j) The standard 1/7 FRC resolution criteria illustrates 25 nm Fourier space cutoff. Objective lens: 80x/0.6 NA. Exposure time: 500 ms. Total frame: 500.
Figure 4. Venus-CAAX tagged Cos-7 cell membrane images. a–d) Conventional wide-field image. e–h) Super-resolution images. Scale bar: 2 μm.

Figure 5. Venus-lifeact tagged Cos-7 cell actin images. a) Conventional wide-field images. b) Super-resolution images. Scale bar: 2 μm. c) FRC resolution criteria curves generated from two reconstructed images based on two independent subsets of data with the number of subframes, N. The standard 1/7 FRC resolution criteria illustrates 34 nm Fourier space cutoff for N = 200. d) The standard 1/7 FRC resolution criteria illustrate the N-dependent resolution of the reconstructed images. e) Reconstructed images generated from N subframes.

significantly reduced photobleaching rates by hyperbolic optical modes supported by the OHM due to the large resultant Purcell effect.[27]

After the optical imaging, the samples were dried and random locations were selected to confirm that the OHM is a chemically stable and biocompatible substrate for cell transfection. The OHM is a conductive substrate to uniquely enable scanning electron microscopy (SEM) imaging of Cos-7 cell without sealing additional layers on top of the cells. An SEM image of Cos-7 cells cultured on a glass coverslip (Figure S6a, Supporting Information) shows overly bright and unstable intensity due to accumulated charges on the top surface of the glass. In contrast, SEM images of Cos-7 cells cultured on the electrically conductive and biocompatible OHM substrate (Figure S6b–f, Supporting
Information) show outstanding resolution and contrast over the entire field of view.

3. Conclusion

In summary, we demonstrated an OHM-based super-resolution imaging technique based on the speckle structured illumination microscopy. The achieved precision of lateral resolution reaches down to 30 nm, which is more than 17 times spatial resolution improvement beyond the diffraction limit. The developed OHM offers an alternative path to controlling light at the nanoscale, leading to a significant resolution improvement in biological cell imaging. We observed that our OHM substrate is highly stable for cellular imaging after cell transfusion and no protection layer was required for cell culture, well surpassing the capability of traditional hyperbolic materials. Our OHM-based imaging technique will open up a new paradigm to utilize the OHM films instead of glass coverslips for microscope slides to achieve enhanced fluorophore stability, biosample friendly, easy to fabricate, and low cost for high-resolution imaging applications.

4. Experimental Section

**OHM Film Fabrication:** 98% regioregular P3HT (molecular weight average $M_w \approx 87$ 000, Sigma-Aldrich) 100 mg were dissolved in 1 mL chlorobenzene by heating the solution to 50 °C for 3 h and resting at room temperature for 2 h, resulting in a dark orange-colored solution. Thin films were spin-coated on 0.17 mm thick clean glass coverslips at 5000 rpm for 60 s. The film thickness of 182 nm was measured by spectroscopic ellipsometry. Please see Supporting Information S2.

**Electron Energy Loss Spectroscopy (EELS) Measurement:** Thin films of rr-P3HT (10 and 100 mg mL$^{-1}$ concentrations) were prepared on glass substrates. Argon reactive-ion etching was performed on an Oxford Instruments Plasmalab 80 Plus to reduce the thickness of the films. The etching condition was set to be 40 sccm Argon with 100W RF power. Film thicknesses of 28 and 26 nm were measured with complex permittivity by VASE. These thin films were collected onto carbon-coated TEM grids after they were floated off by the dissolution of the substrates in water. High EELS measurement was carried out using transmission electron microscopy (TEM) on the FEI monochromated F20 UT Tecnai, operating at 200kV, located in the National Center for Electron Microscopy (NCEM) at Molecular Foundry, Lawrence Berkeley National Laboratory (LBNL). This microscope was equipped with Gatan Imaging Filter (GIF) along with a monochromator incorporated into the gun, therefore, it permitted electron energy loss spectroscopy to be performed with an energy resolution of $\approx 0.2$ eV. This high spectral resolution enabled the characterization of the rr-P3HT film in the electron energy-loss range close to the zero-loss peak with minimum overlapping. Please see Supporting Information S3.

**Cos-7 Cell Transfection:** Cos7 cells were cultured in Dulbecco modified Eagle medium (DMEM; Gibco) containing 4.5 g L$^{-1}$ 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. 24 h prior to transfection, cells were seeded onto OHM substrate and grown to 50–70% confluence. Cells were then transfected with 100 ng of pcDNA3-Lifeact-Venus (87613, Addgene) for actin-labeling or 100 ng of pcDNA3-Venus-CAAX (87612, Addgene) for plasma membrane labeling 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 five min each with PBS on a shaker.

**Imaging Reconstruction:** All of the image processing and reconstruction were performed in MATLAB. An iterative reconstruction algorithm, blind-SIM[8], which does not require exact knowledge of the illumination pattern, was used to retrieve the object information. For blind-SIM, an assumption was made that all illumination patterns add up to a uniform pattern, both the object and the illumination patterns were treated as unknowns in real space and were solved using a cost-minimization strategy. Each super-resolution frame was reconstructed from multiple subframes. The GPU-based reconstruction typically takes 10–30 min on a Nvidia GTX 1080 Ti.[14]

**Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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**Conflict of Interest**

The authors declare no conflict of interest.

**Data Availability Statement**

The data that support the findings of this study are available from the corresponding author upon reasonable request.
