Multi-color live-cell super-resolution volume imaging with multi-angle interference microscopy

Youhua Chen¹,², Wenjie Liu¹, Zhimin Zhang¹, Cheng Zheng¹, Yujia Huang¹, Ruizhi Cao¹, Dazhao Zhu¹, Liang Xu¹, Meng Zhang³, Yu-Hui Zhang³, Jiannan Fan⁴, Luhong Jin⁴, Yingke Xu⁴, Cuifang Kuang¹,⁵ & Xu Liu¹,⁵

Imaging and tracking of near-surface three-dimensional volumetric nanoscale dynamic processes of live cells remains a challenging problem. In this paper, we propose a multi-color live-cell near-surface-volume super-resolution microscopy method that combines total internal reflection fluorescence structured illumination microscopy with multi-angle evanescent light illumination. We demonstrate that our approach of multi-angle interference microscopy is perfectly adapted to studying subcellular dynamics of mitochondria and microtubule architectures during cell migration.
Observe three-dimensional (3D) volumetric subcellular structures and functions is essential for biological research. For this reason, a variety of powerful super-resolution fluorescence imaging techniques have been proposed. One approach for overcoming the diffraction limit in lateral dimension is activating and localizing single molecule1–2. Techniques like modulating the point spread function (PSF)1–3, increasing the numerical aperture (NA) of the system4–6, or detecting supercritical light7 can improve the axial resolution. Similarly, 3D nanometer details can be resolved by combining stimulated emission depletion microscopy (STED) with an NA increasing technique8.

However, although these 3D super-resolution approaches are appropriate to the study of relatively static structures, such as stationary microtubules and nuclear pore complexes, they are difficult to use for imaging and tracking the real-time evolution of live cells in a large field of view because of the requirements of high excitation intensities of $10^3–10^6$ W cm$^{-2}$ (far beyond the safe intensity of 0.1 W cm$^{-2}$ for live-cell imaging), specialized fluorophores and labeling strategies, and a long imaging capturing process.

The aforementioned shortcomings are overcome by 3D structured illumination microscopy (3D SIM)9–11, which can image 3D living samples labeled by conventional dyes with an excitation intensity of $1–10$ W cm$^{-2}$. However, the limited spatial resolution, which is typically improved twofold or less in lateral and axial resolutions, limits its biological applications12.

The evanescent wave is widely used in optical microscopy to illuminate near-surface cell structures on an axial scale down to a few hundred nanometers. By leveraging the exponential decay of the evanescent field from the coverslip, microscopy based on total internal reflection fluorescence (TIRF) can further improve the axial resolution by recording a series of images at different incident angles13 or different wavelengths14. Nevertheless, the immense difference between the wide-field lateral resolution and 10–50-nm axial resolution of TIRF engenders its poor performance in volumetric super-resolution imaging.

Inspired by these existing challenges, we designed and implemented a technique for 3D multi-color live-cell super-resolution volume imaging by incorporating SIM with multi-angle evanescent light illumination. We refer to the proposed approach as multi-angle interference microscopy (MAIM) (Supplementary Figs. 1–3). The introduction of evanescent wave improves the signal-to-noise ratio (SNR) and optical sectioning capacity of the structured illumination methodology (Supplementary Fig. 4). Compared to other 3D super-resolution techniques1–8, MAIM has an advantage with respect to temporal resolution because of the smaller number of required raw images. Moreover, a lower requirement for the surface energy density of the illumination light, which is even lower than that of 3D SIM, enables its use for probing specific long-term dynamics in living cells without photodamage or photobleaching. Furthermore, the portability of the sample preparation protocols and experimental implementation allows it to be easily adapted to a wide range of biological studies.

**Results**

**MAIM theory and demonstration.** In MAIM, we used an established TIRF-SIM algorithm to achieve lateral super-resolution15. Following the robust procedure for breaking the lateral optical diffraction limit by down-shifting high-frequency components, we first recorded nine TIRF-SIM raw images (Fig. 1a). To enhance the axial resolution, we collected a set of multi-angle TIRF (MA-TIRF) images by azimuthal averaged over different evanescent wave propagation directions (so-called “MA-Ring-TIRF”), which contain different depth information of the sample (Fig. 1b–d, Supplementary Fig. 5). The lateral TIRF-SIM super-resolution image is then thresholded, and serves as a binary spatial mask to subtract the local background, abandon the diffraction-limited information, and segment the interesting sample information from the MA-Ring-TIRF raw images. The resulting segmentation image stack is taken as the input of the 3D super-resolution reconstruction. By treating them as a convex optimization problem, a precise knowledge of our system parameters, such as the quantum efficiency, absorption rate, detection efficiency, and PSF, was not required. We then solved the problem with our previously reported fast and efficient reconstruction algorithm based on distributed optimization acceleration16 (Supplementary Fig. 3). With these procedures, MAIM enabled us to map the surface morphology with a sub-100-nm lateral resolution and axial architectures with an approximately 40-nm axial resolution over a 600-nm depth volume. The typical acquisition time of one volume stack in MAIM is 1–2 s. We demonstrated this by tracking the fast temporal evolution of Atto 647N-labeled mitochondria and SiR-tubulin-labeled microtubules in live U2OS cells.

We analyzed the parameters affecting the reconstruction by conducting simulations of 3D microtubule structures incorporating Gaussian white noise while changing the axial position, incident angle range, and angle number. Supplementary Fig. 6a, b present the 3D ground-truth data and the reconstructed depth data (SNR = 30, 20 angles from 61.5° to 71°). We found that the root mean square error of the calculated depth from the true position heavily depends on the SNR, and can be below 5 nm when a fluorescent emitter is bright and near the coverslip (Fig. 2a). Further, to reduce the influence of the refractive index mismatch on the reconstruction (Supplementary Fig. 6e), we experimentally determined the refractive index of each sample before the reconstruction (Supplementary Note 1). Other reconstruction parameters, for example, the angular steps and incident angle varying range, have relatively less effects on the reconstruction (Supplementary Fig. 6c, d). From the analysis, a typical 10–20 angle number from 61.5° to 71° is the optimal choice for MAIM reconstruction to provide the highest spatial and temporal resolutions. Furthermore, we used gradually spaced lines (Argo-HM) and silica microsphere (Bangs Laboratories, Inc.) ground-truth samples to experimentally validate MAIM (Fig. 2b and Supplementary Fig. 7). The lateral resolution enhancement was confirmed by separating 100-nm distance in the spaced line sample (Supplementary Fig. 7a–c). The measured z-positions of the spheres distributed well around the theoretical z-profile. The final fitted averaged sphere diameter of 4.98 μm was slightly larger than the theoretical value of 4.86 μm possibly reflecting the physical uniformity of the spheres (±0.47 μm) (Supplementary Fig. 7d–f). The experimental axial root mean square error had the same descending trend with the simulation result when the depth increases, and was ~20–60 nm in the depth range from 0 to 500 nm (Fig. 2b).

**Fixed cell imaging.** To demonstrate the performance of MAIM in cell imaging, we conducted a series of experiments utilizing our MAIM setup (Supplementary Figs. 1 and 2). Firstly, we imaged the microtubule networks labeled with Alexa Fluor 488 in U373 cells grown in a two-well glass chamber (Fig. 3a–d). After immunofluorescence staining, the cell membrane became permeable for antibodies to enter; therefore, the refractive index inside the cell could be matched with the surrounding aqueous buffer solution. A blue laser ($λ = 488$ nm) was used to excite fluorophore under the TIRF configuration. The excitation power was maintained constant throughout the whole imaging process.
Multi-azimuthal-angle and multi-phase raw images were acquired to reconstruct the lateral super-resolution image of the sample (Fig. 3b). The sub-diffraction structures in the TIRF image (Fig. 3a) were now clearly resolved. By acquiring an image stack of 20 incident angles with a 0.50° angular step size starting from 61.64°, we reconstructed the 3D volume distribution of the microtubules with the lateral and axial resolutions of 98 and 46 nm, respectively, up to approximately 400 nm in depth (Fig. 3c, d). The color-coded depth visualization revealed a continuous upward bending of the microtubule network from the leading edge toward the center. Six different algorithms for reconstructing the multi-angle interference image stack were compared on this image series (Supplementary Fig. 8).

A visual reconstruction of peroxisomes (PMP70) labeled with STAR 580 (λ = 561 nm) at 20 incident angles with a 0.50° angular step size starting from approximately 61.91° showed fine 3D details, revealing distinct topographical features (Fig. 3e–h). The lateral resolving ability of MAIM was confirmed by discriminating the circular shape of the peroxisomes (Fig. 3f, g), which was not straightforward in the conventional TIRF image and multi-angle total internal reflection fluorescence (MA-TIRF) reconstructed image (Fig. 3e and Supplementary Fig. 9). The histogram of several peroxisomes showed a 72-nm axial resolution and a 126-nm mean depth (Fig. 3h).

We next applied MAIM to image the mitochondrial network in fixed bovine pulmonary artery endothelial cells labeled with MitoTracker® Red CMXRos (Cat. no. M7512) (λ = 561 nm) at the illumination power of 1.32 W cm⁻² (Fig. 4). The MAIM image reconstructed from one volume of 30 incident angles with a 0.33° angular step size starting from approximately 61.51° showed fine 3D details, revealing distinct topographical features.
showed more significantly improved lateral and axial resolutions over the conventional diffraction-limited TIRF image. We observed three distinct forms of mitochondrial morphology: many interconnected tubular networks, intermediate mitochondrial structures, and a few dispersed globular structures, which likely indicated the interphase stage of the cell\(^\text{17}\) (Fig. 4b). A series of 3D super-resolution images revealed the hollow shape and morphology change of individual mitochondrial outer-membranes spanning multiple axial slices from a depth of 0 nm to 180 nm relative to the coverslip (Fig. 4c). The change of the whole imaging area was also given (Supplementary Fig. 10). Such process of continuous change in the mitochondrial hollow structure was also previously reported using 3D STED\(^\text{9}\) and 3D STORM\(^\text{13}\). The reconstructed images at different depths given by MAIM were converted into a 3D near-surface volume view of the mitochondria (Fig. 4d and Supplementary Movie 1).
Multiple mitochondria, such as migration, retraction, fusion, and lateral evolution of individual mitochondrion and mitochondria architecture within a 600-nm depth was reconstructed imaged using our method. The result showed that the mitochondria and microtubules (magenta). Scale bars, 4 μm (a-f), and 8 μm (bottom-right of a and b).

Multi-color imaging. At this point, we simultaneously measured the 3D distributions of microtubules stained with Alexa Fluor 488 (λ = 488 nm) and mitochondria stained with Alexa Fluor 568 (λ = 561 nm) in the same U373 cell (Fig. 5). The raw image stacks were all acquired at 16 different incident angles with a 0.50° angular step size starting from 61.64°. As an organelle providing energy to cells, mitochondria generally distribute around the nucleus and near the microtubules, which was in accordance with our co-localization imaging result (Fig. 5d, e). At the axial direction, we quantitatively observed that microtubules were slightly deeper than mitochondria (Fig. 5d–f), verifying the capability of MAIM for multi-color super-resolution volume imaging and co-localization of different cell components.

Live-cell imaging. The fact that MAIM is a powerful tool to study the high dynamics of intracellular architectures was demonstrated by imaging mitochondria and microtubules in different live U2OS cells labeled with Atto 647N (λ = 639 nm) and SiR-tubulin (λ = 639 nm), respectively. Atto 647N has been increasingly used as the optimal labeled dye in live-cell super-resolution imaging because of its high luminous efficiency and excellent photostability. After staining with Atto 647N, U2OS live cells were imaged using our method. The result showed that the mitochondria architecture within a 600-nm depth was reconstructed from a 3D volume corresponding to ten incident angles with a 0.67° angular step size starting from 61.91° corresponding to 250 nm in the depth of the cells (Fig. 6c, d). From the color changes of the reconstructed super-resolution volume images, we observed that their depth tended to be downward (Fig. 6d). In a more quantitative measurement, the average depth of the selected microtubules varied from 138 nm to 32 nm during the 45-s recording time. The lateral real-time movements of extension, contraction, and swirling of the network were also tracked (Fig. 6d).

Discussion
In summary, we demonstrated herein that MAIM is a powerful technique that provides a combination of super-resolution volume imaging, live-cell imaging, multi-color imaging, a long time series, and low photodamage, which other 3D imaging techniques have difficulty providing. The proposed technique achieves this combination without requiring specially-prepared biological samples. Consequently, it is very suitable for the investigation of dynamic molecular behaviors that are proximal to the cell membrane, and can be widely applicable to some critical issues, such as vesicle endocytosis and exocytosis, membrane transport, membrane heterogeneity, and mechanobiology. With respect to the prominent features of our method, it can provide absolute axial sectioning and depth positioning of fluorescent mitochondria, the fission appeared concomitant to constriction/protrusion events (Fig. 6b, purple arrowheads). After fission, we could observe the downward movement of the forming mitochondria because of the high axial resolving capacity of our approach. The 3D dynamics and volume rendering of mitochondrial membranes in the image sequence were both consistent with similar data from the COS-7 cell.
structures relative to the coverslip (Fig. 4 and Supplementary Fig. 10).

Moreover, in real-time live-cell imaging, the MAIM imaging speed depends only on the camera image acquisition time. Theoretically, its speed can reach up to approximately 100 Hz, which means the temporal resolution of our MAIM system can be 3–5 volume per second. However, in practice, the local concentration, quantum efficiency, photostability and brightness of dyes, as well as the laser excitation power, field of view size, and system detection efficiency must be comprehensively considered when recording images. With the rapid development of faster cameras, brighter and more stable fluorescent dyes, alternative cell-permeable organic fluorescent probes, and more mature labeling strategies, the simultaneous multi-color super-resolution imaging of different, but correlative dynamic components in a living system is possible to analyze the interactions among these components. This is difficult to achieve otherwise with other single molecule localization-based techniques.

Methods

Multi-angle interference microscopy instrumentation. All optical elements used in our technique were mounted on a 750 × 750 × 60 mm optical breadboard (Thorlabs, B7357L) to minimize external mechanical vibrations. Three lasers were equipped for fluorescence excitation: a 300-mW, 488-nm laser (Coherent, Sapphire 488-300 CW CDRH); a 300-mW, 561-nm laser (Coherent, Sapphire 561-300 CW CDRH); and a 1-W, 639-nm laser (Coherent, Genesis MX 639–1000). The illumination wavelength and the intensity of the lasers were controlled by an acousto-optical tunable filter (AOTF) (AA Quanta Tech, AOTF3c-400.650-TN). After the AOTF, beams were coupled into a single mode polarization maintaining fiber (Oz Optics, core size of 3 μm, NA 0.11), then collimated by a beam collimator (Thorlabs, ZC618FC). Parallel beams were separated into two paths with orthogonal polarization directions by a polarization beam splitter (PBS) (Thorlabs, CCM1-PBS513/M). Before that point, the intensity ratio of two paths was controlled by a half-wave plate (Thorlabs, AQWP05M-600) to ensure the high modulation contrast of the interference pattern. The two light beams were then directed into two construction-symmetric paths, which consisted of two reflecting mirrors to alter the propagation direction of the beam, a galvanometer set (Cambridge Technology, CTI 8310k) that serves to change the incident angle and rotate the orientation of the interference pattern, and a scanning lens (Thorlabs, SLS-CLS2) to rectify and focus the light. The difference was that a half-wave plate (Thorlabs, AQWP05M-600) and a piezoelectric stage (PI, 753.1, 5 kHz) were inserted into one path to adjust the polarization and shift the interference pattern phase. Two focusing beams were combined by a beam splitter (BS) (Thorlabs, CCM1-BS013) and reimaged in opposite form near the edges of the BFP of a high-NA oil-immersion objective (Nikon, 100 ×/1.49 TIRF) by a 4f configuration constructed from two tube lenses (Thorlabs, TT1200, f = 200 mm) and a periscope. A polarization rotator was placed behind the BS consisting of a liquid crystal cell (LC, Meadowlark, SWIFT) and an achromatic quarter-wave plate (Thorlabs, SAQP05M-700) to simultaneously rotate the polarization of the two focused beams to s-polarization. After entering the TIRF input port of an inverted microscope (Nikon, Eclipse Ti) equipped with input and output side ports and an automated XY translation stage, the beams sequentially passed through a dichroic mirror cube (DM) (Chroma, C174298) and the objective. The fluorescence emitted by the sample was collected by the objective, reflected with the DM, focused by a f = 200 mm tube lens that was internal to the microscope and detected with an electron-multiplying charged-coupled device (EMCCD) camera (Andor, iXon Ultra 888) mounted on the microscope output side port. Supplementary Figs. 1 and 2 present all the elements described earlier. The relationship between the galvanometer voltage and the incident angle was calibrated by measuring light positions and imaging the objective BFP (Supplementary Fig. 11, Supplementary Note 1, and Supplementary Movie 4).
The total magnification between the sample and EMCCD was 250, leading to an imaging pixel size of 52 nm. Although somewhat oversampled, the raw images were binned from 1024 × 1024 to 512 × 512 to meet the Nyquist sampling limit and reduce the acquisition time. The experimental laser power varied from 0.66 W cm⁻² to 1.32 W cm⁻² with reliance on the particular sample and exposure time. Power measurements were performed at the output of the high-NA objective lens by guiding the light into the microscope silicon avalanche power probe (Thorlabs, S170C). The typical exposure time of EMCCD was 50–100 ms with the electron-multiplying gain of 150–300, resulting in a 1–2-s acquisition rate per 3D near-surface volume for a field of view of 312 × 512 pixels. The time consideration among the illumination, exposure, and acquisition caused less photodamage and photobleaching while enabling an optimal SNR and temporal resolution for the given light dose to the sample.

For the multi-color imaging, the MAIM system did not need to correct the chromatic aberration that a spatial light modulator (SLM)-based SIM must consider because of its constraints on pattern generation resulting from the discrete pixel arrangement and the wavelength-dependent nature of SLM. These fixed comparison images were automatically acquired.

Supplementary Figs. 12 and 13 present an example of the control voltages for one frame of the live image sequence shown in Fig. 6. Several features of these voltage signals should be noted, as outlined below:

1. Galvo 1 and 2 comprise Scanning Unit 1 and, control the beam position in one route, while Galvo 3 and 4 comprise Scanning Unit 2 and control the other route. When recording the first half of the image sequence, the TIRF-SIM mode operates, and three pairs of voltages make the orientation of illumination patterns at 0°, 60°, and 120°.
2. We let the piezo stage move forward instead of back and forth to shift the phase to improve the speed of the lateral super-resolution imaging. This is because the push/pull force capacity of the piezo stage is 100/20 N in the operation direction. Moreover, the 12-μm close-loop travel range and 1-nm close-loop resolution are sufficient for the phase shift application.
3. Only Scanning Unit 1 was needed to generate the Ring-TIRF illumination when capturing the second half of the image sequence. Thus, we loaded a relatively high voltage on Galvo 3 and 4 to ensure that the corresponding beam could not enter into the objective. This scheme avoided the mechanical vibration and low switching speed that would have occurred if a shutter was used to control the beam.

Data availability. All data are available from the corresponding authors upon reasonable request.

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References
1. Huang, B., Wang, W., Bates, M. & Zhuang, X. Three-dimensional super-resolution imaging by stochastic optical reconstruction microscopy. Science 319, 810–813 (2008).
2. Pavan, S. R. P. et al. Three-dimensional, single-molecule fluorescence imaging beyond the diffraction limit by using a double-helix point spread function. Proc. Natl. Acad. Sci. USA 106, 2999–2999 (2009).
3. Fia, S., Vaughan, J. C. & Zhuang, X. Isotropic three-dimensional super-resolution imaging with a self-bending point spread function. Nat. Photonics 8, 302–306 (2014).
4. Xu, K., Babcock, H. P. & Zhuang, X. Dual-objective STORM reveals three-dimensional filament organization in the actin cytoskeleton. Nat. Methods 9, 185–188 (2012).
5. Aquino, D. et al. Two-color nanoscopy of three-dimensional volumes by 4Pi detection of stochastically switched fluorophores. Nat. Methods 8, 353–359 (2011).
6. Huang, F. et al. Ultra-high resolution 3D imaging of whole cells. Cell 166, 1028–1040 (2016).

Supplementary methods, technical details, and procedures are given in the Electronic Supplementary Information.

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5. Aquino, D. et al. Two-color nanoscopy of three-dimensional volumes by 4Pi detection of stochastically switched fluorophores. Nat. Methods 8, 353–359 (2011).
6. Huang, F. et al. Ultra-high resolution 3D imaging of whole cells. Cell 166, 1028–1040 (2016).
7. Bourg, N. et al. Direct optical nanoscopy with axially localized detection. Nat. Photonics 8, 387–393 (2015).
8. Schmidt, R. et al. Spherical nanosized focal spot unravels the interior of cells. Nat. Methods 5, 539–544 (2008).
9. Gustafsson, M. G. L. et al. Three-dimensional resolution doubling in wide-field fluorescence microscopy by structured illumination. Biophys. J. 94, 4957–4970 (2008).
10. Siao, L. et al. Rego, E. H. & Gustafsson, M. G. Super-resolution 3D microscopy of live whole cells using structured illumination. Nat. Methods 8, 1044–1046 (2011).
11. Fiolka, R., Shao, L., Rego, E. H., Davidson, M. W. & Gustafsson, M. G. L. Time-lapse two-color 3D imaging of live cells with doubled resolution using structured illumination. Proc. Natl Acad. Sci. USA 109, 5311–5315 (2012).
12. Kraus, F. et al. Quantitative 3D structured illumination microscopy of nuclear structures. Nat. Protoc. 12, 1011–1028 (2017).
13. Boulanger, J. et al. Fast high-resolution 3D total internal reflection fluorescence microscopy by incidence angle scanning and azimuthal averaging. Proc. Natl Acad. Sci. USA 111, 17164–17169 (2014).
14. Stabley, D. R., Oh, T., Simon, S. M., Mattheyses, A. L. & Salaita, K. Real-time fluorescence imaging with 20 nm axial resolution. Nat. Commun. 6, 8307 (2015).
15. Cragg, G. E. & So, P. T. Lateral resolution enhancement with standing evanescent waves. Opt. Lett. 25, 46–48 (2000).
16. Zheng, C. et al. Three-dimensional super-resolved live cell imaging through polarized multi-angle TIRF. Opt. Lett. 43, 1423–1426 (2018).
17. Taguchi, N., Ishihara, N., Jofuku, A., Oka, T. & Mihara, K. Mitotic phosphorylation of dynamin-related GTPase Drp1 participates in mitochondrial fission. J. Biol. Chem. 282, 11521–11529 (2007).
18. Huang, B., Jones, S. A., Brandenburg, B. & Zhuang, X. Whole-cell 3D STORM reveals interactions between cellular structures with nanometer-scale resolution. Nat. Methods 5, 1047–1052 (2008).
19. Han, Y., Li, M., Qiu, F., Zhang, M. & Zhang, Y.-H. Cell-permeable organic fluorescent probes for live-cell long-term super-resolution imaging reveal lysosome-mitochondrion interactions. Nat. Commun. 8, 1307 (2017).
20. Li, D. et al. Extended-resolution structured illumination imaging of endocytic and cytoskeletal dynamics. Science 349, aaa3500 (2015).
21. Winterflood, C. M., Ruckstuhl, T., Verdes, D. & Seeger, S. Nanometer axial resolution by three-dimensional supercritical angle fluorescence microscopy. Phys. Rev. Lett. 105, 108103 (2010).
22. Paszek, M. J. et al. Scanning angle interference microscopy reveals cell dynamics at the nano-scale. Nat. Methods 9, 825–827 (2012).
23. Chizhik, A. L., Rother, J., Gregor, L., Janshoff, A. & Enderlein, J. Metal-induced energy transfer for live cell nanoscopy. Nat. Photonics 8, 124–127 (2014).
24. Dos Santos, M. C., Déturche, R., Vézy, C. & Jaffiol, R. Topography of cells revealed by variable-angle total internal reflection fluorescence microscopy. Biophys. J. 111, 1316–1327 (2016).
25. Arganda-Carreras, I. et al. Trainable Weka Segmentation: a machine learning tool for microscopy pixel classification. Bioinformatics 33, 2424–2426 (2017).

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Author contributions
C.K. and X.L. conceived and initiated the project. Y.C, W.L. and L.X. designed and built the optical system. W.L. and Y.C. designed and performed experiments. W.L., C.Z., Y.H. and R.C. wrote the algorithm. Y.C. and W.L. analyzed the data. Z.Z. and D.Z. wrote the acquisition software. J.F., L.J., Y.X., M.Z. and Y.Z. prepared the experimental samples. C.K. and X.L. guided and supervised the whole project. All authors wrote the paper and contributed to the scientific discussion and revision of the article.

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