

Interferometric fluorescent super-resolution microscopy resolves 3D cellular ultrastructure

Gleb Shtengel^a, James A. Galbraith^b, Catherine G. Galbraith^c, Jennifer Lippincott-Schwartz^{d,1}, Jennifer M. Gillette^d, Suliana Manley^d, Rachid Sougrat^d, Clare M. Waterman^e, Pakorn Kanchanawong^e, Michael W. Davidson^f, Richard D. Fetter^a, and Harald F. Hess^{a,1}

^aHoward Hughes Medical Institute, Janelia Farm Research Campus, Ashburn, VA 20147; ^bNational Institute of Neurological Disorders and Stroke; ^cNational Institute of Dental and Craniofacial Research; ^dCell Biology and Metabolism Branch, National Institute of Child Health and Human Development; and ^eLaboratory of Cell and Tissue Morphodynamics, National Heart Lung and Blood Institute, National Institutes of Health, Bethesda, MD 20892; and ^fNational High Magnetic Field Laboratory and Department of Biological Sciences, Florida State University, Tallahassee, FL 32310

Contributed by Jennifer Lippincott-Schwartz, December 25, 2008 (sent for review November 19, 2008)

Understanding molecular-scale architecture of cells requires determination of 3D locations of specific proteins with accuracy matching their nanometer-length scale. Existing electron and light microscopy techniques are limited either in molecular specificity or resolution. Here, we introduce interferometric photoactivated localization microscopy (iPALM), the combination of photoactivated localization microscopy with single-photon, simultaneous multiphase interferometry that provides sub-20-nm 3D protein localization with optimal molecular specificity. We demonstrate measurement of the 25-nm microtubule diameter, resolve the dorsal and ventral plasma membranes, and visualize the arrangement of integrin receptors within endoplasmic reticulum and adhesion complexes, 3D protein organization previously resolved only by electron microscopy. iPALM thus closes the gap between electron tomography and light microscopy, enabling both molecular specification and resolution of cellular nanoarchitecture.

fluorescence microscopy | interferometry | PALM | photoactivated localization microscopy | single molecule imaging

A fundamental question in biomedical research is how specific, nanometer-scale biomolecules are organized into multicomponent micron-scale structural and signaling ensembles that facilitate cell function. For example, microtubules are built of 8-nm tubulin subunits that incorporate on the ultrastructural level into polymers 25 nm in diameter and $>10\ \mu\text{m}$ in length that serve as the building blocks of superstructures such as mitotic spindles and flagella. However, key challenges remain for determining cellular ultrastructure with high molecular specificity. Because cellular structures are organized on the nanoscale, nanometer resolution is required. Immunoelectron microscopy (EM)-based approaches provide the necessary resolution, but they lack robust molecular specificity because the large size of the antibodies hampers their penetration into dense structures and the specificity of the antibody can be compromised by cross-reactivity and epitope masking caused by the harsh fixation often used for high-resolution EM. Fluorescence microscopy coupled with fluorescent protein (FP) fusion technology enables imaging cellular structure with exquisite molecular specificity, but the resolution of 3D images is diffraction-limited to $\approx 200\ \text{nm}$ in the lateral and $\approx 500\ \text{nm}$ in the axial direction, limiting conventional fluorescence to the characterization of cellular superstructure. Some of the recent fluorescence-based super-resolution microscopy techniques (1–5) demonstrated a resolution of $<100\ \text{nm}$ in the vertical direction; however, this is still insufficient to bridge the resolution gap between cellular ultrastructure and superstructure. To achieve near-ultrastructural 3D resolution even for the limited photon outputs of high-molecular-specificity FPs, we have developed a single-photon multiphase interferometric scheme and integrated it with a lateral photoactivated localization microscopy (PALM) (6),

resulting in a highly photon-efficient system called interferometric PALM (iPALM).

Quantitative high-precision positional measurements are commonly made with phase-based interferometry (7), where coherent light waves from 2 optical paths (a reference path and a measured path) are combined to create a signal that modulates on the length scale of an optical wavelength. Interferometry is widely applied to position objects to 10^{-10} -meter accuracy, measure optical surfaces, and even scout for gravity waves with 10^{-18} -meter sensitivity. Interferometric microscopes (8, 9) when combined with phase-shifting techniques (10) can form sub-nanometer-height images and are routinely used in industry to qualify semiconductor processing or hard disk drive head profiles. However, the benefit of such a quantitative interferometric method has so far been elusive for biological systems, despite the wealth of open questions that hinge on the 3D molecular organization at the nanometer scale. Most biological imaging applications use Nomarski differential interference contrast, Zernike phase contrast, or interference reflection microscopy, which harness qualitative interferometric contrast that depends on the refractive index variations within biological materials and forgoes the molecular specificity afforded by fluorescent labeling. Other more recent concepts, such as 4π (11) and I^2M (12), use interference to create a more precise vertical focus (point-spread function) but do not extract explicit phase values from the interfered signal, a critical advantage that is exploited by phase-shifting techniques. One quantitative approach (13) measures the phase of a fluorescent source but requires multiple replicas of the same sample structure over a patterned silicon substrate, a very limiting requirement for bioimaging.

The obstacles to the precise measurement of biological samples by phase-shifting interferometry techniques may seem fundamental. Cells, tissues, or other biological materials typically lack single well-defined reflecting surfaces and interact with visible light weakly and with low molecular specificity. On the other hand, fluorescence provides superb molecular specificity but poses a new problem for interferometry: how to form 2 path-dependent phase coherent beams that can undergo interference. The brightness of fluorescent labels is also highly variable because of photobleaching and other photophysical

Author contributions: G.S., J.A.G., C.G.G., J.L.-S., J.M.G., S.M., R.S., C.M.W., P.K., and H.F.H. designed research; G.S., J.A.G., C.G.G., J.M.G., S.M., R.S., P.K., R.D.F., and H.F.H. performed research; G.S., J.A.G., C.G.G., J.M.G., S.M., P.K., M.W.D., and H.F.H. contributed new reagents/analytic tools; G.S., J.A.G., C.G.G., J.M.G., S.M., P.K., and H.F.H. analyzed data; and G.S., J.A.G., C.G.G., J.L.-S., J.M.G., S.M., C.M.W., P.K., M.W.D., R.D.F., and H.F.H. wrote the paper.

The authors declare no conflict of interest.

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¹To whom correspondence may be addressed. E-mail: hessh@janelia.hhmi.org or jlippin@helix.nih.gov.

This article contains supporting information online at www.pnas.org/cgi/content/full/0813131106/DCSupplemental.

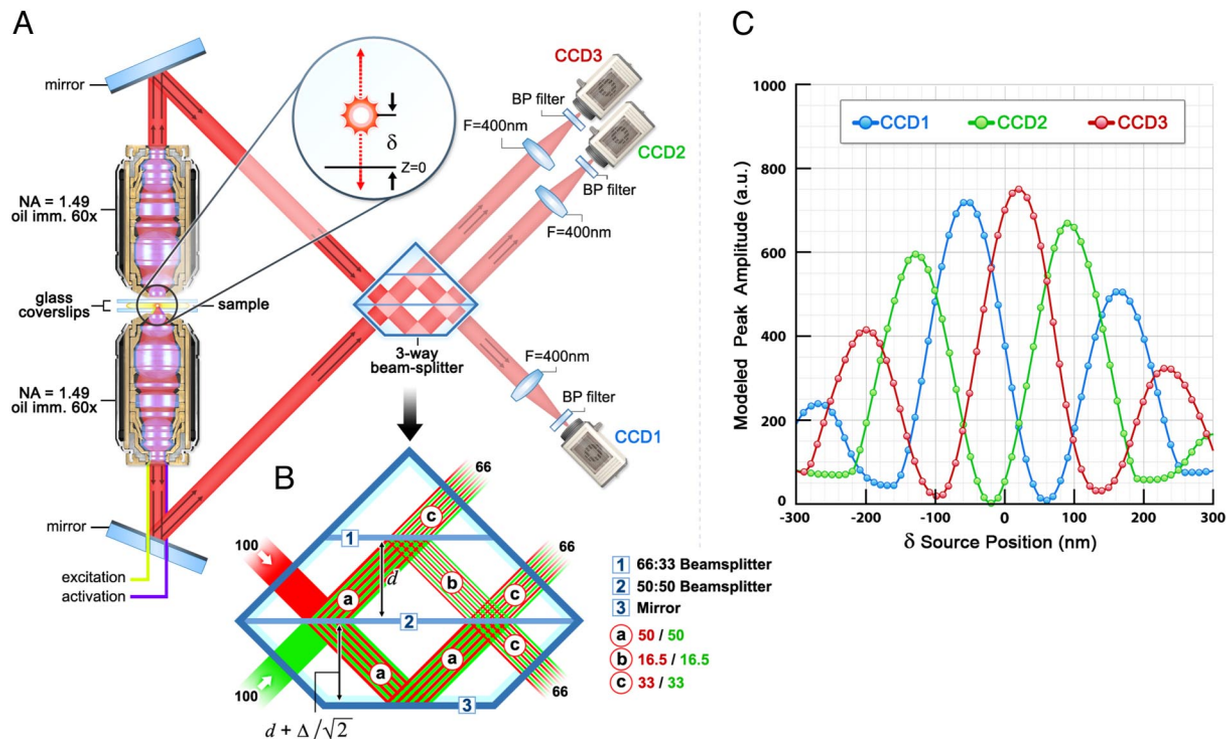


Fig. 1. Schematics and operating principle of multiphase interferometric microscope illustrating how Z-position is resolved. (A and B) Schematic of the single-photon multiphase fluorescence interferometer. A point source with z-position δ emits a single photon both upwards and downwards. These 2 beams (color coded as red and green in B) interfere in a special 3-way beam splitter. (C) The self-interfered photon propagates to the 3 color-coded CCD cameras with amplitudes that oscillate 120° out of phase as indicated.

processes, factors that undermine the ability to take multiple sequential measurements and reliable calibrations that form the basis of phase-shift interferometry.

We have configured an optical system to overcome these obstacles using a combination of insights. First, single fluorescent molecules such as FPs are intrinsic quantum sources. Thus, wave-particle duality allows a single fluorescent photon to form its own coherent reference beam. An emitted photon can simultaneously travel 2 distinct optical paths, which are subsequently recombined so that the photon interferes with itself. The position of the emitter directly determines the difference in the path lengths, hence the relative phase between the 2 beams. This principle has been used in fluorescence interferometry (13, 14) and spectral self-interference fluorescence microscopy (15). Second, the optical system can be configured so that interference can take place over a wide range of lateral source positions to form interference images on an area array detector, such as a CCD for rapid parallel acquisition. Finally and most importantly, *simultaneous multiphase* detection must be implemented for each photon. Such detection methodology is tolerant to intensity variations and short fluorescent lifetimes; it ensures proper self-calibration, and provides complete information to extract a position-dependent interference phase angle.

Results and Discussion

The essence of the single-photon interferometric fluorescence imaging concept is illustrated in Fig. 1, and core to this concept is a custom 3-way beam splitter. Three-way or even higher-order beam splitters are commonly used in a single-mode context (16–18) such as fiber optic gyroscopes, planar waveguide devices for telecommunications, or precision metrology of an object's position. Our device extends this multiphase interference beyond the single-mode, single-point case to the multiphase, and multiple-source case necessary for imaging biological samples. It is composed of 3

parallel planes of a 66:33 beam splitter, a 50:50 beam splitter, and a mirror (Fig. 1B). Self-interference of a single-photon source (such as a fluorescent protein in a sample) placed at the focal plane of 2 opposed objectives enables a precise determination of its axial position. Upon radiation, a fluorescent photon simultaneously enters both the upper and lower objectives, as illustrated in Fig. 1A. The difference in path lengths of the upper and lower beams directly depends on the axial position of the source. Because the photon self-interferes in the 3-way beam splitter, this position-dependent phase difference modulates the relative intensities of the 3 output beams as shown in Fig. 1C. Therefore, the axial position of the source molecule can be determined from the relative amplitudes of the source images from the 3 cameras. Localized lateral positions of a source molecule can be extracted by finding their centers from the same image set so that no extra data acquisition is needed to obtain the full 3D data coordinates. The mathematical description is presented below. The experimental validation of the performance of iPALM, as well as a tutorial demonstrating the interferometry concept are presented in [supporting information \(SI\)](#).

The resolution capability of iPALM for determining the 3D location on the length scale of biomolecules is presented in Fig. 2. The localization accuracy of a single fluorescent source imaged repetitively for 25,000 frames is shown in Fig. 2A. Even with only 1,500 photons into a 4π solid angle (1,200 photons detected), iPALM resolves the position of the source with full width at half maximum (FWHM) of 9.8 nm axially. The lateral position is derived from the centroid of the image spot (19) and the lateral localization accuracy of this same source has FWHM of 22.8 nm. Fig. 2B compares the dependence of localization accuracy on source brightness for iPALM compared with superresolution defocusing approaches (2, 3, 20, 21). Localization accuracy depends critically on source brightness, and scales approximately as the inverse square root of the number of

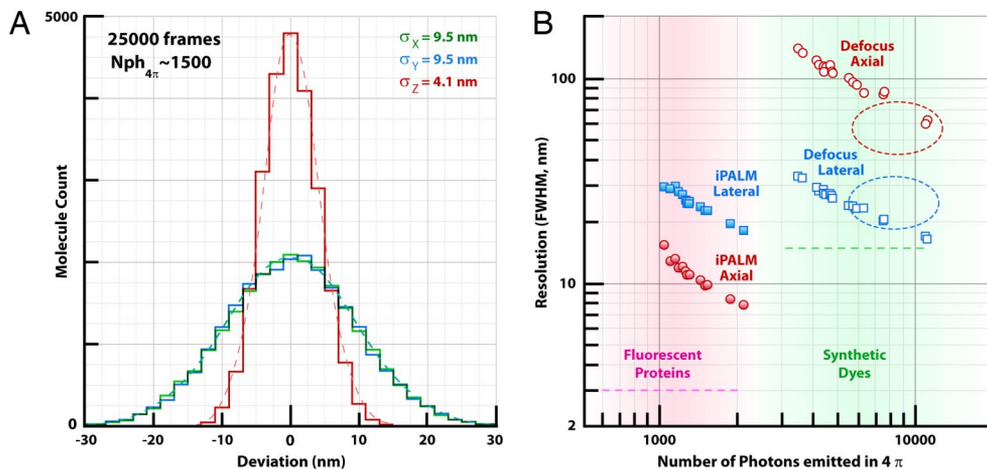


Fig. 2. X, Y, Z resolution of iPALM and its dependence on source brightness, illustrating iPALM's sub-20-nm 3D resolution with endogenous FP labels. (A) A histogram of experimentally determined positions from repeatedly sampling (25,000 frames) a source where $\approx 1,200$ photons are detected per frame from $\approx 1,500$ photons emitted into a 4π solid angle. (B) Axial (solid red circles) and lateral (solid blue squares) resolution of iPALM determined from FWHM of localization of Au beads of different brightness. Note that the positional FWHM number is 2.4 times larger than σ the variance that is also used to characterize resolution. Axial (empty red circles) and lateral (empty blue squares) resolution of the defocusing method determined from FWHM of localized position of Au beads of different brightness. Large ovals indicate approximately the published results for axial (red) and lateral (blue) resolutions of 3D STORM (2) and BP PALM (3). The typical photon output of fluorescent protein tags and synthetic fluorophores are depicted as pink and green gradients. Also shown (horizontal dashed lines) are additional uncertainties resulting from the displacement between the target protein and the fluorescent probes for different imaging methods.

photons detected. This illustrates that iPALM provides 10-fold improvement in axial resolution and 100-fold improvement in photon efficiency compared with the defocus-based techniques (Fig. 2B). Thus, where defocusing methods require antibody-based or cytochemical methods for labeling proteins with brighter organic fluorochromes, iPALM allows accurate localization of FPs even with their modest photon output (typically 500 detected photons for a 2π solid angle). In addition to the advantages of a genetically encoded fluorescent label for specificity, the small size of FPs (3–4 nm) compared with antibodies (>10 nm) confers improvements in spatial sampling density and significantly reduces the localization uncertainty because of probe size, which, in the case of antibody, can introduce up to 15 nm in additional FWHM. These additional uncertainties are illustrated by dotted lines in Fig. 2B. An additional benefit of iPALM compared with defocus-based methods is a factor of 2 better lateral resolution because of the factor of 2 (dual objectives) better photon collection efficiency and the optimal image focusing (2, 3).

We demonstrate the 3D superresolution capability of iPALM on fixed cells expressing photoswitchable or photoactivatable FP fusion proteins. The multiphase interferometric microscopy (the *i* in iPALM) recovers the *z*-axis position, and simultaneously PALM (6, 22, 23) recovers the *x*-*y* coordinates on high densities of FP-fusion molecules that occur in cellular structures. Typically, >20,000 frame triplets of individually imaged photoactivatable FPs are acquired, which capture different proteins with 10–100 labeled molecules per frame. In this manner, a compiled list of 100,000–2,000,000 molecular locations can be combined to form the basis for a 3D volume rendering of protein distributions. iPALM thereby achieves 3D protein-specific contrast images at the size scale associated with electron tomography.

To demonstrate the resolution and sensitivity of iPALM, we imaged well-characterized cellular ultrastructure. Electron microscopy has established that microtubules, which serve as a polarized structural scaffold within cells, have a 25 nm diameter (24). Previous fluorescent-based superresolution 3D imaging approaches (2–4) have lacked the resolution to demonstrate this size. With iPALM, we have resolved the diameter of the microtubules to nearly their known dimension along the *z* axis.

This is illustrated in the iPALM image Fig. 3 of a PtK1 cell expressing human α -tubulin fused to a monomeric variant of the fluorescent protein KikGR (25). The color-coded height image of multiple microtubules crossing each other vertically shows individual microtubules as distinct colors (Fig. 3C). A *z* axis histogram (Fig. 3D) for 2 of the microtubules crossing within 70 nm of each other (boxed region in Fig. 3A) shows a FWHM of 25–30 nm for each microtubule. This size is in good agreement with the dimension of the microtubule itself plus the size of the

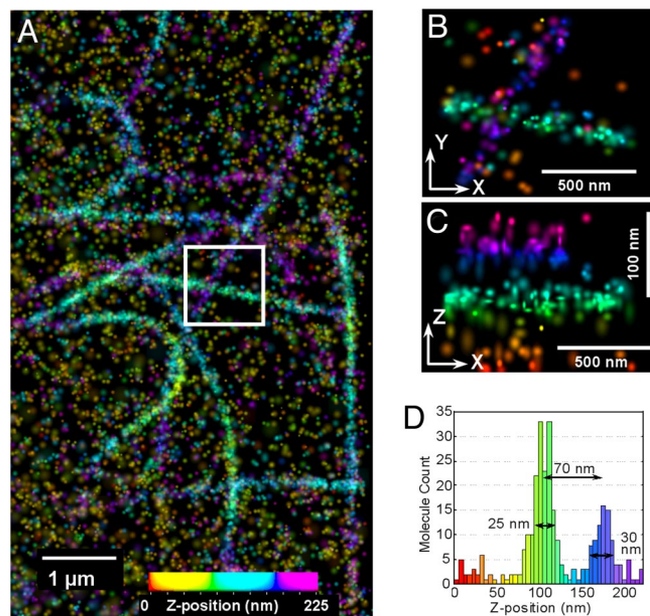


Fig. 3. Superresolution iPALM image of microtubules in a PtK1 cell expressing human α -tubulin fused to *m*-KikGR, rendered with *z* axis color-coding. (A) Large area overview. (B and C) Zoom-in of the area bound by the white box in X-Y (B) and Z-Y (C) projections (*z*-scale is magnified 5 \times). (D) Histogram of *z*-positions of molecules in the boxed region. Each microtubule has a FWHM of 30 nm, and the separation distance of 70 nm between the cyan and purple microtubules is easily resolved.

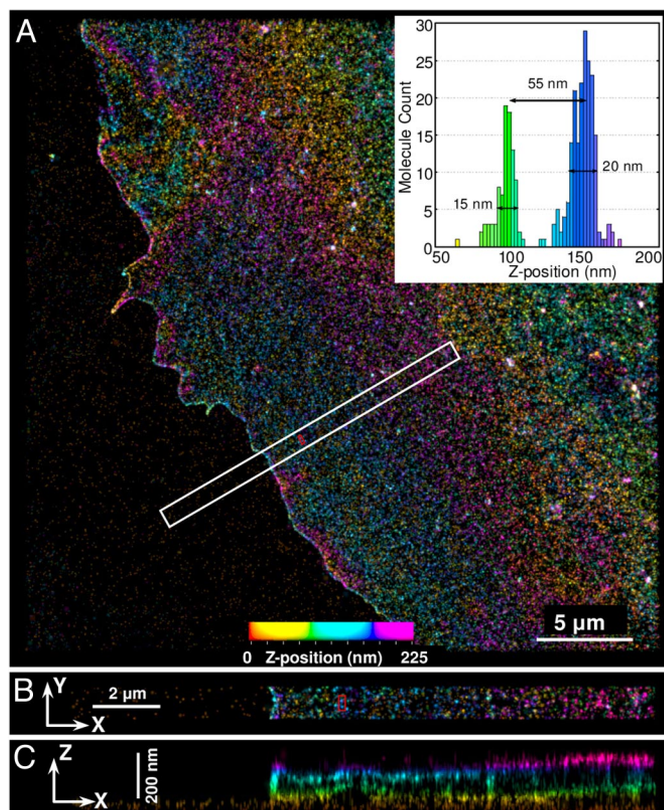


Fig. 4. Superresolution iPALM image of COS7 cell expressing the membrane protein VSVG fused to td-EosFP, rendered with z axis color-coding. (A) Large area overview. (B and C) Area outlined in white is enlarged in B and shown in C as a z cross-section. Shown in the *Inset* is the histogram of vertical distribution of fluorescent molecules in the area limited by the red rectangle.

FP (24). This illustrates the power of iPALM to reveal the 3D morphology with high molecular specificity on a scale relevant for understanding how these proteins are organized and function with other molecular partners within the cell.

Another class of biological structures where iPALM can provide insights is the plasma membrane system, which serves as the boundary between the cytoplasm and extracellular environment. To visualize the plasma membrane, COS-7 cells were transiently transfected with the transmembrane vesicular stomatitis virus G protein (VSVG) tagged with tandem-dimer-Eos FP (tdEosFP) (26). The vertical position of VSVG molecules in the plasma membrane over a 225-nm range is represented by a color hue scale (Fig. 4). A cross-section of the area bound by the white rectangle separately resolves both the dorsal and ventral plasma membrane, even for spacings ≈ 60 nm and shows a cell surface protrusion with an average thickness of 110–160 nm (Fig. 4C), consistent with electron microscopy (27). The lateral dimensions of the displayed area are an order of magnitude wider than the vertical scale so the vertical variations because of membrane roughness manifest as increased apparent membrane thickness in the cross-section. Atomic force microscopy (AFM) measures membrane roughness with the same degree of precision (28), however, with iPALM, we are able to access the topology of the membrane opposing the coverslip with equal precision, which AFM cannot. With the axial resolution provided by iPALM, issues such as plasma membrane polarity and endocytic protein organization can now be investigated at the molecular level.

The molecular precision and spatial resolution of iPALM are especially useful for probing the endomembrane system and its interaction with other cellular structures, particularly in the axial

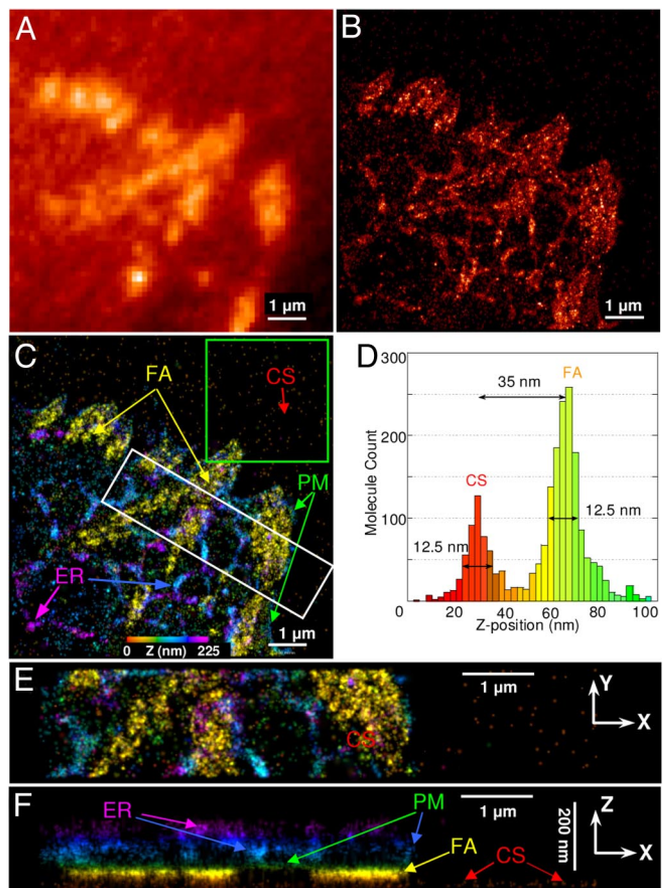


Fig. 5. U2OS cell expressing td-EosFP- α_v -integrin. (A–C) Widefield (A) and PALM (B) images, and z color-coded iPALM image (C). The coverslip surface (CS), focal adhesion (FA), cell plasma membrane (PM), and endoplasmic reticulum (ER) can be identified. (D) Z-position histogram of area limited by the green box in C with peaks corresponding to CS and FA. (E and F) X–Y (E) and X–Z (F) projections of the area bound by the white box in C. Z-scale is magnified by 4 in F.

dimension. The 3D mapping of another membrane protein, α_v integrin fused with tdEos, coexpressed with untagged β_1 integrin in U2OS cells, demonstrates this capability (Fig. 5). Integrins are $\alpha\beta$ heterodimeric transplasma-membrane receptors that cluster in the membrane to form focal adhesions (FAs), where they bind to the extra extracellular matrix and serve as mechanical and signaling linkages between the cells and their environment. The iPALM image (Fig. 5C) with color-encoded vertical position clearly identify those integrin α_v molecules that are incorporated into FAs (yellow in Fig. 5 C, E, and F), whereas a sparser distribution of α_v integrins outline the contour of the plasma membrane. Furthermore, a network of tubular structures is observed at >70 nm above the coverslip plane, which reveals the endoplasmic reticulum (ER) decorated by newly synthesized α_v integrin. The iPALM image clearly identifies finger-like ER structures juxtaposing FA, suggestive of their involvement in integrin delivery (see SI for a movie clearly resolving the ER lumen). Within an FA, α_v integrin occupies a very narrow height distribution of <15 -nm FWHM at a vertical displacement of ≈ 35 nm above the coverslip surface, in good agreement with the cell–substratum separation of ≈ 15 –20 nm, known from electron or interference reflection microscopy (27, 29). Together with the plasma membrane thickness and the position of the tdEos FP tag at the cytoplasmic domain of the integrin α_v , it is also in agreement with the distance between integrin-based adhesions and coverglass determined by interference reflection micros-

the derivations in ref. 19, we obtain the expression for the standard deviation of localization along z axis:

$$\sigma_{z-iPALM} = \frac{s_z - iPALM}{\sqrt{N_{tot}}}, \text{ where } s_z - iPALM = \frac{\lambda(1 + NA/8)}{4\pi n}. \quad [7]$$

For aqueous sample medium with $n \approx 1.33$, $\lambda = 590$ nm, and for $NA = 1.4$, Eqs. 5 and 7 give the localization accuracies of $\sigma_{x,y} \approx 100$ nm/ $\sqrt{N_{tot}}$ and $\sigma_z \approx 40$ nm/ $\sqrt{N_{tot}}$.

It should be pointed out that the only approximation made in deriving Eq. 7 is the omission of the background noise, so when compared with localization accuracy in x - y plane it should be compared with the first 2 terms in Eq. 4.

The contribution of the background noise (third term in Eq. 4) scales as s^4 , so its relative contribution is smaller for z axis localization than that x - y -plane localization.

Biological samples do have index variations that can give additional phase shifts to the radiating photons that depend on

direction and the optical index along that path. This gives rise to slowly varying (over micron lateral scale) distortions. For example the position of molecules on the coverslip surface appear not flat but bowed down by 2–5 nm under a cell of Fig. 4 or Fig. 5. This results from a higher index cell in a lower index aqueous media. However, a reference plane such as fluorescent layer localized on a glass surface allows one to monitor the error from index changes and offers a reference by which to correct for it.

The details of experimental setup and sample preparation are given in SI.

ACKNOWLEDGMENTS. We thank J. Wiedenmann and U. Nienhaus (Ulm University, Ulm, Germany) for the gift of the EosFP, A. Miyawaki (RIKEN, Wako, Saitama, Japan) for the *m*-KikGRFP, Patrick Lee for mechanical design of the iPALM, Hari Shroff and Eric Betzig for help and discussions, Ann McEvoy and Derek Greenfield (University of California, Berkeley) for intriguing 3D samples, and Kevin McGowan and Helen White for help with samples. This work was supported by the Howard Hughes Medical Institute and the National Institutes of Health intramural programs of the National Institute of Neurological Disorders and Stroke, the National Institute of Dental and Craniofacial Research, the National Institute of Child Health and Human Development, and the National Heart, Lung, and Blood Institute.

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