Imaging cellular ultrastructures using expansion microscopy (U-ExM)

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Determining the structure and composition of macromolecular assemblies is a major challenge in biology. Here we describe ultrastructure expansion microscopy (U-ExM), an extension of expansion microscopy that allows the visualization of preserved ultrastructures by optical microscopy. This method allows for near-native expansion of diverse structures in vitro and in cells; when combined with super-resolution microscopy, it unveiled details of ultrastructural organization, such as centriolar chirality, that could otherwise be observed only by electron microscopy.

Cells comprise organelles, large macromolecular assemblies displaying specific structures that for decades could be visualized only by electron microscopy. Although super-resolution fluorescence microscopy has evolved as a powerful tool for subdiffraction-resolution fluorescence imaging of cells, the visualization of ultrastructural details of macromolecular assemblies remains challenging.

Recently, an innovative method called expansion microscopy (ExM) emerged in which immunolabeled samples are physically expanded, and thus can undergo super-resolution imaging by standard fluorescence microscopy (Supplementary Fig. 1a). Alternative ExM protocols such as protein-retention ExM and magnified analysis of the proteome (MAP) have been developed to obtain isotropic macromolecular expansion. Moreover, the tubulin signal appeared inhomogeneous, probably because of epitope masking when stained with both antibodies recognizing epitopes on the central region of the centriole (Supplementary Fig. 1c and 2, and Methods), suggesting an anisotropic macromolecular expansion. As a consequence, the ninefold symmetry of the PolyE-labeled samples was not apparent (Fig. 1c, Supplementary Fig. 3e). However, we observed a reduction in antibody competition (Fig. 1c).

Next, we expanded centrioles with both ExM and MAP protocols and imaged the samples by confocal microscopy followed by HYvolution (Fig. 1b,c). The gels expanded ~4.2-fold (ExM) and ~3.5-fold (MAP). We noticed that the diameter of the centriole in ExM-expanded samples was markedly larger than expected from the determined expansion factor. Indeed, the PolyE signal showed a 1.4x enlargement with an average centriole diameter of 308 ± 42 nm after expansion, compared with the diameter of 216 ± 17 nm determined from non-expanded centrioles imaged by direct stochastic optical reconstruction microscopy (dSTORM) (Fig. 1c, Supplementary Figs. 1c and 2, and Methods).

On the basis of these results, we set out to develop a new method of ExM that could preserve the overall ultrastructure of isolated organelles. Capitalizing on the MAP protocol, we found that avoiding fixation and using a combination of low concentrations of formaldehyde (FA; 0.3–1%) and acrylamide (AA; 0.15–1%) resulted in anisotropic macro-molecular expansion. Moreover, we were able to alleviate the antibody competition, as demonstrated by central core decoration of the PolyE signal with retention of complete tubulin decoration of the centriolar wall.
Fig. 1 | Centriole expansion with U-ExM. a–d. Non-expanded (a) and expanded (b–d) isolated centrioles stained for PolyE (green; Alexa Fluor 488) and α-tubulin (magenta; Alexa Fluor 568) and imaged by confocal microscopy followed by Hyvolution. Centrioles were expanded by ExM (b), MAP (c), or U-ExM (d). Scale bars, 100 nm (a) or 450 nm (b–d). Representative images from 2 (a) or 3 (b–d) independent experiments are shown. e, Diameter of the centrioles in the different conditions. Green and magenta dots represent PolyE and α-tubulin diameters, respectively. Averages and s.d. are as follows. PolyE: 308 ± 42 nm, 133 ± 27 nm, 225 ± 15 nm, and 216 ± 17 nm for ExM, MAP, U-ExM, and non-expanded dSTORM, respectively. n = 30 centrioles for each condition (data from 3 independent experiments) except dSTORM, where n = 15 non-expanded centrioles (1 experiment). α-tubulin: 279 ± 29 nm, 130 ± 32 nm, and 195 ± 12 nm for ExM (n = 29 centrioles), MAP (n = 20 centrioles), and U-ExM (n = 29 centrioles), respectively. Data from 3 independent experiments. Statistical significance was assessed by one-way ANOVA; ****P < 0.0001, ns (nonsignificant) = 0.77. f, Isotropic expansion measured as the ratio of the centriole length and diameter. Average ratios and s.d. are as follows: ExM, 1.8 ± 0.6 (n = 30 centrioles); MAP, 1.9 ± 0.9 (n = 30 centrioles); U-ExM, 2.6 ± 0.3 (n = 29 centrioles); non-expanded structured illumination microscopy (SIM), 2.6 ± 0.2 (n = 22 centrioles). Data from 3 independent experiments except for SIM, where they are from a single experiment. Statistical significance was assessed by one-way ANOVA. ****P < 0.0001; ***P = 0.0002; ns (non-significant) = 0.84 for ExM versus MAP and 0.99 for U-ExM versus non-expanded SIM.
B-tubule facing the flagellar lumen of *Chlamydomonas* axonemes, as previously proposed for both *Chlamydomonas* and *Tetrahymena* (Supplementary Fig. 11g–q).

We then asked whether other dynamic cellular structures can be successfully expanded with U-ExM. We first tested different fixation conditions combined with U-ExM on isolated centrioles to assess structural preservation. We found that centrioles fixed with FA or methanol had good overall structural preservation but a reduced centriole diameter, while fixation with paraformaldehyde (PFA) and glutaraldehyde (GA) did not allow full expansion of centrioles (Supplementary Fig. 12). Then we analyzed the effects of the different fixation conditions followed by incubation in AA–FA solution on human cells. We found that all fixation conditions tested preserved microtubules, and that fixation with both PFA and GA was best suited for structural preservation of mitochondria (Supplementary Fig. 13). Thus, we carried out U-ExM with fixed mammalian cells and analyzed microtubules, mitochondria, and clathrin–coated pits as another membrane-bound structure. We found that U-ExM nicely expanded methanol-fixed microtubules with a full width at half-maximum (FWHM) of 46 nm, in agreement with results from previous ExM methods (Supplementary Fig. 14, 15). Similarly, mitochondria fixed with PFA and GA could be expanded via U-ExM with good structural organization, with the outer mitochondrial membrane translocase TOMM20 surrounding the overall MitoTracker signal (Fig. 3e–h and Supplementary Fig. 16). For both microtubules and mitochondria, U-ExM showed uniform expansion with minimal distortions of 1.6% (microtubules) and 5% (mitochondria), similar to observations from other expansion methods (Supplementary Fig. 17). Finally, we found that FA-fixed clathrin-coated pits could also be visualized as hollow vesicles with U-ExM (Supplementary Fig. 18).

Our results show that ExM protocols have to be carefully optimized to enable isotropic expansion of molecular assemblies. We have demonstrated that U-ExM preserves ultrastructural details and can thus be used successfully to visualize the molecular architecture of diverse multiprotein complexes. In comparison with standard ExM protocols, U-ExM alleviates antibody competition
and prevents fluorophore loss due to post-expansion labeling. By avoiding chemical fixation of isolated protein complexes, U-ExM improves structural integrity, as demonstrated with isolated centrioles. Notably, in standard ExM approaches, the relative distance of the fluorophore to the epitope stays unchanged, whereas post-expansion labeling approaches led to a relatively smaller antibody size compared with that in the expanded sample. Thus, U-ExM coupled with STED imaging can unveil the chirality of the centriole, a structural feature that previously was revealed by super-resolution microscopy only in appendage proteins radiating 50–100 nm out of the centriole18. We are convinced that in the near future U-ExM will be combined with single-molecule localization microscopy to enable fluorescence imaging of molecular details with unsurpassed spatial resolution.

Fig. 3 | U-ExM applied to human cells. a, Representative HyVolution (HyV) confocal image of a U2OS cell fixed with methanol, expanded with U-ExM, and stained for α-tubulin (magenta) and PolyE (green). Scale bar, 10 μm. b–d, Magnified views of the centriolar pair visible within the dotted square in a. P, procentriole; M, mature centriole. Scale bar, 2 μm. Representative images from 3 independent experiments. e–h, Representative HyVolution confocal images of a U2OS cell fixed with PFA + GA and stained for MitoTracker (orange) and the outer membrane mitochondrial translocase TOMM20 (cyan). Scale bars, 12 μm (e) or 3 μm (f–h). The dotted square in e outlines the region shown at higher magnification in f–h. Note that as expected, the TOMM20 signal surrounds the MitoTracker signal. Representative images from 1 experiment.

Online content
Any additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at https://doi.org/10.1038/s41592-018-0238-1.

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D.G., F.U.Z., M.S., V.H., and P.G. conceived and designed the project. M.S., V.H., and P.G. supervised the project. D.G. and M.S.-C. are supported by the European Research Council (ERC; PC66 ReceptorLight (projects A04 and B04)). M.U. is supported by the ERC (GA No. 692726 GlobalBioIm).
Methods

Reagents. FA (36.5–38%; F8775), sodium acrylate (SA; 97–99%; 408220), guanidine hydrochloride (8 M; G7294), AA (40; A0658), N2, N3-methylenebisacrylamide (BIS; 2%; M1533), PIPES (P6757), and poly-d-lysine (A-003-E) were purchased from Sigma-Aldrich, Proteinase K (>1000 U/ml; E00491), ammonium persulfate (APS; 18774), tetramethylethylenediamine (TEMED; 17919), DMAP supplemented with Glutamax (61965), FBS (10270), and penicillin–streptomycin (15140) were obtained from Thermo Fisher. We used DMAP/HA’s F12 with l-glutamine (Sigma; D8062) supplemented with 10% FBS (Sigma; F7524), penicillin (100 U/ml) and streptomycin (100 μg/ml; Sigma; R7587) and GA (25%; 16200) were purchased from Electron Microscopy Sciences. Sodium dodecyl sulfate (SDS), Triton X-100, and Tween-20 were obtained from AppliChem, and Tris was obtained from Biosolve. Nuclease-free water (U-ExM protocols and kept as aliquots at –20°C. Mposts were washed with absolute ethanol and subsequently dried. Next, coverslips were coated with poly-d-lysine (0.1 mg/ml) and GA (2%; 15700) and GA (25%; 16200) were purchased from Electron Microscopy Sciences. NaCl in 1× TAE buffer, 0.2% SDS, 20% (wt/wt) GA (25%; 16200) were purchased from Electron Microscopy Sciences. Nuclease-free water was added last, was placed on the chilled Parafilm, and coverslips were carefully mounted on a glass slide with 6 μl of DABCO containing mounting medium.

Expansion microscopy (ExM) protocol. Centrioles were processed as indicated above for immunofluorescence. After the last PBS wash, coverslips were incubated for 10 min at RT in 0.25% GA in PBS in a six-well plate, washed in PBS three times for 5 min, and then processed for gelation. A small plastic box was covered with Parafilm and put on ice to create a flat hydrophobic surface for gelation. A drop of 35 μl of ExM MS (8.625% (wt/wt) SA, 20% (wt/wt) AA, 0.075% (wt/wt) BIS, 2 M NaCl in 1× PBS) supplemented with 0.2% APS and 0.2% TEMED, with the initiator (APS) added last, was placed on the chilled Parafilm, and coverslips were carefully mounted on the coverslips facing the gelling solution. Gelation proceeded for 1 min on ice, and then samples were incubated at 37°C in the dark for 1 h. Then coverslips with attached gels were transferred into a six-well plate for incubation in 2 ml of digestion buffer (1x TAE buffer, 0.5% Triton X-100, 0.8 M guanidine hydrochloride, pH 8.3) supplemented with fresh primary antibody solution diluted in 2% PBS/BSA at 45 min at 37°C. Finally, gels were removed with tweezers from the coverslips and placed in beakers filled with ddH2O for expansion. Water was exchanged at least twice every 30 min, and then samples were incubated in ddH2O overnight at RT. Gels expanded between 4x and 4.2x according to SA purity.

MAP protocol. Coverslips with isolated centrioles were incubated in a solution of 4% PFA with 30% AA in PBS for 4–5 h at 37°C, without a fixation step. Incubation time in PFA–AA was shortened compared with that in the original ‘cultured cell’ MAP protocol to allow the approach to be adapted to smaller specimens such as isolated centrioles. Immediately after PFA–AA incubation, gelation was carried out as described above for the ExM protocol. Coverslips with centrioles facing down were placed on 35 μl of MAP MS (7% (wt/wt) SA, 20% (wt/wt) AA, 0.1% (wt/wt) BIS in 1x PBS) supplemented with 0.5% APS and 0.5% TEMED, with the initiator (APS) added last, on Parafilm in a precooled humid chamber. Gelation proceeded for 1 min on ice and then shifted to 37°C for 10 min in the dark. Coverslips with gels were placed in 2 ml of denaturation buffer (200 mM SDS, 200 mM NaCl, and 50 mM Tris in nuclease-free water, pH 9) in a six-well-plate for 15 min at RT. Gels were then removed from the coverslips with tweezers, moved into a 1.5-ml Eppendorf centrifuge tube filled with fresh denaturation buffer, and incubated at 95°C for 30 min. After denaturation, gels were placed in beakers filled with ddH2O for 45 min, and water was exchanged once. Gels were then washed in PBS with 0.1% Tween 20 (PBST) three times for 10 min with shaking, and then incubated with secondary antibody solution diluted in 2% PBS/BSA for 6–7 h at RT with gentle shaking. Gels were then washed in PBST three times for 10 min with shaking, and finally were placed in beakers filled with ddH2O for the final expansion. Water was exchanged at least twice every 30 min, and then gels were incubated in ddH2O overnight. Gels expanded between 3.3x and 3.5x according to SA purity.

U-ExM protocol. In U-ExM, the sample was not fixed or was mildly fixed prior to expansion. First, coverslips with unfixed isolated centrioles were incubated in a solution of 0.7% FA with 0.15% or 1% AA in PBS for 4–5 h at 37°C. Next, similar to the ExM and MAP protocols, gelation was carried out via incubation of coverslips with centrioles facing down with 35 μl of U-ExM MS composed of 19% (wt/wt) SA, 10% (wt/wt) AA, 0.1% (wt/wt) BIS in 1x PBS supplemented with 0.5% APS and 0.5% TEMED, on Parafilm in a pre-cooled humid chamber. Note that APS was added last. Importantly, the MS was adapted specifically for U-ExM to achieve an expansion factor of approximately fourfold. Briefly, to find the best expansion conditions, we increased SA and reduced AA concentrations in the MS. We tested the following combinations to compare gel expansion: 20% AA and 7% SA (original MAP MS), 10% AA and 7% SA, 10% AA and 19% SA (U-ExM MS), 5% AA and 7% SA, and 5% AA with 19% SA (Supplementary Fig. 19). Isolated centrioles were then embedded in gels made with MSs with the different AA–SA combinations and expanded to check their quality. The shape of expanded centrioles looked preserved only in gels made with 10% AA (Supplementary Fig. 19). Three independent experiments were performed for each condition.

Gelation proceeded for 1 min on ice, and then samples were incubated at 37°C in the dark for 1 h. Coverslips with gels were then transferred into ~2 ml of denaturation buffer (200 mM SDS, 200 mM NaCl, and 50 mM Tris in ultrapure water, pH 9) in a six-well-plate for 15 min at RT. Gels were then removed from the coverslips with flat tweezers, moved into a 1.5-ml Eppendorf centrifuge tube filled with fresh denaturation buffer, and incubated at 95°C for 30 min. After denaturation, gels were placed in beakers filled with ddH2O for expansion. Water was exchanged at least twice every 30 min at RT, and then gels were incubated with fresh primary antibody solution diluted in 2% PBS/BSA for 1 h at RT in a humid chamber. Coverslips were then washed in PBS three times for 5 min and subsequently incubated for 1 h at RT with 75 μl of the secondary antibody solution diluted in 2% PBS/BSA in a humid chamber, protected from light. Finally, coverslips were washed in PBS three times for 5 min. Coverslips were then mounted on a glass slide with 6 μl of DABCO containing mounting medium.

Chlamydomonas reinhardtii isolation and centrifugation on coverslips. Chlamydomonas reinhardtii strain CW15 and span on coverslips as previously described. Coverslips were then processed either for regular immunofluorescence or for ExM protocols.

Immunofluorescence of non-expanded isolated centrioles. Coverslips with isolated centrioles were fixed with 4% FA in PBS for 10 min at RT and washed in PBS. Coverslips, with centrioles facing down, were then placed on 75 μl of the primary antibody solution diluted in 2% PBS/BSA for 1 h at RT in a humid chamber. Coverslips were then washed in PBS three times for 5 min and subsequently incubated for 1 h at RT with 75 μl of the secondary antibody solution diluted in 2% PBS/BSA in a humid chamber, protected from light. Finally, coverslips were washed in PBS three times for 5 min. Coverslips were then mounted on a glass slide with 6 μl of DABCO containing mounting medium.

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poly-d-lysine-coated coverslip, already inserted in the metallic chamber, and gently pressed with a brush to ensure adherence of the gel to the coverslip. This step is crucial to completely avoid gel drift during imaging. After a few seconds, a couple of drops of dH2O were slowly added on top of the sample until the gel was completely covered with water, to avoid shrinking of the polymer. Confocal microscopy was performed on a Leica TCS SP8 with a 63×/1.4-NA (numerical aperture) oil-immersion objective, using the HyVolution mode to generate deconvoluted images, with the following parameters: ‘HyVolution Grade’ at max resolution, ‘HyPyrens Essential’ as Approach, water as ‘Mounting Medium’, and Best Resolution as ‘Strategy’. 3D + stacks at 0.12-µm intervals were acquired with a pixel size of 35 nm.

For post-U-ExM imaging of clathrin, microtubules, and DNA, we used a re-scanning confocal microscope (RCM), which is based on the image-scanning principle whereby pixel reassignment is achieved purely optomechanically. The RCM unit (Cordiavision) is attached to a side port of the microscope and connected to the RCM unit. The laser unit has four laser lines (405, 488, 561, and 640 nm) with 50 mW each (free beam). Various OD filters are introduced in the excitation path to attenuate the laser power. The TIE is equipped with a motorized stage (Nikon) and a 60× water-immersion objective (CEI Plan APO, 1.27-NA; Nikon). The setup is fully controlled with NIS-Elements version 4.6 on Windows 8.

Post-U-ExM imaging was performed on a commercial STED microscope (Expert Line, Abbevor-Instruments, Germany) working at a repetition rate of 40 MHz (ref. 2). Centrioles were immuno-stained with secondary antibodies conjugated to STAR RED and/or Star 580 dye (Abberior, Germany). STAR RED was imaged with excitation at a wavelength of 640 nm and time-gated fluorescence detection between 650 and 720 nm. STAR 580 was excited at 561 nm with time-gated detection between 580 and 630 nm. The STED laser had a wavelength of 775 nm and a pulse width of roughly 500 ps. The pinhole was set to 0.75 Airy units (AU). For imaging, a water-immersion objective lens was used (UPLSAPO 60XW, Olympus, Japan). To arrive at high fluorescence signals and STED resolutions for a clear structure representation, we used the recently published adaptive-illumination scan technique DyMIN11. Non-expanded centrioles were imaged with a 63×/1.4-NA oil-immersion objective.

dSTORM imaging was conducted on an inverted microscope (Zeiss Axio Observer.Z1, Carl Zeiss Microscopy) equipped with a 100× oil-immersion objective (alpha Plan-Apochromat 100×/1.4 Oil DIC, Carl Zeiss Microscopy) and a 63× water-immersion objective lens (LD C-Apochromat 63×/1.5 W Corr M27, Carl Zeiss Microscopy). For illumination of the sample, a 640-nm diode laser (iBeam smart, Toptica Photonics) was used. The laser beam was adjusted to quasi-TIRF mode for use with the oil-immersion objective and to epifluorescence configuration with the water-immersion objective. Between 20,000 and 40,000 frames were collected on an EM-CCD (electron-multiplying charge-coupled device) camera (Andor Ixon Ultra DU897U-CSO) at a frame rate of 50–80 Hz. An autofocus system (Zeiss Definite Focus) kept the focus stable during image acquisition. For 3D imaging, a cylindrical lens (f = 250 mm) was placed in the detection path of the microscope setup. Samples were placed in freshly prepared photoswitching buffer consisting of 100 mM cysteine hydroxychloride (Sigma), PBS (1×), pH 7.5, supplemented with an scavenger system (2% glucose (w/v), 2 U/ml glucose oxidase (Sigma), and 200 U/ml catalase (Sigma)). 2D super-resolution images were reconstructed with the ImageJ plugin ThunderSTORM14, and for 3D images the open-source software rapidSTORM 3.3 (ref. 15) was used.

Measurements of centriole diameter. We selected only nearly perfectly top-view centrioles for measurement of centriole diameter (Supplementary Fig. 2). Briefly, we included in the analysis only centrioles for which the most distal and most proximal regions were aligned. We used the line scan and plot profile tools of Fiji to determine the diameter. For each centriole, a single z plane (the same z plane used to quantify the centriole diameter) was transformed to polar coordinates with the Polar Transformer plugin, generating ‘unwrapped’ images (Supplementary Fig. 3c). Then a straight line wide enough to cover the whole signal was drawn to obtain the profile. For each centriole, a Nikon S8000 from three independent experiments were merged to create a unique averaged curve as follows. A sinusoidal model was generated to represent a theoretical plot profile of a polar-transformed centriole. This model is described by the following formula:

$$f(x) = 0.5 + 0.5 \sin \left( \frac{2 \pi x}{\text{Period}} \right)$$

where the period can be calculated by dividing the length of the experimental plot profile by 9. To compare an experimental centriole plot profile to this model, we performed the experiment between 0 and 1. This value represented the diameter of the centriole progressively moved along its x axis. For each shift, the cross-correlation between the experimental data and the model was calculated. The shift value giving the best cross-correlation was conserved, and the plot profile was moved according to it. Once all plot profiles had been aligned on the sinusoidal model, an average profile was generated. The model and the cross-correlation search were done using the language R.

Analysis of centriolar shape. We used the shape descriptor tool of Fiji to analyze centriole shape quality. The single z plane already used to measure centriole diameter was also used to analyze centriole shape. Using Polyf, we drew a polygon with nine vertexes around the circumference, joining the nine microtubule triplets when visible. The shape descriptor tool gives several parameters, among them the roundness value, defined as the ratio of minor axis to major axis for the figure. A value of 1 represents a perfectly round shape. For each condition, we analyzed 30 centrioles from three independent experiments.

Comparison between dSTORM and U-ExM. Isolated centrorioles, either non-expanded or expanded via the U-ExM protocol, were stained for PolE and imaged with dSTORM or confocal microscopy, respectively. A straight line that bisected the microtubule triplets of procentrioles was drawn with the line tool in Fiji. Then we used the line scan and plot profile tools to measure the fluorescence profile along this line and obtain the FWHM of the curve for dSTORM (4 non-expanded procentrioles, total of 24 triplets, one experiment), U-ExM + confocal microscopy (9 non-expanded procentrioles, total of 72 triplets, 3 independent experiments) and U-ExM + HyVolution (9 expanded procentrioles, total of 81 triplets, 3 independent experiments). Analysis was performed on the maximum-intensity projection of 3D stack acquisitions. To monitor the ninefold symmetry of the PolE signal, we used the Polar Transformer plugin in Fiji as described above.

Isotropic 3D averaging. We obtained particle averaging results for isolated C. reinhardtii centrioles by using the method described in ref. 15. We used the StackReg plugin18 for data preprocessing to correct drifts between slices of the image stacks. We averaged 14 centrioles from one experiment that we selected with the software ImageJ. To model the point spread function (PSF), we acquired images of 0.1-µm fluorescent beads embedded in a U-ExM gel. We obtained the final PSF volume by registering and averaging 15 images of beads.

The reconstruction was realized in two steps. The first step used as input the PSF model and a restricted number L of particles representing the main orientations to create a coarse initial reconstruction without reference. Because of the cyclical symmetry of the centriole, most of the information can be captured from top and side views, which corresponds to L = 2. We downsampled the input volumes by a factor of 2 to accelerate computations. We used the bi-level and block-coordinate optimization approach described in ref. 22. To further speed up the computations, we replaced the stochastic optimization approach for the estimation of pose parameters by a deterministic search with coarse discretization. The second step refined the result of the first step by considering all the available data and a more accurate model. The volume and particle poses were alternately updated until convergence. In both steps, we applied a C9 symmetry constraint to the reconstructed volumes. The custom Matlab source code is available at https://github.com/dfortu22/U-ExM. The reconstruction code with known angles uses the inverse problem library GlobalBoltz22. The parameters used for the reconstruction of Supplementary Fig. 6 are the default parameters accessible in the code.
Sub-triplet localization analysis. Isolated cilia were expanded via the U-ExM protocol, stained for PolyE and α-tubulin, and imaged with a confocal microscope. Before analysis, deconvolution was applied on images. A straight line that bisected the microtubule triplets from the inside toward the outside was drawn with the line tool in Fiji. Then, the line scan and plot profile tools were used to measure the fluorescence profiles of PolyE and α-tubulin along the same line and normalize them on the highest value (Supplementary Fig. 7a–d). Both curves were aligned on the peak of PolyE as a reference point. N = 8 cilia from 3 independent experiments, for a total of 61 microtubule triplets.

To model the position of the fluorescence signal in the centriole, we scaled a 2D image extracted from cryo-EM data to obtain a centriole with the expanded diameter after expansion. The resulting image gave a centriole with a diameter of 1,125 nm (Supplementary Fig. 7e) (centriole diameter × expansion factor = 250 nm × 4.5 = 1,125 nm). We repeated the same operation for A-, B-, and C-microtubules (100-nm diameter) in Fiji to filter the resolution of HyVolution (140 nm) with a bandpass filter in Imagej (Supplementary Fig. 7j–l). We generated the final images (Supplementary Fig. 7m–o) by merging each specific microtubule signal (Supplementary Fig. 7j–l) on the initial microtubule triplet image (Supplementary Fig. 7i). Sub-triplet localization analysis was done as described above.

Cryo-electron microscopy of Chlamydomonas centrioles. Isolated Chlamydomonas centrioles were applied to lacey carbon film grids (300Mesh, EMS), vitrified in liquid ethane. Grids were transferred to a JEM 2200FS cryo-electron microscope (JEOL) operating at 200 kV and equipped with a field emission gun. Images were collected with a 2,048 × 2,048 CCD camera (Gatan).

Quantification of microtubule-triplet angle. We used the angle tool in Fiji to measure the angle between the center of the centriole and the microtubule triplets. N = 77 microtubule triplets for electron microscopy images and n = 65 for U-ExM. Data from one experiment.

Chlamydomonas culture and expansion. Chlamydomonas cells were cultured in Tris-acetate phosphate medium for 3 d at 23 °C (ref. 23). Cells were allowed to adhere to 12-mm coverslips and incubated overnight at 37 °C with 5% CO₂. Next, coverslips were incubated with 50 nM PolyE-α-tubulin diluted in antibody solution (PBS with 1% BSA and 0.05% Tween) for 1 h at RT in a humid chamber. For condition (1), cells were rapidly pre-extracted in BRBB80 solution with 0.5% Triton and fixed with 3% PFA and 0.1% GA in PBS for 15 min at RT. After fixation, cells were quickly washed in PBS and then incubated for 5 h in AA–FA U-ExM solution at 37 °C. Next, coverslips were incubated with primary antibody diluted in antibody solution (PBS with 1% BSA and 0.05% Tween) for 1 h at RT in a humid and dark chamber. Finally, coverslips were washed in PBS three times for 5 min, quickly dried, and mounted on a glass slide with 3 μl of DABCO containing mounting medium.

Similarly, we tested the effects of fixation on the preservation of mitochondria under four different conditions. Cells were first incubated for 15 min with Mitotoxin Red CMXRos (100 nM, diluted in the culturing medium) and then were stained with rat anti-α-tubulin (YLI2/1) and secondary Cy3 antibodies (Supplementary Fig. 7c) as described above and imaged with a confocal microscope. These experiments were analyzed with Imagej, where the differences in signal intensity between pre- and post-U-ExM images were calculated. To examine the effect of fixation on the preservation of mitochondria, we used mouse anti-α-tubulin (DM1a) and Alexa Fluor 488.

To compare mitochondria pre- and post-U-ExM, we incubated cells for 15 min with Mitotoxin Red CMXRos (100 nM, diluted in the culturing medium), fixed them in 3% PFA and 0.1% GA in PBS for 15 min at RT, and then incubated them in AA–FA U-ExM solution for 5 h at 37 °C. Next, we acquired pre- and post-expansion images. Then the U-ExM protocol was applied similarly as for microtubules, but with denaturation allowed to proceed for 1 h at 70 °C (1.5 h of denaturation completely destroyed mitochondria (data not shown)). For post-expansion staining, rabbit anti-TOMM20 and Alexa Fluor 488 were used. Note that the Mitotoxin signal was retained after expansion, and thus was acquired and used to calculate the r.m.s. error between pre- and post-expansion images.

To recognize the region of the coverslip where cells were cultured, we applied a mark to the opposite side of the coverslip where cells were present. This allowed us to cut the piece of gel including only the cells acquired pre-expansion and facilitated their acquisition post-expansion.

Post-U-ExM analysis of clathrin, microtubules, and DNA. For post-U-ExM labeling of clathrin-coated pits, microtubules, and DNA, we cultured COS-7 African green monkey kidney cells (purchased from CLS Cell Line Service GmbH) in DMEM/HAM's F12 with L-glutamine supplemented with 10% FBS, penicillin (100 U/ml), and streptomycin (0.1 mg/ml) at 37 °C and 5% CO₂. We seeded 30,000 cells per well on round 18-mm high-precision coverslips (12.5 mm × 12.5 mm) in MatTek wells (TPP 30212). Cells were grown for 24 h at 37 °C and 5% CO₂ and subsequently fixed in 4% FA in cytoskeleton buffer at 37 °C for 10 min. After fixation, cells were briefly washed with PBS and then transferred into AA–FA U-ExM solution for 5 h at 37 °C. Next, we acquired pre- and post-expansion images. Then the U-ExM protocol was applied similarly as for microtubules, but with denaturation allowed to proceed for 1 h at 70 °C (1.5 h of denaturation completely destroyed mitochondria (data not shown)). For post-expansion staining, rabbit anti-TOMM20 and Alexa Fluor 488 were used. Note that the Mitotoxin signal was retained after expansion, and thus was acquired and used to calculate the r.m.s. error between pre- and post-expansion images.

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For post-U-ExM labeling of clathrin-coated pits, microtubules, and DNA, we cultured COS-7 African green monkey kidney cells (purchased from CLS Cell Line Service GmbH) in DMEM/HAM's F12 with L-glutamine supplemented with 10% FBS, penicillin (100 U/ml), and streptomycin (0.1 mg/ml) at 37 °C and 5% CO₂. We seeded 30,000 cells per well on round 18-mm high-precision coverslips (12.5 mm × 12.5 mm) in MatTek wells (TPP 30212). Cells were grown for 24 h at 37 °C and 5% CO₂ and subsequently fixed in 4% FA in cytoskeleton buffer at 37 °C for 10 min. After fixation, cells were briefly washed with PBS and then transferred into AA–FA U-ExM solution for 5 h at 37 °C. Gelation and denaturation of the sample were carried out for 3 h in AA–FA U-ExM protocol, with the following differences: gelation was allowed to initiate on ice and denaturation allowed to proceed for 1.5 h (a longer time was chosen for cells than for isolated centrioles to ensure maximal expansion in this complex specimen). For post-expansion staining, we used mouse anti-α-tubulin (DM1a) and Alexa Fluor 488.

Typically, we observed MTDs or doublets and central-pair microtubules to measure the plot profile of the microtubules. Briefly, a cryo-EM image of a cilium was drawn, and the fluorescence profiles of both PolyE and α-tubulin were measured along this line. A curve for each cilium was obtained from the average of two measurements and normalized on the highest value, for both stainings. All curves were then aligned with the center between the two peaks of intensity used as a reference point. N = 23 flagella from 3 independent experiments were analyzed.

The modeling of PolyE on the B-microtubule (half or full) was done similarly to the sub-triplet localization on centrioles. Briefly, a cryo-EM image of a cilium cross-section was scaled to obtain a cilium with a diameter of 900 nm (cilium diameter × expansion factor = 200 nm × 4.5 = 900 nm). These images were then filtered at the resolution of HyVolution (140 nm) with a bandpass filter in Imagej.

Quantification of PolyE signal in Chlamydomonas flagella. Polylglutamylation of doublets and central-pair microtubules of Chlamydomonas flagella was analyzed on sprayed flagella (Supplementary Fig. 11a–f). For each flagellum, using the software Fiji, we drew a line scan of a few micrometers across the nine microtubule doublets (MTDs) and two central-pair microtubules to measure the plot profile of the PolyE signal. Next, we obtained the average of the nine intensity peaks of the MTDs and the average of the two intensity peaks of the central-pair microtubules. Finally, we calculated the ratio between the PolyE signal at the MTDs and that at the central-pair microtubules for each flagellum and obtained the average for four sprayed flagella from one experiment.
The s.e. of the diameter was calculated from the square root of the sum of the squared errors from the center values of the single Gaussian fits.

**Distortion analysis.** To estimate the sample deformation after expansion, we calculated the r.m.s. error between two images of the same structure before and after expansion, following the protocol described by Chozinski et al. This protocol also provides the scale factor between the images, thus giving the expansion factor of the experiment. For both microtubules and mitochondria, the data from three independent experiments were used.

**Statistics and reproducibility.** All experiments were carried out three times independently, unless indicated otherwise in the figure legends. All data are expressed as the average (mean) ± s.d. The n values, which represent the number of centrioles or the number of triplets analyzed, are stated in figure legends and in the Methods. Statistical one-way ANOVA and unpaired two-tailed t-tests were used as indicated in the figure legends.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Code availability.** The custom Matlab source code is available at https://github.com/dfortun2/U-ExM.

**Data availability**

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

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Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

- [ ] The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement.
- [ ] An indication of whether measurements were taken from distinct samples or whether the same sample was measured repeatedly.
- [ ] The statistical test(s) used AND whether they are one- or two-sided.

  - Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- [ ] A description of all covariates tested.

- [ ] A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons.
- [ ] A full description of the statistics including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals).
- [ ] For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted.

  - Give P values as exact values whenever suitable.
- [ ] For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings.
- [ ] For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes.
- [ ] Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated.

- [ ] Clearly defined error bars.

  - State explicitly what error bars represent (e.g. SD, SE, CI)

Software and code

Policy information about availability of computer code.

Data collection

- Centrioles were imaged using a Leica TCS SP8 using a 63x 1.4 NA oil objective, with the HyVolution mode2 to generate deconvolved images, with the following parameters. 'HyVolution Grade' at max Resolution, Huygens Essential as 'Approach', water as 'Mounting Medium' and Best Resolution as 'Strategy'.
- STED imaging was performed on a commercial STED microscope (Expert Line, Abberior-Instruments, Germany) working at repetition rate of 40 Mhz.
- dSTORM imaging was conducted on an inverted microscope (Zeiss Axio Observer.Z1, Carl Zeiss Microscopy) equipped with a 100x oil-immersion objective (alpha Plan-Apochromat 100x/1.46 Oil DIC, Carl Zeiss Microscopy) and a 63x water objective lens (LD C-Apochromat 63x/1.15 W Corr M27, Carl Zeiss Microscopy).
Data analysis

The extraction of individual particles in input volumes was realized manually with the software ImageJ, version 1.51s. A drift correction was applied in the acquired stacks of images with the ImageJ plugin StackReg: http://bigwww.epfl.ch/thevenaz/stackreg/. The code for particle averaging was developed by the authors and is available on the GitHub repository: https://github.com/dfortun2/U-ExM. This is a Matlab code, with a graphical interface for each step of the reconstruction. We refer to the README file of the repository for more details.

2D Super-resolution images were reconstructed using the ImageJ plugin ThunderSTORM 5 and for 3D images the open source software rapidSTORM 3.36 was used.

The Intensities of clathrin-coated pits were normalized to the maximum intensity value and double gaussian fits were fitted to the Intensity profiles using the software Origin (OriginLab, Northampton, MA).

2D Super-resolution images were reconstructed using the ImageJ plugin ThunderSTORM 5 and for 3D images the open source software rapidSTORM 3.36 was used.

Field-specific reporting

Please select the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☑ Life sciences  ☐ Behavioural & social sciences  ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/authors/policies/ReportingSummary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size

We analyzed in most experiments around 30 isolated centrioles (10 centrioles per each three independent experiments). Note that we stated in the manuscript how many centrioles were analyzed in each experiment. We did not use a predetermined sample size. Considering the time of acquisition, we acquired 10 centrioles per experiment.

Data exclusions

To measure centriole diameter, we solely quantify nearly perfect top view centrioles to avoid any bias due to tilted centrioles. We stated this in the online methods section (Measurements of centriole diameter) and made a supplementary figure to explain this choice (Supplementary Fig.2).

For the in cellulo flagella analysis, we analyzed only fully expanded flagella and we specified this in the online methods section.

Replication

All experiments, imaging and analysis were carried out 3 times independently, unless specified otherwise for Figures 2b, 3b-f, 4c, which were performed once. This is reported in the Online method (statistics and reproducibility).

Randomization

This is not relevant for our study because we selected only nearly perfect top and lateral views of centrioles. In a given gel, centrioles are found in many orientations.

Blinding

Blinding is not relevant for our study for the same reasons as specified above.

Reporting for specific materials, systems and methods
Materials & experimental systems

| Involved in the study | n/a |
|-----------------------|-----|
| Unique biological materials | x |
| Antibodies | x |
| Eukaryotic cell lines | x |
| Palaeontology | x |
| Animals and other organisms | x |
| Human research participants | x |

Methods

| Involved in the study | n/a |
|-----------------------|-----|
| ChIP-seq | x |
| Flow cytometry | |
| MRI-based neuroimaging | |

Antibodies

| Antibodies used |
|-----------------|
| - rabbit polyclonal anti-polyglutamate chain (PolyE, IN105), reference AG-25B-0030-C050, Adipogen, dilution 1/500 |
| - mouse monoclonal anti-alpha tubulin (DM1alpha), reference T6199, Sigma, dilution 1/500 |
| - rat monoclonal anti-alpha tubulin (YL1/2), Abcam, ab6160, dilution 1/500 |
| - goat anti-rabbit Alexa Fluor 488 IgG (H+L), Invitrogen A11008, dilution 1/400 |
| - goat anti-mouse Alexa Fluor 488 IgG (H+L), Invitrogen A11029, dilution 1/400 |
| - goat anti-mouse Alexa Fluor 568 IgG (H+L), Invitrogen A11004, dilution 1/400 |
| - anti-rabbit STAR 580, Aberrior, S-12-2015Hp, dilution 1/400 |
| - anti-mouse STAR RED, Aberrior, S-08-2016Hp, dilution 1/400 |
| - A1647 conjugated F(ab')2 fragment of goat anti-rabbit, reference A-21246, ThermoFisher, dilution 1/200 |
| - MitoTracker red CMXRos, M7512, Invitrogen, 100nM |
| - rabbit monoclonal anti-TOMM20 (EPR15581-39), ab186734 Abcam, dilution 1/200 |
| - anti-rat Cy3, Jackson ImmunoResearch, 712-166-153, dilution 1/400 |
| - mouse anti-alpha tubulin (B-5-1-2) Sigma T5168, dilution 1/500 (6.7mg/ml) |
| - rabbit anti-clathrin heavy chain, Abcam, dilution 1/500 |
| - Alexa Fluor 488 F(ab')2 of goat anti rabbit IgG (2mg/ml, 1:200, A11070, ThermoFisher) |
| - Se Tau-647-NHS (K9-4149, Seta BioMedicals) conjugated to F(ab')2 of goat anti-Rabbit IgG (SAS-10225, ThermoFisher), 1.5mg/ml, 1/200 |
| - DNA-dye Hoechst 3342, C10340, Invitrogen, 10mg/ml, 1/1000 |
| - Alexa Fluor 647 F(ab')2 of goat anti rabbit IgG, A-21246, ThermoFisher, 2mg/ml, 1/200 |

Validation

- PolyE antibody (IN105) recognizes specifically glutamate chains of four or more glutamates. [https://adipogen.com/ag-25b-0030-anti-polyglutamate-chain-polye-pab-in105.html](https://adipogen.com/ag-25b-0030-anti-polyglutamate-chain-polye-pab-in105.html)
- DM1A antibody recognizes the following epitope in alpha tubulin: aa 426-450. [https://www.sigmaaldrich.com/catalog/product/sigma/T6199?lang=fr&region=CH](https://www.sigmaaldrich.com/catalog/product/sigma/T6199?lang=fr&region=CH)

Eukaryotic cell lines

Policy information about cell lines

| Cell line source(s) | U2OS: cell line provided by Erich Nigg (Biozentrum, Basel, Switzerland). COS-7 were purchased from CLS Cell Line Service GmbH. |
|---------------------|---------------------------------------------------------------------------------------------------------------------|

Authentication

None of the cell lines used were authenticated

Mycoplasma contamination

U2OS cell lines were negative for mycoplasma contamination. COS-7 were not tested for mycoplasma.

Commonly misidentified lines

(See [ICCLAC register](https://icclac.org/))

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

| Laboratory animals | The cell wall-less Chlamydomonas strain CW15- used in this study was grown in liquid TAP (Tris-acetate-phosphate) buffer at ~22°C or on solid TAP plates with 1.5% agar. |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------|
| Wild animals | no wild animals were used in this study. |
| Field-collected samples | no field-collected samples were used in this study. |
Imaging cellular ultrastructures using expansion microscopy (U-ExM)

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Supplementary Figure 1. Expansion methods and non-expanded centrioles in dSTORM. (a) Schematic illustration of two methods of expansion microscopy, ExM and MAP. (b) Schematic representation of a centriole seen either in top view (top) or lateral view (bottom). (c) 2D dSTORM image of isolated Chlamydomonas centrioles stained with PolyE (Alexa 647). Scale bar: 400 nm. (1-3) Magnified views of boxed regions of (c) Scale bar: 100 nm. Examples of centrioles used to quantify the diameter of non-expanded centriole in Fig.1. Representative images from 1 experiment.
Supplementary Figure 2. Centriole diameter and length measurements. (a, b) Maximum intensity projection of a 3D image of an U-ExM *Chlamydomonas* centriole stained with tubulin (magenta) and PolyE (green) to illustrate a top view centriole in a good orientation (a) or a tilted orientation (b). Note that we excluded from our measurements tilted centrioles. (c) Illustration of centriole diameter measurement. Dotted lines represent the lines used to perform the measurements in Image J. An average of the two values was taken as the final diameter. Scale bar: 400nm. (d, e) Top view 3D image of an U-ExM *Chlamydomonas* centriole stained with tubulin (magenta) and PolyE (green) taken with confocal (d) or after Hyvolution (e). Steps every 240nm. Scale bar: 400nm. (f, g) Lateral view 3D image of an U-ExM *Chlamydomonas* centriole stained with tubulin (magenta) and PolyE (green) taken with confocal (f) or after Hyvolution (g). Steps every 240 nm. Scale bar: 400nm. Representative images from 3 independent experiments.
Supplementary Figure 3. Expansion conditions. (a-b) Representative confocal images (HyVolution) of isolated centrioles in ExM stained with PolyE (green, Alexa 568) and α-tubulin (magenta, Alexa 488) (a) or only α-tubulin (magenta, Alexa 488) (b). Note the 9-fold symmetry visible with tubulin in absence of the PolyE staining. Representative images from 3 independent experiments (a) and 1 experiment (b). Scale bar: 400nm. (c) Steps explaining how the 9-fold symmetry was analyzed starting from the confocal image to the polar transformation and the actual plot profile. Scale bar: 400nm. (d, e) Plot profiles of the polar transform showing the 9-fold symmetry for ExM (d) and MAP (e). The center line represents the average and the shaded region the standard deviation. n= 30 centrioles for each condition. Data from 3 independent experiments. (f) Representative confocal images (deconvolved by HyVolution) of isolated Chlamydomonas centrioles incubated in the reported
solutions before expansion in a MAP gel and stained with PolyE (green, Alexa 488) and α-tubulin (magenta, Alexa 568). Representative images from 2 experiments. Scale bar: 800nm. (g) Quantification of PolyE (green) and α-tubulin (purple) diameters of isolated Chlamydomonas centrioles incubated in the reported FA solutions. Average and standard deviation are as follows. PolyE: 0.3%: 213nm +/- 12; 0.5%: 206nm +/- 10; 0.7%: 213nm +/- 11 and 1%: 200nm +/- 11. Tubulin: 0.3%: 179nm +/- 13; 0.5%: 179nm +/- 9; 0.7%: 184nm +/- 11 and 1%: 173nm +/- 12. N=25 centrioles for each condition from 2 independent experiments. Statistical significance was assessed by ordinary one-way ANOVA test. Only statistically significant differences are shown: ***=0.0004 for 0.3% vs 1% and 0.0008 for 0.7% vs 1%. **=0.0049. (h) Quantification of PolyE (green) and α-tubulin (purple) diameters of isolated Chlamydomonas centrioles incubated in 0.7%FA without AA or in 0.7%FA + different concentration of AA (as reported in the graph). Average and standard deviation are as follows. PolyE: NO AA: 213nm +/- 11; 0.15% AA: 227nm +/- 12; 1%AA: 222nm +/- 14, 5%AA: 220nm +/- 7 and 10%AA: 219nm +/- 8. Tubulin: NO AA: 184nm +/- 11; 0.15% AA: 195nm +/- 12; 1%AA: 193nm +/- 14, 5%AA: 192nm +/- 11 and 10%AA: 182nm +/- 8. Data in condition NO AA are the same used in (g) as 0.7%. N=30 centrioles for conditions 0.15% AA and 1% AA from 3 independent experiments. For condition 0.7%FA + 5% AA, n=20 centrioles from 2 experiments and for condition 0.7%FA + 10% AA, n=10 centrioles from 1 experiment. Statistical significance was assessed by ordinary one-way ANOVA test. Only statistically significant differences are shown: *=0.018 for PolyE in NO AA vs 1% AA, 0.014 for Tubulin in NO AA vs 0.15% AA, 0.044 for Tubulin in NO AA vs 1% AA, and 0.046 for Tubulin in 0.15% AA vs 10% AA. ***=0.0002.
Supplementary Figure 4. U-ExM preserves the ultrastructure of centrioles. (a) Cryo-electron microscopy image representing a lateral view of an isolated *Chlamydomonas* centriole. (b) Same confocal image (deconvolved by HyVolution) shown in Fig. 1d of an isolated centriole expanded with U-ExM (0.7%+ 1% AA) and stained with PolyE (green, Alexa 488) and α-tubulin (magenta, Alexa 568). Representative lateral view from 3 independent experiments. Scale bars in (a): 100nm and (b): 400nm. (c) Plot profile of the polar transform showing the 9-fold symmetry for U-ExM. The center line represents the average and the shaded region the standard deviation. n= 30 centrioles. Data from 3 independent experiments. (d) Roundness, shape of the centriole for the three expansion methods. Averages and standard deviation are as follows. For ExM: 0.7=16.67 ± 11.55, 0.8= 43.33 ± 25.17, 0.9=30 ± 10, 1=10 ± 17.32. For MAP, 0.7=0 ± 0, 0.8=26.67 ± 5.77, 0.9= 50 ± 10, 1=23.33 ± 15.28. For U-ExM: 0.7=0 ± 0, 0.8=6.67 ± 5.77, 0.9=73.33 ± 15.28, 1=20 ± 10. Note that for all the quantifications provided in this figure, we included data from U-ExM performed with (0.7%FA + 0.15%) and (0.7%FA + 1% AA). N= 30 centrioles for each condition. Data from 3 independent experiments.
Supplementary Figure 5. Comparison of U-ExM to dSTORM. (a) Montage of a Z-stack through *Chlamydomonas reinhardtii* isolated centrioles. Note the presence of procentrioles highlighted with the black arrowheads. Representative images from 3 independent experiments. (b, c) Examples of 3D projections of isolated centrioles fixed in FA and stained for PolyE using Al647. Imaging was performed using dSTORM in 3D mode using a water objective. Representative images from 1 experiment. Scale bar: 500nm. (d, e) Two examples of isolated *Chlamydomonas* centrioles imaged using 2D dSTORM with a oil objective. Representative images from 1 experiment. Scale bar: 200nm. (f-h) Plot profile of the polar transformation calculated from 2D-dSTORM images of non-expanded procentrioles (f, n=4 procentrioles from one experiment), from confocal images of procentrioles expanded
with U-ExM (g, 30 precentrioles from 3 independent experiments) and from deconvolved images of the same procentrioles used in (g) using HyVolution (h, 30 precentrioles from 3 independent experiments). The center line represents the average and the shaded region, the standard errors. (i-k) Plot profile of the full width at half maximum (FWHM) calculated from procentrioles triplets: dSTORM $\approx$ 44nm (n=4 procentrioles, and 24 triplets from one experiment), U-ExM (confocal)$\approx$ 59nm (n=9 procentrioles, and 76 triplets from 3 independent experiments), U-ExM (HyVolution)$\approx$ 21nm (n=9 procentrioles, and 81 triplets from 3 independent experiments). Line connects the mean of each x value and error bars represent the standard deviation.
Supplementary Figure 6. Polyglutamylated tubulin localization in an isotropic reconstruction of a U-ExM centriole. (a) Gallery of a Z stack from distal to proximal of a mature centriole expanded using U-ExM (0.7%FA+1%AA) and stained with α-tubulin (magenta, Alexa568) and PolyE (green, Alexa488). Scale bar: 200nm. Arrowhead points to PolyE surrounding the tubulin signal. Representative data from 3 independent experiments. (b) 3D volume reconstruction of a mature *Chlamydomonas* centriole stained with α-tubulin (magenta, Alexa568) and PolyE (green, Alexa488). Scale bar: 200nm. Data obtained from 14 centrioles averaged (one experiment). (c-f) Sections through the centriole spanning the distal (c) and the central core (d-f) regions. Scale bar: 200 nm. (g-i) Quantification of PolyE and α-tubulin diameter for the distal (g), central (h) and proximal parts of the central core region of centrioles (i) corresponding to (d), (e) and (f), respectively. Data obtained from 14 centrioles averaged (one experiment). Note that the polyglutamylation diameter signal is larger than that of tubulin.
Supplementary Figure 7. PolyE sub-triplet localization revealed by U-ExM. (a–c) Representative image taken from a Z-stack of the distal most-part of the central core of a deconvolved mature centriole stained for α-tubulin (magenta, Alexa568) and PolyE (green, Alexa488). Scale bar: 200nm. Green arrowheads highlight the PolyE signal (b, c). The dotted arrow in (a) illustrates how the fluorescence intensity was measured in (d). (e) Schematic representation of microtubule triplets superimposed
onto the fluorescent signal shown in b. (d) Quantification of the fluorescence intensity shift between the magenta and the green signal. Note that the shift is of 21 nm (n=61 triplets from 8 centrioles). Data from 3 independent experiments. Line connects the mean of each x value and error bars represent the standard deviation. (e-h) Model images of a *Chlamydomonas* centriole with entire triplets (e) only A-tubule (f) only B-tubule (g) and only C-microtubule (h). Images were scaled to mimic an expanded centriole. Scale bar: 500 nm. (i-l) Images were bandpass filtered to obtain a resolution of 140 nm typical of a confocal microscope in HyVolution mode. Colors are added to mimic fluorescent signal. (m-o) Model representation of PolyE signal onto specific microtubule blades, A- (m), B- (n) and C- (o). Note that in this model, the entire triplet is stained with α-tubulin (magenta). Below is represented the fluorescence peak shift between the two colors in each condition. Scale bar: 200nm.
Supplementary Figure 8. Coupling STED to U-ExM. (a) 3D DyMIN images of U-ExM (0.7% FA+1% AA)-treated centrioles stained with PolyE (green, STAR 580) and α-tubulin (magenta, STAR RED). Maximum intensity projection. Data from one experiment. (b) Stack slices thought the procentrioles. Scale bar in (a): 1µm, (b): 200nm. (c) Electron microscopy (EM) image of a centriole pair comprising two
mature centrioles and two procentrioles (black arrowheads) interconnected by striated fibers. Scale bar: 200nm (d) A similar centriole pair stained for α-tubulin (magenta, STAR RED) after U-ExM (1% FA) and imaged using DyMIN. Note that the overall ultrastructure of this organelle resembles the EM image. White arrowheads points to procentrioles and the arrow points to the mature centriole. Data from 2 independent experiments. Scale bar: 800nm.

(e, f) Two representative DyMIN images of non-expanded *Chlamydomonas* isolated centrioles stained for α–tubulin (magenta). Insets show the same centriole imaged at confocal. White arrowheads indicate procentrioles. Data from one experiment. Scale bar: 200nm.
Supplementary Figure 9. In cellulo expanded *Chlamydomonas* centrioles. (a, b) Representative Hyvolution confocal image of an *in cellulo* *Chlamydomonas* centriole stained for PolyE (green) and α−tubulin (magenta). The dotted square highlight the centriole shown in the inset (b). Arrowheads indicate the nine-fold symmetry of the centriole. Scale bar: 250nm. Representative images from 3 independent experiments. (c) Diameter in nm of U-ExM centrioles *in cellulo* or from purified centrioles. Averages and standard deviation are as follows. PolyE: 236 nm +/- 18 nm for U-ExM *in cellulo* (n=31 centrioles) and 225 nm +/- 15 nm for U-ExM isolated centrioles (n=30 centrioles). α-tubulin: 212 nm +/- 22 nm for U-ExM *in cellulo* (n=31 centrioles) and 195 nm +/- 12 nm for U-ExM isolated centrioles (n=29 centrioles). Data from 3 independent experiments. Note that centrioles *in cellulo* are statistically slightly larger. Unpaired two-sided t-test: for PolyE: p= 0.0145 and tubulin: p= 0.0004. (d) Isotropic expansion calculated by the ratio of centriolar length/diameter. Average and standard deviation are as follows: 2.6 +/- 0.3 (n=23 centrioles from 3 independent experiments). Unpaired two-sided t-test = 0.68 (non significative) compared to U-ExM isolated centrioles (Fig. 1f). Note that centrioles *in cellulo* expanded isotropically.
Supplementary Figure 10. U-ExM applied on *Chlamydomonas* cells. (a) *In cellulo* U-ExM expansion of a *Chlamydomonas* cell stained with PolyE (green) and α–tubulin (magenta) imaged using confocal microscopy coupled to Hyvolution. Scale bar: 13µm. Representative image of a top-view across a flagellum. Note that the nine doublets are highly polyglutamylated (green) while the central pair is weakly polyglutamylated. Representative images from 3 independent experiments. (b) Plot profile of the polar transform showing the 9-fold symmetry of the axoneme. The center line represents the average and the shaded region, the standard errors. N=23 axoneme from 3 independent experiments. (c) Quantification of the fluorescence intensity shift between the magenta and the green signal showing that the PolyE signal is more internal than the tubulin signal with a shift of -10 nm. Line connects the mean of each x value and error bars represent the standard deviation. N= 23 axonemes from 3 independent experiments.
Supplementary Figure 11. Analysis of the Polyglutamylation profile in axonemes using U-ExM. (a-d) Opened axoneme stained with PolyE (green, Alexa 488) and α-tubulin (magenta, Alexa 568). (b-d) Inset of the region boxed in (a). Scale bar: 1µm. (e) Plot profile of the PolyE signal over the 9 microtubule doublets and the 2 central...
pairs. Numbers represent single microtubules doublets (MTD) while CP1 and CP2 are individualized microtubules of the central pair. Representative images and plot profile from one experiment (n=4 sprayed flagella). (f) Fluorescence intensity (A.U) of MTD and CP. Note that CPs display a weak PolyE signal. Average and standard deviation is as follows: Average intensity of CP= 17% +/- 7 unpaired t test two-tailed, p<0.0001. Data from one experiment (n= 4 sprayed flagella). (g) Schematic representation of an axoneme composed of nine outer microtubule doublets and a central pair of single microtubules (9+2). Microtubules are in magenta; inner dynein arms (IDA), outer dynein arm (ODA), nexin and radial spokes are in grey; polyglutamylated tubulin is in green. Note that we draw two models: one with polyglutamylated tubulin restricted to the inner side of the B-tubule (half B) and one with the entire B-tubule decorated (Full B). (h-j) Model images of a *Chlamydomonas* flagellum adapted from a cryoEM map with the entire microtubule doublets (h), only half B-tubule (i) and only full B-tubule (j). Images were scaled to mimic an expanded axoneme. (k-m) Bandpass filtered images to obtain a resolution of 140 nm typical of a confocal microscope in HyVolution mode. Colors are added to mimics fluorescent signal. (n-o) Model combining the α-tubulin and PolyE signals to show how the PolyE signal in either half B-microtubule or full B-microtubule would be positioned in relation to the α-tubulin signal of the entire doublet. (p, q) Fluorescent peak shift between the magenta and the green signal for the Half B (p) and the Full B (q) models. The shifts are -7 nm and +6 nm, respectively. Note that the experimental data mimics the Half B model (p).
Supplementary Figure 12. Effect of different fixations on U-ExM-isolated Chlamydomonas centrioles. (a-d) 2 examples of top (left) and one example of lateral (right) views of Hyvolution images of centrioles unfixed (a) or fixed with FA (b), methanol (c) or PFA/GA (d) stained with α-tubulin (magenta) and PolyE (green). Scale bar: 400nm. (e) Diameter of the centrioles in the indicated conditions. Averages of PolyE diameter and standard deviations are as follows: no fixation: 225 nm +/- 15 nm; FA: 201 nm +/- 7 nm (n=11); methanol: 200 nm +/- 18 nm (n=16); PFA/GA: 160 nm +/- 9 nm (n=13). For ‘No fixation’ condition, same data from Fig. 1e was used. For methanol fixed condition, data from 2 independent experiments. For FA and PFA/GA conditions, data from 1 experiment. Statistical significance was assessed by one-way ANOVA test: ****<0.0001, ns (non significant)=0.996.
Supplementary Figure 13. Effect of different fixations on cellular structures in human cells. (a-d) Representative Hyvolution images (from 2 independent experiments) of U2OS cells unfixed (a) or fixed with FA (b), methanol (c) or PFA+GA (d) and stained with α-tubulin (green). Scale bar: 10µm. Insets (shown with the dotted square) show a magnified region. Scale bar: 0.5µm. (e-h) Representative Hyvolution images (from 2 independent experiments) of U2OS cells unfixed (e) or fixed with FA (f), methanol (g) or PFA+GA (h) and stained with mitotracker (fire). Scale bar: 10µm. Insets (shown with the dotted square) show a magnified region. Scale of insets: 500nm. Note that only PFA/GA preserves the mitochondrial organization.
Supplementary Figure 14. U-ExM applied on human cells. (a) Schematic illustration of the pre and post-U-ExM expansion imaging experiment. U2OS cells were first methanol fixed and immunostained with a rat anti-tubulin (tubulin 1, green) followed by imaging. Then, cells were processed for U-ExM. Post staining was performed with mouse anti-tubulin (tubulin 2, magenta). (b-d) Inset of a U-ExM-U2OS cell processed for pre- (b) and post expansion imaging (c, confocal; d, Hyvolution) as depicted in (a). Scale bars: 1.5 µm (a) and 6 µm (c, d). Representative images from 1 experiment, n=4 cells. (e, f) Plot profiles of the regions indicated with a dotted white line with the corresponding full width at half maximum (FWHM): confocal, U-ExM microtubule: 54 nm (n=47); Hyvolution, U-ExM microtubule: 46 nm (n=50). Line connects the mean of each x value and error bars represent the standard deviation. Data from 3 independent experiments. (g) Two examples of human centrioles expanded using U-ExM in cells and imaged at confocal (left) or Hyvolution (right). Data from one experiment. Scale bar: 800nm.
Supplementary Figure 15. U-ExM microtubules. Hyvolution image of microtubules from a U2OS cell fixed with methanol and processed for U-ExM. Color code indicates the Z-position. Representative images from 3 independent experiments. Scale bar: 4µm.
Supplementary Figure 16. U-ExM mitochondria. Hyvolution image of mitochondria from a U2OS cell fixed with PFA/GA, processed for U-ExM and stained for the outer membrane mitochondrial translocase TOMM20. Color code indicates the Z-position. Representative images from 3 independent experiments. Scale bar: 4µm.
Supplementary Figure 17. Evaluation of distortions over length scales for microtubules and mitochondria staining. (a, b) Hyvolution images of a U2OS cell treated for pre- (a) (tubulin 1, green) and post-expansion imaging (tubulin 2, magenta) (b). (c) Overlay of pre- and post-expansion images of microtubules with distortion vector field. Representative images from 1 experiment. (d, e) Hyvolution images of a U2OS cell treated for pre- (d) (mitotracker 1, green) and post-expansion imaging (mitotracker 2, magenta) (e). (f) Overlay of pre- and post-expansion images of mitochondria with distortion vector field. Representative images from 2 experiments. (g) Quantification of root mean square (RMS) error over distance (micrometers) comparing pre- (a, d) and post-U-ExM (b, e) expansion for both microtubules (n=4 cells from 1 experiment) and mitochondria (n=3 cells from 2 experiments). Line connects the mean of each x value and error bars represent the standard deviation. (h) Calculated expansion factor based on the RMS graph for both microtubules (n=4 cells from 1 experiment) and mitochondria (n=3 cells from 2 experiments). Red dotted lines correspond to the averages: 4.1 for microtubules and 3.9 for mitochondria.
Supplementary Figure 18. Clathrin Coated Pits visualized after U-ExM.

(a) Representative image of a human cell stained for tubulin (magenta), clathrin coated pits (yellow) and DNA (blue) with a 10x objective. Scale bar: 40µm. Data from 3 independent experiments. (b) Same cell imaged with a 60X objective highlighting the clathrin coated pits. Scale bar: 4µm. (c) Zoom in of the region shown with the squared dotted line. White arrowheads indicate clathrin-coated pits. Scale bars: 1µm. (d) Gallery of U-ExM clathrin-coated pits. Note the hollow shape of the clathrin pits. Scale bar: 400nm. (e) Plot profiles along the clathrin-coated pits, as indicated in (d) with a dotted line, highlighting the hollow center in the clathrin pits. Double gaussian fits (blue line) of the normalized intensity profiles were used to calculate the diameter of the pits with standard errors deriving from the fits. Using an expansion factor of 4x, the values translate to: 72nm, 110nm, 178nm, 90nm and 102nm.
Supplementary Figure 19. Optimization of the U-ExM expansion factor. (Online Methods). Monomer solutions of different composition were tested for gel expansion. Averages expansion factor with their corresponding standard error of the mean are indicated (data from 2 independent experiments). Centriolar shape was analyzed: ✓ indicates proper shape and ✗ deformed centrioles. Below are representative top and side views from 2 independent experiments of Hyvolution-images of isolated *Chlamydomonas* centrioles treated in the indicated conditions. Centrioles were stained for tubulin (magenta) and PolyE (green). Scale bar: 450nm. Note that U-ExM gave the best structural preservation of the specimen.

| Conditions   | 20%AA/7%SA | 10%AA/7%SA | 10%AA/19%SA | 5%AA/7%SA | 5%AA/19%SA |
|--------------|------------|------------|-------------|-----------|-----------|
| Expansion Factor | 3.40 ± 0.08 | 3.60 ± 0.04 | 4.00 ± 0.02 | 3.90 ± 0.06 | 4.30 ± 0.00 |
| Correct shape | ✓          | ✓          | ✓           | ✗         | ✗         |