Correlated cryogenic photoactivated localization microscopy and cryo-electron tomography

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Cryo-electron tomography (CET) produces three-dimensional images of cells in a near-native state at macromolecular resolution, but identifying structures of interest can be challenging. Here we describe a correlated cryo-PALM (photoactivated localization microscopy)-CET method for localizing objects within cryo-tomograms to beyond the diffraction limit of the light microscope. Using cryo-PALM-CET, we identified multiple and new conformations of the dynamic type VI secretion system in the crowded interior of Myxococcus xanthus.

CET allows large macromolecular complexes to be visualized directly within intact cells1. Because cells are preserved in a near-native state, and reconstructions exhibit ‘macromolecular’ resolution (~4 nm), some complexes can be identified simply by their known shapes2. Unfortunately, this requires prior knowledge of a macromolecule’s morphology, association with other macromolecules and/or subcellular location. It is therefore of great interest to expand this technique to characterize structures that are smaller, dynamic or less well characterized.

One way to identify objects of interest in cryo-tomograms is through correlated fluorescence light microscopy (FLM) and electron microscopy (EM)3,4. In this approach, proteins of interest are fused to fluorophores and coarsely localized by FLM. The same samples are then imaged by CET—an EM technique—and the two images superposed. Although powerful, correlated FLM-CET approaches have limitations. FLM is best carried out at room temperature to allow use of oil-immersion lenses with high numerical aperture (NA = 1.4), but movement can prevent correlation of subcellular structures. As a result, only relatively static structures such as bacterial stalk cross-bands5 or eukaryotic focal adhesions6 can be studied this way. Alternatively, cells can be chemically fixed before imaging, but this can introduce artifacts7,8. Cells can also be plunge frozen before FLM and imaged cold (at ~80 K), but this severely limits fluorescence resolution because long-working-distance air objectives (NA = 0.7) are then required. Intriguingly, several super-resolution microscopy techniques have been described recently, including stochastic optical reconstruction microscopy (STORM)9, PALM10 and fluorescence PALM11. These techniques have dramatically improved fluorescence resolution, allowing single-molecule localization, but none has been performed under cryogenic conditions. Here we report the development of a super-resolution method that can be applied to frozen samples (cryo-PALM), allowing dynamic cellular objects to be localized within cryo-tomograms.

PALM relies on sequential laser-induced activation and inactivation of photoactivatable fluorophores10. To perform PALM on frozen EM grids, we added a multicolor laser to our cryo-FLM-CET system3. Normally, methods such as PALM rely on total-internal-reflection illumination to reduce background, but as imaging frozen EM grids requires an air objective lens, we used epi-illumination. Fortunately, because CET samples are thin (<500 nm), there was little background fluorescence, and the signal-to-noise ratio was sufficient for single-molecule localization after sample vibration and drift were minimized (Online Methods).

In CET, living cells are preserved by vitrification without chemical fixation or permeabilization. This limits the possible cryo-PALM probes to those that can be incorporated into living cells, such as genetically encoded photoactivatable fluorescent proteins (PA-FPs). Because low temperature drastically reduces the photobleaching rate of fluorescent proteins12, and might also alter their photoconversion behaviors, we screened PA-FPs for activation and inactivation at 80 K. We chose only monomeric candidates in order to avoid PA-FP polymerization13. Surprisingly, most PA-FPs tested did not respond, or responded very weakly, to illumination by the activation/conversion laser (Supplementary Fig. 1). Only PA-GFP maintained its photoactivatable behavior at 80 K. As at room temperature, photoconversion was reliably triggered by a 488-nm laser, and photoactivation could be accelerated by adding a 405-nm laser. With the long-working-distance air objective (NA of 0.7) and the presumably slower photobleaching rate at 80 K (ref. 12), the average number of photons collected per PA-GFP molecule was 206 (Supplementary Fig. 2).

One concern with cryo-PALM is that exposure to the intense laser might warm the sample above 135 K, causing vitreous ice to crystallize and denature the sample14. To examine this, we exposed vitrified samples on EM grids to a laser with sufficient energy for PA-GFP photoconversion (300 W/cm²) and then imaged the grids by CET. The laser exposure caused small crystals to form.

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within the amorphous ice surrounding cells (Supplementary Fig. 3a), but cytoplasmic structures were not affected perceptibly, likely because the crowded cellular interior served as a natural cryoprotectant. Adding 5% Ficoll PM 70 as a cryoprotectant and reducing the total laser exposure time to <300 s per target reduced ice crystallization considerably (Supplementary Fig. 3b). Residual ice crystallization and devitrification were fully overcome by adding 10% Ficoll PM 70 and 10% ethylene glycol and by pulsing the exposure, allowing 60 s of heat dissipation between each 60-s period of laser illumination (Supplementary Fig. 3c).

Widespread in Gram-negative bacteria, the type VI secretion system (T6SS) is a virulence-associated nanomachine that translocates inhibitors into prey cells\(^\text{15-16}\). In *Vibrio cholerae* the T6SS forms highly dynamic intracellular tubes\(^\text{17}\). Through cycles of sheath extension, contraction and disassembly, these tubes deliver protein effectors into adjacent target cells. Time-lapse FLM has revealed that the T6SS assembles within tens of seconds in a wide range of subcellular locations and then contracts and disassembles in a similar time frame. Identification of T6SS structures in cryo-tomograms is therefore complicated by their highly dynamic nature and random localization within the cell. In our previous study characterizing the T6SS in *V. cholerae* by CET, mutant strains and purified sheaths needed to be imaged in order to identify which ultrastructure was the T6SS\(^\text{17}\).

To characterize the T6SS in a second species, we chose *M. xanthus*, a model bacterial predator whose genome has been reported to encode all 13 core proteins of the T6SS\(^\text{18}\). We first tried to identify the T6SS in *M. xanthus* through a conventional correlated cryo-FLM-CET approach using a mutant strain containing a deletion of the gene encoding the sheath protein VipA (also known as TssB) (ΔvipA) and expressing a functional VipA-GFP fusion protein (Supplementary Figs. 4 and 5a).

Figure 1 | Correlated cryo-PALM-CET used to visualize extended and contracted conformations of the T6SS sheath in *M. xanthus*. (a,d) Low-resolution EM images (grayscale background), cryo-PALM images (red and yellow foreground), slices from high-resolution three-dimensional cryo-tomograms (grayscale foreground), and segmentations of cellular structures (blue, tubular structures; green, filament bundles; white, spherical granules) superposed. The cryo-PALM images reveal VipA–PA-GFP localization (red, low precision; yellow, high precision), identifying the tubular structures (grayscale foreground), and segmentations of cellular structures (blue, tubular structures; green, filament bundles; white, spherical granules) superposed. (b,e) Tomographic slices through the tubular structures (blue) in a,d showing extended and contracted T6SS sheaths, respectively. (c,f) Cross-sectional views of b and e, respectively. Scale bars, 400 nm (d; applies to a,d) and 50 nm (f; applies to b,c,e,f).

Figure 2 | Correlated cryo-PALM-CET identifies new T6SS structures. (a) A very short loaded T6SS structure with a ‘baseplate’ attached to the membrane. Superposed low-resolution EM image, cryo-PALM signal, high-resolution cryo-tomographic slice and object segmentations are shown as in Figure 1. (b) Tomographic slice through the tubular structure corresponding to the segmentation model shown in a. Features of the T6SS are highlighted in blue (sheath) and cyan (baseplate). (c) Tomographic slice in b without highlights. (d) Cross-section of the T6SS sheath in c. (e) A bent T6SS structure with an additional sheet, distinguished from other tubular structures in the vicinity. Segmentation of the cryo-tomogram shows different tubular structures (individually colored blue, pink and green) and spherical granules (white). (f,j,m) Segmented models of tubular structures shown in e. (g,k,n) Tomographic slices through the tubular structures corresponding to the segmentation models shown in f, j and m, respectively. White arrows in g indicate a sheet adjacent to the hollow tube. (h,i) Tomographic slices of cross-sections h’ and i’ of the tubular structure shown in g. (l,o) Cross-sectional views of k and n, respectively. Diameters of the tubules shown in d, h, l and o are 13, 14, 20 and 33 nm, respectively. Scale bars, 400 nm (e; applies to a,e) and 50 nm (o; applies to b,d,g,i,k,l,n,o).
Unfortunately, the limited resolution of cryo-FLM gave a broad fluorescence signal that, in the crowded interior of the bacterial cell, often ambiguously spanned a variety of candidate structures in cryo-tomograms (Supplementary Fig. 5b). To overcome this problem, we performed correlated cryo-PALM-CET using the mutant expressing VipA–PA-GFP. We localized VipA–PA-GFP molecules in frozen-hydrated cells on EM grids by cryo-PALM (Supplementary Fig. 5c). We then imaged locations with high cryo-PALM signal density at high resolution by CET. The overall imaging process is demonstrated in Supplementary Video 1 (with 5% Ficoll PM 70) and Supplementary Video 2 (with 10% Ficoll PM 70 and 10% ethylene glycol and pulsed laser exposure). In nearly every case (17/20), we observed a tubular structure in the location of VipA–PA-GFP signals. The three signals that did not correlate with a tubular structure likely corresponded to just-disassembled T6SSs whose fluorescent tags were still colocalized. We found that the structure of the T6SS sheath in _M. xanthus_ is similar to that reported for _V. cholerae_, and we observed both extended (‘loaded’) and contracted (‘fired’) conformations (Fig. 1). Extended tubes exhibited filled lumens (Fig. 1c) and had an average length of 608 ± 75 nm and a diameter of 12.6 ± 0.6 nm (n = 12; mean ± s.d.). Contracted tubes had clear lumens (Fig. 1f) and an average length of 287 ± 62 nm and diameter of 14.7 ± 0.7 nm (n = 9).

One cryo-PALM focus identified a very short (60-nm) filled tube, which was likely a T6SS in an early stage of assembly (Fig. 2–d and Supplementary Video 2). The width of this tube matched that of the other extended tubes, a result suggesting that the inner rod and outer sheath of the T6SS form concomitantly rather than sequentially. Another cryo-PALM focus superposed on a contracted tube bent approximately one-quarter length from its membrane-proximal end (Fig. 2e). Interestingly, we observed an additional layer associated with the tube on one side of the bend (Fig. 2f–i). The tube diameter was identical to that of contracted T6SS sheaths. It therefore likely represents an intermediate in the disassembly process of the sheath after contraction, raising the question of whether the additional layer is ClpV, the AAA-ATPase known to disassemble T6SS sheaths.19,20 The fact that this bent sheath was identifiable as a T6SS despite being surrounded by a variety of other tubular structures (Fig. 2j–o) demonstrates the utility of correlated cryo-PALM-CET.

Correlated cryo-PALM-CET brings together two of the most powerful light microscopy and EM techniques. By precisely localizing the fluorescent tag on a cellular object by cryo-PALM and then resolving the higher-resolution molecular structure of the object itself by CET, correlated cryo-PALM-CET should allow numerous dynamic molecular machines to be structurally characterized _in vivo_—and, notably, without fixation artifacts. Once the typical structures of an object are known, many will then be recognizable by their morphologies alone in cryo-tomograms of fully wild-type cells (without any tags at all). The cryoprotectant properties of different cytoplasms will likely vary, however, so the laser pulsing may need to be optimized for each species. The cryo-PALM setup described here had a maximum lateral position error of ~160 nm (Supplementary Table 1), easily allowing us to identify T6SS structures (11–15 nm wide and 200–500 nm long in two-dimensional projections). Our identification of an unusually short T6SS sheath just 12 nm wide and 60 nm long demonstrates the power of this method to identify smaller structures as well. Future development of brighter and longer-lasting cryo-photoactivatable fluorophores, higher-NA cryo-FLM objectives, more stable cryo-stages and better tools for transfer of samples and correlation of images could further increase the power of this approach. The use of a cryo-fluorophore ten times brighter and lasting ten times longer before bleaching at 80 K than PA-GFP, imaged with a 100× long-working-distance air objective with an NA of ~0.8, would increase the precision to better than 2 nm, sufficient to pinpoint the location of almost any macro-molecule within the cell.

**METHODS**

Methods and any associated references are available in the online version of the paper.

**Note:** Any Supplementary Information and Source Data files are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

Y.-W.C. and G.J.J. conceived the cryo-PALM idea. Y.-W.C. and S.C. configured the optical system. Y.-W.C., S.C. and E.I.T. tested fluorophores for photoactivatability at low temperatures. Y.-W.C. improved stability of cryo-FLM stage, prepared samples, overcame laser-induced ice crystallization on the sample, acquired and analyzed cryo-PALM data and conducted correlated cryo-PALM-CET. A.T.-L., S.L. and L.S.-A. generated the optical system. Y.-W.C. and G.J.J. wrote the paper with input from all authors.

**COMPETING FINANCIAL INTERESTS**

The authors declare competing financial interests: details are available in the online version of the paper.

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M. xanthus strain construction. The in-frame deletion mutations were generated in the WT strain DK1622 (ref. 22) using standard methods and plasmids pSl02 (to generate strain SA5716 (ΔvipA = ΔMXAN4807)) and pAK88 (to generate SA5707 (ΔT6SS = ΔMXAN4800–4813)). Strains SA4137 (vipA/P_pilA-vipa-GFP) and SA5718 (ΔvipA/P_pilA-vipa-PA-GFP) were generated by electroporation of plasmids pMAT36 and pSl05, which encode Vipa-GFP and Vipa–PA-GFP, respectively, and integrate into the Mx8 attB site, into SA5716. Assays for motility and development were carried out as described.

Plasmid construction. Plasmid pSl02 was generated by amplification of the vipA gene without its stop codon using primers oVipA1 and oVipA3 (see Supplementary Table 3 for primer sequences) and M. xanthus chromosomal DNA as template. The product was ligated into pTP100 by XbaI-BamHI digestion. The plasmid was amplified from pCrGFp (BioCat) using primers oGFP1 and oGFP2. PA-GFP was amplified from pRSETA-PA-GFP (Addgene 11911) using primers oPAGFP1 and oPAGFP2. The GFP and PA-GFP fragments were cloned into pSl02 by BamHI-KpnI digestion to generate pMAT36 and pSl05, which encode Vipa-GFP and Vipa–PA-GFP, respectively, and integrate into the pilA attB site. In the two fusion proteins, GFP and PA-GFP are separated from Vipa by an Ala-Ala-Ala-Ala-Gly-Gly-Gly linker.

To generate the in-frame deletion of vipA, we constructed pSl04 by cloning the upstream (primer pair mxan4807AB) and downstream (primer pair mxan4807CD) regions of vipA into vector pBJ114 (ref. 25). This deletion extends from nucleotide 31 to 465 of vipA (vipA has a total length of 495 bp). To generate the in-frame deletion of all T6SS-encoding genes MXAN4800–4813 (ref. 18), plasmid pAK88 was generated by cloning the upstream (primer pair mxan4800AB) and downstream (primer pair mxan4813CD) regions of the T6SS gene cluster into vector pBJ114 (ref. 25). The resulting deletion extends from codon 26 of MXAN4800 to codon 867 of MXAN4813.

Immunoblot analysis for M. xanthus T6SS activity. To determine whether the M. xanthus T6SS is active, we monitored by immunoblot analysis the accumulation of (i) hemolysin-coregulated protein (Hcp) in the culture supernatant of an exponentially growing WT strain, (ii) a mutant containing a deletion of the entire T6SS gene cluster including the gene encoding Hcp (ΔT6SS) and (iii) a mutant containing a deletion of the gene encoding the core protein Vipa (ΔvipA). The SDS-PAGE gel for the anti-Hcp immunoblot was loaded with total cell extracts from 2.5 × 10^7 cells and concentrated cell-free supernatants from 1.5 × 10^10 cells. To prepare cell-free supernatants, we used cultures of exponentially growing M. xanthus cells at OD550 of 0.9–1.1. Cells were harvested by centrifugation, and the supernatant was filtered through a 0.22-µm sterile filter. Precipitation of the proteins in the filtered supernatant was performed using TCA-DOC precipitation. Briefly, 1/100 volume of 2% sodium deoxycholate (DOC) was added, and the solution incubated at 4°C for 30 min. Afterward, TCA was added to a final concentration of 10%, and the solution was incubated at 4°C for another 16 h. Protein precipitate was sedimented by centrifugation (15 min, 10,000g). The collected pellet was washed twice with acetone and then resuspended in SDS-PAGE loading buffer. Immunoblotting was performed using standard procedures with rabbit anti-Hcp (Eurogentec, SY2563, 1:500) and peroxidase-conjugated goat anti-rabbit IgG secondary antibodies (Sigma-Aldrich, A8275, 1:10,000) following the recommendations of the manufacturer. Rabbit antisera against Hcp was generated by Eurogentec, using a mixture of the peptides REAGSLATGRQYEG and ITQGGVTHEDTWDTQR, corresponding to residues 44–59 and 148–163 of the Hcp protein, respectively. The blot was developed using Luminata Western HRP substrate (Millipore, WBLUF100). Hcp was detected in total cell extract as well as in the cell-free supernatant of WT cells, in only the total cell extract of the ΔvipA mutant, and in neither fraction in the ΔT6SS mutant (Supplementary Fig. 4), indicating that the M. xanthus T6SS is active. The ΔT6SS and ΔvipA mutants were indistinguishable from WT with respect to growth, motility, fruiting-body formation and sporulation. To detect Vipa–PA-GFP and Vipa–GFP, we stripped the anti-Hcp blot (Restore Plus western blot stripping buffer; Thermo Scientific, 46430), and the blot was probed with mouse anti-GFP primary (Roche, 1181446001, 1:2000) and anti-mouse IgG secondary antibodies (DakoCytomation, P0260, 1:2000) and developed as described.

Enhancing cryo-FLM stage stability. In the cryo-FLM setup, liquid nitrogen (LN) is continuously pumped into the stage in order to maintain the sample at 80 K. The flow and boiling of LN cause vibrations in the stage. To control this problem, we improved the insulation of the cryo-FLM stage by introducing a 1-cm-thick poly(methyl) methacrylate plate as a lid to reduce heat loss through the condenser hole while preserving transparency for light traveling through the condenser aperture into the cryo-stage (Supplementary Fig. 6). The resulting slower heat exchange suppressed LN boiling and reduced its flow. The slight residual stage drift during data recording could be corrected by registering fluorescent fiducial beads, as in the room-temperature PALM method.

Photoactivation/photoconversion test of PA-FPs in cells at 80 K. E. coli BL21(DE3) cells harboring plasmid pRSETA-PA-GFP, pRSETA-Dendra2, pRSETA-Dronpa, pRSETA-mEosFP2, pRSETB-PA-mRFP1, pET28A-PA-CFP2 or pBAD/HisB-PA-mCherry1 were first grown to an OD_600 of 0.6, and expression of PA-FPs was then induced with 1 mM isopropyl β-D-1-thiogalactopyranoside or 0.2% (w/v) L-arabinose for 2 h. The cells were applied to coverslips, and photoactivation/photoconversion of the PA-FPs was confirmed with room-temperature FLM using a 100× oil-immersion objective lens. Exposures of 1–3 s using a mercury lamp with a DAPI filter set (Semrock DAPI-1160A-NTE) were used to trigger the photoactivation/photoconversion. Exposures of 1–3 s through a FITC (Semrock FITC-3540B-NTE) or a TxRed (Semrock TxRed-4040B-NTE) filter set were used to record green or red fluorescence signals, respectively. Aliquots from the
same batches of cells were then plunge frozen on EM grids and loaded into the cryo-FLM setup (detailed procedure is described in the next section) for imaging at 80 K. Exposures of 0.1–10 s using a 405-nm laser with 100–1,000 W/cm² of power at the sample plane were used in an attempt to trigger photoactivation/ photoconversion. Exposures of 0.1–1 s using a 488-nm or 561-nm laser with 100–1,000 W/cm² of power at the sample plane were used to record green or red fluorescence signals.

Correlated cryo-PALM-CET. *M. xanthus* SA5718 cells were grown to an OD₅₅₀ of 0.4. 10-nm colloidal gold (Sigma-Aldrich) pretreated with bovine serum albumin was added to the cells for fiducial markers during tomogram reconstruction. 200-nm orange fluorescent Fluospheres (Life Technologies) were added to the cells for stage-drift correction in cryo-PALM and also for use as landmarks for correlating FLM and EM images. 5% Ficoll PM 70 (Sigma-Aldrich) or 10% Ficoll PM 70 with 10% ethylene glycol (Sigma-Aldrich) was added as a cryoprotectant for laser exposure. 3 µl of the resulting sample were pipetted onto a freshly glow-discharged Quantifoil gold London-finder EM grid (Quantifoil Micro Tools) and plunge frozen in a liquid ethane propane mixture using an FEI Vitrobot. Frozen grids were then loaded into Polara EM cartridges, transferred into a ethane propane mixture using an FEI Tecnai G2 Polara 300-keV FEG transmission electron microscope equipped with a Gatan energy filter and a Gatan K2 Summit direct detector. Tabs on the finder-grid were used to locate the cells imaged by cryo-PALM. Energy-filtered tilt-series of images on individual cells were collected automatically from −60° to +60° at 1° intervals using the UCSF Tomography data collection software with a total dosage of 180 e/Å² and a defocus of −10 µm. Three-dimensional reconstructions and segmentations were produced with IMOD.

To overlay the cryo-PALM signals onto cryo-tomograms, we acquired low-magnification EM images of the target cells to locate surrounding fluorescent polystyrene beads. Cryo-PALM images were then rescaled to match the EM images according to the locations of the beads by using the GNU image manipulation program (GIMP) (http://www.gimp.org/). The precision of the superimposition of the beads between FLM/EM in Supplementary Figure 2 is shown in Supplementary Table 1. Central slices of cryo-tomograms were superposed onto low-resolution EM images by aligning the edges of cells and the holes of the EM grid via GIMP.

\[
\frac{(\sigma_{x,y})^2}{N} = \frac{s^2 + a^2/12}{N} + \frac{8\pi s^4 b^2}{a^2 N^2}
\]

where \(N\) is the number of photons recorded in the fitted point spread function (PSF), \(a\) is the pixel size, \(b\) is the background noise and \(s\) is the s.d. of the PSF.

After cryo-PALM imaging, EM cartridges containing frozen grids were stored in LN and maintained at 123 K or less throughout transfer into and imaging in an FEI Tecnai G2 Polara 300-keV FEG transmission electron microscope equipped with a Gatan energy filter and a Gatan K2 Summit direct detector. Tabs on the finder-grid were used to locate the cells imaged by cryo-PALM. Energy-filtered tilt-series of images on individual cells were collected automatically from −60° to +60° at 1° intervals using the UCSF Tomography data collection software with a total dosage of 180 e/Å² and a defocus of −10 µm. Three-dimensional reconstructions and segmentations were produced with IMOD.

In Supplementary Figure 2, the acquisition time for each frame of the image stack (\(\tau_{frame}\)), total number of acquired frames in the image stack (\(F_{total}\)), total number of molecules localized in the image stack (\(M_{total}\)), maximum acceptable position error for inclusion in the final image (\((\sigma_{x,y})_{max}\)), and number of localized molecules composing the final image (\(M_{image}\)) for Figures 1 and 2 are summarized in Supplementary Table 1. The position error \(\sigma_{x,y}\) for each single molecule was estimated as previously reported, with

\[
\frac{(\sigma_{x,y})^2}{N} = \frac{s^2 + a^2/12}{N} + \frac{8\pi s^4 b^2}{a^2 N^2}
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