Eliminating effects of particle adsorption to the air/water interface in single-particle cryo-electron microscopy: Bacterial RNA polymerase and CHAPSO

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Abstract

Preferred particle orientation presents a major challenge for many single particle cryo-electron microscopy (cryo-EM) samples. Orientation bias limits the angular information used to generate three-dimensional maps and thus affects the reliability and interpretability of the structural models. The primary cause of preferred orientation is presumed to be due to adsorption of the particles at the air/water interface during cryo-EM grid preparation. To ameliorate this problem, detergents are often added to cryo-EM samples to alter the properties of the air/water interface. We have found that many bacterial transcription complexes suffer severe orientation bias when examined by cryo-EM. The addition of non-ionic detergents, such as NP-40, does not remove the orientation bias but the Zwitter-ionic detergent CHAPSO significantly broadens the particle orientation distributions, yielding isotropically uniform maps. We used cryo-electron tomography to examine the particle distribution within the ice layer of cryo-EM grid preparations of *Escherichia coli* 6S RNA/RNA polymerase holoenzyme particles. In the absence of CHAPSO, essentially all of the particles are located at the ice surfaces. CHAPSO at the critical micelle concentration coats the air/water interface and eliminates particle absorption there, allowing particles to randomly orient within the vitreous ice layer. We find that CHAPSO reduces orientation bias for a number of bacterial transcription complexes containing *E. coli* or *Mycobacterium tuberculosis* RNA polymerases. Findings of this study confirm the presumed basis for how detergents can help remove orientation bias in cryo-EM samples and establishes CHAPSO as a useful tool to facilitate cryo-EM studies of bacterial transcription complexes.

1. Introduction

Advances in single particle cryo-electron microscopy (cryo-EM) now allow structure determination of biological macromolecular complexes to near atomic-resolution (Cheng, 2018). Nevertheless, a major complication for many biological cryo-EM specimens is particle orientation bias (Glaeser and Han, 2017; Glaeser, 2016). Specimens that suffer particle orientation bias can have an anisotropic distribution of angular projection directions leading to under-sampling of Fourier components in the final three-dimensional reconstruction. This under-sampling leads to an overall loss of structural information parallel to the axis of preferred orientation, giving the maps an anisotropic point spread function leading to a "smearing effect" artifact (Barth et al., 1989; Naydenova and Russo, 2017; Tan et al., 2017), which affects the interpretability of the cryo-EM maps.

In examining a complex of *Escherichia coli* (Eco) 6S RNA bound to RNAP $\sigma^{70}$-holoenzyme (6S-E$\sigma^{70}$) for single-particle cryo-EM structure determination, we encountered a severe orientation bias problem (Chen et al., 2017). We explored a range of solution conditions and detergent additives to solve the orientation bias problem. We discovered that the detergent 3-([3-Cholamidopropyl]dimethylammonio)-2-hydroxy-1-propanesulfonate (CHAPSO) was uniquely effective. We hypothesized that the preferred orientation was due to adsorption and orientation of the particles at the air-water interface (Taylor and Glaeser, 2008) and that CHAPSO mitigated this problem by preventing adsorption at the interface. We used fiducial-less cryo-electron tomography (cryo-ET) on the single-particle specimens to visualize particle distributions within the vitreous ice (Noble et al., 2018a; Noble and Stagg, 2015). The results confirmed our hypothesis; the orientation bias arises from interactions of the particles with the surfaces of the ice layer. In the presence of a sufficient concentration of CHAPSO, the particles were excluded from the ice surfaces and distributed within the ice layer with nearly...
random orientations. We show that CHAPSO solves preferred orientation problems for a number of single-particle samples comprising bacterial transcription complexes.

2. Results

2.1. 6S-Eσ70 particles show severe orientation bias which is significantly relieved with CHAPSO

Initial single particle cryo-EM analysis of the 6S-Eσ70 complex was performed using a potassium L-glutamate buffer (KGlu, Supplemental Table 1). Micrographs showed uniform, homogenously dispersed particles that yielded detailed high-resolution 2D classes (Fig. 1A, Supplemental Fig. 1A). However, 3D alignment resulted in “smeared” maps (Supplemental Fig. 1B), suggesting that the particles in the vitreous ice layer exhibited orientation bias. The angular distribution plot of the particles aligned to a low-pass filtered X-ray crystal structure of Eco core RNAP (PDB ID 4LJZ with σ70 removed; (Bae et al., 2013) as a 3D template revealed a distribution corresponding to essentially one orientation (Fig. 1A). A rough characterization of the particle distribution as a Gaussian yielded a peak at about rotation angle (rot) −19°, tilt angle (tilt) −13°, and a standard deviation (±) of 15°.

To overcome this particle orientation bias, we prepared samples in a number of alternative conditions, such as a buffer containing KCl instead of KGlu (KCl), and the addition of non-ionic detergents n-dodecyl-β-D-maltoside (DDM), Triton X-100, Tween 20, and Nonidet P40 substitute (NP40S), and the Zwitterionic detergent CHAPSO (Supplemental Table 2). All of the detergents were added at their critical micelle concentration (CMC). We screened the samples for complex stability using an electrophoretic mobility shift assay, and particle homogeneity/distribution using negative stain EM. Triton X-100 and Tween 20 (at CMC) affected the stability of the 6S-E σ70 complex and were not explored further. The 6S-E σ70 complex appeared to be stable with DDM, but negative stain EM images showed protein aggregation. From these results, we proceeded with Cryo-EM analysis of the four sample conditions listed in Supplemental Table 1.

Single particle cryo-EM analysis of the 6S-E σ70 particles in the KCl condition also showed severe orientation bias (Fig. 1B), with one peak at an orientation and spread very similar to the KGlu condition (rot −12°, tilt −8°, ± 14°), but with an additional peak at approximately (rot 168°, tilt 8°), corresponding to a mirror image projection of the first orientation. Since the mirror image projection does not contribute any new information to the 3D reconstruction, having two mirror image orientations is equivalent to having one orientation.

Preparation of the particles in KGlu + NP40S (Fig. 2A) also yielded two mirror image peaks [(rot −130°, tilt −47°) and (rot 50°, tilt 47°)]. The distribution was broadened with respect to the KGlu and KCl distributions, with standard deviation ± 20°. Thus, the addition of the detergent still gave only one effective orientation, but the bias was slightly ameliorated. In contrast to KGlu, KCl, and KGlu + NP40S, particles prepared in KCl + CHAPSO did not exhibit peaks of preferred orientation; instead the particle orientations were spread over a large fraction of Euler angles (Fig. 2B), resulting in isotropically uniform 3D reconstructions (Supplemental Fig. 1B).
For each individual sample, the particles were grouped into 163 bins according to their Euler angles (corresponding to 20° increments in rotation and tilt angles) and ranked according to the percentage of particles in each bin (bin #1, highest %; bin #2, next highest %; so on). The orientation distribution of the particles prepared in each condition was compared by plotting the histograms of the % particles in each bin according to bin # (Fig. 2C). A completely random distribution of particle orientations would yield a flat distribution, with 0.61% particles in each bin (dashed horizontal line in Fig. 2C). Visualizing the orientation distributions this way highlights the bias of the KGlu, KCl, and KGlu + NP40S samples. The KCl + CHAPSO sample, while not completely randomized, approaches the random orientation distribution more closely. The inset of Fig. 2C plots the cumulative % of particles across the bins. This plot reveals that 50% of the particles are binned into only 5, 9 and 12 bins for the KGlu, KCl, and KGlu + NP40S conditions, respectively, while 50% of the KCl + CHAPSO particles are spread out over 36 bins (Fig. 2C).

The particle orientation distributions and their effects on resulting reconstructions were also analyzed using cryoEF (Naydenova and Russo, 2017). A cross-section through the middle of the expected PSFs (calculated using cryoEF (Naydenova and Russo, 2017) are superimposed, illustrating the anisotropy for the KGlu (red), KCl (blue), and KGlu + NP40S (orange) samples, while the KCl + CHAPSO sample yields an isotropic PSF (green). Parameters further characterizing the orientation distributions (the orientation efficiency, $E_{od}$, and the fraction of unsampled Fourier space, $f_{empty}$ (Naydenova and Russo, 2017) are also tabulated. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
efficiency ($E_{cd}$) and the fraction of unsampled Fourier space ($f_{	ext{empty}}$) (Naydenova and Russo, 2017), illustrating the dramatic improvement with CHAPSO.

2.2. Cryo-electron tomography shows that orientation bias corresponds to adsorption to an ice surface

We employed fiducial-less cryo-electron tomography (cryo-ET) on the single-particle specimens in order to visualize the locations of the particles in the vitreous ice layer (Noble et al., 2018a; Noble and Stagg, 2015). Tilt series of cryo-grids of the 6S-Eσ70 complex were collected for each solution condition (Fig. 3). The 6S-Eσ70 particles prepared in KGlu and KCl were restricted to a thin layer at one of the ice surfaces (Fig. 3A, B). The complexes were prepared with excess 6S RNA and free 6S RNA molecules could be visualized distributed throughout the vitreous ice layer (Fig. 3A, B, Supplementary videos). The 6S-Eσ70 particles prepared in KGlu + NP40S were restricted to two thin layers corresponding to both ice surfaces (Fig. 3C). By contrast, particles prepared in KCl + CHAPSO were excluded from the air/water interfaces and were evenly distributed throughout the middle of the ice layer (Fig. 3D).

2.3. Effect of CHAPSO on particle orientations is concentration dependent

Our results thus far indicate that adsorption of the 6S-Eσ70 particles at air/water interfaces gives rise to the severe orientation bias seen in the KGlu, KCl, and KGlu + NP40S samples (Figs. 1, 2). The addition of CHAPSO at the critical micelle concentration (CMC, 8 mM) completely eliminates surface interactions and significantly randomizes the particle orientations (Fig. 2B-D, 3D). To investigate if CHAPSO at CMC was required for the full effect, we compared particle orientations for datasets collected with 0, 4 mM (0.5XCMC), and 8 mM (1XCMC) CHAPSO.

Fig. 3. Cryo-ET reveals mechanism for preferred orientation. A – D. (Left Panel) Surface tomographic cross-section of the vitreous ice layer. (Middle Panel) Middle tomographic cross-section of the vitreous ice layer. (Top Right Panel) Schematic diagram of particle distribution in vitreous ice. Top and bottom surfaces of the ice are shown with a solid blue line. Thickness of ice is indicated on the bracket right of the cartoon. 6S-Eσ70 and free 6S RNA particles are shown as grey volumes in the cartoon. (Bottom Right Panel) Spatial plot of particles in vitreous ice layer, oriented orthogonal to the ice surface. Each 6S-Eσ70 particle is represented as a blue point and graphed based on 3D position in the ice layer. A. Tomogram of 6S-Eσ70 particles in KGlu. B. Tomogram of 6S-Eσ70 particles in KCl. C. Tomogram of 6S-Eσ70 particles in KGlu + NP40S. D. Tomogram of 6S-Eσ70 particles in KCl + CHAPSO. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Fig. 4. Effect of CHAPSO on particle orientations is concentration dependent. A – C. (Top Panel) 3D distribution plot of particle orientations. Particles were 3D classified into one class using Eco core RNAP (PDB ID 4LJZ (Bae et al., 2013); σ70 was deleted and the structure was low-pass filtered using EMAN2) (Tang et al., 2007) as a 3D template in RELION (Scheres, 2012). The resulting density is shown as a solid grey volume and the angular distribution from this alignment is shown as red spheres. Each sphere represents a particular Euler angle and the sphere volume represents the absolute number of particles at that particular angle. (Bottom Panel) 2D distribution plot of particle orientations. Particles are plotted on a tilt angle vs rotation angle graph. Areas of the points represents the percentage of particles at that particular orientation. A. Angular distribution of TEC particles in TEC buffer (20 mM Tris-HCl, pH 8.0, 150 mM KCl, 5 mM MgCl2, 5 mM DTT) without CHAPSO. B. Angular distribution of TEC particles in TEC buffer + 4 mM CHAPSO (0.5XCMC). C. Angular distribution of TEC particles in TEC buffer + 8 mM CHAPSO (1XCMC). D. Particles for each dataset were grouped into Euler angle bins (20° rotation angle × 20° tilt angle bind) and then the bins were ranked according to the number of particles populating that bin (bin #1 has the most particles, so on). Plotted on a semi-log scale is the percent of total particles in each dataset by bin #. The horizontal dashed line represents a totally random particle orientation distribution (equal number of particles in each bin). (Inset) Plotted is the cumulative percent particles by bin #. The random distribution is denoted by the dashed line. E. Cross-sections through the middle of the expected PSFs (calculated using cryoEF (Naydenova and Russo, 2017) are superimposed. Parameters further characterizing the orientation distributions (the orientation efficiency, $E_{od}$, and the fraction of unsampled Fourier space, $f_{empty}$ (Naydenova and Russo, 2017) are also tabulated. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
of a different bacterial transcription complex, an Eco RNAP ternary elongation complex (TEC) (Kang et al., 2017). In the absence of CHAPSO, the particles again exhibited severe orientation bias (Fig. 4A). The particle orientations were significantly spread by the presence of 4 mM CHAPSO (Fig. 4B), but there is a clear difference in the particle spread between 4 mM and 8 mM CHAPSO (Fig. 4B, C) and 8 mM CHAPSO induces a distribution of particle orientations much closer to a random distribution (Fig. 4D, E).

2.4. Cryo-EM maps reveal CHAPSO interacts with specific sites on the Eco RNAP surface

Examination of the nominal 3.5 Å resolution cryo-EM map of an Eco RNAP transcription elongation complex bound to RfaH (Kang et al., 2018a), the highest resolution cryo-EM map available for an Eco RNAP transcription complex, revealed three CHAPSO molecules bound to the RNAP surface (Fig. 5A). Retrospective analysis of previously published cryo-EM structures of Eco RNAP transcription complexes where 8 mM CHAPSO was used to prevent orientation bias revealed CHAPSO molecules consistently bound at the same sites (Fig. 5B). In each case, the cholic acid-derived amphifacial moiety of CHAPSO was bound to the RNAP while the attached flexible chain and hydrophilic head group were disordered.

3. Discussion

From the results of this study, we conclude that:
1) Transcription complexes containing Eco RNAP adsorb strongly to air/water interfaces during cryo-EM grid preparation, even in the presence of non-ionic detergents such as NP40S (Fig. 3A-C).

2) The complexes adsorbed to the air/water interface are oriented, confounding single particle reconstruction approaches (Figs. 1, 2, 4).

3) The addition of CHAPSO at the CMC (8 mM) completely prevents adsorption of the complexes to air/water interfaces (Fig. 3D) and dramatically broadens the distribution of particle orientations (Figs. 2B-D, 3D, 4C-E), allowing for the determination of isotropically uniform maps (Figs. 2D, 4E; Supplemental Fig. 1B) (Naydenova and Russo, 2017).

Although CHAPSO at 0.5 CMC (4 mM) significantly broadens the particle orientation distribution compared to no CHAPSO (Fig. 4A, B, D) and the calculated PSF and orientational parameters indicate that a detailed reconstruction would be achieved at this condition (Fig. 4E) (Naydenova and Russo, 2017), the full effect on the relative randomization of the particle orientation distribution requires CHAPSO at its CMC (8 mM; Fig. 4). At the same time, due to the high CHAPSO concentration, we observed CHAPSO molecules consistently bound to three sites on the surface of Eco RNAP (Fig. 5). Our cryoET results showed that in the presence of 8 mM CHAPSO, the RNAP complexes are completely excluded from the air/water interfaces and found in the middle of the ice layer (Fig. 3D) where rotational diffusion allows for nearly randomized particle orientations. This effect of CHAPSO on eliminating RNAP adsorption at the air/water interface could be due to: 1) CHAPSO at or above the CMC coating the air/water interfaces with a monolayer, altering the interfacial surface properties to prevent RNAP adsorption, and/or 2) CHAPSO molecules binding to the surface of RNAP (Fig. 5), altering the RNAP surface properties to prevent adsorption at the air/water interface. Our result that 8 mM CHAPSO is required for the full effect (Fig. 4) strongly supports hypothesis (1) as the primary factor, since at 4 mM CHAPSO we still observe CHAPSO molecules bound to the RNAP (Supplemental Fig. 2).

In the single particle analyses under conditions where the RNAP complexes are adsorbed to air/water interfaces and oriented, we sometimes observe an orientation distribution comprising one peak (Fig. 1A), while in other cases we observe two peaks, one corresponding to the mirror image projection of the other (Figs. 1B, 2A, 4A). The cryoET analysis revealed some samples where the complexes were adsorbed to only one air/water interface (Fig. 3A, B), while in other samples particles were adsorbed to both the top and bottom interfaces (Fig. 3C), explaining the observation of mirror image projection orientations. While we have not studied this phenomenon specifically, our data suggest that whether the complexes are adsorbed to one or both interfaces is a random occurrence dependent on grid preparation rather than properties of the solution conditions.

A number of cryo-EM structures of bacterial RNAP transcription complexes have benefited from CHAPSO (Chen et al., 2017; Kang et al., 2017; Kang et al., 2018a,b; Boyaci et al., 2018). The observation of specifically-bound CHAPSO molecules on the Eco RNAP surface (Fig. 5) raises the potential for the introduction of structural artifacts. For this reason, it is important to examine the complexes under investigation with sensitive and quantitative functional assays to show that the biochemical function is not altered by the presence of CHAPSO, as was shown in each of the cases listed in Fig. 5B. We believe the very high concentration of CHAPSO (8 mM) allows binding at the observed sites on the RNAP (Fig. 5) but that the binding energy for these sites is very low and is insufficient to alter the conformational/functional properties of the RNAP. For instance, a conformational change in the HisPEC altered the relationship between β i9 and the rest of the RNAP, disrupting CHAPSO site 3 (Fig. 5), and no complexes were observed with CHAPSO bound at this site. In biochemical assays, 8 mM CHAPSO had a minimal (less than 2-fold) effect on the pause lifetime (Kang et al., 2018b).

Orientation bias has long been an issue confounding single particle cryo-EM structure determination (Glaeser and Han, 2017; Glaeser, 2016). Many of the recent advances in high-resolution structure determination of macromolecular complexes by cryo-EM do not address this potential obstacle (Naydenova and Russo, 2017). Maps generated from oriented samples are limited in their interpretability due to a “smearing” artifact in the final 3D construction (Supplemental Fig. 1B). 2D classification can be an effective method for diagnosing whether a sample suffers from preferred orientations (Figs. 1A, B, 2A, B). A common solution to mitigate particle orientation bias has been to add surfactants during grid preparation (Taylor and Glaeser, 2008). It has often been presumed that particle orientation bias arises from adsorption to the air/water interface and that surfactant additives mitigate the orientation bias by reducing the propensity for interface adsorption. Our cryoET analysis establishes that this is indeed the case for samples comprising Eco RNAP transcription complexes and the surfactant CHAPSO (Fig. 3).

Recent studies have established that severe particle orientation bias can be overcome by cryo-EM data collection from tilted grids (Tan et al., 2017) or by rapid grid freezing after sample application (Noble et al., 2018b). These are important advances that are potentially generally applicable. Nevertheless, imaging on tilted grids presents many technical obstacles to optimal high-resolution data collection. Moreover, tilting the grid does not address the issue of particle adsorption, which has recently been suggested to cause denaturation for most particles (D’Imprima et al., 2018). The specialized plunge-freezing devices necessary for sufficiently rapid plunge-freezing to ‘outrun’ some of the air/water interface adsorption effects are not yet widely available (Noble et al., 2018b). We show here that the addition of CHAPSO during cryo-EM grid preparation of samples comprising Eco RNAP transcription complexes is relatively functionally inert, completely eliminates interaction and orientation of the particles at air/water interfaces (Fig. 3), and significantly broadens the particle orientation distributions to allow determination of high-resolution cryo-EM maps with isotropic PSFs (Figs. 2D, 4E). These properties greatly facilitate high-resolution structure determination of these complexes using modern cryo-EM approaches (Chen et al., 2017; Kang et al., 2017; Kang et al., 2018a,b; Boyaci et al., 2018).

4. Materials and methods

4.1. Protein expression and purification

Eco RNAP TEC was assembled using Eco RNAP core lacking the α C-Terminal Domain (ΔαCTD) and was prepared as previously described (Kang et al., 2017; Twist et al., 2011). The 6S-Δσ70 complex was prepared as described previously (Chen et al., 2017).

4.2. Preparation of 6S-/Δσ70 for single particle Cryo-EM

Frozen aliquots of Eco RNAP ΔαCTD-core and Δσ70Δ1.1 were mixed in a 1:2 M ratio and incubated for 15 mins at 37 °C. 6S RNA was added in a 1:1.5 M ratio and incubated for 15 mins at room temperature. The samples were diluted and detergents added (if used) immediately before grid preparation. 3.5 μL of sample were deposited on glow discharged Quantifoil R 1.2/1.3, 400 mesh, copper grids (EMS) and plunged frozen into liquid ethane using a Cryoplunge 3 system (Gatan).

4.3. Single particle cryo-EM of 651-Δσ70 complex

Cryo-EM grids of 6S-Δσ70 were imaged using a 300 kV Tecnai G2 Polara (FEI) equipped with a K2 Summit direct electron detector (Gatan). Dose-fractionated images were collected using UCSFImage4 (Li et al., 2015) in super-resolution mode with a nominal magnification of 31,000 ×, corresponding to a calibrated pixel size of 1.23 Å on the specimen level (0.615 Å for super-resolution). The dose rate on the
camera was 8 counts/physical pixel/second using Digital Micrograph (Gatan). The exposure time per movie was 6 s (30 frames) leading to a total dose of 31.7 electrons/Å². Movies were collected using a defocus range from 1 µm to 2.5 µm. Movies were 2 × 2 binned using IMOD (Kremer et al., 1996) and then drift corrected using MotionCor (Li et al., 2009). Imaging conditions are summarized in Supplemental Table 3.

4.4. Preparation of 6S-Eσ70 for Cryo-ET

For KGlu and KGlu + NP40S conditions (Supplemental Table 1), Eσ70 was purified in KGlu buffer using a Superose6 INCREASE column (GE Healthcare). For KCl and KCl + CHAPSO, Eσ70 was purified in KCl buffer. Peak fractions corresponding to Eσ70 were pooled and concentrated by centrifugal filtration (VivaScience) to 4 mg/mL protein concentration. 6S RNA was added in 1.2 M excess compared to holoenzyme and incubated at room temperature. Immediately prior to grid freezing, samples of KGlu or KCl were diluted 1:10 with their respective buffers while NP40S was added to the KGlu + NP40S to CMC (0.06 mM) and CHAPSO was added to the KCl + CHAPSO sample to CMC (8 mM). After centrifugation to remove aggregates, 3.5 µL of sample were deposited on glow discharged Quantifoil R 1.2/1.3, Au, 400 mesh grids (EMS) and plunged frozen into liquid ethane using a Vitrobot Mark IV (FEI).

4.5. Acquisition of cryo-electron tomograms of 6S-Eσ70

Tilt-series were collected at NYSBC using Titan Krios #1 (FEI Company, Hillsboro, OR) with a Gatan K2 (Gatan, Inc., Pleasanton, CA). Tilt-series were collected using Leginon (Suloway et al., 2009) with 100 ms frames for each tilt image, which were full-frame aligned using MotionCor2 (Zheng et al., 2017). Tilt-series were collected bi-directionally with a tilt range of −45˚ to 45˚ and a tilt increment of 3˚. Most tilt-series were collected at a nominal defocus between 4 and 6 µm. Tilt-series were collected with a dose rate around 8 e-/pixel/s and an incident dose of 3.29 e-/Å² for the zero-degree tilt image, with increasing dose for higher tilt angles according to the cosine of the tilt angle, resulting in a total dose of 120 e-/Å². Most tilt-series were collected at a pixel size of 1.33 Å. Imaging conditions are summarized in Supplemental Table 4.

4.6. Cryo-ET data analysis

Tilt-series were aligned using Appion-Protomo (Noble and Stagg, 2015; Winkler and Taylor, 2006; Lander et al., 2009). Tilt-series were first dose compensated using equation 3 in Grant, Grigorieff (Grant and Grigorieff, 2015a), coarsely aligned, manually fixed if necessary, refined using a set of alignment thicknesses, then the best aligned iteration was reconstructed for visual analysis using Tomo3D SIRT (Agulliero and Fernandez, 2011; Agulliero and Fernandez, 2015). CTF correction was not performed. Subtomogram analysis was performed with Dynamo (Castano-Diez et al., 2012; Navarro et al., 2018). First, about 10 representative particles were manually picked with two defined Euler angles, averaged together to make a template, then the tomograms were template picked, and the picks were cleaned manually. Data processing is summarized in Supplemental Table 4.

4.7. Preparation of Eco RNAP TEC for single particle cryo-EM

Purified RNAP ΔαCTD-core was buffer-exchanged over the Superose 6 INCREASE (GE Healthcare Life Sciences) column into 20 mM Tris-HCl, pH 8.0, 150 mM KCl, 5 mM MgCl₂, 5 mM DTT. At a molar ratio of 1.3:1, template DNA/RNA hybrid was mixed into the eluted RNAP core and incubated for 15 min at room temperature. Subsequently non-template DNA was added and incubated for an additional 10 min (Kang et al., 2017). The complex was concentrated by centrifugal filtration (VivaScience) to 3 mg/mL RNAP concentration before grid preparation.

CHAPSO was added to the samples to give a final concentration of 0xCMC, 0.5xCMC, or 1xCMC. C-flat CF-1.2/1.3 400 mesh copper grids (EMS) were glow-charged for 15 s. 3.5 µL of sample (~2.0–3.0 mg/mL protein concentration) was absorbed onto the grid, blotted, and plunge-frozen into liquid ethane using a Vitrobot Mark IV (FEI).

4.8. Single particle Cryo-EM of RNAP TEC

Grids of RNAP TEC were imaged using a 300 keV Krios (FEI) (for 0xCMC CHAPSO and 1xCMC CHAPSO datasets) or a 200 keV Talos Arctica (FEI) (for 0.5xCMC CHAPSO). Both microscopes were equipped with a K2 Summit direct electron detector (Gatan). Imaging parameters were as outlined previously (Kang et al., 2017). Dose-fractionated images were recorded with Serial-EM (Mastronarde, 2005) in super-resolution mode with a super-resolution pixel size of 0.65 Å (nominal magnification 22,500x and a calibrated pixel size of 1.3 Å) on the Titan Krios and with a super-resolution pixel size of 0.75 Å (nominal magnification 28,000 × and a calibrated pixel size of 1.5 Å) on the Talos Arctica. The dose rate at the camera level was 10 electrons/physical pixel/second in Digital Micrograph (Gatan). Images were recorded in dose-fractionation mode with subframes of 0.3 s over a total exposure of 15 s (50 frames). Images were collected over a defocus range of 0.8 µm to 2.6 µm. Movies were gain-normalized and 2X2 binned using ‘mag_distortion_estimate’ (Grant and Grigorieff, 2015b). Images were drift-corrected and summed using Unblur (Grant and Grigorieff, 2015a). Imaging conditions are summarized in Supplemental Table 3.

4.9. Single particle cryo-EM data analysis

The Cryo-EM data analysis pipeline used for all single particle cryo-EM datasets is illustrated in Supplemental Fig. 1A. CTF estimations were calculated for each dataset using Getf (Zhang, 2016). Particles were picked using Gautomatch (developed by K. Zhang, MRC Laboratory of Molecular Biology, Cambridge, UK, http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch) without a 2D template. Picked particles were extracted from the dose-weighted images in RELION (Scheres, 2012). Particles were curated by 2D classification (N classes = 50) and 3D classification (N classes = 3) using a crystal structure of Eco RNAP (PDB ID 4LJZ) (Baé et al., 2013) with σ70 removed. The crystal structure was converted to an EM map and low-pass filtered to 60 Å using EMAN2 (Tang et al., 2007) before classification and refinements. Particles were coarsely aligned to the 3D template using RELION (Scheres, 2012) 3D classification (N class = 1). Histograms of particle orientations (BILD format) were graphically represented as spheres. The PSFs and orientation distribution parameters (f₆₀ and f₀₆₀) were calculated using cryoEF (Naydenova and Russo, 2017). Data processing of single particle cryo-EM datasets is summarized in Supplemental Table 3.

5. Accession numbers

Single particle cryo-EM micrographs, cryo-ET tilt-series, cryo-ET tilt-series alignment runs with Appion-Protomo, and cryo-ET tomograms have been deposited to the Electron Microscopy Pilot Image Archive (EMPIAR) with accession code EMPIAR-10214.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jysb.2019.100005.

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