

# Imaging protein conformational space in liquid water

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## Abstract

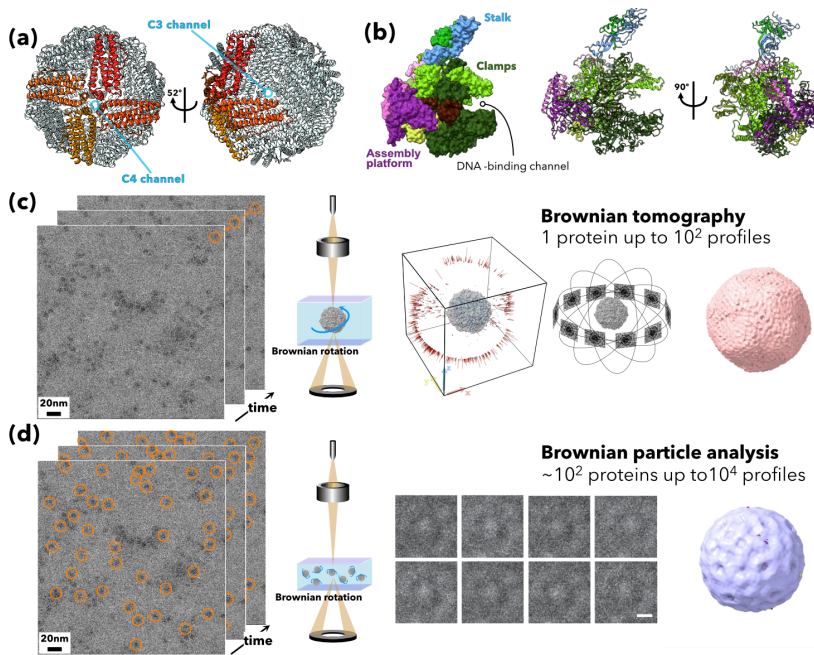
Recent developments in electron-transparent materials have paved the way for liquid-phase electron microscopy (LPEM) leading to an unprecedented understanding of the structure and dynamics of specimens in their liquid environment. Image reconstruction in liquid-state poses several challenges, and most importantly, it undermines the single-particle analysis assumption that the three-dimensional objects captured on the image sensor are identical. We propose the combination of all-atom simulations with LPEM to complement structural studies with dynamic investigations. In this work, we exploited LPEM to image the dynamics of particles undergoing Brownian motion, using their natural rotation to access the particle structural landscape for reconstructing its architecture in 3D using tomographic techniques. We have selected two test cases for our approach according to prior data accessibility and physiological environment factors: apoferritin and archaeal RNA polymerase. We show that the adopted approach allows to achieve sub-nanometer spatial resolutions of protein structures, either imaging proteins one by one and assessing different conformational states or combining several proteins into one statistical conformational ensemble.

## Introduction

Electron microscopy (EM) is one of the most powerful techniques for structure determination with the ability to image matter down to the subatomic level. The short wavelength associated with electrons is pivotal for achieving such a high resolution. However, electrons interact strongly with matter making imaging possible only under ultrahigh vacuum conditions. Such a necessity limits EM imaging to solid-state or non-volatile specimens. Thus samples dispersed in most common liquids or containing liquids, especially water, require special preparation techniques involving either controlled drying or cryogenic treatments with consequent artefacts. Such alterations become particularly critical for biological samples whose meso- and nano-scale structures comprise water as one of the building blocks [1]. Some of these limitations can be overcome using fast vitrification processes to solidify liquid samples [2]. In the last three decades, cryogenic electron microscopy (cryoEM) has radically changed biology [3-7]. A critical step forward in the tremendous advancement of cryoEM has been the development of single-electron detectors and image analysis techniques, such as electron tomography (ET) and single-particle analysis (SPA), for resolving specimens' three-dimensional

(3D) structure. Structure determination has enabled biologists to shed light on the hierarchical organisation and associated functions of complex biological machinery [8, 9]. In ET, many specimen two-dimensional (2D) projections or profiles are collected by rotating the samples to different angles [8, 9]. In SPA the specimen profiles are collected over large number of images containing many particles per field-of-view with the general assumption that all particles are identical or very similar to each other [4, 10]. For both techniques all the profiles are classified, aligned, refined, and collated together to give form to a 3D structure [11]. A large number of profiles provides high spatial resolution thus making SPA considerably superior to ET, which often is limited to the analysis of large complexes or sub-cellular sections [8, 9] which are too large and/or heterogenous for SPA. In the case of proteins, SPA enables atomic reconstruction [12] with tremendous consequences in molecular and chemical biology, and drug discovery [13]. Cryogenic and standard dry TEM provide snapshots of three-dimensional (3D) structural information that can be correlated with function and more recently with close to single angstrom resolution [14-16]. Yet, most biomolecules such as proteins, nucleic acids, or sugars are naturally soft and subjected to thermal fluctuations and internal motion over a broad range of length and time scales. Biomolecules' conformational space is unique to each temperature and environmental conditions and includes several micro-states. Shedding light on the dynamics of biomolecules is critical to understand molecular recognition patterns, enzymatic conversions, and allostery. Thus, it would appear evident that the new frontier of microscopy is the temporal dimension, and the ability to access dynamic changes near physiological temperatures where the molecule of interest operates. While four-dimensional electron microscopy has been proposed with temporal resolutions approaching nanoseconds [17], accessing the biomolecule conformational landscape under physiological conditions is still a major challenge. In cryoEM, rapid sampling [18, 19] or appropriate classification during SPA refinement to separate different configurations [20, 21] can probe protein conformational states, but the time resolution is limited and the molecules are vitrified at -180°C. Physiological conditions can also be achieved using NMR spectroscopy which grants access to the biomolecule structure and dynamics with atomic resolution. However, unlabelled structure determination via NMR is very much restricted to small proteins, with the requirement of incorporating stable isotopes for structures larger than ~100kDa [22], and often in combination with other methods such as X-ray crystallography or cryoEM imaging [20]. More importantly structure and dynamics determination by NMR require time-consuming chemical shift assignment [22]. These challenges call for a completely new approach to probe biological molecules and materials at physiological temperature and with relevant timescales. The recent development of electron-transparent materials and liquid-phase electron microscopy (LPEM) have enabled the imaging of samples at room temperature and in liquid state [23, 24]. LPEM imaging has been applied to study electrochemical reactions [25], nanocrystal growth [26, 27], whole cells [28, 29], and to extract atomistic structure of inorganic particles in liquid state [30, 31] to name a few applications. Examples of 3D reconstruction of particles in liquid have been reported for inorganic particles [30, 31] and proteins immobilised on the viewing window of the liquid cell [32].

In the present investigation, we employ LPEM to image protein complexes undergoing Brownian motion. In this way, we exploit the proteins natural rotation in liquid in order to access most of the proteins profiles with the aim of reconstructing their 3D structure using tomographic techniques. We imaged proteins dispersed in phosphate buffer solution (PBS) using two liquid cell holders from DENSSolutions, the Ocean and Stream holders (**Figure S1**). The imaging set-up comprised a JEOL JEM2200 FS TEM equipped with a metal-oxide-semiconductor (CMOS)-based direct electron detection camera (Gatan K2 IS) which allows low-dose and high speed imaging with high detective quantum efficiency (DQE) [33]. We chose two different proteins to



**Figure 1. Description of Brownian tomography and particle analysis.**

Structure of apoferritin (a) showing both C3 and C4 channels, and archaeal RNA polymerase (RNAP) showing the different domains (b). Schematics of the Brownian tomography (c) and Brownian particle analysis (d) workflows to extract the 3D structure of a protein dispersed in water and undergoing Brownian motion. Reconstructed 3D models of apoferritin via Brownian Tomography and Brownian particle analysis shown at the far right side of (c) and (d) respectively.

establish our imaging and analysis methodologies: apoferritin (**Figure 1a**) and archaeal RNA polymerase (RNAP) (**Figure 1b**). Apoferritin quaternary structure is an octahedral shell comprising 24  $\alpha$ -helical subunits with 432 point symmetry, forming a hollow spherical shell [34] with an external and internal diameter of 12 and 8nm respectively. The three- and four-fold symmetry axes traverse the protein structure across eight C3 and six C4 channels respectively. The C3 channels emerge from the contact of three different subunits pairs and are circa 2 nm in diameter and negatively charged. The C4 channels result from the contact of four different subunits pairs and are

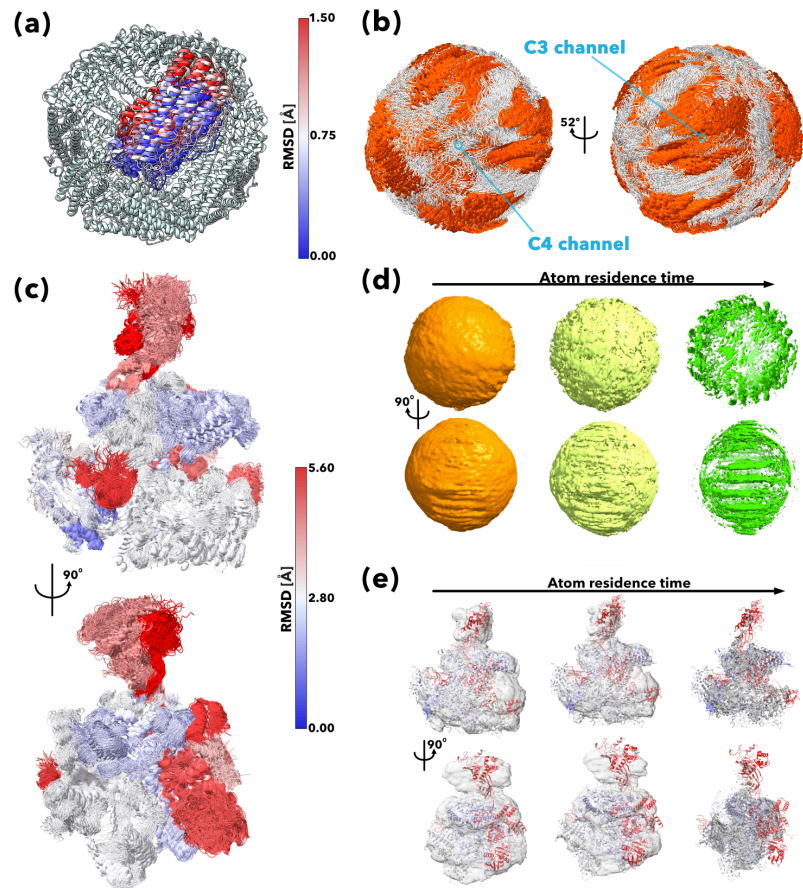
narrower than the C3 channels, and to a large extent hydrophobic [36].

Double-psi beta barrel (DPBB) RNA polymerases (RNAPs) carry out transcription in all cellular life forms as well as some viruses and bacteriophages [35]. Archaeal and eukaryotic RNA polymerases are closely related in terms of subunit composition, structure, molecular mechanisms, utilisation of basal transcription factors, promoter DNA sequence motifs, and histone-based chromatin templates [36]. As shown in **figure 1b**, we can identify four prominent structural features of RNAP, the DNA-binding channel, the clamp that closes over the DNA-binding channel and stabilises RNAP-DNA complexes, the assembly platform, and the stalk that interacts with nascent RNA chain and serves as docking point for accessory RNA-processing factors. [36] Both RNAP clamp and stalk are flexible motifs, and their exact conformation is influenced by the binding of general factors (TFE and Spt4/5), and the DNA/RNA scaffold of transcription complexes as RNAP progresses repeatedly through the transcription cycle [36][37, 38].

We imaged apoferritin in liquid phase (**Figures S2.a-b**) obtaining contrast similar to that afforded by cryogenic fixation (**Figure S2.b**) under the same imaging conditions. We achieve similar results with RNAP proteins (**Figures S2.c-d**) and most importantly we can collect a sufficient number of frames maintaining the total electron dose well below the damage lethal dose threshold  $<10$  electron per  $\text{\AA}^2$  of cumulative dose, as recognised from cryoEM work [39]. We imaged protein dispersions free to diffuse in water over time, and we propose two approaches to analyse their structure: Brownian tomography (BT) (**Figure 1c**) and Brownian particle analysis (BPA) (**Figure 1d**). In BT we select and track a single protein long enough to access most of its profile views which are then aligned and converted into a 3D structure. In BPA, we image a given population of proteins captured in the field of view over time and create a

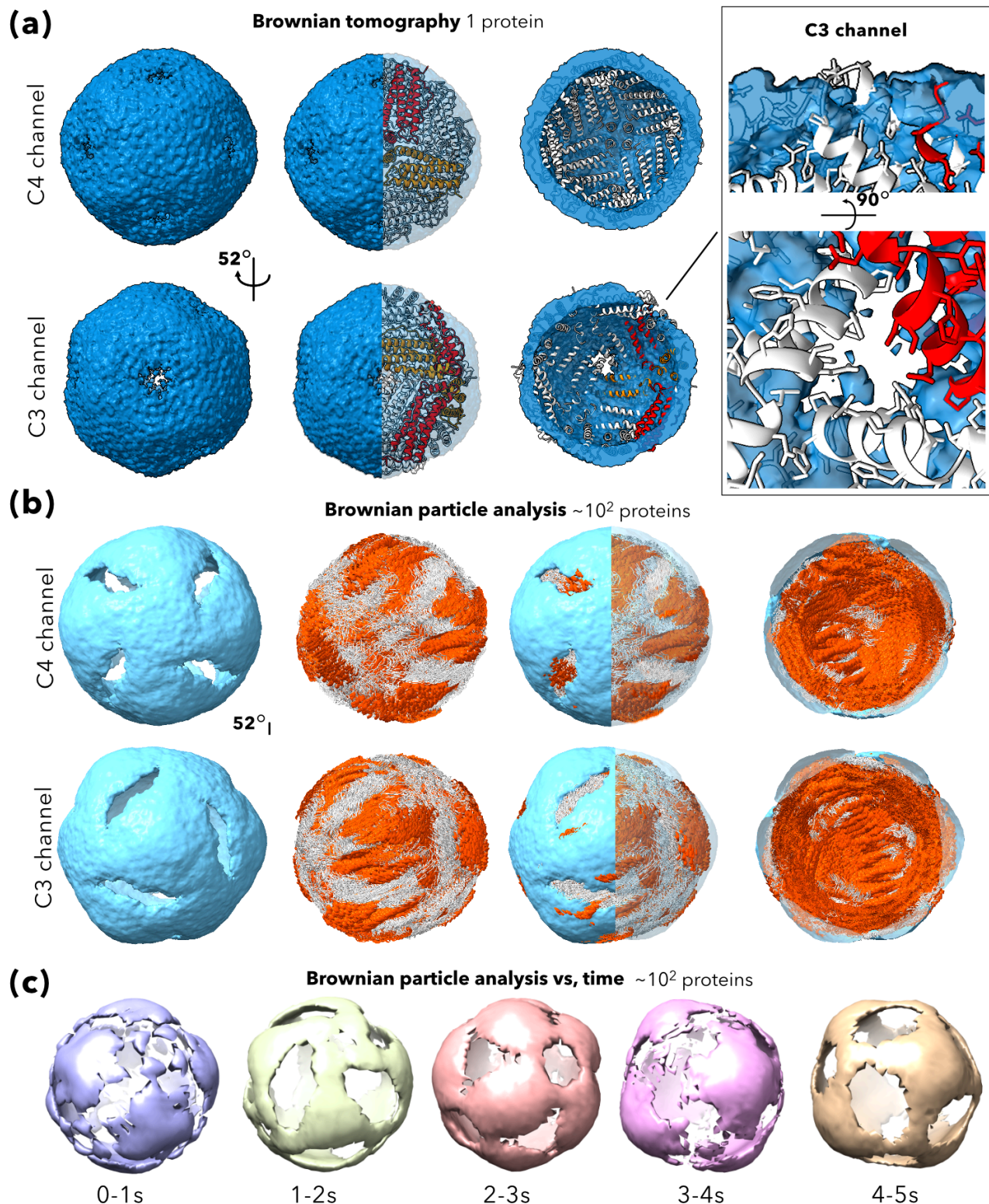
data set of different profiles which are sifted and sorted into different 2D classes to reconstruct the relative 3D structure. Such a process is iterative and involves, first an alignment and averaging step to create an initial volume that becomes the reference for the next iteration.. This initial model is further refined through an iterative algorithm either unbiased or introducing an initial reference model increasing spatial resolution across iterations [40]. Such a reference model should be accurate enough to improve data refinement but not too accurate in order not to bias the data. Hence, it is common practice to use a low-pass filtered map of a known similar structure. This latter can be either a structure or a sub-structure obtained by other experimental techniques (e.g. x-ray or NMR), or computationally generated homology or even de-novo models. In **Figure S3**,

we show the electron density (ED) maps for apoferritin generated by means of both BT and BPA using either unbiased refinement or a low-pass filtered ED map calculated from the deposited x-ray apoferritin PDB structure. The generated maps show that templating the refinement improves the resolutions considerably for both reconstruction methods allowing the visualisation of the core-shell structure of apoferritin. However, only the BT maps show the expected octahedral symmetry (**Figure S3.a**) while the BPA maps show a spherical structure with undistinguishable features (**Figure S3.b**). Most importantly, the resolution achieved for none of the generated ED maps is sufficient to visualise unique apoferritin structural elements such as C4 or C3 channels. The lack of detail is very likely due to the fact that proteins in liquid at room temperature can explore wider conformational spaces than in frozen vitrified state. Our time resolution ( $\sim 100\text{ms}$ ) is not close to the protein rotational correlation time  $\tau_R \sim 1\mu\text{s}$  nor the protein internal structure vibrations with correlation times in the order of ns. Whilst protein rotational dynamics being faster than our sampling aid to capture more profiles from one frame to another, we expect our reconstructions independently of the intervals used, to be time-averaged. In this fashion, we anticipate that the LPEM generated reconstructions represent the volume occupied by all the



**Figure 2. Molecular dynamics of apoferritin and archaeal RNAP.**

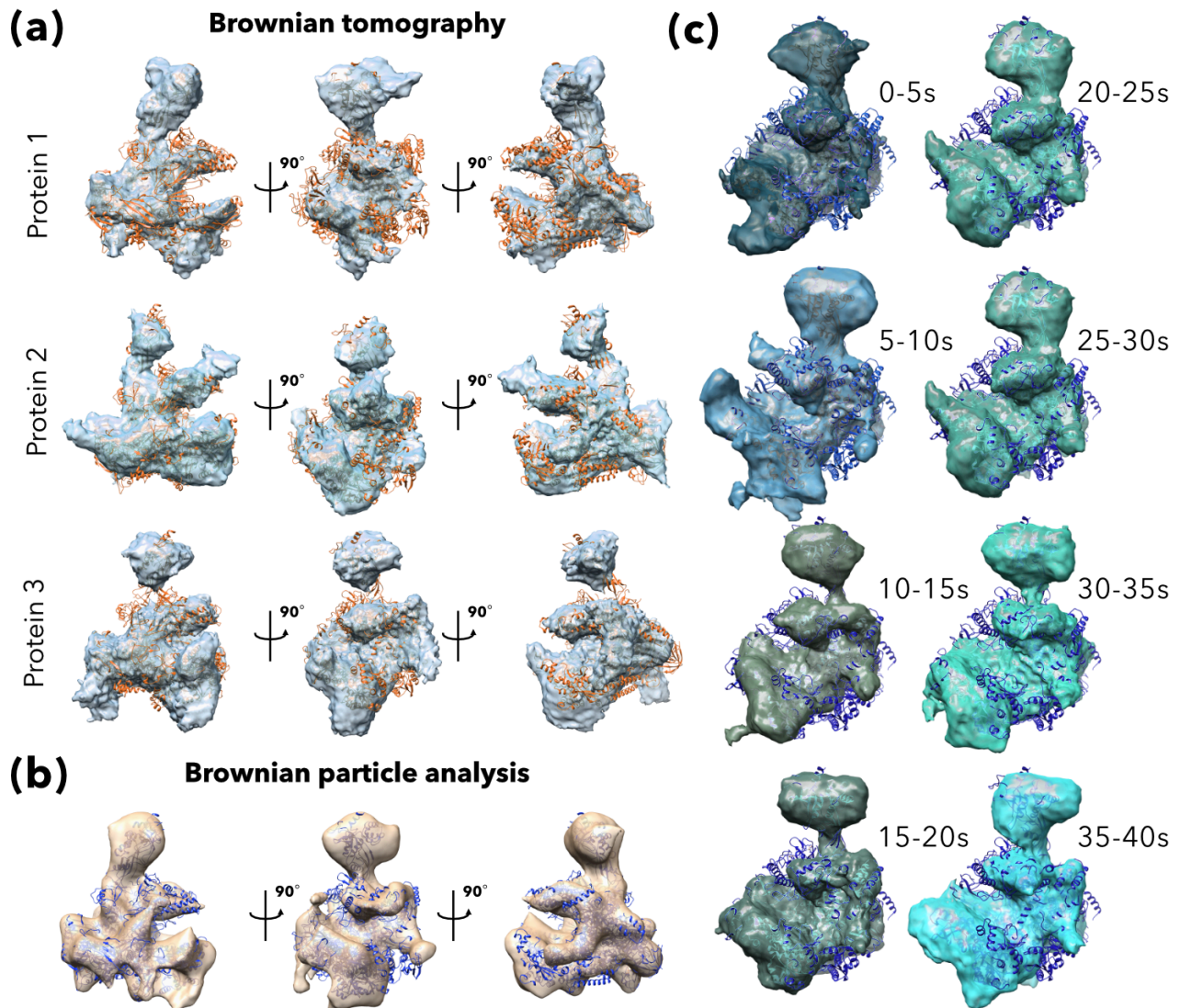
Apoferritin structure showing sixty over-imposed conformations of a single unit coloured as a function of the root-mean-square deviation of atomic positions RMSD (a). C4 and C3 orientation of sixty superimposed conformations of apoferritin where  $\alpha$ -helixes and coils are coloured in orange and white respectively (b). Two different orientations of sixty superimposed conformations of archaeal RNAP coloured as a function of the RMSD (c). Using the molecular dynamics datasets obtained for the two different proteins, we can create Apoferritin (d) and RNAP (e) fourth-dimensional dynamical maps shown at two orientations (top and bottom) and across three orders of magnitude of atomic residency time.



**Figure 3. Apoferritin structure.** Apoferritin brownian tomography (BT) electron density (ED) map generated from LPEM displayed at both C4 and C3 channel orientations as stand alone density (left hand side), as well as overlapped with the apoferritin crystallographic structure both clipped (centre) or as equatorial cross section (right hand side); detailed view of the C3 channel at the far right side (a). Apoferritin brownian particle analysis (BPA) ED map generated from LPEM (light blue) displayed alongside one hundred overlapping configurations from molecular dynamics simulations ( $\alpha$ -helices and coils coloured in orange and white respectively) The BPA ED map is overlapped with MD simulations both clipped or as equatorial cross section at the right and far right side respectively (b). Temporal evolution of the Apoferritin density map reconstructed by BPA. Profiles of 32 particles were extracted from each frame and were split into 10-frame sets of 1s long duration, these were then used for separate refinements, generating five different density maps shown here (c).

protein conformations within the time scale used for imaging.

Multiple 3D density maps in distinct conformational or compositional states can sometimes be obtained [20, 21] however these states maps required large number of profiles and adequate classifications. Herein, we leverage long molecular dynamics (MD) simulations of both apoferritin and archaeal RNAP in solution (150mM NaCl) at room temperature. During long MD simulations the system under investigation explores a conformational ensemble accessible for the starting state at a given temperature. In **Figure 2a** we depicted the apoferritin structure showing one unit with sixty overlapped conformations and coloured as a function of the root-mean-square deviation from the reference atomic positions (RMSD) during MD. We also show the whole apoferritin protein with the sixty conformations superimposed and coloured



**Figure 4. BT and BPA reconstruction of archaeal RNAP.** BT electro density (ED) map generated from LPEM employing the dynamical map as refinement model showing three different RNAP proteins at different orientations. The published RNAP structure is overlapped to show correlation **(a)**. BPA ED map generated from 5600 profiles of 14 particles over a 40s (400 frame) time period, with published structure aligned **(b)**. Temporal evolution of the RNAP density map reconstructed by BPA. The set of 5600 profiles was broken up into 50 frame (5s) groups, each with 700 profiles, yielding eight individually refined models **(c)**.

according to its secondary structure, the  $\alpha$ -helices and coils in orange and white respectively

(**Figure 2b**). We performed a similar analysis for RNAP and we show two orientations of sixty superimposed RMSD coloured conformations in **Figure 2c**. To shed light on the effect of dynamics on the local structure, we extracted a fourth-dimensional residence time matrix from the all-atom trajectories. For each frame of the trajectory, the system is roto-translated into a reference template (initial molecular dynamics structure). The rotation matrix of the system snapshot onto the template is calculated via a least-square rigid motion transformation using single value decomposition. This algorithm finds the best-fitting rigid transformation that aligns two sets of corresponding points. Then an occupation fourth-dimensional matrix is calculated and exported as a volumetric map. The resulting maps for apoferritin and RNAP are displayed in **Figures 2d** and **2e** respectively for different atom residency times showing dynamical information at different time-scales. The MD simulations show a relative rigid structure for apoferritin with maximum RMSD of 1.5 Å. The RNAP structure is considerably more flexible with several units within the stalk and the DNA binding cleft reaching RMSD of 5.4 Å. The 'dynamical maps' (**Figures 2d-e**) calculated from molecular dynamics encodes thermal fluctuations and the accessible conformational micro-states of the structure at the given temperature. Importantly, the conformational dynamics of the RNAP stalk and clamp are in good agreement with structural changes that have been partially characterised by comparing different molecular structures determined by cryoEM or crystallography, or by using single-molecule FRET of double-labelled RNAP in solution [38][41]. The different levels of the map i.e. from left to right correspond to the dynamical information at different time-scales and associated structural features. In **Figure S4** we show the difference between a low-pass filtered map, we refer to as 'static map', generated from x-ray of the all-atom structure for the archaeal RNAP polymerase and its 'dynamical map'. It is interesting to note that the correlation between the two maps increases up to a certain spatial resolution, without being a perfect match, see **Table S1**.

In **Figure 3**, we show the reconstructed ED map from both BT and BPA methods for apoferritin using the 'dynamical map' as initial model for the refinement. As shown in **Figure 3a**, the resolution is considerably improved when employing the BT method, as opposed to BPA, to generate the maps as both the C3 and C4 channels become detectable in the generated BT ED map. The C3 appears as a hole in the ED map (**Figure 3a**) indicating that the local density is the same as bulk water while the C4 appears to be solid in accordance with the channel reported polarity. The C3 pore is notably not visible in the input dynamical map and therefore cannot be a result of model bias. Interestingly, the C3 pore feature is critical to define our spatial resolution; indeed, the biological features suggest an ED map with 9Å resolution.

As for the BPA ED maps, shown in **Figure 3b**, the results are rather different; here, the reconstructions are the result of averaging thousands of profiles across several proteins, precisely 2340 profiles of 32 ferritin proteins, and 5600 profiles of 14 RNAP particles. Thus, the reconstructions obtained from BPA represent the conformational space conserved across the different proteins. Interestingly the BPA ED map density matches with close correlation the secondary structure distribution of the proteins. In this fashion the densest areas in the BPA ED map corresponds to the areas where the conformation is mostly an  $\alpha$ -helix arrangement, while the lowest density corresponds to the areas with random coil arrangements, depicted in orange and white respectively in **Figure 3b**. BPA reconstructions can also be performed from a smaller data set and the video can be thus broken down into periods of 1s. Profiles of 32 particles were extracted from each frame generating 468 total profiles and were split into 10 frames (1s) sets. The profiles contained in each 1s set were then used for separate refinements, generating five different sequential density maps over 5 seconds, as shown in **Figure 3c**. While these maps do not show notable differences from the cumulative map we can grasp some fluctuations between the different secondary arrangements of the protein.

The symmetrical nature of apoferritin is advantageous for introducing our methodology, however the real impact of the dynamical analysis can be better illustrated when applying it to an asymmetrical protein such as as archaeal RNAP. In **Figure 4a**, we show the reconstructed BT ED maps for three different RNAP proteins followed during the video along frames, using the 'dynamical map' as initial model. All three particles yield different reconstructions, showing effectively different conformations of the RNAP stalk and clamp, the two most flexible sections. Furthermore, BPA, yields an average reconstruction (**Figure 4b**) that smooth the conformational information encoded in the experimental data, yet the structure is not different from the single BT ED maps suggesting that the defying RNAP features are conserved across long time scales. We broke down the BPA profiles into smaller cohorts each representing a 5s long period, and the relative ED maps are shown in **Figure 4c** as well as morphed together in **video S1**. **Figure 4c** shows the fluctuations of the main RNAP features with the stalk exploring a large volume and the DNA binding channel expanding and contracting. This structural evolution demonstrates that while the different parts of the protein fluctuate around the structure, our technique can capture their dynamics over 40s periods.

In conclusion, we report a completely new way to image proteins in liquid phase exploiting their natural brownian dynamics with the aim of accessing the proteins' conformational repertoire. We show that a single protein particle can be followed using the so called Brownian tomography (BT) method, where we can reconstruct independently every particle, losing 'spatial resolution' due to the fact that single reconstructions are made with less frames, hence, profiles. Nevertheless, the loss of spatial resolution when employing the BT method is compensated by the access to dynamical resolution. We also showed that by combining single particle analysis with brownian dynamics, the so-called Brownian Particle Analysis (BPA), we can access more profiles than when using BT; yet, these profiles are shared across several protein units. Hence, the resulting structure is the average conformational space of the investigated proteins across several micro-states. Most importantly, every particle will evolve independently, stochastically, following a Brownian dynamics as well as being free to explore different conformational arrangements. Imaging proteins in liquid state allows us to use Brownian dynamics to capture and assess the different sides of the protein in a fast manner and probe its conformational dynamics. Finally, we can observe that despite the fact that our reconstruction do not offer atom level information, we do image the entire particle and we have access to structural information using BT coupled with molecular dynamics. Both inter- and intra- particle dynamics are lost by the need to vitrify the water in cryo-EM or crystallise the proteins in x-ray crystallography losing important structural information associated with molecular dynamics. Moreover, it is important to notice that the best cryo-EM reconstructions are based on  $10^6$  independent particles across as a large number of proteins. Herein, we limit the protein number to several hundreds in BPA dictated by the number of particles in the field of view, and to one in BT with  $10^3$  and  $10^2$  number of profiles respectively. These number of profiles are three and four orders of magnitudes less than current cryo-EM approaches, yet they achieve spatial resolution of about six times worst than that achieved by cryo-EM. While we are years away from filling such a resolution gap, we believe that our LPEM BT and BPA approach can be very well combined with cryo-EM and simulations to access the 4th dimension in structural biology accessing important dynamic information so far out of reach.



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## Experimental methods

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**Protein sources and production.** RNAP polymerase was prepared from *Sulfolobus acidocaldarius* cells using the method reported by Blombach *et al.* [42]. Equine spleen apoferritin was purchased from Merck (previously Sigma-Aldrich) and diluted with phosphate buffer solution (PBS) at a concentration of 2mg/ml.

**Liquid holders methodology.** Two liquid holders were used for the experiments, the OCEAN and STREAM holders, both manufactured by DENSsolutions. The first generation Ocean holder was employed for the experiments on apoferritin. The last generation Stream holder was employed for the experiments on RNA polymerase. The use of the different holders does not influence the results of the BPA and BT techniques. The most important variable is always the liquid thickness, which will directly affect the electrons transmitted through the specimen, hence specimen contrast and ultimately the resolution of the acquired images and the magnification that can be used.

For the OCEAN holder preparation, the ferritin in solution was sandwiched in a liquid cell formed by two chips made of silicon nitride (SixNy), with 50nm thick rectangular observation windows at the centre measuring 20mmx200mm. Silicon nitride prevents the solution from leaking from the closed cell and allows the passage electrons through the sample as it is electron transparent. The liquid cell is comprised of a spacer-chip placed at the bottom and a spacer-less chip placed at the top. 200nm spacer chips were used for the reported investigation. The SiN chips were rinsed in HPLC-graded acetone followed by isopropanol for five minutes each, to eliminate their protective layer made of poly(methyl methacrylate) (PMMA). Then, the chips were plasma cleaned for thirteen min to increase their hydrophilicity. Soon after, the chip with the 200nm spacer was fitted at the bottom of the liquid cell and 1.5 $\mu$ L of ferritin solution was deposited onto the observation window of the chip. It was important to not allow the ferritin solution dry at this stage as the film thickness generated by dried ferritin solution cannot be easily controlled and impedes good imaging conditions. The chip without spacer was then positioned on top of the liquid layer closing the liquid cell. The observing windows at the top and bottom chips were placed in a perpendicular configuration. The liquid holder was sealed, and 300 $\mu$ L of the ferritin solution at 2mg/mL was flushed in at 20  $\mu$ L/min with a peristaltic pump. A volume of 300  $\mu$ L of solution ensured that the holder tubing system and cell were filled with

ferritin solution. Although the liquid thickness is dictated by the spacer used in the liquid cell, the observation windows of the chips tend to experience some bulging, mainly at their centre point. This bulging effect is due to the differential pressure encountered by the holder once it is inserted into the vacuum of the microscope. The bulging phenomenon of the windows consequently adds some extra thickness to the observation chamber. The bulging effect is at minimum however at the corners of the rectangular windows. Thus, ferritin imaging was performed at the areas close to the corners of the SixNy window.

For the STREAM holder preparation, the chips did not require cleaning with organic solvent as they were supplied without protective layer. The chips were plasma cleaned to increase their hydrophobicity and hence wettability of the protein solution. The observing windows of the top and bottom chips were aligned manually using an optical microscope and in a perpendicular configuration. As with the Ocean, the top and bottom chips were separated by a 200nm spacer. The liquid holder was sealed by slowly increasing the torque of the screws to a maximum of 1cN while checking the windows were aligned. Then, 1 mg/mL of RNAP solution was used to fill the tubing system. A volume of 300  $\mu$ L of RNAP solution was flushed in the sealed holder at 1 $\mu$ L/min with a liquid nitrogen pump.

In both liquid holders, we stopped the pumps after collecting the respective sample solution from the outlet tube (Fig.S1). In the case of the STREAM holder the liquid thickness was reduced by applying a negative pressure in the outlet tube, effectively extracting some specimen sandwiched between the observing windows. The negative pressure did also help to reduce the bulging of the observing windows. A reduced liquid thickness and window bulging allowed to acquire images of RNAP at 30Kx magnification with good contrast and SNR. Then, we waited five minutes in order to guarantee that convection effects from the flowing process were not affecting the Brownian movement of the samples in solution. The inlet and outlet tubes of the pumps were sealed to ensure a closed liquid circuit.

The liquid holders were then introduced into the microscope. Imaging was performed with the holder in static conditions i.e. the pumps were stopped during imaging. Fluid dynamics simulations were performed on the OCEAN liquid holder and are reported in the SI section addressing Fluid dynamics simulation.

**Imaging methodology.** The experiments were performed using a JEOL JEM-2200FS TEM microscope equipped with a field emission gun (FEG) at 200 kV, and an in-column Omega filter. The camera used was the direct detection device (DDD) in-situ K2-IS camera from Gatan. The ultra-high sensitivity of the K2 allows low-dose imaging modes limiting considerably electron dose damage and facilitating high spatial (3838 $\times$ 3710 pixels) and temporal (up to 1600 fps in in-situ mode) resolution. The microscope was used in transmission electron microscopy mode. In order to limit the beam dose on the specimen, images were collected at the minimum spot size (5) with a small condenser lens aperture (CLA 2). The K2 can be used in two recording modes, linear and counted. The former is fast, but it generates the output image by capturing the integrated signal from each pixel, in a similar way to a standard charge coupled device (CCD). Counted mode processes the image and assigns one count to the pixel or sub pixel of the landing site of that electron. For our investigations dose fractionation videos were recorded in counted mode. Further details on the quality of the videos are provided in the Video analysis section. The apoferritin video used was a single 5s dose fractionation video at 10fps resulting in a total of 50 frames. The electron flux of this video was circa 0.9 e-/A<sup>2</sup> per frame and a total cumulative electron dose of 4.5e-/A<sup>2</sup>. The magnification used for apoferritin was 20Kx. The RNAP video used was composed by 400 frames, generated from four 10s dose fractionation videos at 10fps under identical imaging conditions. The electron flux of these videos were

0.038e-/Å<sup>2</sup> per frame, giving a total cumulative dose of 15.2e-/Å<sup>2</sup>. The magnification used for RNAP was 30Kx. Every image was recorded in the format of 4-byte grayscale and required the full size of the detector.

**MD.** We simulate apoferritin starting from the biological assembly of a 1.43Å resolution x-ray crystal structure (PDB-id: 6MSX) containing Cd<sup>2+</sup>. Only the positions of the protein coordinated Cd<sup>2+</sup> atoms were retained and simulated as Fe<sup>2+</sup>. The protein complex presented no mutations or structural gaps; therefore, it was protonated at pH 7.4 and solvated (150mM NaCl) using a TIP3P[43] water model and Charmm36m[44] protein and ions force field. The final simulated system consisted of 336806 atoms, resulting in a truncated octahedron simulation cell of 16.14nm x 15.21nm x 13.17nm (5.37, -5.37, 7.6). All simulations were performed with Gromacs v2019.2 [45] with a 2fs integration step. We performed 2000 steps of minimization (steepest descent algorithm), followed by 5000 steps of isotropic NPT at 300K, using the velocity-rescaling thermostat[46] and Parrinello-Rahman barostat [47]. After 50000 steps of NVT the system was fully equilibrated and ready for production, which was performed in the NVT ensemble for 1 microsecond.

The simulation of archaeal RNA polymerase was started from the cryo-EM structure of the apo-RNA polymerase (manuscript submitted at the time of submission of the present work). PDBidX. Missing residues and loops were modelled using MODELLER[48]. The protein complex, including the structural Zn<sup>2+</sup>, Mg<sup>2+</sup>, was protonated at pH 7.4 and solvated (150mM NaCl) using a TIP3P[43] water model and Charmm36m[44]. The final simulated system consisted of 539642 atoms, resulting in a truncated octahedron simulation cell of 18.94nm x 17.86nm x 15.47nm (6.31, -6.31, 8.93). The same equilibration protocol mentioned above was followed and 1 microsecond of NVT production was generated.

**Dynamical map calculation.** We calculated a fourth-dimensional residence time matrix for both systems, apoferritin and RNAP, from their all-atom MD trajectories. For each frame of the trajectory, the system is roto-translated into a template (initial molecular dynamics structure). The rotation matrix of the system snapshot onto the template is calculated using a least-squares rigid motion transformation using single value decomposition. Then an occupation fourth-dimensional matrix is calculated and exported as a volumetric map.

**Reconstruction methodology.** The individual frames of the dose-fractionation videos were exported and processed individually. The apoferritin frames were first denoised using a noise2void neural network [49], with a network trained on sets of 200 similar apoferritin video frames, this procedure did not prove beneficial for RNAP. Reconstructions were performed using Relion 2.0 methods [50, 51] within the Scipion framework [52]. For Brownian tomography, the particles were tracked and picked manually across the consecutive frames to render 400 profiles per particle for RNAP, and 50 profiles per particle for apoferritin. For Brownian particle analysis in RNAP all 400 profiles of 14 particles were combined to render a total of 5600 effective profiles. These profiles were split into groups of 700 particles for each of the 5s time-resolved reconstructions. For apoferritin, many particles within each frame were manually picked, rendering a total of 2340 particles. The time-resolved structures for apoferritin used profiles of 32 particles across the 10 frame (1s) intervals, making 320 profiles per reconstruction. The particle sets were then used with "relion\_refine" to refine models of the respective protein, with initial models originating either from atomic models low-pass filtered to 60Å or from dynamic models representing the residence time of atoms from a 1µs long simulation of the proteins. Octahedral symmetry was applied for ferritin refinement.

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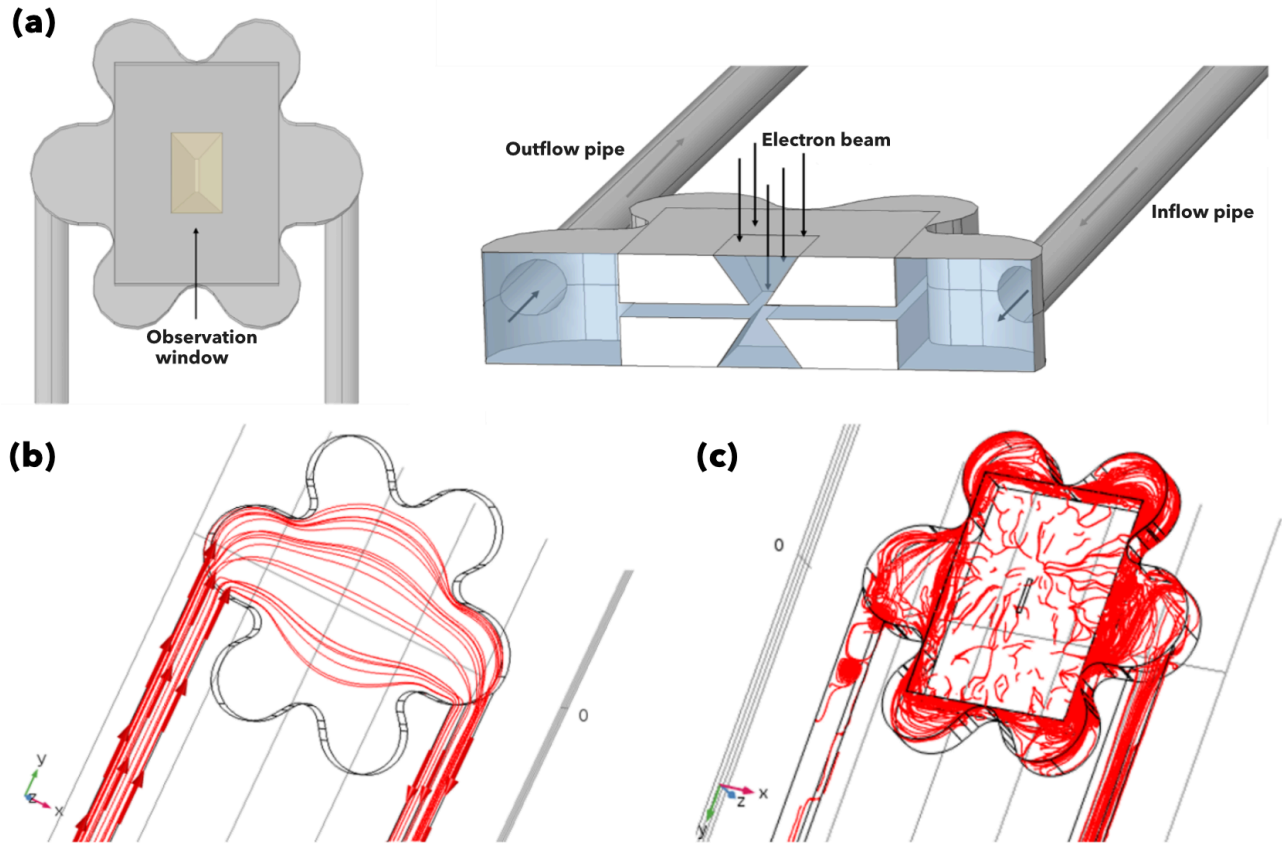
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## Supporting information

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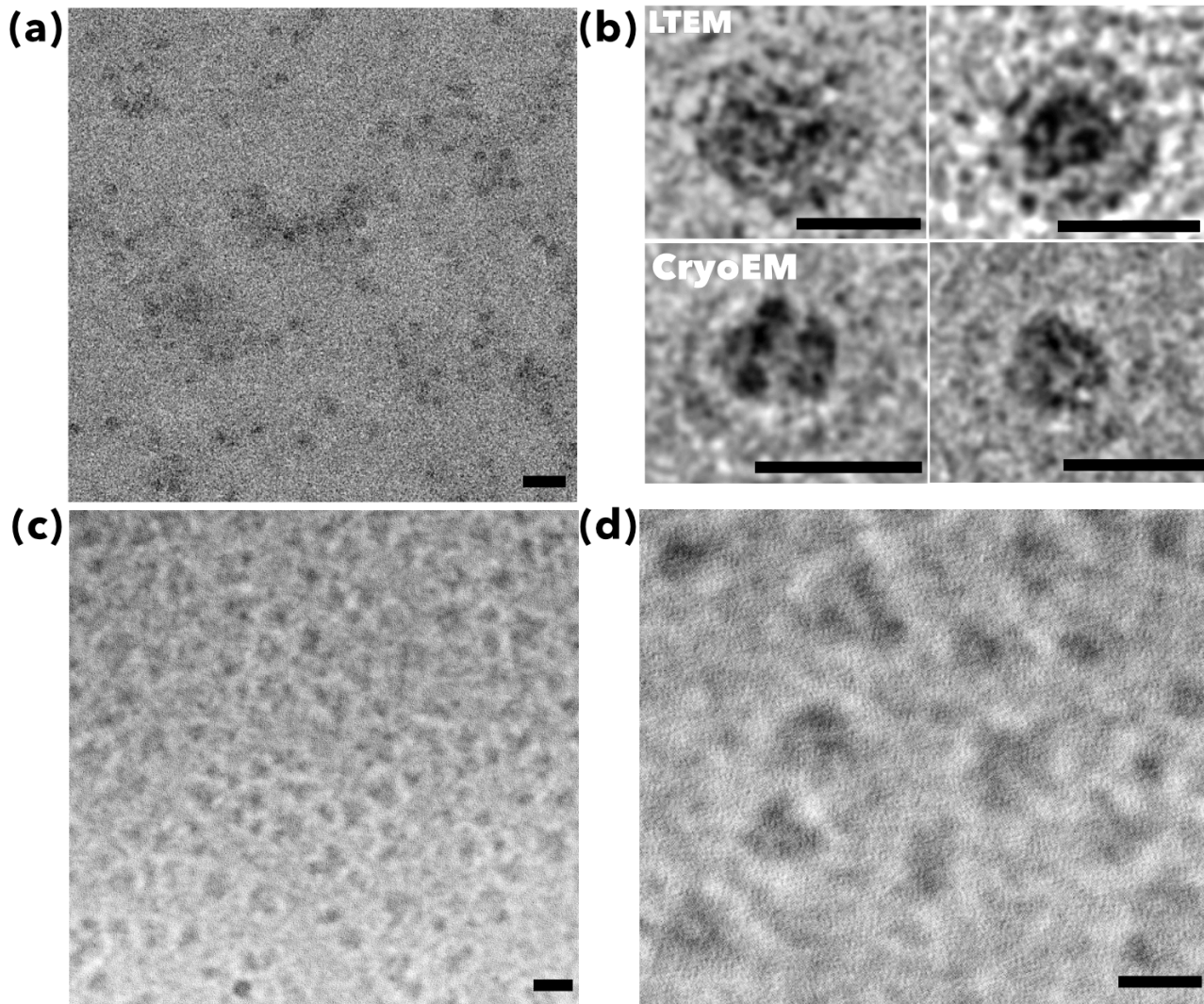
**Fluid dynamics simulation.** Section Liquid cell preparation and assembly shows how demanding this technique can be, and it is not surprising that the first attempts to image samples

in liquid state failed due to the high complexity of the system. Therefore, a fluid dynamics simulation was performed by using COMSOL Multiphysics in order to understand the functioning of the liquid cell in “in-flow” conditions and obtain the best performance from the imaging process. The liquid holder was modelled in COMSOL using the actual dimensions of the device, the values of which will not be reported in this paper so as not to violate the confidentiality agreement signed between the authors and the manufacturing company. However, the simplified geometry of the holder is shown in Fig. S1a. The tip of the holder is flower-shaped and is connected to two in-plane parallel pipes, an inlet and outlet pipe for flowing in and out of the solution respectively. The holder hosts on the inside two SiN chips separated by a variable gap, creating a microfluidic channel to enable inflow analysis and keep the sample hydrated. The bulging effect on the observation windows occurring when imaging the sample and mentioned in section Liquid cell preparation and assembly of SI has been excluded from the simulation. This is because the simulation has been performed at atmospheric pressure conditions, i.e. when the sample is flowed in the liquid holder outside the microscope. Thus, the windows are treated as flat, rigid surfaces trapping the solution on the inside. Input and output ports were set to the end of the pipes, guaranteeing good fluid circulation. At the beginning of the simulation the chamber was modelled as empty, both chips were removed, and an incompressible Newtonian fluid was pumped in with an initial velocity of 5  $\mu\text{L}/\text{min}$ . The analysis was performed at steady state, imposing laminar flow, planar symmetry and no-slip condition on the walls of the holder. The velocity of the fluid in these conditions was analyzed and the results are reported in Fig. S1b. This first preliminary simulation was mainly performed to validate the goodness of the model, and especially to analyze the trajectory of the flow. The first simulation confirmed that the fluid linearly went from the inlet to the outlet pipes, slowly expanding in the whole volume of the chamber. The results showed that the trajectory of the injected particles were a combination of both convection and Brownian motion experienced by nanoparticles in liquid. Then the complexity of the system was increased by adding the two chips inside the flower-shaped tip, separated by a 200nm spacer. The inclusion of the chips significantly influenced the liquid flow inside the chamber, modifying the liquid trajectory and narrowing it down to the small microfluidic channel between the chips. Consequently, most of the fluid went into the petals of the structure, generating a turbulent flow. Furthermore, the volume between the two chips was filled with only a small portion of the incoming fluid, the motion of which was hardly moved by the effect of the pump (Fig. S1c). In a first approximation model, these results can be extended to the particles in solution pumped in the inflow port; mirroring the actual experimental conditions. As a consequence, most of the particles would not flow between the observation window, but instead be trapped in the flower petal reservoir. These findings strongly suggest that parameters such as sample concentration and velocity at which the sample is injected into the holder have to be carefully chosen, in order to maximize the number of the particles in the viewing window area.

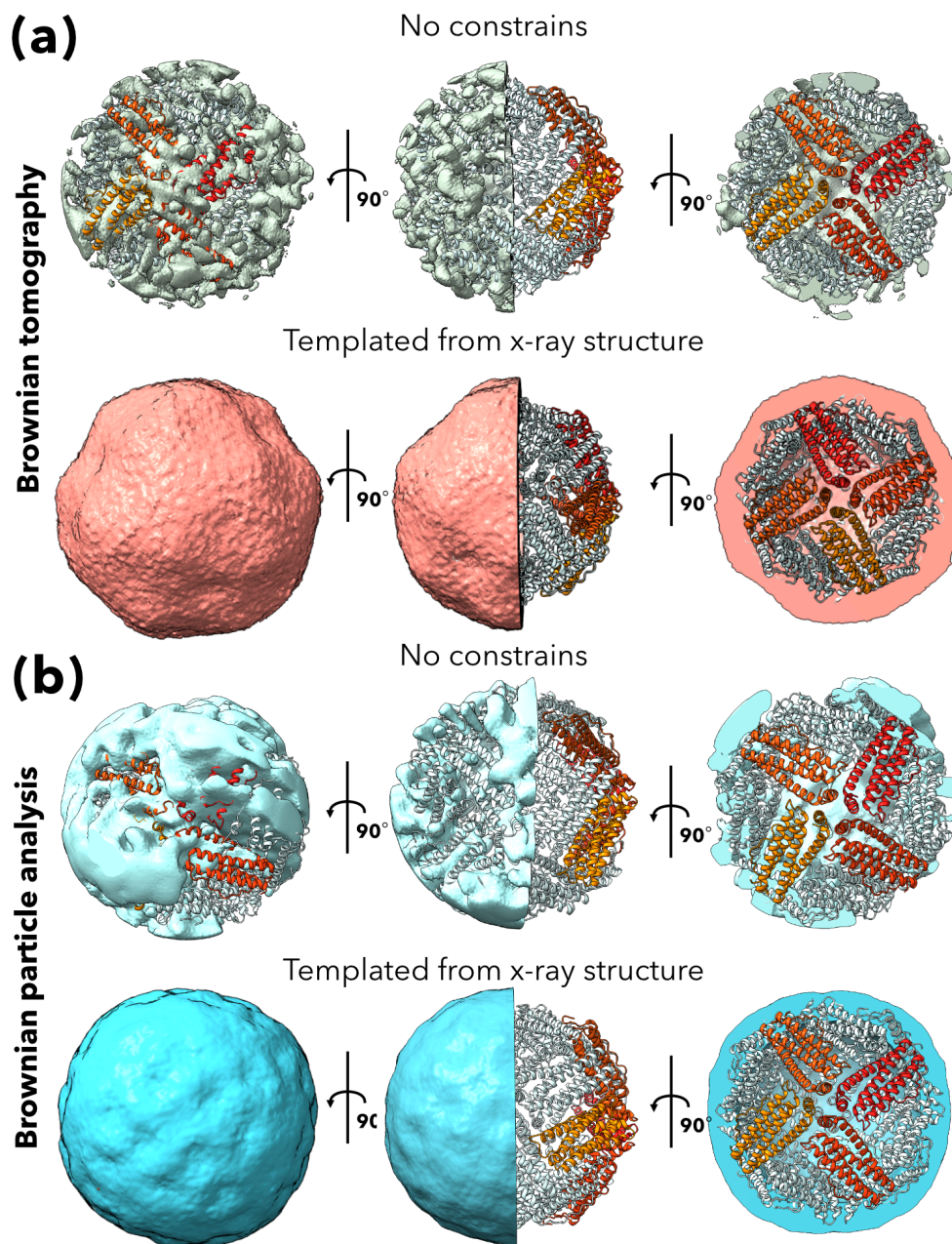


**Figure S1. Liquid cell holder.** Schematics of the liquid holder used for imaging liquid sample by transmission electron microscopy (a). Fluid dynamics simulations of the holder, modelled without the SiN chips (b). A Newtonian fluid was pumped in the inflow pipe (on the left) and out from the outflow pipe (on the right). The flow expands towards the flower shape reservoir perpendicular to the liquid main trajectory while flowing through the chamber. Fluid dynamics simulations of the holder comprising two chips inside (c). Both simulations run applying laminar flow, planar symmetry and no slip conditions; the analysis was then performed at steady state.



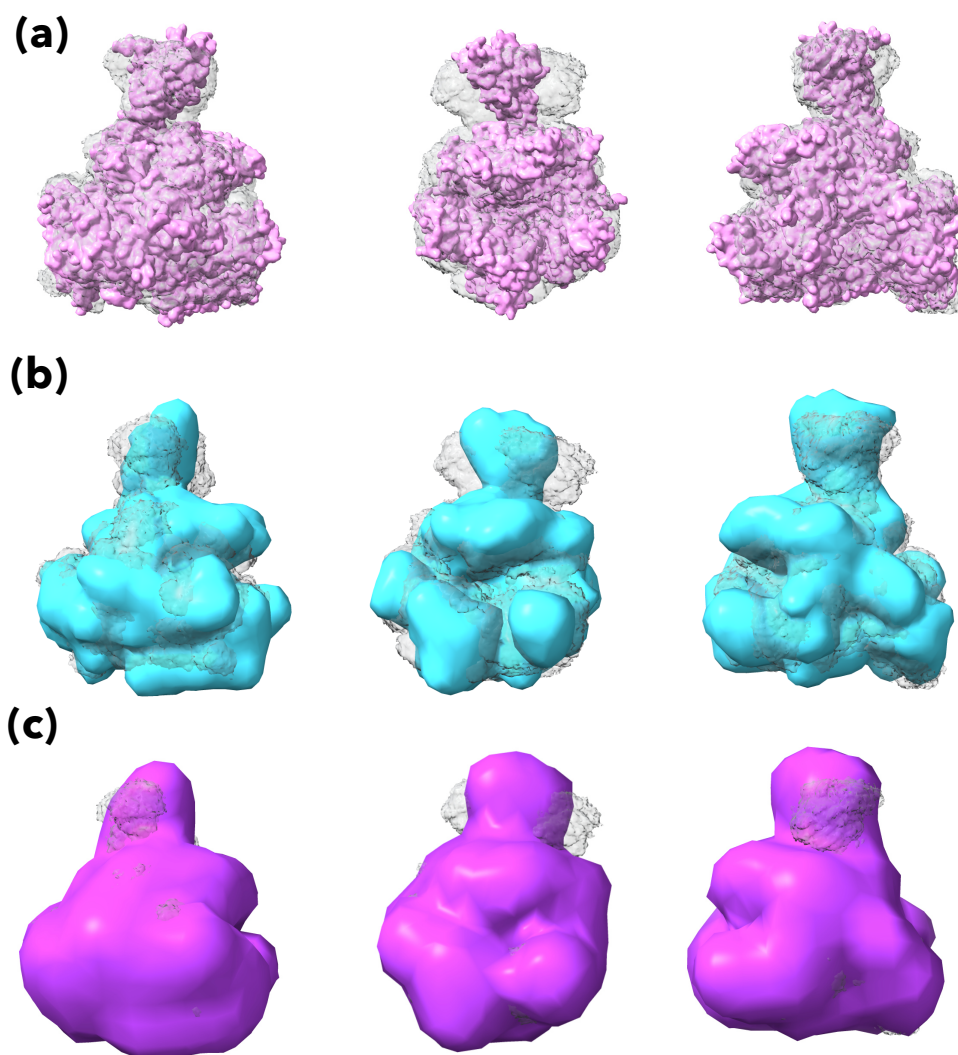


**Figure S2. Liquid phase transmission electron microscopy (LPEM) imaging of proteins.** LPEM micrographs of apoferritin particles collected at 20 frame per second (fps) and electron cumulative dose  $<10 \text{ e}/\text{\AA}^2$  (a). High magnification transmission electron micrographs (b) of apoferritin imaged in liquid and vitrified water. LPEM micrographs of archeal RNA polymerase particles collected at 10 frame per second (fps) in liquid water shown at low (c) and high (d) magnification with cumulative dose  $<10 \text{ e}/\text{\AA}^2$ . Scale bars: 20nm



**Figure S3. Brownian tomography (BT) versus Brownian particle analysis (BPA).**

Comparison between electron density (ED) maps for apoferritin generated by means of both a) BT and b) BPA using either unbiased refinement or a low-pass filtered ED map calculated from the deposited x-ray apoferritin PDB structure, shown at different orientation.



**Figure S4. Static versus Dynamical maps.** Three different low-pass-filtered maps of the all-atom structure of the Archeal RNA polymerase generated from Cryo-EM are depicted as surface, overlaid onto the 'dynamical map', and shown at different orientations (left, center and right hand side). Each map corresponds to the following resolutions: a) 5Å, b) 20Å and c) 40Å. show three different orientations

Resolution (Å)	Correlation
5	0.7479
20	0.8662
40	0.8523
60	0.8122
80	0.7714

**Table S1.** Correlation between 'dynamical map' and 'static map' at different resolutions