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Heritable yeast prions have a highly organized three-dimensional architecture with interfiber structures

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Yeast prions constitute a “protein-only” mechanism of inheritance that is widely deployed by wild yeast to create diverse phenotypes. One of the best-characterized prions, [PSI⁺], is governed by a conformational change in the prion domain of Sup35, a translation-termination factor. When this domain switches from its normal soluble form to an insoluble amyloid, the ensuing change in protein synthesis creates new traits. Two factors make these traits heritable: (i) the amyloid conformation is self-templating; and (ii) the protein-remodeling factor heat-shock protein (Hsp)104 (acting together with Hsp70 chaperones) partitions the template to daughter cells with high fidelity. Prions formed by several other yeast proteins create their own phenotypes but share the same mechanistic basis of inheritance. Except for the amyloid fibril itself, the cellular architecture underlying these protein-based elements of inheritance is unknown. To study the 3D arrangement of prion assemblies in their cellular context, we examined yeast [PSI⁺] prions in the native, hydrated state in situ, taking advantage of recently developed methods for cryosectioning of vitrified cells. Cryo-electron tomography of the vitrified sections revealed the prion assemblies as aligned bundles of regularly spaced fibrils in the cytoplasm with no bounding structures. Although the fibers were widely spaced, other cellular complexes, such as ribosomes, were excluded from the fibril arrays. Subtomogram image averaging, made possible by the organized nature of the assemblies, uncovered the presence of an additional array of densities between the fibers. We suggest these structures constitute a self-organizing mechanism that coordinates fiber deposition and the regulation of prion inheritance.

P Prions are self-replicating protein conformations found in yeast, fungi, and mammals. A nearly universal feature of prions is a conformational switch that converts a normally soluble, monomeric protein into a fibrous assembly with a cross- β -amyloid structure. Prions have the additional property of “infectivity,” with the prion amyloid conformation catalyzing conformational conversion of the normal soluble form of the protein.

In mammals, conversion of a prion protein, PrP, to its infectious PrP^{Sc} conformation is the basis of transmissible spongiform encephalopathies, in which spreading of PrP^{Sc} kills neurons by mechanisms that still remain elusive (1). Heightening interest in this process, proteins involved in several other neurodegenerative diseases have recently been shown to propagate in a prion-like manner, and this activity is likely to play an important role in determining the pathogenic properties of these proteins (2).

Remarkably, prions can play important roles in normal biological processes (3, 4). In metazoans, prion-like conformational switches function in processes as diverse as the maintenance of neuronal synapses (5, 6) and antiviral immunity (7). In fungi, prions provide a paradigm-shifting mechanism for the inheritance of biological phenotypes: the prion conformational change produces

a change in function that is inherited through the transmission of a self-perpetuating protein conformation rather than through transmission of altered nucleic acids (3, 4, 8).

Nearly a dozen yeast proteins are now known to switch into prion states. Most of them are hubs of cell regulation, and their prion states have a major impact on cell phenotype (3, 4, 9–11). Many more potential prion proteins have been discovered recently (12). Several of these, as well as the factors that regulate them, have been conserved for hundreds of million years (5, 13–15). Moreover, it has been discovered that yeast prions are widespread in nature, where they confer many phenotypes that have clear adaptive value (8). One of the most remarkable features of prion biology is their ability to link biological stresses to heritable changes in phenotype. This is a natural consequence of the effect of such stresses on protein homeostasis (3, 16–18). We are still, however, at an early stage in understanding the mechanisms by which these protein-based elements of inheritance are propagated. Here, we apply recently developed imaging methods to one of the best-characterized yeast prion proteins, Sup35.

Sup35 forms the prion known as [PSI⁺]. (The brackets denote its nonchromosomal, cytoplasmic inheritance, and the capital letters denote its dominant behavior in genetic crosses.) The protein has a C-terminal GTPase domain that normally functions as a translation termination factor. Two other domains govern the ability of Sup35 to switch between a stable functional state and a stable nonfunctional prion state. The middle region, M, is highly charged and helps maintain the soluble nonprion state. The N-terminal region, N, that is strongly enriched in asparagine and glutamine, is amyloidogenic, and promotes prion formation (5, 11, 19, 20). Importantly, the conformational transitions of

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the rings contained long fibers, and the dots were formed by a mosaic of short fibril bundles. Both ring and dot fibrils were packed in aligned bundles, with a center-to-center spacing of ~ 34 nm. In this stained material, the diameter of the fibril itself was ~ 25 nm.

To trace the fibril paths, we used serial tomography covering most or all of several dot structures and most of one ring. Strikingly, in all images, we found no discontinuities in hundreds of ring fibers examined, except for those located at the artificial boundaries between successive cut sections. That is, the fibers of ring cells were continuous. Typical views of fibril packing are shown in a cell section with NM-GFP rings (Fig. 1*A*), and a rendered model of the cell is shown in Fig. 1*B*. A section and rendered model of a cell containing an NM-GFP dot are shown in Fig. 1*C* and *D*. A movie of the dot fiber model, constructed from seven serial sections of one cell is shown in [Movie S1](#).

The width of fiber bundles in cells with NM-GFP rings was ~ 200 nm. Dot bundles ranged up to ~ 700 nm in width. The individual fibers in rings were very long, averaging several microns in length. Dot fibrils were much shorter, with many fiber ends. The fibrils of ring and dot cells have the same biochemical properties (35). Indeed, lysates made from such cells have equivalent capacities to seed the polymerization of purified NM, and such preparations, in turn, have equivalent capacities to transform [*psi*⁻] cells to [*PSI*⁺] by protein-only transformation. Biologically, ring and dot cells are distinguished only by the fact that ring cells cannot transfer the prion state to [*psi*⁻] mating partners and are, therefore, not “infectious.” Thus, tomography provides independent support for the hypothesis that fiber ends are the source of infectious propagons in vivo (41).

Notably, the lengths of fibrils in dot structures fit to a normal distribution, with a peak at 110 nm (Fig. S1*A* and *B*). Thus, it seems that the infectious dot arrays are not simply formed by random cleavage of fibers in the ring bundles. Rather, to account for their relatively uniform length, there must be a concerted mechanism for growth, lateral interaction, and fiber cleavage in these cells.

Fibril Arrays in Ring and Dot Assemblies Are Attributable to the Prion Domain. GFP retains its fluorescence in ring and dot assemblies (29, 33, 35, 39, 41). It must, therefore, be natively folded when incorporated in these fibrils. Because our interest was in discerning architectural elements characteristic of prion-based mechanisms of inheritance, it was important to ensure that the spatial organization and structural features of these assemblies were dictated by the prion domain and not by GFP. We prepared cells expressing NM from the same overexpression plasmid with an HA, myc or FLAG tag, but no GFP, downstream of NM ([SI Materials and Methods](#)). Freeze-substituted sections of the HA-tagged fibrils showed the same interfibril spacing (~ 31 nm; Fig.

S24). Thus, the observed fibril packing is dictated by the NM prion domain and any associated cellular factors, not by GFP.

NM-GFP Fibrils Do Not Maintain Associations with the Preautophagosomal Structure. Previously, the NM-GFP plasmid used in our studies was used to envision early events in prion induction, which are too rare to capture in the absence of overexpression. Conversion events were associated with an ancient system for the asymmetric inheritance of damaged proteins at the rim of the vacuole, at a site adjacent to the preautophagosomal structure (PAS) (35) known as the Insoluble Protein Deposit (IPOD) (46). This site is also associated with accumulations of Rnq1 (35, 46), a factor that promotes prion induction (42, 47). We, therefore, examined established NM fibril bundles for associations with the vacuolar rim. In some fibril dot structures, a minimal contact site was seen, at which one or a few fibrils at the edge of the dot were directly adjacent to the vacuole surface, excluding the otherwise ubiquitous ribosomes from the putative contact region. In serial reconstructions of other dots, no contact with the vacuole was seen. No direct contact with the vacuole was observed in the ring cell shown in Fig. 1*A* and *B*. Therefore, we conclude that direct contacts with the vacuolar rim are not essential for maintenance of the dots or rings, although they are observed in some cases.

Cryo-Electron Tomography to Characterize Native, Hydrated Fibril Assemblies in Situ. To examine the fibril assembly in the absence of potential artifacts caused by staining and dehydration, we used cryo-electron tomography of vitrified sections. In this approach, the unfixed, high-pressure frozen cells are sectioned in the vitreous state and viewed by cryomicroscopy with liquid nitrogen cooling and without thawing, fixation, or staining (48). Therefore, the material remains in the native hydrated state. Under these conditions, fibrils appeared thinner than in the stained, freeze-substituted sections (10 nm), and their center-to-center distance was more variable in the cryosections (25–50 nm) (Fig. 2). This may be attributable to mechanical section compression, but we did not observe a direct relationship between the sectioning direction and the interfiber spacing. An alternative explanation for the observed variable spacing is that the apparent separation distance depends on the oblique orientations of the fibrils in these 50- to 80-nm-thick sections. The core fibril diameter was consistent with EM studies of full-length, NM- and N-domain fibrils in vitro, which showed that the middle and C-terminal domains are flexible and can extend considerably from the fibril backbone (49). By analogy, it is likely that the globular GFP domains are also mobile in our fibrils. As noted above, a similar fibril spacing was observed with constructs lacking GFP, demonstrating that these properties were attributable to

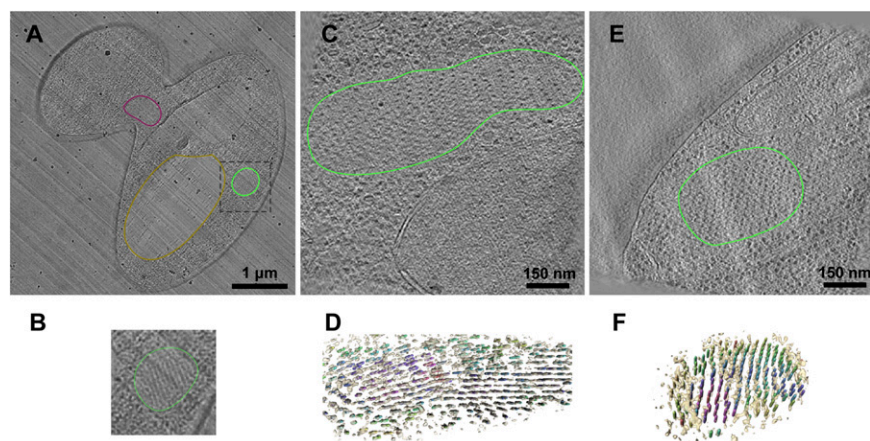


Fig. 2. Cryosections of dot and ring cells. (A) Overview section of a budding yeast cell with a dot aggregate, outlined as in Fig. 1. The cell is compressed along the direction of sectioning, indicated by the knife marks (diagonal streaks). (B) Enlargement of the dot region. (C) Tomogram slice of a ring, adjacent to the nucleus. (D) Rendered views of the fibers in the ring, colored by their cross-correlation values, with the most ordered regions in purple and least ordered in yellow. (E) Tomogram slice of a dot cell. (F) Rendered view of the fibril array in the dot, colored as in D.

fundamental organizing principles of the prion domain and its association with other cellular factors (Fig. S2A).

Subtomogram Averaging Reveals Additional Structures Intercalated Between the Fibrils. Two striking features of the assemblies revealed by electron tomography are: (i) the exclusion of other cytoplasmic contents, such as the normally ubiquitous and abundant ribosomes, despite the wide separation between fibrils; and (ii) the alignment of fibrils in arrays (Fig. 1). These observations suggest the presence of additional cellular components between the fibril cores, factors that might act to exclude ribosomes and to align the fibrils into small arrays. To investigate the 3D structural organization of these assemblies in more detail, we used subtomogram averaging, a single-particle approach in which small 3D subregions of the fibril lattice are extracted from a tomogram, aligned, classified, and averaged in 3D, to improve the signal to noise ratio and reveal more structural detail.

Overviews of a ring and dot lattice analyzed in this manner are shown in Fig. 3A and B, respectively. The fibrils are arranged in a quasicrystalline lattice, with several shells of neighboring fibrils revealed by the subtomogram averaging, as seen in the resin-

embedded sections. Remarkably, the averaging clearly revealed additional structures between the fibrils. Elongated densities surrounded and extended from the fibrils (Fig. 3C–F). The additional features were each ~8- to 9-nm in length. The most detailed average was obtained from the ring assemblies, in which the fibrils were aligned more nearly parallel to the section plane. However, similar structures were also seen, at lower resolution, after image averaging of dot assemblies (Fig. 3G–J). Although the interfibrillar structures do not appear to be strongly connected to the fibers, the level of defocus needed to give sufficient contrast in cryosections causes a halo of lower density around high-density features such as the fibrils. Therefore, a direct contact between the individual fibers and the newly identified interfibrillar structures is depicted in the illustration in Fig. 3K. The presence of six to seven discrete densities surrounding the fiber in the subtomogram average indicates that they are ordered, repeating structures in the fiber array (Fig. 3A and B). The reliability of the averaged maps is supported by the appearance of two to three shells of neighboring fibers, because these surrounding structures were not present in the reference image that was generated from the central fiber and used for alignment. These additional structures could explain the wide fiber spacing and exclusion of cytoplasmic components in between the fibers. It is likely that interfiber contacts would underlie the observed, ordered arrangement of fibers into arrays.

Molecular Chaperones Are Abundant Components of the Fiber Assemblies and Can Influence Their Structural Organization. As an initial approach to identifying the interfibrillar structures, we probed factors known previously to interact with prions and influence their inheritance. The model derived from our subtomogram averaging predicts that molecules forming these interfibrillar structures should be present at a similar level as NM-YFP. In a quantitative study, Bagriantsev et al. (34) found the Hsp70 proteins Ssa1 and Ssa2 in a ~1:1 ratio, with Sup35 in prion aggregates from $[PSI^+]$ cell lysates, along with other less abundant proteins, such as additional molecular chaperones implicated in prion biology (26, 31, 32, 44, 50–53), for example Sis1 and Hsp104, and a protein known to regulate actin dynamics during endocytosis, Sla2 (33, 36).

Therefore, we asked whether molecular chaperones and protein remodeling factors are components of the assembly. We tagged the genomic copies of Ssa1 and Ssa2 and the Hsp40/J protein Sis1, as well as Hsp104 and Hsp110 (a member of the Hsp70 family known as Sse), with CFP in cells that were expressing NM-YFP (*SI Materials and Methods*). The shift from NM-GFP to YFP was made simply to allow separation of its fluorescent signal from that of CFP. All of these proteins colocalized with both rings and dots (Fig. 4).

To further investigate the role of Hsp70 proteins and to determine whether genetic manipulations of yeast might provide tools for future analyses of prion cytoarchitecture, we used a $[PSI^+]$ strain deficient in Hsp70. Hsp70 proteins are redundant in all eukaryotes, and in yeast, four genes encode the family of Hsp70s in the cytosolic and nuclear compartments (known as Ssa proteins). We used a strain in which three of the four Ssa proteins were deleted (Ssa2–4 Δ) and the single remaining Ssa member, Ssa1, was replaced by a temperature-sensitive (ts) mutant (54). Because of the number of markers involved in the construction of this strain, the wild-type copy of Sup35 was retained, and we examined only newly assembled NM dot structures, induced with a galactose-inducible NM-YFP construct (18). Gold-labeled anti-YFP antibodies were used to identify prion dot assemblies in freeze-substituted EM sections with and without the additional ts mutation in Ssa1, both at 25 °C and after a shift to 37 °C. The organization of fiber arrays was disturbed in the strain with the Ssa1-ts mutation, compared with cells with wild-type SSA1. The fibril bundles were more disorganized, especially after the shift to 37 °C (Fig. S2B–D).

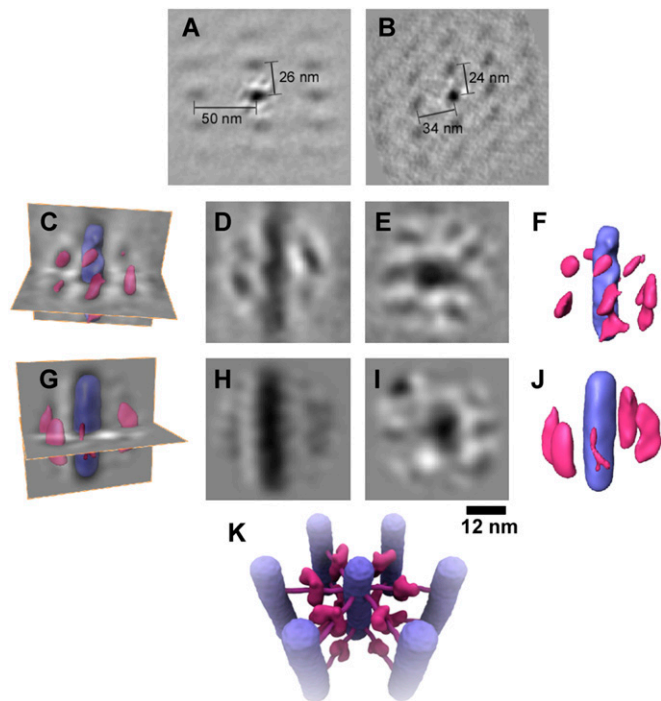


Fig. 3. Subtomogram averages of the fiber arrays. Subregions of the cryotomograms were aligned and averaged to reveal additional structures surrounding the fibers. (A) Section through the array of ring fibers, with the region in the center showing the averaged density surrounded by the array of fibers. (B) Equivalent view of the dot fiber array. The variable spacing might arise from sectioning artifacts and/or oblique fibril orientations. (C) Rendered views of the ring average superposed on density sections. (D and E) Long- and cross-sections of the averaged ring fiber. (F) Rendered view of the averaged fiber with the surrounding structures. The fiber is shown in purple and the additional structures in pink. (G) Rendered view of the dot average superposed on density sections as above. (H and I) Long- and cross-sections of the dot fiber average. (J) Rendered view of the averaged fiber with the surrounding structures, colored as in F. The differences in dot and ring structures most likely arise from the lower resolution obtained for the dot average. The defocus needed to obtain sufficient contrast in cryosections causes a white halo around high-density (dark) features; therefore, obscuring regions of contact between the central fiber and surrounding structures. (K) Illustration of the deduced arrangement of cross-bridges between the fibers. Fibers are in purple, and cross-bridge structures are in pink.

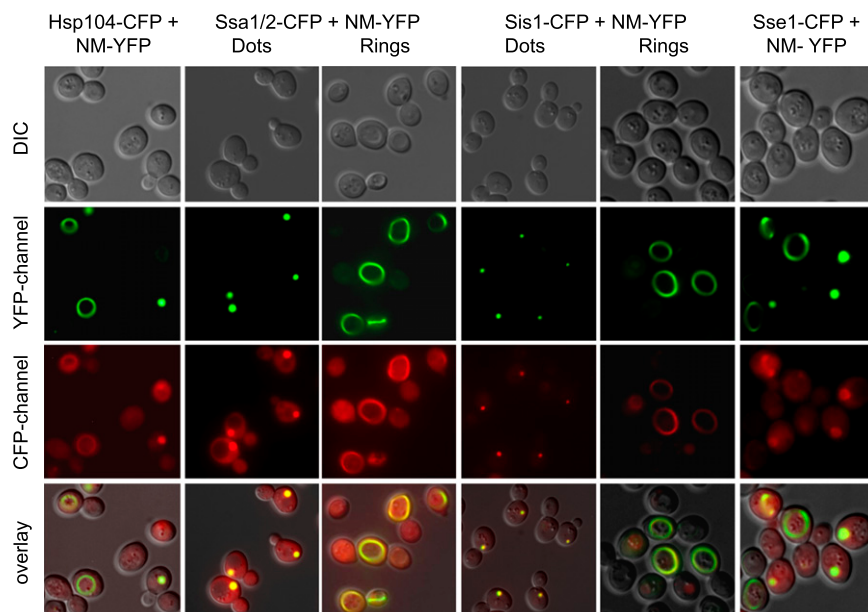


Fig. 4. Fluorescence microscopy showing colocalization of chaperones with NM-YFP in ring and dot structures. DIC, differential interference contrast images. The prion fibers are labeled with YFP (NM-YFP), and the chaperones are labeled with CFP. The results show that Hsp104, the Hsp70 proteins Ssa1/2 and Sse1, and the Hsp40 protein Sis1 are all found in dots and rings.

Discussion

The yeast prion phenomenon is of much broader importance in biology than was initially anticipated (3, 4, 8, 10, 12, 17). Despite recent advances in characterizing prions and the biochemistry of their propagation (4, 5, 11, 19, 20), little is known about the cellular organization of amyloid fibers in these protein-based elements of inheritance. A growing body of evidence suggests that prion amyloid assemblies, like other amyloids, are deposited at distinct protein quality control sites, particularly for abundant species (12, 35–37, 46). Such deposition could serve to sequester potentially damaging aggregates for retention in the mother cell, a common principle also observed for oxidatively damaged proteins during cell division, with implications for cellular aging (55). However, despite retention of prion aggregates at the IPOD, sufficient amounts of seeds are transmitted to daughter cells to faithfully propagate the prion state (35, 40, 41). Therefore, deposition and transmission of prion aggregates must be regulated, a process likely to involve specific organization of prion fibrils by cellular factors.

How can a cell achieve such a high degree of order for amyloid deposits as observed previously (35, 41) and in greater detail in this study? NM fibers have never been observed to form such arrays in vitro (22, 27, 49, 52). Therefore, we reasoned that other cellular factors must dictate the arrangement of the fibril arrays. The 10-nm diameter of the fibers observed in our cryo-electron tomography studies is consistent with that of fibers generated in vitro from NM or N alone (22, 49). Subtomogram averaging identified elongated structures intercalated at semiregular positions along and between the fibrils that could explain their wide, parallel spacing.

What could be forming these 8- to 9-nm densities? Various studies have shown molecular chaperones coprecipitating with $[PSI^+]$ prion aggregates (32, 34, 35). Indeed, we found colocalization of several molecular chaperones including Hsp104, Ssa1/2, and Sis1 with the prion assemblies by fluorescence microscopy (Fig. 4). Ssa1 appeared particularly abundant in the assemblies, consistent with previous copurification of Ssa1/2 with $[PSI^+]$ prion aggregates in a nearly 1:1 ratio (34). The presence of molecular chaperones is likely related to their roles in remodeling of prion aggregates and propagation of the prion state (31, 50, 56). Indeed, Hsp104 (26) and Sis1, a cochaperone of Ssa, play essential roles in severing of $[PSI^+]$ prion fibers during the prion-propagation cycle, likely in cooperation with Ssa (44, 51). Given the high abundance of Ssa1/2 in prion aggregates relative to other factors required for

fiber severing, e.g., Hsp104 (34), it may serve an additional function. A recent study of a ribosome-associated Hsp70-40 complex, Ssz-zuotin, suggests an end-to-end binding of these two components, which could generate an extended assembly (57). By analogy, elongated structures could be formed by Ssa1/2 and Sis1, and either serve a scaffolding function and/or regulate access of other factors, thereby influencing the processing of prion fibrils. Furthermore, overexpression of Ssa1/2 in $[PSI^+]$ cells expressing a construct with GFP incorporated into the endogenous copy of SUP35 between the N and M domains causes a shift from multiple small aggregates to fewer, larger ones (58). This phenotype may also hint at a role of Ssa1/2 in organization of prion aggregates.

Indeed, in an initial approach, deleting all functional copies of Ssa (54) from cells displaying dots caused some disruption to the prion fibril assemblies (Fig. S2 C and D). At this point, we cannot distinguish whether the effect observed in cells with reduced Ssa is directly or indirectly attributable to the loss of Ssa function. Thus, the present results raise the possibility that the Hsp70 system organizes fibril deposition in NM dots. However, it is not currently possible to directly identify the interfibril structures, which are packed too closely to be discriminated from the fibrils by immuno-EM.

Further analysis is needed to confirm this hypothesis and to establish how these ordered prion fiber assemblies are formed in yeast cells. The appearance of similar ring and dot structures for other prion-like proteins (12, 36, 47) raises the possibility that the amyloid fiber organization in highly ordered arrays is a common principle for units of protein-based inheritance.

Materials and Methods

Yeast Strains, Constructs, Media, and Fluorescence Microscopy. Yeast strains, constructs, media, and fluorescence microscopy were performed as described in refs. 18, 35, and 54 and in *SI Materials and Methods*.

EM Sample Preparation and Data Collection. Cells were high-pressure-frozen in an EMPACT 2 freezer and freeze-substituted into Lowicryl HM20 (Polysciences) following the protocol of Hawes et al. (59) with 0.2% or 2% (wt/vol) uranyl acetate, or with 0.2% glutaraldehyde, 0.1% uranyl acetate and 1% (vol/vol) water in acetone. The electron micrographs were recorded on a 2,000 × 2,000 pixel CCD camera with a pixel size at the specimen level between 0.6 and 1 nm. The total electron dose applied on the cryosamples was around 70 e/Å².

Image Processing. Alignment of the room temperature tomograms was done with etomo (60) using the gold particles as fiducial markers, whereas the cryotomograms were aligned by cross-correlation (61).

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