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In situ structure of trypanosomal ATP synthase dimer reveals a unique arrangement of catalytic subunits

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We used electron cryotomography and subtomogram averaging to determine the in situ structures of mitochondrial ATP synthase dimers from two organisms belonging to the phylum euglenozoa: Trypanosoma brucei, a lethal human parasite, and Euglena gracilis, a photosynthetic protist. At a resolution of 32.5 Å and 27.5 Å, respectively, the two structures clearly exhibit a noncanonical F1 head, in which the catalytic (αβ)3 assembly forms a triangular pyramid rather than the pseudo-sixfold ring arrangement typical of all other ATP synthases investigated so far. Fitting of known X-ray structures reveals that this unusual geometry results from a phylum-specific cleavage of the α subunit, in which the C-terminal αC fragments are displaced by ~20 Å and rotated by ~30° from their expected positions. In this location, the αC fragment is unable to form the conserved catalytic interface that was thought to be essential for ATP synthesis, and cannot convert γ-subunit rotation into the conformational changes implicit in rotary catalysis. The new arrangement of catalytic subunits suggests that the mechanism of ATP generation by rotary ATPases is less strictly conserved than has been generally assumed. The ATP synthases of these organisms present a unique model system for discerning the individual contributions of the α and β subunits to the fundamental process of ATP synthesis.

F1F0 (F-type) ATP synthases are ancient, energy-converting nanomachines that generate ATP by rotary catalysis (1). In mitochondria, the ATP synthases form rows of dimers along the highly curved ridges of the inner membrane cristae (2, 3). All known F1F0 ATP synthases consist of a catalytic hydrophilic (αβ)3 hexamer, a hydrophobic membrane-embedded region containing a rotor ring driven like a turbine by the proton gradient, and a pair of stalks connecting the two. The central stalk transmits the torque of the rotor ring to the (αβ)3 hexamer to power catalysis, whereas the peripheral stalk acts as a stator to prevent idle rotation of the catalytic subunits with the rotor assembly. The (αβ)3 subunits and the central stalk make up the membrane-extrinsic F1 subcomplex. The membrane-intrinsic subunits, including the rotor ring and the peripheral stalk, form the F0 subcomplex.

The F0 subcomplex has three catalytic sites, each located at an interface of the α and β subunits. The two subunits alternate in (αβ)3 hexamer, which forms a ring of near-sixfold symmetry (4). ATP synthesis is powered by the proton-driven rotation of the central stalk, which induces conformational changes in the α and β subunits (5, 6). In the course of these changes, a conserved α subunit arginine inserts into the nucleotide binding pocket of the β subunit (7). Substitution of the arginine residue inhibits ATP production, highlighting the essential role of the α subunit in catalysis (8, 9).

Trypanosomes belong to a group of parasitic protists, which cause severe and widespread insect-borne human and animal diseases. Biochemistry and mass spectrometry have indicated that the α subunit in the mitochondrial ATP synthase of euglenozoa (which include trypanosomes) exists as two separate fragments: an N-terminal 14-kDa fragment (αN) and a C-terminal 44-kDa fragment (αC) (10–14). Even though the α subunit is cleaved, the complex still hydrolyzes and synthesizes ATP, which is critical for the survival of Trypanosoma brucei, the sleeping sickness parasite (10, 11, 14–19). How this cleavage affects the structure and molecular mechanism of the enzyme is unknown.

Here we report the in situ structure of the mitochondrial ATP synthase from the “procyclic” insect stage of the lethal human parasite T. brucei and the related Euglena gracilis, a photosynthetic protist. The structures were determined by electron cryotomography and subtomogram averaging to a resolution of 32.5 Å and 27.5 Å, respectively. Both structures reveal an unexpected arrangement of the catalytic subunits, which form a triangular pyramid rather than a hexameric ring of near-sixfold symmetry, as is seen in all other known rotary ATPases. Rigid body fitting of X-ray structures showed that the unusual arrangement of catalytic subunits in the F1 head results from the phylum-specific cleavage of the α subunit. The subtomogram averages from both organisms indicate that the αC fragment is displaced from the central stalk by ~2 nm and forms a link between neighboring β subunits. Consequently, the typical catalytic interfaces of rotary ATPases cannot form, and therefore the mechanism of ATP synthesis must be different.

Significance

The mitochondrial F1F0 ATP synthase is an essential membrane protein machine that supplies all eukaryotic cells with ATP. The proton-driven rotation of the rotor assembly in the membrane transmits energy to the catalytic F1 head, where ATP is generated by rotary catalysis. We determined the in situ structures of ATP synthase dimers from the lethal human parasite Trypanosoma brucei and its free-living relative Euglena gracilis. In both ATP synthases, the catalytic subunits form a threefold pyramid rather than the usual near-sixfold ring. This unexpected finding indicates that the structure of the F1 head, and therefore its catalytic action, is less highly conserved than previously thought, and provides insight into the fundamental mechanism of ATP production in higher organisms.

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The authors declare no conflict of interest.

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Data deposition: The structures of the mitochondrial ATP synthase dimers from E. gracilis and T. brucei have been deposited in the Electron Microscope Data Bank, emdbdata bank. org (EMDB ID codes EMDB-3559 and EMDB-3560, respectively).

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Results
Structure of Euglenozoan ATPSynthase Dimers in Situ. Mitochondrial membranes were isolated from procaryotic (insect stage) T. brucei or light-grown E. gracilis cells and imaged by electron cryotomography. Tomographic volumes revealed that cristae were discs decorated with double rows of 10-nm particles. The structure of the particles obtained by subtomogram averaging confirmed that they were ATP synthase dimers. A total of 1,294 dimers from E. gracilis and 925 dimers from T. brucei were averaged to generate maps at a resolution of 27.5 Å and 32.5 Å, respectively (Fig. 1, Fig. S1, and Movie S1). The ATP synthase dimers of both species are twofold-symmetric V-shaped structures, with distances between the catalytic F1 heads of 24.5 nm in T. brucei and 22 nm in E. gracilis (Fig. 1A and D) and an angle between the long axes of the monomers of 65° in T. brucei and 50° in E. gracilis. The peripheral stalks are positioned on opposite sides of the dimer and are structurally unrelated to other peripheral stalk subcomplexes (Fig. 1B and E) (3, 20). On the luminal side of the membrane, the monomers are connected by several interlinked protein densities (Fig. 1C and F). In the complex from E. gracilis, two protein domains near the dimer axis tie the monomers together on the matrix side.

Cleavage of the α Subunit Results in an Unusual Arrangement of Catalytic Subunits. The most conspicuous feature of the two new ATP synthase dimer maps is the catalytic F1 head (Fig. 2 and Fig. S2). In all known rotary ATPases, the F1 head is a hexagonal prism of near-sixfold symmetry, consisting of alternating α and β subunits or their equivalents (4, 21–24). In the T. brucei and E. gracilis ATP synthases, the F1 head resembles a triangular pyramid (Fig. 2A and D and Fig. S2 A and E). A comparison with the bovine F1 reveals that the subtomogram average map accommodates only every other subunit of the αβ, hexamer (Fig. 2B and E and Fig. S2 B and C and S3). The densities of the three other intervening subunits are arc-shaped and look strikingly different from those in the bovine or yeast F1 heads.

Previous mass spectrometry and biochemical studies indicated that the α subunit of Euglenozoa ATP synthases exists as two separate fragments, αN and αC (10–13). To investigate whether the cleavage of the α subunit might account for the unusual structure of the F1 head, we fitted a cleaved model of the bovine F1 structure into the subtomogram average densities (Fig. 2 B and E and Fig. S2 C and G). The α subunit is cleaved by an unidentified protease in a sequence found only in Euglenozoa (residues 145–169 in T. brucei) (10, 13). This cleavage site maps to a loop that connects the N-terminal β barrel to the central domain of the bovine α subunit (Fig. 2 G and H). We severed the α subunit of the bovine (αβ), atomic model (4) at this location and positioned the resulting subcomplex (β + αN subunits) into the EM maps as a rigid body. In both subtomogram averages, the atomic model closely matched the density, with the N-terminal β barrels from both the αN fragment and the β subunit occupying a narrow ring of density below the oligomycin sensitivity-conferring protein (OSCP), known as the crown region. The remainder of the β subunits occupied the three densities adjacent to the central stalk (Fig. 2 B and E and Fig. S2 C and G).

The fit of the αN and β subunits did not occupy the volumes of the arc-shaped densities that form the vertices of the triangular pyramid (blue densities in Fig. 2 B and E and Fig. S2 C and G). These densities, which span the gap between neighboring β subunits, are 10 nm long and ~2 nm wide and enclose a volume of ~49 × 103 Å3 in E. gracilis and 58 × 103 Å3 in T. brucei. Based on an average partial specific volume of 1.2 Å3/Da (25), each arc-shaped density should accommodate a protein with a molecular mass of 41–48 kDa. This is very similar to the size of the αC fragment (44 kDa), which in T. brucei has been shown to remain bound to the F1 subcomplex, as demonstrated by αC-specific antisera (26) and mass spectrometry (10).

To assess whether the cleaved αC fragments could be located in the arc-shaped densities, we performed rigid body fitting (Fig. 2 C and F, Fig. S2 D and H, and Movie S2). The αC fragments fitted the profiles of the arc-shaped density well, indicating that they do indeed constitute the vertices of the triangular F1. Despite the good overall match, rigid body fitting permits several possible orientations of the αC fragment (Fig. S4). We consider
the αε fragment position that involves an outward displacement of ~20 Å and a rotation by 30° to be the most likely orientation (Fig. 2 C and F and Fig. S2 D and H), because it requires the minimal offset from the canonical structure. The accuracy of the fit is limited by the map resolution. Nevertheless, based on our difference map (Fig. S3 B and D), the rigid body fits (Fig. 2 C and F and Fig. S2 D and H), and the fact that the αε fragment remains attached to the F₁ subcomplex after cleavage (10, 26), we can unambiguously assign the αε fragments to the arc-shaped densities. Thus, our results show that in euglenozoa, cleavage of the mitochondrial α subunit results in an unusual arrangement of catalytic subunits in a triangular F₁ pyramid. We conclude that the structure of the catalytic F₁ head in euglenozoa is substantially different from that of any other known rotary ATPase.

To assess whether the in situ structures obtained by subtomogram averaging are of an active complex, we solubilized samples of the E. gracilis mitochondrial membranes used for electron cryotomography with n-Dodecyl β-D-maltoside (DDM) and performed blue native gel electrophoresis (BN-PAGE). Two bands reacted to an in-gel ATPase activity assay, one in the megadalton range, which is the ATP synthase dimer, and the other at ~700 kDa, which is the ATP synthase monomer (Fig. S5). These results indicate that the ATP synthases averaged in the tomograms of isolated mitochondrial membranes from E. gracilis are catalytically active.

**Dimer Rows in Euglena and Trypanosomes Indicate an Unexpected Arrangement of ATP Synthase Monomers.** To determine the macromolecular arrangement of ATP synthase dimers from E. gracilis and T. brucei in the membrane, the subtomogram averages were placed into the original tomographic volumes in the positions and orientations determined in the averaging process. In both species, the isolated mitochondrial membranes were decorated with multiple ATP synthase dimer rows that are arranged in short left-handed helix segments around the edges of discoid vesicles (Fig. 3 A–H and Movie S3). Each row contained between three and six dimers. The short dimer rows encompass the entire circumference of the disk-shaped membranes (Fig. 3 D and H).

In each row, the two F₁ monomers from one dimer interdigitate with monomers from the next dimers along the row. As a result, the nearest-neighbor monomers, both across and along the row, belong to different dimers (Fig. 3 I and L). This contrasts with the arrangement of ATP synthase dimers in metazoans and fungi, where the nearest monomers across the row belong to the same dimer (3). When viewed from the lumen, the dimers form a ladder-like assembly (Fig. 3 C, G, K, and N). Examination of each row in isolation shows that the principal curvature is caused by the association of dimers into rows, rather than by the dimers themselves (Fig. 3 J and M). The resulting membrane curvature is stronger in T. brucei than in E. gracilis (Fig. 3 J and M), most likely due to the different dimer angles in the two species (Fig. 1 A and D).

**Discussion**

**Structural Variation of Mitochondrial ATP Synthase Dimers and Dimer Rows.** The assembly of the mitochondrial ATP synthase into dimers, and of dimers into rows, is a feature common to mitochondria of all species examined to date (2, 3, 27–29). The principal role of dimer rows appears to be in the formation of mitochondrial inner membrane cristae. Tomographic analysis has revealed two different row architectures that bend the membrane in different ways. The first type of row architecture, which we call the metazoa type, is found in metazoans, plants, and fungi. In this group, the ATP synthases form V-shaped dimers with a dimer angle of ~90° between monomers (3). Dimers of the metazoa type assemble into straight rows along the edges of lamellar cristae, without any lateral offset between adjacent dimers (3). The V-shape of individual dimers induces a sharp, local 90°-membrane curvature that is responsible for driving the self-association of dimers into rows and shaping cristae membranes (30, 31).

In ciliates (29), euglenozoa, and green algae (20), the ATP synthase dimers form a left- or right-handed helix along the curved ridges of the cristae or along the outer perimeter of helical tubular cristae. In these organisms, the principal membrane curvature is caused by the macromolecular association of dimers into helical arrays, rather than by the individual dimers as in metazoans (Fig. 3 J and M). It will be interesting to investigate the architecture of ATP synthase complexes in the bloodstream stage of T. brucei, which lacks oxidative phosphorylation and contains sparse, if any, mitochondrial cristae (32) and uses the enzyme as an ATP-powered proton pump to generate the mitochondrial membrane potential (16, 18).

A striking difference between the metazoa-type ATP synthase dimers and ATP synthase dimers of protozoans or unicellular algae, which we refer to as the protozoan type, is the structural diversity of the peripheral stalk and dimer interfaces (33). The architectures of all metazoa-type ATP synthase dimers studied so far are virtually identical (3), and sequence analysis and proteomics have identified homologous subunits for nearly all of the subunits that constitute the mitochondrial ATP synthase in these organism (34, 35). In contrast, there seem to be no subunits in the protozoan-type ATP synthases that correspond to the peripheral stalk or dimer interface in the metazoa type. Instead, proteomics and biochemical analysis have identified several unique subunits, none of which are homologous among the
various phyla (10, 11, 36, 37). This variation in subunit composition correlates with the observed diversity in structure, which may reflect adaptations to different energetic requirements or environmental conditions. Apparently, these adaptations occurred before trypanosomes adopted their parasitic lifestyle.

Unusual Arrangement of Catalytic Subunits on the F1 Head of Euglenozoan ATP Synthase Dimers. Despite the diversity of ATP synthase dimer architectures observed in different species, the catalytic subunits that are involved in ATP synthesis and form the F1 subcomplex are structurally conserved in all previously reported structures (36, 38, 39). Here we show that the catalytic F1 head of the mitochondrial ATP synthase dimers from *E. gracilis* and *T. brucei* exhibits an unprecedented pyramidal structure (Fig. 2 and Fig. S2). This unusual arrangement of catalytic subunits is surprising, given the otherwise strictly conserved pseudo-sixfold symmetry and catalytic mechanism of the (αβ)3 hexamer in all other known F-type ATPases, including those of bacteria and chloroplasts (4, 24, 40). The observed alterations in the *T. brucei* and *E. gracilis* F1 region are in line with previous biochemical studies reporting a complete cleavage of the α subunit into two fragments, both of which remain bound to the ATP synthase (10–14). Furthermore, it has been shown that despite the α-subunit fragmentation, the *T. brucei* ATP synthase remains catalytically active throughout the life cycle and is essential for parasite survival in both the procyclic and bloodstream forms (10, 16, 18, 19); as mentioned above, it is the reverse function as a proton-pumping ATPase that is essential for survival in the mammalian host. In addition, the isolated mitochondrial membranes from *E. gracilis* isolated in this study contained catalytically active ATPase, as demonstrated by in-gel activity (Fig. S5).

In the bovine ATP synthase, the catalytic sites are located at the interface of the α- and β-subunits (Fig. 4C). The nucleotide-binding sites are formed largely by the central domain of the β subunits, whereas the α subunits contribute a conserved arginine (the arginine finger) that is essential for catalysis (8) (Fig. 4C and F). In euglenozoa, the nucleotide-binding site in the β subunit is conserved (13), but the α/β interface is not (Fig. 2B and E and Fig. S2 C and G); however, in the unusual arrangement of catalytic subunits, it appears unlikely that the arginine in the α subunit, which is present in *T. brucei* and *E. gracilis*, can reach the bound nucleotide from its remote position in the central domain of the α fragment (Fig. 4D and E). Because the arginine finger is critical for catalysis in both the ATP synthase (8) and in many other ATPase domains in general (e.g., AAA+ ATPases) (41), it must be located at the active site and thus must be contributed by a different part of the structure. Apart from a suitably positioned residue in the α fragment, a prime candidate is the euglenozoan-specific F1 subunit p18 (10–13) (systematic ID in *T. brucei* Tb927.5.1710). Knockdown of p18 in procyclic *T. brucei* by RNAi has been shown to result in slowed growth and disappearance of F1, indicating that p18 is required for F1 integrity (42). Although functional data on the role of p18 in other trypanosomatids has not been reported to date, p18 has been identified as a subunit of mitochondrial ATP synthase in other trypanosomatids, including *Leishmania tarentolae* (13) and *Crithidia fasciculata* (12), as well as in the euglenoid *E. gracilis* (11), strongly suggesting that this subunit is generally conserved in euglenozoa. The location of p18 in the unusual F1 structure is currently unknown; however, an unassigned density region between αN and αC and the likely presence of more than one copy of p18 per F1 molecule (10) suggest that this region may contain p18 (Fig. S6). Resolving the question of how the euglenozoan ATP synthase dimers produce ATP will require a higher-resolution structure.

The in Situ Structure of Euglenozoan ATP Synthase Dimers Provides Insight into Rotary Catalysis. ATP synthase by rotary catalysis involves three key stages: (i) proton movement across the membrane-embedded F0 sector drives the rotation of the central stalk; (ii) rotation of the central stalk causes conformational changes at the catalytic αβ interface, which triggers phosphate bond formation; and (iii) conformational changes are cooperatively transmitted between catalytic sites. The exact molecular mechanism of this process is not known. A central question is the extent to which each of the α and β subunits contributes to this process. In the euglenozoan ATP synthase, the contributions of each subunit to rotary catalysis can be dissected. The displacement of the αC fragments away from the central stalk means that the γ subunit is unable to interact with the α subunits directly (Fig. 4A and B). Thus, at least in the euglenozoan ATP synthase, but possibly also in the conserved F1 structure, interactions between the β and γ subunits are sufficient for the conversion of γ-subunit rotation into conformational changes in the process of mechanocompelling coupling. The different position of the αC fragments in the euglenozoan ATP synthase also rules out involvement of the γ subunit in the transmission of conformational changes between catalytic sites, because this subunit can interact with only one β subunit at a time. This observation is consistent with results of high-speed atomic force microscopy demonstrating that in *Bacillus PS3*, conformational transmission between catalytic sites does not require the central stalk, but the loss of a single α subunit stops the propagation of conformational changes (43). In the *T. brucei* and *E. gracilis* structures, the central and C-terminal domains of the α subunit are displaced from their canonical position but remain bound to the complex, indicating an essential role of the fragment in enzymatic activity. In these complexes, the αC fragment connects the C-terminal domain of one β subunit to the central domain of the neighboring subunit (Fig. 2C and Fig. S2D). In the bovine complex, the distance between these two contact points in one β subunit changes significantly during nucleotide binding and release (4). Therefore, the arc-shaped αC fragments in the euglenozoan complexes likely transmit conformational changes from one β subunit to the next, ensuring cooperativity among the three catalytic sites in the F1 head.

![Fig. 4. Domain arrangement in the catalytic F1 region. (A and B) Cross-section of catalytic domain of *E. gracilis* and *T. brucei* with fitted atomic models as indicated in Fig. 2C, contoured in steps of 0.8 e. (C) Cross-section of the conserved bovine structure (PDB code: 1BMF) (4) for comparison. Blue, β subunits; yellow αC fragment; gray, central stalk. Green asterisks indicate nucleotide-binding sites in the β subunits. (D–F) Enlarged view of the catalytic interface from *E. gracilis* (D), *T. brucei* (E), and *B. taurus* (F), with the arginine finger circled in red and the nucleotide-binding site in green. (Scale bar: 5 nm.)](https://www.pnas.org/content/114/5/995)
Conclusion

Electron cryotomography of mitochondria from *E. gracilis* and insect-stage *T. brucei* shows that the ATP synthases of both organisms form dimers, which assemble into short left-handed helical arcs around the edge of disk-shaped cristae. Subtomogram averaging of the ATP synthase dimers in these species reveals a noncanonical F₁ head structure. This unusual structure is the result of a protelytic cleavage of the α subunit, which causes a displacement of the central and C-terminal domains of the α subunit from their canonical position. The structure of the euglenozoan F₁ head provides insight into the mechanism of rotary catalysis, in particular how rotation of the central stalk induces conformational changes and how these changes are transmitted between catalytic sites. The unique structure of the F₁ head of the euglenozoan ATP synthases represents a potential model system for establishing future in-depth studies on the mechanism of ATP synthesis, a fundamental and essential life process.

Materials and Methods

Culture Conditions and Sample Preparation. *T. brucei* procyclic form cells (strain 29.13 (44)) were grown at 27 °C and 5% (vol/vol) CO₂ in SDM-79 medium (5 mM Hepes, 55 μM MgCl₂, and 10% (vol/vol) FBS (Gibco)). Cells were counted with a 22 Cell Counter (Beckman Coulter) and harvested by centrifugation at 1,300 × g for 5 min. Then 600 × 10⁶ cells were resuspended in 1.5 mL of homogenization buffer (20 mM Tris pH 7.4, 250 mM sucrose, and 5 mM MgCl₂) and homogenized by 350 passages in a ball-bearing cell homogenizer with 4-μm clearance (Isobiotec). The cell lysate was centrifuged at 500 × g for 10 min at 4 °C. Mitochondrial membranes were pelleted by centrifugation at 16,000 × g for 10 min at 4 °C.

*E. gracilis* (Lebendkulturen Helbig) cultures were grown in Euglena medium (46) under a cycle of 10 h of light at 10,000 lux and 14 h of darkness at 21 °C. Then 100-mL cultures were harvested by centrifugation at 800 × g for 10 min at 4 °C, followed by resuspension in 10 mL of homogenization buffer (20 mM Tris pH 7.4, 250 mM sucrose, and 5 mM MgCl₂). Cells were homogenized by 30 passages in a cell homogenizer with 8-μm clearance, and the homogenate was centrifuged at 1,500 × g for 5 min at 4 °C. The supernatant was collected and centrifuged at 6,000 × g for 10 min at 4 °C. The resulting pellet contained an upper green layer, which was carefully removed and discarded, and a brownish pellet containing the mitochondrial membranes.

**BN-PAGE of Solubilized Membrane Complexes and In-Gel ATPase Activity Assay.** *E. gracilis* mitochondrial membranes were resuspended in solubilization buffer (final concentration, 50 mM Tris·HCl pH 8.0, 1 mM MgCl₂ with Roche Complete Protease Inhibitor Mixture added), mixed with DDM in a final detergent-to-protein ratio (DPR) of 1:1 (wt/wt) or 5:1 (wt/wt), and incubated for 30 min on ice. Samples were centrifuged at 15,000 × g for 15 min at 4 °C, after which solubilized protein complexes were collected from the supernatant. Solubilized *E. gracilis* mitochondrial membrane protein complexes were separated by BN-PAGE as described by Wittig et al. (47), using 3–12% NativePAGE Novex Bis-Tris gradient gels (Life Technologies), which were subsequently stained with Coomassie blue R-250. In-gel ATPase activity assays were performed as described previously (48), with slight modifications. After gel electrophoresis, native gels were rinsed twice in 1 μL H₂O and once in 1 M Tris·HCl pH 7.8, followed by incubation in activity buffer (35 mM Tris·HCl pH 7.8 and 14 mM Mg(SO₄)). ATP, lead (III) nitrate, and N,N-Dimethyl-n-dodecylamine N-oxide were added in trace amounts, whereupon white lead phosphate precipitate formed inside the protein bands with ATPase activity during overnight incubation.

**Electron Cryotomography and Tomogram Processing.** Mitochondrial membrane pellets were resuspended in 50 μL of freezing buffer (20 mM Tris pH 7.4 and 250 mM trehalose) and mixed 1:1 with 6-nm colloidal gold fiducial markers (Aurion). Then 3 μL was applied to a glow-discharged 22/2 Cu 300-mesh holey carbon-coated support grid (Quantifoil). Excess liquid was removed by blotting (#4 Whatman paper; Sigma-Aldrich), and the grid was rapidly frozen in liquid ethane using a home-built gullotine. Frozen-hydrated specimens of *T. brucei* mitochondrial membranes were imaged in a Titan Krios electron microscope (FEI) at a nominal magnification of 42,000× (pixel size 3.35 Å). Tomographic tilt series were collected at 4.5-μm defocus to ±60° in 2° increments, starting at +24°.

*E. gracilis* mitochondrial membranes were imaged with an FEI Tecnai Polara transmission electron microscope at a nominal magnification of 90,000× (pixel size 2.28 Å), and tilt series were collected at 3.5-μm defocus from -50° to +60° in 2° increments, starting at +24°. In both cases, tilt series were collected using the program LATITUDE (Digital Micrograph; Gatan). The cumulative dose of a tilt series was ~100 electron/A². Both microscopes were operated at 300 kV and equipped with a Quantum Energy filter and a K2 summit direct electron detector (Gatan) operated in counting mode. Tomographic volumes were reconstructed using IMOD (49). The handedness of tomographic volumes was determined by evaluating the tilt axis rotation angle in tilt series of a sample of known handedness. For 3D visualization, tomographic volumes were contrast-enhanced by nonlinear anisotropic diffusion filtering (51) and segmented manually with AMIRA (FEI). Subtomogram averages were placed into original tomographic volumes using the EMPerick module (52) for AMIRA.

**Subtomogram Averaging.** ATP synthase dimers were averaged as described previously (3). In brief, particle pairs were manually identified in tomographic volumes, and initial rotation angles were assigned based on the F₁ head positions relative to the membrane in SPIDER (53). Particles were extracted, rotated, and averaged to create an initial reference. Particle alignment was optimized iteratively in PEET (54), using a restricted search range. Resolution of final maps was estimated by Fourier shell correlation (FSC). Particle sets were randomly split into two halves after alignment. The resulting half-maps were multiplied with a Gaussian mask before FSC. To prevent mask bias, tomographic volumes were phase-randomized beyond 40 Å using PEET, and two half-maps were generated by extracting, aligning, and averaging the same particle subsets. The FSC curves of the phase-randomized half-sheets indicated a drop at the randomization frequency. 3D visualization, rigid body fitting, and difference map generation were performed in UCSF Chimera (55).

**Sequence Alignment.** Sequence alignment was performed with ClustalW2 (56). Sequences were obtained from TriTrypDB (57), UniProt (58), and Perez et al. (11).

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