1	An ancestral interaction module promotes oligomerisation in divergent
2	mitochondrial ATP synthases
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18	Abstract
19	Mitochondrial ATP synthase forms stable dimers arranged into oligomeric assemblies that
20	generate the inner-membrane curvature essential for efficient energy conversion. Here, we
21	report cryo-EM structures of the intact ATP synthase dimer from trypanosomes in 10 different
22	rotational states. The model consists of 25 subunits, including 11 lineage-specific, as well as
23	36 lipids. The rotary mechanism is influenced by the divergent peripheral stalk, conferring a
24	greater conformational flexibility. Proton transfer in the lumenal half-channel occurs via a
25	chain of five ordered water molecules. The dimerization interface is formed by subunit-g that
26	is critical for interactions but not for the catalytic activity. Although overall dimer architecture
27	varies among eukaryotes, we find that subunit- g and $-e$ form a common ancestral
28	oligomerisation motif, which is shared between the trypanosomal and mammalian lineages.
29	Therefore, our data defines the subunit-g/e module as a structural component determining ATP
30	synthase oligomeric assemblies.
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34 Main

Mitochondrial ATP synthase consists of the soluble F₁ and membrane-bound F₀ subcomplexes, 35 and occurs in dimers that assemble into oligomers to induce the formation of cristae folds. The 36 cristae folds are the sites for oxidative phosphorylation and energy conversion in eukaryotic 37 cells. Dissociation of ATP synthase dimers into monomers results in the loss of native cristae 38 architecture and impairs mitochondrial function^{1,2}. While the cristae morphology varies 39 substantially between organisms from different lineages, ranging from flat lamellar in 40 opisthokonts to coiled tubular in ciliates and discoidal in euglenozoans³, the ATP synthase 41 42 dimers represent a universal occurrence to maintain the membrane shape⁴.

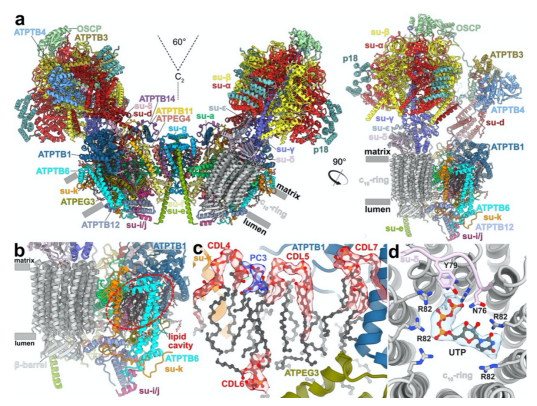
- ATP synthase dimers of variable size and architecture, classified into types I to IV have 43 recently been resolved by high-resolution cryo-EM studies. In the structure of the type-I ATP 44 synthase dimer from mammals, the monomers are only weakly associated^{5,6}, and in yeast 45 46 insertions in the membrane subunits form tighter contacts⁷. The structure of the type-II ATP synthase dimer from the alga *Polytomella* showed that the dimer interface is formed by 47 phylum-specific components⁸. The type-III ATP synthase dimer from a ciliate *Tetrahymena* is 48 characterized by parallel rotary axes, and a substoichiometric subunit, as well as multiple lipids 49 were identified at the dimer interface, while additional protein components that tie the 50 51 monomers together are distributed between the matrix, transmembrane, and lumenal regions⁹. The structure of the type-IV ATP synthase with native lipids from Euglena also showed that 52 specific protein-lipid interactions contribute to the dimerization, and that the central and 53 peripheral stalks interact with each other directly¹⁰. Finally, a unique apicomplexan ATP 54 55 synthase dimerises via 11 parasite-specific components that contribute ~7000 Å² buried surface area¹¹, and unlike all other ATP synthases, that assemble into rows, it associates in higher 56 oligomeric states of pentagonal pyramids in the curved apical membrane regions. Together, the 57 available structural data suggest a diversity of oligomerisation, and it remains unknown 58 59 whether common elements mediating these interactions exist or the dimerization of ATP synthase occurred independently and multiple times in evolution⁴. 60
- The ATP synthase of Trypanosoma brucei, a representative of kinetoplastids an established 61 medically important model organism causing the sleeping sickness, is highly divergent, 62 exemplified by the pyramid-shaped F_1 head containing a phylum specific subunit^{12,13}. The 63 dimers are sensitive to the lack of cardiolipin¹⁴ and form short left-handed helical segments 64 that extend across the membrane ridge of the discoidal cristae¹⁵. Uniquely among aerobic 65 eukaryotes, the mammalian life cycle stage of *T. brucei* utilizes the ATP synthase as a proton 66 pump maintaining the mitochondrial membrane potential at the expense of ATP^{16,17}, whereas 67 68 the insect stage of the parasite employs the forward ATP-producing mode of the enzyme^{18,19}.
- Given the conservation of the core subunits, the different nature of oligomerisation and the
 ability to test structural hypotheses biochemically, we reasoned that investigation of the *T*. *brucei* ATP synthase structure and function would provide the missing evolutionary link to
 understand how the monomers interact to form physiological dimers. Here, we address this
- 72 question by combining structural, functional and evolutionary analysis of the *T. brucei* ATP
- 74 synthase dimer.
- 75

76 Results

77 Cryo-EM structure of the *T. brucei* ATP synthase

We purified ATP synthase dimers from cultured *T. brucei* procyclic trypomastigotes by affinity chromatography with a recombinant natural protein inhibitor TbIF_{1}^{20} , and subjected to cryo-EM analysis (Supplementary Fig. 1 and 2). Using masked refinements, maps were obtained for the membrane region, the rotor, and the peripheral stalk. To further describe the conformational space of the *T. brucei* ATP synthase, we resolved 10 distinct rotary substates, which were refined to 3.5-4.3 Å resolution. Finally, particles with both monomers in rotational state 1 were selected, and the consensus structure of the dimer was refined to 3.2 Å resolution

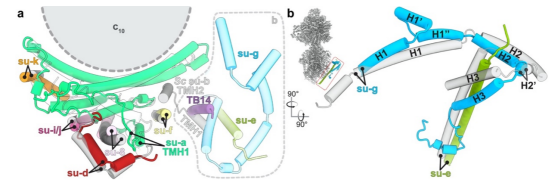
- 85 (Supplementary Table 1, Supplementary Fig. 2).
- 86 Unlike the wide-angle architecture of dimers found in animals and fungi, the T. brucei ATP
- 87 synthase displays an angle of 60° between the two F₁/*c*-ring subcomplexes. The model of the
- 88 *T. brucei* ATP synthase includes all 25 different subunits, 11 of which are lineage-specific (Fig.
- 89 1a, Supplementary Video 1, Supplementary Fig. 3). We named the subunits according to the
- 90 previously proposed nomenclature $^{21-23}$ (Supplementary Table 2). In addition, we identified and
- 91 modeled 36 bound phospholipids, including 24 cardiolipins (Supplementary Fig. 4). Both
- 92 detergents used during purification, n-dodecyl β -D-maltoside (β -DDM) and glyco-diosgenin
- 93 (GDN) are also resolved in the periphery of the membrane region (Supplementary Fig. 5).
- In the catalytic region, F_1 is augmented by three copies of subunit p18, each bound to subunit-94 $\alpha^{12,13}$. Our structure shows that p18 is involved in the unusual attachment of F₁ to the peripheral 95 stalk. The membrane region includes seven conserved F_0 subunits (d, f, 8, i/j, k, e, and g) 96 arranged around the central proton translocator subunit-a. We identified those subunits based 97 98 on the structural similarity and matching topology to their yeast counterparts (Fig 2). Surprisingly, the long helix-2 of subunit-b (bH2), which constitutes the central part of the 99 peripheral stalk in other organisms and associates with subunit-*a* in the membrane, is absent in 100 101 T. brucei. By contrast, bH1 from the yeast structure superposes well with the single transmembrane helix of ATPTB14, which anchors the newly identified subunit-e and -g to the 102 F_o (Fig 2a). Therefore, ATPTB14 may represent a highly reduced homolog of subunit-*b*. 103
- 104 The membrane region contains a peripheral subcomplex, formed primarily by the phylum-
- specific ATPTB1,6,12 and ATPEG3 (Fig. 1b). It is separated from the conserved core by a membrane-intrinsic cavity, in which nine bound cardiolipins are resolved (Fig. 1c), and the
- 107 C-terminus of ATPTB12 interacts with the lumenal β -barrel of the c_{10} -ring. In the cavity of the
- 108 decameric c-ring near the matrix side, 10 Arg66_c residues coordinate a ligand density, which
- is consistent with a pyrimidine ribonucleoside triphosphate (Fig. 1d). We assign this density as
- uridine-triphosphate (UTP), due to its large requirement in the mitochondrial RNA metabolism
 of African trypanosomes being a substrate for post-transcriptional RNA editing²⁴, and addition
- 112 of poly-uridine tails to gRNAs and rRNAs 25,26 , as well as due to low abundance of cytidine
- 113 triphosphate $(CTP)^{27}$. The nucleotide base is inserted between two Arg82_c residues, whereas
- 114 the triphosphate region is coordinated by another five Arg82_c residues, with Tyr79_{δ} and Asn76_{δ}
- 115 providing asymmetric coordination contacts. The presence of a nucleotide inside the *c*-ring is
- surprising, given the recent reports of phospholipids inside the c-rings in mammals^{5,6} and
- 117 ciliates⁹, indicating that a range of different ligands can provide structural scaffolding.





119 Fig. 1: The *T. brucei* ATP synthase structure with lipids and ligands.

a. Front and side views of the composite model with both monomers in rotational state 1. The 120 121 two F_1/c_{10} -ring complexes, each augmented by three copies of the phylum-specific p18 subunit, are tied together at a 60°-angle. The membrane-bound F_o region displays a unique architecture 122 and is composed of both conserved and phylum-specific subunits. b, Side view of the Fo region 123 showing the lumenal interaction of the ten-stranded β -barrel of the *c*-ring (grey) with ATPTB12 124 125 (pale blue). The lipid-filled peripheral F_o cavity is indicated. c, Close-up view of the bound lipids within the peripheral F_o cavity with cryo-EM density shown. d, Top view into the 126 decameric *c*-ring with a bound pyrimidine ribonucleoside triphosphate, assigned as UTP. Map 127 density shown in transparent blue, interacting residues shown. 128



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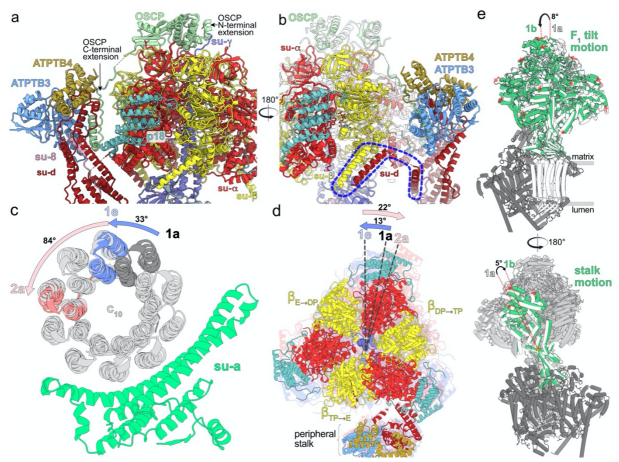
130 Fig. 2: Identification of conserved F₀ subunits.

a, Top view of the membrane region with *T. brucei* subunits (colored) overlayed with *S. cerevisiae* structure (gray transparent). Close structural superposition and matching topology
allowed the assignment of conserved subunits based on matching topology and location.
b, Superposition of subunits-*e* and -*g* with their *S. cerevisiae* counterparts (PDB 6B2Z)
confirms their identity.

136 Peripheral stalk flexibility and distinct rotational states

The trypanosomal peripheral stalk displays a markedly different architecture compared to its 137 yeast and mammalian counterparts. In the opisthokont complexes, the peripheral stalk is 138 organized around the long bH2, which extends from the membrane ~ 15 nm into the matrix and 139 attaches to OSCP at the top of $F_1^{5,7}$. By contrast, *T. brucei* lacks the canonical *b*H2 and instead, 140 helices 5-7 of divergent subunit-d and the C-terminal helix of extended subunit-8 bind to a C-141 terminal extension of OSCP at the apical part of the peripheral stalk (Fig. 3a). The interaction 142 between OSCP and subunit-d and -8 is stabilized by soluble ATPTB3 and ATPTB4. The 143 144 peripheral stalk is rooted to the membrane subcomplex by a transmembrane helix of subunit-8, wrapped on the matrix side by helices 8-11 of subunit-*d*. Apart from the canonical contacts 145 at the top of F_1 , the peripheral stalk is attached to the F_1 via a euglenozoa-specific C-terminal 146 extension of OSCP, which contains a disordered linker and a terminal helix hairpin extending 147 between the F_1 -bound p18 and subunits -d and -8 of the peripheral stalk (Fig. 3a, 148 149 Supplementary Videos 2,3). Another interaction of F_1 with the peripheral stalk occurs between the stacked C-terminal helices of subunit- β and -d (Fig. 3b), the former of which structurally 150 belongs to F₁ and is connected to the peripheral stalk via a flexible linker. 151

To assess whether the unusual peripheral stalk architecture influences the rotary mechanism, 152 we analysed 10 classes representing different rotational states. The three main states (1-3) result 153 from ~120° rotation of the central stalk subunit- γ , and we identified five (1a-1e), four (2a-2d) 154 and one (3) classes of the respective main states. The rotor positions of the rotational states 1a, 155 156 2a and 3 are related by steps of 117°, 136° and 107°, respectively. Throughout all the identified substeps of the rotational state 1 (classes 1a to 1e) the rotor turns by $\sim 33^{\circ}$, which corresponds 157 approximately to the advancement by one subunit-c of the c_{10} -ring. While rotating along with 158 159 the rotor, the F_1 headpiece lags behind, advancing by only ~13°. During the following transition from 1e to 2a, the rotor advances by $\sim 84^\circ$, whereas the F₁ headpiece rotates $\sim 22^\circ$ in the opposite 160 direction (Fig. 3c,d). This generates a counter-directional torque between the two motors, 161 which is consistent with a power-stroke mechanism. Albeit with small differences in step size, 162 this mechanism is consistent with a previous observation in the *Polytomella* ATP synthase⁸. 163 However, due to its large, rigid peripheral stalk, the *Polytomella* ATP synthase mainly displays 164 rotational substeps, whereas the *Trypanosoma* F_1 also displays a tilting motion of ~8° revealed 165 by rotary states 1 and 2 (Fig. 3e, Supplementary Video 2). The previously reported hinge 166 motion between the N- and C-terminal domains of OSCP⁸ is not found in our structure, instead, 167 the conformational changes of the F_1/c_{10} -ring subcomplex are accommodated by a 5° bending 168 169 of the apical part of the peripheral stalk. (Fig. 3e, Supplementary Videos 2,3). Together, the structural data indicate that the divergent peripheral stalk attachment confers greater 170 conformational flexibility to the T. brucei ATP synthase. 171



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Fig. 3: A divergent peripheral stalk allows high flexibility during rotary catalysis. a, N-173 terminal OSCP extension provides a permanent central stalk attachment, while the C-terminal 174 extension provides a phylum-specific attachment to the divergent peripheral stalk. b, The C-175 176 terminal helices of subunits $-\beta$ and -d provide a permanent F₁ attachment. **c**, Substeps of the *c*ring during transition from rotational state 1 to 2. d, F₁ motion accommodating steps shown in 177 (c). After advancing along with the rotor to state 1e, the F_1 rotates in the opposite direction 178 when transitioning to state 2a. e, Tilting motion of F₁ and accommodating bending of the 179 180 peripheral stalk.

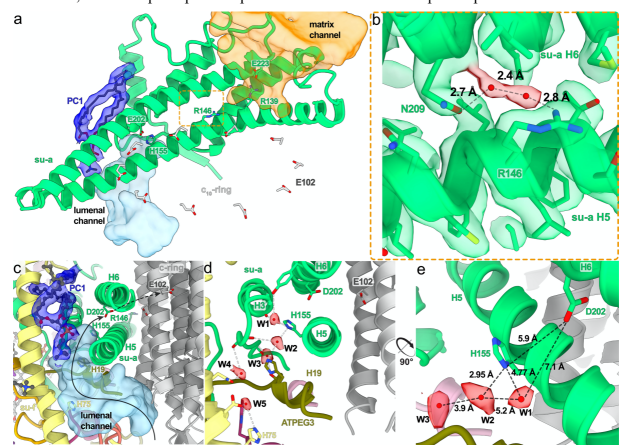
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182 Lumenal proton half-channel is insulated by a lipid and contains ordered water molecules

183 The mechanism of proton translocation involves sequential protonation of E102 of subunits-c, 184 rotation of the c_{10} -ring with neutralized E102c exposed to the phospholipid bilayer, and release 185 of protons on the other side of the membrane. The sites of proton binding and release are

- 186 separated by the conserved R146 contributed by the horizontal helix H5 of subunit-a and are
- accessible from the cristae lumen and mitochondrial matrix by aqueous half channels (Fig. 4a).
- 188 Together, R146 and the adjacent N209 coordinate a pair of water molecules in between helices
- 189 H5 and H6 (Fig. 4b). A similar coordination has been observed in the Polytomella ATP
- 190 synthase⁸. The coordination of water likely restricts the R146 to rotamers that extend towards
- 191 the *c*-ring, with which it is thought to interact.
- In our structure, the lumenal half-channel is filled with a network of resolved water densities,ending in a chain of five ordered water molecules (W1-W5; Fig. 4c,d,e). The presence of

- 194 ordered water molecules in the aqueous channel is consistent with a Grotthuss-type mechanism
- 195 for proton transfer, which would not require long-distance diffusion of water molecules⁵.
- 196 However, because some distances between the observed water molecules are too large for 197 direct hydrogen bonding, proton transfer may involve both coordinated and disordered water
- molecules. The distance of 7 Å between the last resolved water (W1) and $D202_a$, the conserved
- residue that is thought to transfer protons to the *c*-ring, is too long for direct proton transfer.
- Instead, it may occur via the adjacent $H155_a$. Therefore, our structure resolves individual
- 201 elements participating in proton transport (Fig. 4d,e).
- 202 The lumenal proton half-channel in the mammalian^{5,6} and apicomplexan¹¹ ATP synthase is
- 203 lined by the transmembrane part of bH2, which is absent in *T. brucei*. Instead, the position of
- bH2 is occupied by a fully ordered phosphatidylcholine in our structure (PC1; Fig. 4a,c).
- 205 Therefore, a bound lipid replaces a proteinaceous element in the proton path.



207 Fig. 4: The lumenal half-channel contains ordered water molecules and is confined by an 208 F_0 -bound lipid. a, Subunit-a (green) with the matrix (orange) and lumenal (light blue) channels, and an ordered phosphatidylcholine (PC1; blue). E102 of the c_{10} -ring shown in grey. 209 **b**, Close-up view of the highly conserved R146 $_a$ and N209 $_a$, which coordinate two water 210 molecules between helices H5- 6_a . c, Sideview of the lumenal channel with proton pathway 211 (light blue) and confining phosphatidylcholine (blue). d, Chain of ordered water molecules in 212 the lumenal channel. Distances between the W1-W5 are 5.2, 3.9, 7.3 and 4.8 Å, respectively. 213 e, The ordered waters extend to $H155_a$, which likely mediates the transfer of protons to $D202_a$. 214

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217 Subunit-g facilitates assembly of different ATP synthase oligomers

Despite sharing a set of conserved F_o subunits, the *T. brucei* ATP synthase dimer displays a 218 markedly different dimer architecture compared to all the previously determined structures. 219 First, its interface of 3,600 Å² is smaller than that of the *E. gracilis* type-IV (10,000 Å²) and 220 221 the *T. thermophila* type-III ATP synthases (16,000 Å²). Second, unlike mammalian and fungal 222 ATP synthase, in which the peripheral stalks extend in the plane defined by the two rotary axes, in our structure the monomers are rotated such that the peripheral stalks are offset laterally on 223 the opposite sides of the plane. Due to the rotated monomers, this architecture is associated 224 225 with a specific dimerization interface, where two subunit-g copies interact homotypically on 226 the C₂ symmetry axis (Fig. 5a, Supplemetary Video 1). Both copies of H1-2_g extend horizontally along the matrix side of the membrane, clamping against each other (Fig. 5c,e). 227 228 This facilitates formation of contacts between an associated transmembrane helix of subunit-e with the neighbouring monomer via subunit-a' in the membrane, and -f' in the lumen, thereby 229 230 further contributing to the interface (Fig. 5b). Thus, the ATP synthase dimer is assembled via the subunit-e/g module. The C-terminal part of the subunit-e helix extends into the lumen, 231 towards the ten-stranded β -barrel of the *c*-ring (Supplementary Fig. 6a). The terminal 23 232 residues are disordered with poorly resolved density connecting to the detergent plug of the c-233 234 ring β -barrel (Supplementary Fig. 6b). This resembles the lumenal C-terminus of subunit-*e* in the bovine structure⁵, indicating a conserved interaction with the c-ring. 235

The e/g module is held together by four bound cardiolipins in the matrix leaflet, anchoring it to the remaining F_o region (Fig. 5c). The head groups of the lipids are coordinated by polar and charged residues with their acyl chains filling a central cavity in the membrane region at the dimer interface (Fig 5c, Supplementary Fig. 4f). Cardiolipin binding has previously been reported to be obligatory for dimerization in secondary transporters²⁸ and the depletion of cardiolipin synthase resulted in reduced levels of ATP synthase in the bloodstream trypanosomes¹⁴.

Interestingly, for yeasts, early blue native gel electrophoresis²⁹ and subtomogram averaging 243 studies² suggested subunit-g as potentially dimer-mediating, however the e/g modules are 244 located laterally opposed on either side of the dimer long axis, in the periphery of the complex, 245 246 ~8.5 nm apart from each other. Because the e/g module does not interact directly within the yeast ATP synthase dimer, they have been proposed to serve as membrane bending elements, 247 whereas the major dimer contacts are formed by subunit-a and $-i/j^7$. In mammals, the e/g248 module occupies the same position as in yeasts, forming the interaction between two diagonal 249 monomers in a tetramer^{5,6,30}, as well as between parallel dimers³¹. The comparison with our 250 251 structure shows that the overall organization of the intra-dimeric trypanosomal and interdimeric mammalian e/g module is structurally similar (Fig. 5d). Furthermore, kinetoplastid 252 parasites and mammals share conserved GXXXG motifs in subunit- e^{32} and -g (Supplementary 253 Fig. 8), which allow close interaction of their transmembrane helices (Fig. 5e), providing 254 255 further evidence for subunit homology. However, while the mammalian ATP synthase dimers are arranged perpendicularly to the long axis of their rows along the edge of cristae³³, the 256 T. brucei dimers on the rims of discoidal cristae are inclined ~ 45° to the row axis¹⁵. Therefore, 257 258 the *e/g* module occupies equivalent positions in the rows of both evolutionary distant groups 259 (Fig. 5f and reference 31).

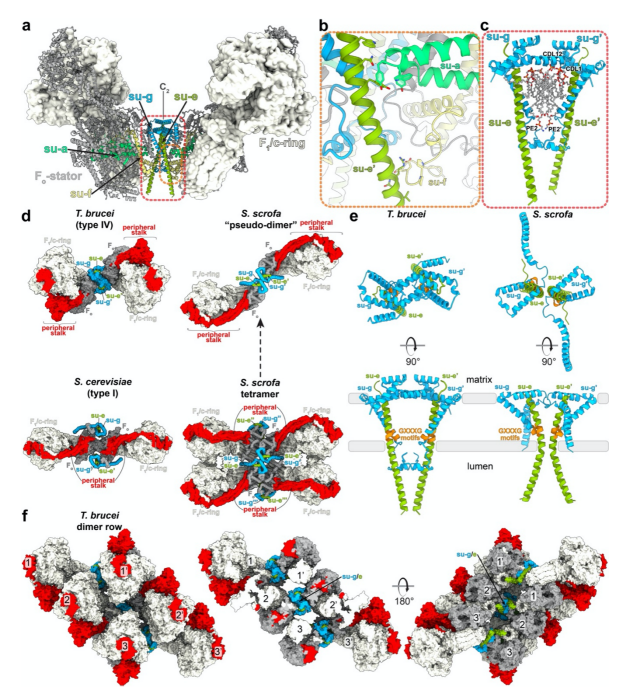
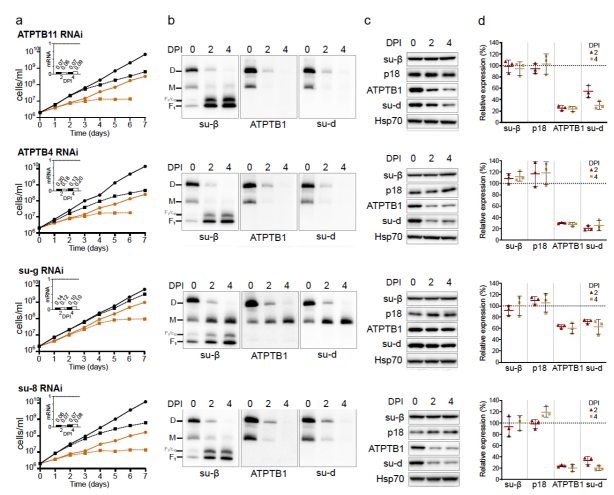


Fig. 5: The homotypic dimerization motif of subunit-g generates a conserved 261 oligomerisation module. a, Side view with dimerising subunits colored. b,c, The dimer 262 interface is constituted by (b) subunit-e' contacting subunit-a in the membrane and subunit-f263 264 in the lumen, (c) subunits e and g from both monomers forming a subcomplex with bound lipids. d, Subunit-g and -e form a dimerization motif in the trypanosomal (type-IV) ATP 265 synthase dimer (this study), the same structural element forms the oligomerisation motif in the 266 porcine ATP synthase tetramer. The structural similarity of the pseudo-dimer in the porcine 267 structure with the trypanosomal dimer suggests that that type I and IV ATP synthase dimers 268 have evolved through divergence from a common ancestor. e, The dimeric subunit-e/g 269 structures are conserved in pig (PDB 6ZNA) and T. brucei (this work) and contain a conserved 270 GXXXG motif (orange) mediating interaction of transmembrane helices. f, Models of ATP 271 synthase dimers fitted into subtomogram averages of short oligomers¹⁵ (EMD-3560). 272



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Fig. 6: RNAi knockdown of subunit-g results in monomerization of ATP synthase. a, 274 Growth curves of non-induced (solid lines) and tetracycline-induced (dashed lines) RNAi cell 275 276 lines grown in the presence (black) or absence (brown) of glucose. The insets show relative levels of the respective target mRNA at indicated days post-induction (DPI) normalized to the 277 278 levels of 18S rRNA (black bars) or ß-tubulin (white bars). b, Immunoblots of mitochondrial 279 lysates from indicated RNAi cell lines resolved by BN-PAGE probed with antibodies against 280 indicated ATP synthase subunits. c, Representative immunoblots of whole cell lysates from indicated RNAi cell lines probed with indicated antibodies. d, Quantification of three replicates 281 of immunoblots in (c). Values were normalized to the signal of loading marker Hsp70 and to 282 283 non-induced cells. Plots show means with standard deviations (SD).

284 Subunit-g retains the dimer but is not essential for the catalytic monomer

To validate structural insights, we knocked down each individual F_o subunit by inducible RNA 285 interference (RNAi). All target mRNAs dropped to 5-20 % of their original levels after two 286 and four days of induction (Fig. 6a and Supplementary Fig. 7a, insets). Western blot analysis 287 288 of whole-cell lysates resolved by denaturing electrophoresis revealed decreased levels of Fo subunits ATPB1 and -d suggesting that integrity of F_0 moiety depends on the presence of other 289 Fo subunits. Immunoblotting of mitochondrial complexes resolved by blue native 290 polyacrylamide gel electrophoresis (BN-PAGE) with antibodies against F₁ and F_o subunits 291 292 revealed a strong decrease or nearly complete loss of dimeric and monomeric forms of ATP 293 synthases four days after induction of RNAi of most subunits (e, f, i/j, k, 8, ATPTB3, ATPTB4, ATPTB6, ATPTB11, ATPTB12, ATPTB14, ATPEG3 and ATPEG4), documenting an 294 increased instability of the enzyme or defects in its assembly. Simultaneous increase in F₁-295 296 ATPase levels demonstrated that the catalytic moiety remains intact after the disruption of the 297 peripheral stalk or the membrane subcomplex (Fig. 6b,c,d and Supplementary Fig. 7b,c,d).

In contrast to the other targeted F_o subunits, the downregulation of subunit-g with RNAi 298 299 resulted in a specific loss of dimeric complexes with concomitant accumulation of monomers 300 (Fig. 6b), indicating that it is required for dimerization, but not for the assembly and stability of the monomeric F1F0 ATP synthase units. Transmission electron microscopy of thin cell 301 sections revealed that the ATP synthase monomerization in the subunit-g^{RNAi} cell line had the 302 same effect on mitochondrial ultrastructure as nearly complete loss of monomers and dimers 303 upon knockdown of subunit-8. Both cell lines exhibited decreased cristae counts and aberrant 304 305 cristae morphology (Fig. 7a,b), including the appearance of round shapes reminiscent of structures detected upon deletion of subunit-g or -e in Saccharomyces cerevisiae¹. These results 306 indicate that monomerization prevents the trypanosomal ATP synthase from assembling into 307 short helical rows on the rims of the discoidal cristae¹⁵, as has been reported for impaired 308 oligomerisation in counterparts from other eukaryotes^{2,34}. 309

- Despite the altered mitochondrial ultrastructure, the subunit-g^{RNAi} cells showed only a very 310 mild growth phenotype, in contrast to all other RNAi cell lines that exhibited steadily slowed 311 growth from day 3 to 4 after the RNAi induction (Fig. 6a, Supplementary Fig. 7a). This is 312 313 consistent with the growth defects observed after the ablation of F_o subunit ATPTB1¹⁹ and F₁ subunits- α and p18¹². Thus, the monomerization of ATP synthase upon subunit-g ablation had 314 only a negligible effect on the fitness of trypanosomes cultured in glucose-rich media, in which 315 ATP production by substrate level phosphorylation partially compensates for compromised 316 oxidative phosphorylation³⁵. 317
- Measurement of oligomycin-sensitive ATP-dependent mitochondrial membrane polarization 318 by safranin O assay in permeabilized cells showed that the proton pumping activity of the ATP 319 synthase in the induced subunit-g^{RNAi} cells is sufficient to generate mitochondrial membrane 320 321 potential, demonstrating that the monomerized enzyme is catalytically functional. By contrast, RNAi downregulation of subunit-8, ATPTB4 and ATPTB11, and ATPTB1 resulted in a strong 322 323 decline of the mitochondrial membrane polarization capacity, consistent with the loss of both monomeric and dimeric ATP synthase forms (Fig. 7c). Accordingly, knockdown of the same 324 325 subunits resulted in inability to produce ATP by oxidative phosphorylation (Fig. 7d). However, upon subunit-g ablation the ATP production was affected only partially, confirming that the 326

327 monomerized ATP synthase remains catalytically active. The ~50 % drop of ATP production 328 in the subunit- g^{RNAi} cells can be attributed to the decreased oxidative phosphorylation 329 efficiency due to the impaired cristae morphology. Indeed, when cells were cultured in the 330 absence of glucose enforcing the need for oxidative phosphorylation, knockdown of subunit-*g* 331 results in a growth arrest, albeit one to two days later than knockdown of all other tested 332 subunits (Fig. 6a). The data show that dimerization is critical when oxidative phosphorylation 333 is the predominant source of ATP.

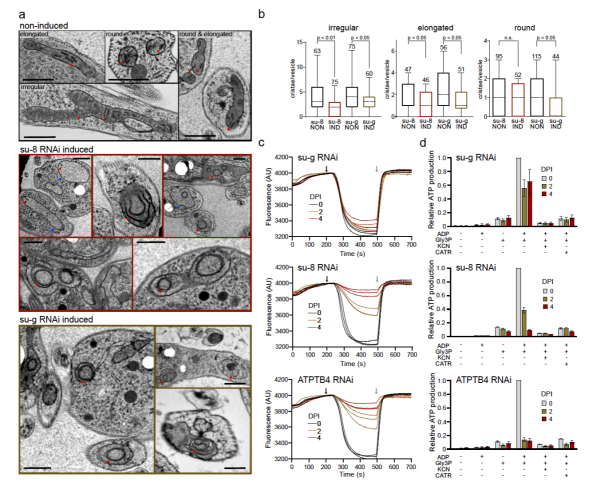
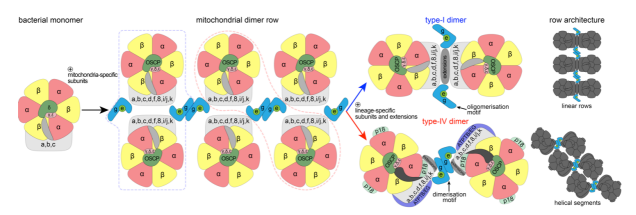


Fig. 7: Monomerization of ATP synthase by subunit-g knockdown results in aberrant 335 mitochondrial ultrastructure but does not abolish catalytic activity. a, Transmission 336 electron micrographs of sections of non-induced or 4 days induced RNAi cell lines. 337 Mitochondrial membranes and cristae are marked with blue and red arrowheads, respectively. 338 Top panel shows examples of irregular, elongated and round cross-sections of mitochondria 339 quantified in (b). b, Cristae numbers per vesicle from indicated induced (IND) or non-induced 340 (NON) cell lines counted separately in irregular, elongated and round mitochondrial cross-341 section. Boxes and whiskers show 25th to 75th and 5th to 95th percentiles, respectively. The 342 numbers of analysed cross-sections are indicated for each data point. c, Safranin O 343 measurement of ability to generate mitochondrial membrane potential in non-induced or 344 345 tetracyline-induced RNAi cell lines 2 and 4 DPI. Black and gray arrow indicate addition of ATP and oligomycin, respectively. d, ATP production in permeabilized non-induced or 346 tetracyline-induced RNAi cells 2 and 4 DPI in the presence of indicated substrates and 347 inhibitors. Error bars represent SD of four replicates. 348

349 Discussion

Our structure of the mitochondrial ATP synthase dimer from the mammalian parasite T. brucei 350 offers new insight into the mechanism of membrane shaping, catalytic rotation, and proton 351 transfer. Considering that trypanosomes belong to an evolutionarily divergent group of 352 353 Kinetoplastida, the ATP synthase dimer has several interesting features that differ from other 354 dimer structures. The subunit-b found in bacterial and other mitochondrial F-type ATP synthases appears to be highly reduced to a single transmembrane helix bH1 represented by a 355 likely homolog ATPTB14. The long bH2, which constitutes the central part of the peripheral 356 357 stalk in other organisms, and is also involved in the composition of the lumenal half proton 358 channel, is completely absent in *T. brucei*. Interestingly, the position of *b*H2 in the proton half channel is occupied by a fully ordered phosphatidylcholine molecule that replaces well-359 conserved proteinaceous element in the proton path. Lack of the canonical bH2 also affects 360 composition of the peripheral stalk in which the divergent subunit-d and subunit-8 binds 361 362 directly to a C-terminal extension of OSCP, indicating a remodeled peripheral stalk architecture. The peripheral stalk contacts the F₁ headpiece at several positions conferring 363 greater conformational flexibility to the ATP synthase. 364

- Using the structural and functional data, we also identified a conserved structural element of 365 366 the ATP synthase that is responsible for its multimerization. Particularly, subunit-g is required for the dimerization, but dispensable for the assembly of the F₁F₀ monomers. Although the 367 monomerized enzyme is catalytically competent, the inability to form dimers results in 368 defective cristae structure, and consequently leads to compromised oxidative phosphorylation 369 370 and cease of proliferation. The cristae-shaping properties of mitochondrial ATP synthase are critical for sufficient ATP production by oxidative phosphorylation, but not for other 371 mitochondrial functions, as demonstrated by the lack of growth phenotype of subunit-g^{RNAi} 372 cells in the presence of glucose. Thus, trypanosomal subunit-g depletion strain represents an 373 374 experimental tool to assess the roles of the enzyme's primary catalytic function and 375 mitochondria-specific membrane-shaping activity, highlighting the importance of the latter for 376 oxidative phosphorylation.
- 377 Based on our data and previously published structures, we propose an ancestral state with 378 double rows of ATP synthase monomers connected by *e/g* modules longitudinally and by other 379 Fo subunits transversally. During the course of evolution, different pairs of adjacent ATP synthase monomer units formed stable dimers in individual lineages (Fig. 8). This gave rise to 380 the highly divergent type-I and type-IV ATP synthase dimers with subunit-*e/g* modules serving 381 382 as dimerization or oligomerization motives, respectively. Because trypanosomes belong to the 383 deep-branching eukaryotic supergroup Discoba, the proposed arrangement might have been present in the last eukaryotic common ancestor. Although sequence similarity of subunit-g is 384 low and restricted to the single transmembrane helix, we found homologs of subunit-g in 385 addition to Opisthokonta and Discoba also in Archaeplastida and Amoebozoa, which represent 386 387 other eukaryotic supergroups, thus supporting the ancestral role in oligomerization (Supplementary Fig. 8). Taken together, our analysis reveals that mitochondrial ATP synthases 388 that display markedly diverged architecture share the ancestral structural module that promotes 389 390 oligomerization.



392

Fig. 8: The subunit-e/g module is an ancestral oligomerization motif of ATP synthase. 393 Schematic model of the evolution of type-I and IV ATP synthases. Mitochondrial ATP 394 synthases are derived from a monomeric complex of proteobacterial origin. In a mitochondrial 395 396 ancestor, acquisition of mitochondria-specific subunits, including the subunit-e/g module resulted in the assembly of ATP synthase double rows, the structural basis for cristae 397 biogenesis. Through divergence, different ATP synthase dimer architectures evolved, with the 398 subunit-e/g module functioning as an oligomerization (type I) or dimerization (type IV) motif, 399 400 resulting in distinct row assemblies between mitochondrial lineages.

401

402 Materials and Methods

403 <u>Cell culture and isolation of mitochondria</u>

404 T. brucei procyclic strains were cultured in SDM-79 medium supplemented with 10% (v/v) 405 fetal bovine serum. For growth curves in no glucose conditions, cells were grown in SDM-80 medium with 10 % dialysed FBS. RNAi cell lines were grown in presence of 2.5 µg/ml 406 phleomycin and 1 µg/ml puromycin. For ATP synthase purification, mitochondria were 407 isolated from the Lister strain 427. Typically, 1.5×10^{11} cells were harvested, washed in 20 mM 408 sodium phosphate buffer pH 7.9 with 150 mM NaCl and 20 mM glucose, resuspended in 409 hypotonic buffer 1 mM Tris-HCl pH 8.0, 1 mM EDTA, and disrupted by 10 strokes in 40 ml 410 Dounce homogenizer. The lysis was stopped by immediate addition of sucrose to 0.25 M. 411 Crude mitochondria were pelleted (15 min at 16,000 xg, 4°C), resuspended in 20 mM Tris-412 HCl pH 8.0, 250 mM sucrose, 5 mM MgCl₂, 0.3 mM CaCl₂ and treated with 5 µg/ml DNase I. 413 414 After 60 min on ice, one volume of the STE buffer (20 mM Tris-HCl pH 8.0, 250 mM sucrose, 2 mM EDTA) was added and mitochondria were pelleted (15 min at 16000 xg, 4°C). The pellet 415 was resuspended in 60% (v/v) Percoll in STE and loaded on six linear 10-35% Percoll gradients 416 in STE in polycarbonate tubes for SW28 rotor (Beckman). Gradients were centrifuged for 1 h 417 418 at 24,000 rpm, 4°C. The middle phase containing mitochondrial vesicles (15-20 ml per tube) 419 was collected, washed twice in the STE buffer, and pellets were snap-frozen in liquid nitrogen 420 and stored at -80°C.

421

422 <u>Plasmid construction and generation of RNAi cell lines</u>

423 To downregulate ATP synthase subunits by RNAi, DNA fragments corresponding to 424 individual target sequences were amplified by PCR from Lister 427 strain genomic DNA using

425 forward and reverse primers extended with restriction sites XhoI&KpnI and XbaI&BamHI, respectively (Supplementary Table 3). Each fragment was inserted into the multiple cloning 426 sites 1 and 2 of pAZ0055 vector, derived from pRP^{HYG-iSL} (courtesy of Sam Alsford) by 427 replacement of hygromycine resistance gene with phleomycine resistance gene, with restriction 428 429 enzymes KpnI/BamHI and XhoI/XbaI, respectively. Resulting constructs with tetracycline inducible T7 polymerase driven RNAi cassettes were linearized with NotI and transfected into 430 a cell line derived from the Lister strain 427 by integration of the SmOx construct for 431 432 expression of T7 polymerase and the tetracycline repressor³⁶ into the β -tubulin locus. RNAi was induced in selected semi-clonal populations by addition of 1 µg/ml tetracycline and the 433 434 downregulation of target mRNAs was verified by quantitative RT-PCR 2- and 4-days post 435 induction. The total RNA isolated by RNeasy Mini Kit (Oiagen) was treated with 2 µg of 436 DNase I, and then reverse transcribed to cDNA with TaqMan Reverse Transcription kit (Applied Biosciences). qPCR reactions were set with Light Cycler 480 SYBR Green I Master 437 438 mix (Roche), 2 µl of cDNA and 0.3 µM primers (Supplementary Table 3), and run on LightCycler 480 (Roche). Relative expression of target genes was calculated using - $\Delta\Delta$ Ct 439 method with 18S rRNA or β-tubulin as endogenous reference genes and normalized to 440 noninduced cells. 441

442

443 Denaturing and blue native polyacrylamide electrophoresis and immunoblotting

444 Whole cell lysates for denaturing sodium dodecyl sulphate polyacrylamide electrophoresis (SDS-PAGE) were prepared from cells resuspended in PBS buffer (10 mM phosphate buffer, 445 130 mM NaCl, pH 7.3) by addition of 3x Laemmli buffer (150 mM Tris pH 6.8, 300 mM 1,4-446 447 dithiothreitol, 6% (w/v) SDS, 30% (w/v) glycerol, 0.02% (w/v) bromophenol blue) to final concentration of 1×10^7 cells in 30 µl. The lysates were boiled at 97°C for 10 min and stored at 448 -20°C. For immunoblotting, lysates from 3x10⁶ cells were separated on 4-20 % gradient Tris-449 glycine polyacrylamide gels (BioRad 4568094), electroblotted onto a PVDF membrane (Pierce 450 88518), and probed with respective antibodies (Supplementary Table 4). Membranes were 451 incubated with the Clarity Western ECL substrate (BioRad 1705060EM) and 452 chemiluminescence was detected on a ChemiDoc instrument (BioRad). Band intensities were 453 quantified densitometrically using the ImageLab software. The levels of individual subunits 454 455 were normalized to the signal of mtHsp70.

456 Blue native PAGE (BN-PAGE) was performed as described earlier¹² with following modifications. Crude mitochondrial vesicles from 2x10⁸ cells were resuspended in 1 M ε-457 aminocaproic acid (ACA) and solubilized with 2% (w/v) dodecylmaltoside (DDM) for 1 h on 458 ice. Lysates were cleared at 16,000 g for 30 min at 4°C and their protein concentration was 459 estimated using bicinchoninic acid assay and diluted to 0.25 μ g/ μ l. 16 μ l of each sample was 460 mixed with 2 µl of 50 % (w/v) glycerol and 1.5 µl of loading dye (500 mM ACA, 5% (w/v) 461 Coomassie Brilliant Blue G-250) and resolved on Native PAGE 3-12% Bis-Tris gels 462 (Invitrogen). After the electrophoresis (3 h, 140 V, 4°C), proteins were transferred by 463 464 electroblotting onto a PVDF membrane (2 h, 100 V, 4°C, stirring), followed by immunodetection with an appropriate antibody (Supplementary Table 4). 465

467

468 <u>Mitochondrial membrane potential measurement</u>

Mitochondrial membrane potential was determined fluorometrically employing safranin O dye 469 (Sigma S2255) in permeabilized cells. For each sample, $2x10^7$ cells were harvested and washed 470 471 with ANT buffer (8 mM KCl, 110 mM K-gluconate, 10 mM NaCl, 10 mM free-acid Hepes, 10 mM K₂HPO₄, 0.015 mM EGTA potassium salt, 10 mM mannitol, 0.5 mg/ml fatty acid-free 472 BSA, 1.5 mM MgCl₂, pH 7.25). The cells were permeabilized by 4 µM digitonin in 2 ml of 473 ANT buffer containing 5 µM safranin O. Fluorescence was recorded for 700 s in a Hitachi F-474 7100 spectrofluorimeter (Hitachi High Technologies) at a 5-Hz acquisition rate, using 495 nm 475 and 585 nm excitation and emission wavelengths, respectively. 1 mM ATP (PanReac 476 AppliChem A1348,0025) and 10 µg/ml oligomycin (Sigma O4876) were added after 200 s and 477 500 s, respectively. Final addition of the uncoupler SF 6847 (250 nM; Enzo Life Sciences 478 479 BML-EI215-0050) served as a control for maximal depolarization. All experiments were performed at room temperature and constant stirring. 480

481

482 <u>ATP production assay</u>

ATP production in digitonin-isolated mitochondria was performed as described previously³⁷. 483 Briefly, 1x10⁸ cells per time point were lysed in SoTE buffer (600 mM sorbitol, 2 mM EDTA, 484 20 mM Tris-HCl, pH 7.75) containing 0.015% (w/v) digitonin for 5 min on ice. After 485 centrifugation (3 min, 4,000 g, 4°C), the soluble cytosolic fraction was discarded and the 486 487 organellar pellet was resuspended in 75 µl of ATP production assay buffer (600 mM sorbitol, 10 mM MgSO₄, 15 mM potassium phosphate buffer pH 7.4, 20 mM Tris-HCl pH 7.4, 2.5 488 mg/ml fatty acid-free BSA). ATP production was induced by addition of 20 mM DL-glycerol 489 490 phosphate (sodium salt) and 67 µM ADP. Control samples were preincubated with the 491 inhibitors potassium cyanide (1 mM) and carboxyatractyloside (6.5 µM) for 10 min at room temperature. After 30 min at room temperature, the reaction was stopped by addition of 1.5 µl 492 493 of 70% perchloric acid. The concentration of ATP was estimated using the Roche ATP Bioluminescence Assay Kit HS II in a Tecan Spark plate reader. The luminescence values of 494 495 the RNAi induced samples were normalized to that of the corresponding noninduced sample.

496

497 <u>Thin sectioning and transmission electron microscopy</u>

The samples were centrifuged and pellet was transferred to the specimen carriers which were 498 completed with 20% BSA and immediately frozen using high pressure freezer Leica EM ICE 499 (Leica Microsystems). Freeze substitution was performed in the presence of 2% osmium 500 tetroxide diluted in 100% acetone at -90°C. After 96 h, specimens were warmed to -20°C at a 501 502 slope 5 °C/h. After next 24 h, the temperature was increased to 3°C (3°C/h). In room 503 temperature, samples were washed in acetone and infiltrated with 25%, 50%, 75% acetone/resin EMbed 812 (EMS) mixture 1 h at each step. Finally, samples were infiltrated in 504 100% resin and polymerized at 60°C for 48h. Ultrathin sections (70 nm) were cut using 505 diamond knife, placed on cupper grids and stained with uranyl acetate and lead citrate. TEM 506 507 micrographs were taken with Mega View III camera (SIS) using a JEOL 1010 TEM operating 508 at an accelerating voltage of 80 kV.

509

510 <u>Purification of *T. brucei* ATP synthase dimers</u>

Mitochondria from 3x10¹¹ cells were lysed by 1 % (w/v) DDM in 60 ml of 20 mM Bis-tris 511 propane pH 8.0 with 10 % glycerol and EDTA-free Complete protease inhibitors (Roche) for 512 513 20 min at 4°C. The lysate was cleared by centrifugation at 30,000g for 20 min at 4°C and adjusted to pH 6.8 by drop-wise addition of 1 M 3-(N-morpholino) propanesulfonic acid pH 514 5.9. Recombinant TbIF₁ without dimerization region, whose affinity to F₁-ATPase was 515 increased by N-terminal truncation and substitution of tyrosine 36 with tryptophan²⁰, with a C-516 terminal glutathione S-transferase (GST) tag (TbIF₁(9-64)-Y36W-GST) was added in 517 approximately 10-fold molar excess over the estimated content of ATP synthase. Binding of 518 TbIF₁ was facilitated by addition of neutralized 2 mM ATP with 4 mM magnesium sulphate. 519 After 5 min, sodium chloride was added to 100 mM, the lysate was filtered through a 0.2 µm 520 521 syringe filter and immediately loaded on 5 ml GSTrap HP column (Cytiva) equilibrated in 20 mM Bis-Tris-Propane pH 6.8 binding buffer containing 0.1 % (w/v) glyco-diosgenin (GDN; 522 Avanti Polar Lipids), 10 % (v/v) glycerol, 100 mM sodium chloride, 1 mM tris(2-523 carboxyethyl)phosphine (TCEP), 1 mM ATP, 2 mM magnesium sulphate, 15 ug/ml 524 cardiolipin, 50 ug/ml 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), 25 ug/ml 1-525 palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (POPE) and 10 ug/ml 1-palmitoyl-2-526 oleoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (POPG). All phospholipids were purchased 527 from Avanti Polar Lipids (catalog numbers 840012C, 850457C, 850757C and 840757, 528 529 respectively). ATP synthase was eluted with a gradient of 20 mM reduced glutathione in Tris pH 8.0 buffer containing the same components as the binding buffer. Fractions containing ATP 530 531 synthase were pooled and concentrated to 150 µl on Vivaspin centrifugal concentrator with 30 kDa molecular weight cut-off. The sample was fractionated by size exclusion chromatography 532 on a Superose 6 Increase 3.2/300 GL column (Cytiva) equilibrated in a buffer containing 20 533 534 mM Tris pH 8.0, 100 mM sodium chloride, 2 mM magnesium chloride, 0.1 % (w/v) GDN, 3.75 ug/ml cardiolipin, 12.5 ug/ml POPC, 6.25 ug/ml POPE and 2.5 ug/ml POPG at 0.03 535 536 ml/min. Fractions corresponding to ATP synthase were pooled, supplemented with 0.05% (w/v) DDM that we and others experimentally found to better preserve dimer assemblies in 537 cryo-EM³⁸, and concentrated to 50 ul. 538

539

540 Preparation of cryo-EM grids and data collection

541 Samples were vitrified on glow-discharged Quantifoil R1.2/1.3 Au 300-mesh grids after 542 blotting for 3 s by plunging into liquid ethane using a Vitrobot Mark IV. 5,199 movies were 543 collected using EPU 1.9 on a Titan Krios (ThermoFisher Scientific) operated at 300 kV at a 544 nominal magnification of 165 kx (0.83 Å/pixel) with a Quantum K2 camera (Gatan) using a 545 slit width of 20 eV. Data was collected with an exposure rate of 3.6 electrons/px/s, a total 546 exposure of 33 electrons/Å² and 20 frames per movie.

547

548 Image processing

549 Image processing was performed within the Scipion 2 framework³⁹, using RELION-3.0 unless 550 specified otherwise. Movies were motion-corrected using the RELION implementation of the

MotionCor2. 294,054 particles were initially picked using reference-based picking in 551 Gautomatch (http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch) and Contrast-transfer 552 function parameters were using GCTF⁴⁰. Subsequent image processing was performed in 553 RELION-3.0 and 2D and 3D classification was used to select 100,605 good particles, which 554 were then extracted in an unbinned 560-pixel box (Fig. S1). An initial model of the ATP 555 synthase dimer was obtained using *de novo* 3D model generation. Using masked refinement 556 with applied C₂ symmetry, a 2.7-Å structure of the membrane region was obtained following 557 per-particle CTF refinement and Bayesian polishing. Following C2-symmetry expansion and 558 signal subtraction of one monomer, a 3.7 Å of the peripheral stalk was obtained. Using 3D 559 classification (T=100) of aligned particles, with a mask on the $F_{1/c}$ -ring region, 10 different 560 rotational substates were then separated and maps at 3.5-4.3 Å resolution were obtained using 561 3D refinement. The authors note that the number of classes identified in this study likely 562 reflects the limited number of particles, rather than the complete conformational space of the 563 564 complex. By combining particles from all states belonging to main rotational state 1, a 3.7-Å map of the rotor and a 3.2-Å consensus map of the complete ATP synthase dimer with both 565 rotors in main rotational state 1. 566

567

568 Model building, refinement and data visualisation

An initial atomic model of the static F_o membrane region was automatically built using 569 Bucaneer⁴¹. Subunits were subsequently assigned directly from the cryo-EM map, 15 of them 570 corresponding to previously identified *T. brucei* ATP synthase subunits²¹, while three subunits 571 572 (ATPTB14, ATPEG3, ATPEG4) were newly identified using BLAST searches. Manual model 573 building was performed in *Coot* using the *T. brucei* F₁ (PDB 6F5D) and homology models⁴² of the E. gracilis OSCP and c-ring (PDB 6TDU) as starting models. Ligands were manually fitted 574 575 to the map and restraints were generated by the GRADE server (http://grade.globalphasing.org) Real-space refinement was performed in PHENIX using auto-sharpened, local-resolution-576 filtered maps of the membrane region, peripheral stalk tip, c-ring/central stalk and F_1F_0 577 monomers in different rotational states, using secondary structure restraints. Model statistics 578 were generated using MolProbity and EMRinger. Finally, the respective refined models were 579 combined into a composite ATP synthase dimer model and real-space refined against the local-580 resolution-filtered consensus ATP synthase dimer map with both monomers in rotational state 581 1, applying reference restraints. Figures of the structures were prepared using Chimera X^{43} , the 582 proton half channels were traced using HOLLOW⁴⁴. 583

584

585 Data availability

The atomic coordinates have been deposited in the Protein Data Bank (PDB) and are available 586 under the accession codes: XXX (membrane-region), XXX (peripheral stalk), XXX (rotor), 587 588 XXX (F1Fo dimer), XXX (rotational state 1a), XXX (rotational state 1b), XXX (rotational 589 state 1c), XXX (rotational state 1d), XXX (rotational state 1e), XXX (rotational state 2a), XXX (rotational state 2b), XXX (rotational state 2c), XXX (rotational state 2d), XXX (rotational 590 state 3). The local resolution filtered cryo-EM maps, half maps, masks and FSC-curves have 591 been deposited in the Electron Microscopy Data Bank with the accession codes: EMD-XXX 592 593 (membrane-region), EMD-XXX (peripheral stalk), EMD-XXX (rotor), EMD-XXX (F₁F₀

594 dimer), EMD-XXX (rotational state 1a), EMD-XXX (rotational state 1b), EMD-XXX

- 595 (rotational state 1c), EMD-XXX (rotational state 1d), EMD-XXX (rotational state 1e), EMD-
- 596 XXX (rotational state 2a), EMD-XXX (rotational state 2b), EMD-XXX (rotational state 2c),
- 597 EMD-XXX (rotational state 2d), EMD-XXX (rotational state 3).
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724 Author contributions

- 725 O.G. prepared the sample for cryo-EM. O.G. and A.M. performed initial screening. A.M.
- processed the cryo-EM data and built the model. B.P., C.H.Y., M.J., M.S., O.G. and A.Z.
- performed biochemical analysis. O.G., A.M. and A.A. analyzed the structure. O.G., A.M., A.A.
- and A.Z. wrote the manuscript. All authors contributed to the analysis and the final version of
- the manuscript.

730	SUPPLEMENTARY INFORMATION
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732	Supplementary Videos:
733	
734	Supplementary Video 1. Overall structure and subunit-e/g module of trypanosomal ATP
735	synthase dimers. Subunits are coloured as in Figure 1. Phospholipids and ligands are shown
736	as sticks.
737	
738	Supplementary Video 2. Rotary cycle of T. brucei ATP synthase. Top view showing
739	flexibility of the peripheral stalk, including bending of its apical part, and rotational and tilting
740	motions of F_1 .
741	
742	Supplementary Video 3. Rotary cycle of T. brucei ATP synthase. Side view showing
743	rotational and tilting motions of the F_1/c -ring subcomplex.
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764 Supplementary Tables:

766 Supplementary Table 1. List of models and refinement statistics

	Mem- brane region	Rotor	Periphe- ral stalk	F1F0 dimer	Rot. 1a	Rot. 1b	Rot. 1c	Rot. 1d	Rot. le	Rot. 2a	Rot. 2b	Rot. 2c	Rot. 2d	Rot. 3
Data collection		•					1	1	ł	ł	ł		1	
Microscope							Titan	Krios						
Voltage (kV)							3	00						
Camera							K2 S	ummit						
Magnification							16	5 kx						
Exposure (e ⁻ /Å ²)							3	33						
Defocus range (µm)							-1.6 t	to -3.2						
Pixel size (Å)							0.	.83						
Movies collected							5,1	199						
Frames per movie							. 2	20						
Data processing														
Initial particles						100,605 (C ₂ symmetr	y-expanded	201,210)					
Final no. particles	100,605	118,683	201,210	36,925	19,764	26,427	23,019	16,991	34,482	12,173	24,096	11,035	17,833	17.312
Symmetry	C ₂	C_1	C_1	C ₂	C 1	C 1	C 1	Cı	C 1	C_1	\mathbf{C}_1	C 1	C 1	\mathbf{C}_1
Map resolution (Å)	2.7	3.7	3.7	3.2	3.7	3.5	3.7	3.8	3.7	4.3	3.5	3.8	3.8	3.7
Sharpening B factor	-46.2	-74.4	-92.5	-49.8	-61.8	-61.1	-57.6	-45.6	-58.0	-73.8	-54.5	-65.2	-54.9	-61.7
EMD ID											-	-		
Model refinement statistics	-													
CC (map/model)	0.86	0.83	0.82	0.71	0.79	0.79	0.82	0.79	0.69	0.71	0.81	0.77	0.77	0.79
Resolution (map/model)	2.65	3.4	3.68	3.13	3.48	3.56	3.36	3.55	3.57	3.94	3.39	3.73	3.64	3.58
No. of atoms	76,690	19,669	12,083	251,552	129,568	129,568	129,568	129,568	129,568	129,563	129,563	129,563	129,563	129,56
No. of residues	4074	1285	767	15,356	7872	7872	7872	7872	7872	7872	7872	7872	7872	7872
No. of lipids	36	0	0	36	21	21	21	21	21	21	21	21	21	21
No. of ATP/ADP	0	0	0	10	5	5	5	5	5	5	5	5	5	5
No. of Mg ions	0	0	0	10	5	5	5	5	5	5	5	5	5	5
B-factor (Ų)														
- protein	54.05	56.13	77.88	84.48	55.65	70.37	80.22	83.27	70.70	112,72	79.93	65.52	66.49	101.5
- ligands	50.57	58.25	-	69.94	40.99	72.29	63.18	78.43	63.76	75,25	74.47	61.79	46.55	83.68
Rotamer outliers (%)	0.44	0.40	0.31	0.22	0.42	0.09	0.18	0.26	0.58	0.18	0.27	0.48	0.42	0.39
Ramachandran (%)														
- outliers	0.00	0.00	0.00	0.01	0.001	0.003	0.004	0.01	0.003	0.01	0.00	0.04	0.04	0.04
- allowed	1.57	1.91	1.59	1.56	1.52	1.65	1.44	1.49	1.49	1.67	1.58	1.47	1.65	1.79
- favored	98.43	98.08	98.41	98.42	98.47	98.34	98.56	98.49	98.48	98.31	98.42	98.49	98.31	98.17
Clash score	1.66	2.44	2.32	2.26	2.60	2.65	2.53	2.67	2.99	2.38	2.30	2.52	2.38	3.57
MolProbity score	0.92	1.03	1.01	1.00	1.05	1.05	1.04	1.05	1.09	1.02	1.01	1.04	1.02	1.15
RMSD														
- bonds (Å)	0.004	0.004	0.02	0.003	0.003	0.003	0.004	0.003	0.003	0.002	0.003	0.003	0.003	0.003
- angles (°)	0.455	0.416	0.386	0.407	0.414	0.424	0.417	0.407	0.412	0.410	0.416	0.419	0.428	0.421
EMRinger score	5.11	3.96	1.61	2.56	3.24	2.95	3.32	2.85	3.32	1.35	2.89	2.32	2.49	2.8
and a score	5.11	3.90	1.01	2.50	3.24	2.95	3.32	2.00	3.32	1.55	2.09	2.32	2.49	2.0

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770 Supplementary Table 2. Composition of *T. brucei* ATP synthase dimer

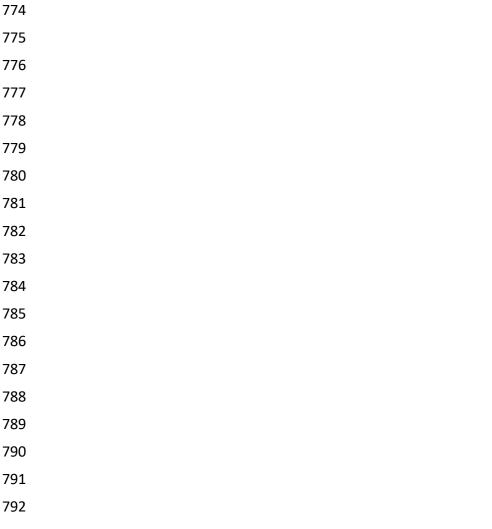
Subunit	TriTrypDB Lister	TriTrypDB	Uniprot	Residues	Residues
name	strain 427 ID	TREU927 strain ID	TREU927 strain		built
			ID		
	1	F1 subcomplex			
α	Tb427_070081800	Tb927.7.7420	Q57TX9	584	45-151,
	Tb427_070081900	Tb927.7.7430			161-584
β	Tb427_030013500	Tb927.3.1380	Q57XX1	519	26-514
γ	Tb427_100005200	Tb927.10.180	B0Z0F6	305	2-301
δ	Tb427_060054900	Tb927.6.4990	Q586H1	182	22-182
Е	Tb427_100054600	Tb427.10.5050	N/A	75	11-75
p18	Tb427_050022900	Tb927.5.1710	Q57ZP0	188	23-188
-		F ₀ subcomplex			
OSCP	Tb427 100087100	Tb927.10.8030	Q38AG1	255	18-202,
	_				208-255
a	mt encoded	mt encoded	N/A	231	1-231
с	Tb427 100018700	Tb927.10.1570	Q38C84 Q385P0	118	41-118
		ТЬ927.11.5280	Q57WQ3		
	Tb427_070019000	Tb927.7.1470			
d	Tb427_050035800	Tb927.5.2930	Q57ZW9	370	17-325,
					332-354
e	Tb427_110010200	Tb927.11.600	N/A	92	1-383
f	Tb427_030016600	Tb927.3.1690	Q57ZE2	145	2-136
g	Tb427_020016900	Tb927.2.3610	Q586X8	144	16-144
i/j	Tb427_030029400	Tb927.3.2880	Q57ZM4	104	2-104
k	Tb427_070011800	Tb927.7.840	Q57VT0	124	20-124
8	Tb427_040037300	Tb927.4.3450	Q585K5	114	29-114
ATBTB1	Tb427_100008400	Tb927.10.520	Q38CI8	396	1-383
ATPTB3	Tb427_110067400	Tb927.11.6250	<u>Q385E4</u>	269	2-269
ATPTB4	Tb427_100105100	Tb927.10.9830	Q389Z3	157	21-157
ATPTB6	Tb427_110017200	Tb927.11.1270	<u>Q387C5</u>	169	2-169
ATPTB11	Tb427_030021500	Tb927.3.2180	Q582T1	156	18-156
ATPTB12	Tb427_050037400	Tb927.5.3090	Q57Z84	101	5-100
ATPTB14	Tb427_040009100	Tb927.4.720	Q580A0	105	26-105
ATPEG3	Tb427_060009300	Tb927.6.590	Q583U4	98	14-98
ATPEG4	N/A	Tb927.11.2245	N/A	62	1-62

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773 Supplementary Table 3. List of oligonucleotides

Primers for amplification of RNAi cassettes e TAATCTCGAGGGTACCGGGAGTACAGAAGGGCTACA TAGATCTAGAGGATCCCGTGACACCATCAGCAG GCGTCTAGAGGATCCAGCACCGACCATCAGCAGCAG GCGTCTAGAGGATCCAGCAGCAGCACCAAACTGC f ATACTCGAGGGTACCAGCAGGAGTACACAAACTGC g ACTGCTCGAGGGTACCAGCAGGAGTACACAAACTGC g ACTGCTCGAGGGTACCAGCGGGGAATTCAAAAGACC g GCGGTCTAGAGGATCCAGTGCGGGGAATTCAAAAGACC g TAATCTCGAGGGTACCGACATGCGGGGATCATGTCACTGCATGA ij TAATCTCGAGGGTACCGGAATATCCGATGCAGCGC g GCGGTCTAGAGGATCCACTTCGCTGACAGGGGGATTTT gCCGTCTAGAGGATCCGGGCATCAGTGCAGGGGAGTTTT GCCGTCTAGAGGATCCGGGCTATGGTGGTGTATTATGC gACGTCTAGAGGATCCCGCGCAAGAAAACTCCCAACGACA ATGACTCGAAGGGTACCGAGGAAAACTCCCAACGACA ATPTB3 ACTGCTCGAGGGTACCCAAAGAGGAGGTGAGGTCTGC GCAGTCTAGAGGATCCCCCTAGGGTCTCCCAATTG GCAGTCTAGAGGATCCCCTCAGGGTTCCAATTG GCAGTCTAGAGGATCCCCTCACGGGCTTCCAATTG GCAGTCTAGAGGATCCCCCTCACGGCGCTTCCCATTTC ATPTB4 ACTGCTCGAGGGTACCAGCGCCCTCACACAGGAGGG ATPTB11 ACTGCTCGAGGGTACCAGCGCCCTCACACATTGCGGGGTGTTAGGGAGGAACGGG ATPTB12 TAATCTCGAGGGTACCAGCGCCACCACACATTACGCGCCACAC ATPTB14 TAATCTCGAGGGTACCAGCGCCTCCACACACACACACACA	Subunit	Primer pair sequences
eTAGATCTAGAGGATCCCGTGCACACCATCAGCTGfATACTCGAGGGTACCGTGAGTACCGCCTTACGCgGCGTCTAGAGGATCCAGCACGCGGAATTCAAAAGACCgGCGGTCTAGAGGATCCCGTTGCGGTGCTGTCATTAijTAATCTCGAGGGTACCACGCGGGAATTCAAAAGACCgGCCGTCTAGAGGATCCCGTTGCGATGCAGGGCGATCAGTGCAGGGAkATTACTCGAGGGTACCGGCGATCAGTGCAGGGGATTTgGCCGTCTAGAGGATCCGCGGCGATCAGTGCAGGGGATTGCkATTACTCGAGGGTACCGGGCTATGGTGGTGGTATTATGCGCCGTCTAGAGGATCCGCAGAAAACTCCCAACGACAAACGCTCGAGGGTACCGGGCTATGGTGGTGGTATTATGCGCGTCTAGAGGATCCCCCAAGAGGAGGTGAGGAATPTB3GCGTCTAGAGGATCCCCCAAGAGGAGGTGAGGAATPTB4GCAGTCTAGAGGATCCCCCAACATGGCAGTACCGGGGATPTB6GCAGTCTAGAGGATCCCAACATGGCAGTATCCGGTGGCAGTCTAGAGGATCCCAGCGCCTCCAACATGGCAGGAGGAGAATPTB1ACTGCTCGAGGGTACCCAACATGGCAGGATGCCGCAGACTTGGAGGATCCCAGGCGCCATCAAAGGAAGACAATPTB12GCAGTCTAGAGGATCCAGCGCCACAAAAACAGAACAATPTB13ACTGCTCGAGGGTACCAACGAGGAGGAACGGGGCAGTCTAGAGGATCCCAGCGCCACAAAAAAAACGAACAGATPTB14GCAGTCTAGAGGATCCAGCGAGGAACAGGAATPTB15TAATCTCGAGGGTACCAACCGAGGAGGAACGGGGCAGTCTAGAGGATCCCACCACCCTTCTCGGCCGCCTGATAfTAACCTCGAGGGTACCAAACCTGAAGGCACCACCfTAACCTCGAGGGTACCACACCTTTATGcCCAGCCTAGAGGATCCCCTCTTCGGCCGCCTGACAgCCAAGCCTTGCACAACCTTTATGgGCAATTGTGTGAGCTGAACGfTTTCTACATACCGCAGCACCfTACCGGCCCACGCATGCAACGjGCAATTGTGTGAGCTGAACGgGCAATTGTGTGAGCTGAACGgGCAATTGCTGGAGA		Primers for amplification of RNAi cassettes
TAGATCTAGAGGATCCCGTGCACACCATCAGCTG f ATACTCGAGGGTACCGTGAGTACCGCCATCAGCACACCATCAGC g ACTGCTCGAGGGTACCAGCAGGAATTCAAAAGACC g ACTGCTCGAGGGTACCAGCGGGAATTCAAAAGACC g GCGGTCTAGAGGATCCCGTGCGGTGCTTGTCATTA <i>ij</i> TAATCCGAGGGTACCGAATATCCGATGCAGCGCG <i>ij</i> TAATCCGAGGGTACCGGGCGATCAGTGCAGGGGGATTTT <i>k</i> ATTACTCGAGGGTACCGGGCGATCAGTGCAGGGGGATTTT GCCGTCTAGAGGATCCTTCCTCGAAAACGCACACA ATTACTCGAGGGTACCGGGCATAGGTGGGGGTATTAGC <i>k</i> ATGACTCGAGGGTACCGGGCAAAAACTCCCAACGACA <i>k</i> ACTGCTCGAGGGTACCGCGAGAAAACTCCCAACGACA <i>ATPTBB</i> ACTGCTCGAGGGTACCTCCTTTTTTCGCGCACCGG <i>GCAGTCTAGAGGATCCCCCAACGGGTGGTGGTCGC</i> GCAGTCTAGAGGATCCTTCTTTTTTTCGCTGCATCGG <i>ATPTBB</i> ACTGCTCGAGGGTACCCAACATGGCAGTATCCGGTG <i>ATPTBB</i> ACTGCTCGAGGGTACCGACGCCTCGTCTTCTCCATTTC <i>GCAGTCTAGAGGATCCCTCACGCGCCTGTCTTTAGGGAGG</i> GCAGTCTAGAGGATCCCAACAACAGGAACGGG <i>ATPTBB</i> ACTGCTCGAGGGTACCGACGCCAACAAAAGGACAA <i>ATPTBB</i> ACTGCTCGAGGGTACCGACGCCACACAAAAGGACAA <i>ATPTBB</i> ACTGCTCGAGGGTACCGACGCCACCACACATTAGGGGAGGTGGTGCGT <i>GCAGTCTAGAGGATCCCAACAGGCCAACAAACGGCAGGCGGCCCACCACACAACA</i>	a	TAATCTCGAGGGTACCGGGAGTACAGAAGGGCTACA
fGCGTCTAGAGGATCCAGCACTGATCACCAAACTGCgACTGCTCGAGGGTACCAGCGGGAATTCAAAAGACC GCGGTCTAGAGGATCCGATGCGGGGATCAATACCGATGCATGAA GCGTCTAGAGGATCCGATAATCCGATGCATGCAA ATTACTCGAGGGTACCAGGGCGATCAGTGCAGGGGAATTTT GCCGTCTAGAGGATCCTTCCTCAAAAACGCAACAAkATTACTCGAGGGTACCGGCGATCAGTGCAGGGGAATTATOCC GCAGTCTAGAGGATCCGCAGAAAACTCCCAACGACAAAATGACTCGAGGGTACCGGCGATCAGTGCAGGGGAATTATOCC GACGTCTAGAGGATCCGCAGAAAAACTCCCAACGACAAATPTB8ACTGCTCGAGGGTACCGAGAAAACTCCCAACGGAGAATPTB8ACTGCTCGAGGGTACCCAAAGAGGAGGTGAGGTCTGC GCAGTCTAGAGGATCCCTCTCTGGGGTTCTCGAAGCAATPTB84CTGACTCGAGGGTACCCAACATGGCAGTATCCGGTG GCAGTCTAGAGGATCCCTCTTCTCCAATTTG GCAGAAGCTTGGATCGAGGGTACCCAACATGGCAGTATCCGGTG 	e	TAGATCTAGAGGATCCCGTGCACACCATCAGCTG
GCGTCTAGAGGATCCAGCACTGATCACCCAAACTGC g ACTGCTCGAGGGTACCACGCGGGAATTCAAAAGACC gCGGGTCTAGAGGATCCCGTTGCGGTGCTTGTCATTA ij TAATCTCGAGGGTACCGAATATCCGATGCATGCCGC gCCGTCTAGAGGATCCACTTCGCTCTACTGCATGCA k ATTACTCGAGGGTACCGAGGGATCAGTGCAGGGGAATTTT GCGTCTAGAGGGTACCGGGCGATCAGTGCAGGGGGATTTT GCCGTCTAGAGGGTACCGGGCGATCAGTGGCAGGGGGAATTATGC gA ATTACTCGAGGGTACCGGGCATAGGTGTGGTATTATGC GACGTCTAGAGGGTACCGGGCTATGGTGTGGTATTATGC GCAGTCTAGAGGGTACCGCGCAGAAAACTCCCAACGACA ATPTB3 ACTGCTCGAGGGTACCCACAAGAGGGAGGTGAGGTCTGC GCAGTCTAGAGGGTACCCCCCTAGGGTTCTCCAATTG GCAGTCTAGAGGGTACCCACACATGGCAGTATCCGGTG GCAGTCTAGAGGGTACCCACACATGGCGGTGTTCCAATTG GCAGTCTAGAGGGTACCGCGCTCGTCTTCTCCATTTC GCAGTCTAGAGGGTACCGACGCCTGCTTTTCCCATTTC GCAGTCTAGAGGGTACCGACGCCTCGTCTTCCCATTCC ATPTB11 ACTGCTCGAGGGTACCGACGCCCCCCACCACAAAAGGAAGG	£	ATACTCGAGGGTACCGTGAGTACCGCCTTTACGC
gGCGGTCTAGAGGATCCCGTTGCGGTGCTTGTCATTAijTAATCTCGAGGGTACCGAATATCCGATGCATGCGCgGCCGTCTAGAGGATCCACTTCGCTCACTGCATGCATGCAkATTACTCGAGCCCGGGCGATCAGTGCAGGGGATTTTgGCCGTCTAGAGGGTACCGGGCTATGGTGGTGTTTTATGCgATGACTCGAGGGTACCGGGGCTATGGTGGTATTATGCgGACGTCTAGAGGATCCGCAGAAAACTCCCAACGACAATPTB3ATGCTCGAGGGTACCAAGAGGAGGGGAGGTGAGGTCGCGCAGTCTAGAGGATCCCCCTAGGGTTCTCGAAGGAATPTB4CTGACTCGAGGGTACCCAACATGGCAGTATCCGGGGCAGTCTAGAGGATCCCTCCTCGGGCTTCCCAATTTGGCAGTCTAGAGGATCCCCCCACGGCTGCTCCCAATTGGCAGTCTAGAGGATCCCCCCCCGGGCTGCGTGGTGGTATPTB6ACTGCTCGAGGGTACCCAACATGGCAGTATCCGGTGGCAGAAGCTTGGATCCAGGTGGGGGGTGTTAGGGAGGATPTB11ACTGCTCGAGGGTACCCACGCCAACAAACAGACAAATPTB12CTAATCTCGAGGGTACCAGCGCCATCAAAGGAATGCCGCAGTCTAGAGGATCCAGCGCCAACAAACAGACAAATPTB14TAATCTCGAGGGTACCAGCGCCAACAAACAGGCAGGGCAGTCTAGAGGATCCAGCTGAGGAGGAACCGGGGCAGTCTAGAGGATCCACTTTCCTTCCACCCACCGCAGTCTAGAGGATCCACTTACCCTTCCACCCACCGCAGTCTAGAGGATCCACTTACCCTCCACCCACCGCAGTCTAGAGGATCCCACTTACCCTCCACCCACCGCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAGCAGCTTAGAAGGATCCCCTCTTCCGACCACCfTACACTCGAGGGTACCAACACTTTATGCCCGCAAAGAAGTACGCCACfTTTTCTACATACCGCAGCAGTfTACCATTGCATGCGCGCATGCAACGgGCAATTGTGTGAGCTGAACGgGCAATTGTGTGAGCTGAACGgCAAGGCTTGCAATAACCGGTGGTGCgCAAGTGCGCCCATTGCATAACgCAAGTTAAAAAGCGCGCCTACGgCAAGTTGGAAAA	J	GCGTCTAGAGGATCCAGCACTGATCACCAAACTGC
ij TAATCTCGAGGGTACCGAATATCCGATGCATGCAGC ij TAATCTCGAGGGTACCGAATATCCGATGCATGCAG k ATTACTCGAGGGTACCGGGCGATCAGTGCAGGGGATTTT GCGGTCTAGAGGGTACCGGGCGATCAGTGCAGGGGGATTATGC ATTACTCGAGGGTACCGGGCGATGGTGGTGGTATTATGC gCGGTCTAGAGGGTACCGGGCTATGGTGTGGTATTATGC GACGTCTAGAGGGTACCGAGGGTAGGGTGGG ATPTB3 ACTGCTCGAGGGTACCAAAGAGGAGGTGAGGTCTGC GCAGTCTAGAGGATCCCCCTAGGGTTCTCGAAGCA CTGACTCGAGGGTACCCACAAGGGAGGTGAGGTCTGC ATPTB4 CTGACTCGAGGGTACCCACACAGGGGTGCGTGGT ATPTB6 CTGACTCGAGGGTACCCAACATGGCAGTATCCGGTG GCAGTCTAGAGGATCCCTCCTTCTGCGTGTGTTCCCATTTC GCAGTCTAGAGGGTACCGACGCCTCGTCTTCCCATTTC GCAGTCTAGAGGGTACCGACGCCTCGTCTTCCCATTC GCAGGCTTGGAGGTACCGACGCCAACAAACAGAAGG ATPTB16 ACTGCTCGAGGGTACCGACGCCAACAAACAGAACAGA ATPTB11 ACTGCTCGAGGGTACCGACGCCAACAAACAGAACAG ATPTB12 TAATCTCGAGGGTACCGACGCCAACAAACAGGAACGGG GCCGTCTAGAGGGTACCAACGCCACCACACAACAAGAGGAACGGG GCAGTCTAGAGGATCCTATCCCTTCCACCACACACACACA	a	ACTGCTCGAGGGTACCACGCGGGAATTCAAAAGACC
i/jGCCGTCTAGAGGATCCACTTCGCTCACTGCATGCAkATTACTCGAGCCCGGGCGATCAGTGCAGGGGATTTT GCCGTCTAGAGGATCCTTCCTCGAAAAACGCACACA8ATGACTCGAGGGTACCGGGCTATGGTGTGGTATTATGC GACGTCTAGAGGATCCGCAGAAAACTCCCAACGACAATPTB3ACTGCTCGAGGGTACCGAGAAAACTCCCAACGAGAATPTB4CTGACTCGAGGGTACCAAAGAGGAGGTGAGGTCGC GCAGTCTAGAGGATCCCCCTAGGGTTCTCGAAGCAATPTB6CTGACTCGAGGGTACCAACATGGCAGTATCCGGTG GCAGTCTAGAGGGTACCAACATGGCAGTATCCGGTG GCAGTCTAGAGGGTACCAACAGGCAGCAACAAACGGCAGATPTB11ACTGCTCGAGGGTACCAACAGGCAGTATCCGGTG GCAGTCTAGAGGATCCTGCTGTCTCTCCATTTC GCAGAAGCTTGGATCCAGGTGGGGTGTTAGGGAGATPTB11ACTGCTCGAGGGTACCAACAGGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCAGCGCCATCAAAGGAATGCC GCCGTCTAGAGGATCCAGCAGCCAACAAACAGACAAATPTB12TAATCTCGAGGGTACCAGCAGCCAACAAACAGACAAATPTB13TAATCTCGAGGGTACCAACGGAGGGAACGGG GCAGTCTAGAGGATCCAGCAGCCAACAAACAGACAAATPTB14TAATCTCGAGGGTACCAAACCTGAAGGCCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAeCCAAGCCTTGCACACACTTTATG CCGCAAAGAAGTACGCCACfTAACACTCGAGGGTACCATCCACCACACTTTATG GCAGTCTAGAGGATCCCTCTTCCACCCACCfTACCATTCCATGCGCGATGfTACCATTCCATGCGCGCAGT TACCATTCCATGCGCGCTGATACgGCAATTGTGTGAGCTGAACG TACCGCGCATTGCATAACi/jAGAGTAAAAGCGCGCCTACG CAGTTGGAAAACCGGTAGCC	8	GCGGTCTAGAGGATCCCGTTGCGGTGCTTGTCATTA
GCCGTCTAGAGGATCCACTTCGCTTACTGCATGCAkATTACTCGAGGCCCGGGCGATCAGTGCAGGGGATTTT GCCGTCTAGAGGATCCTTTCCTCGAAAAACGCACACA8ATGACTCGAGGGTACCGGGCTATGGTGGGTATTATGC GACGTCTAGAGGGTACCACAGAGAAAACTCCCAACGACAATPTB3CTGACTCGAGGGTACCAAAGAGGAGGTGAGGTCGC GCAGTCTAGAGGGTACCCTAGGGTTCTTCGAAGCAATPTB4CTGACTCGAGGGTACCAAAGAGGAGGTGAGGTCGC GCAGTCTAGAGGGTACCCACACAGGGCTGCTCCAATTTG ACTGCTCGAGGGTACCCAACATGGCAGTATCCGGTG GCAGTCTAGAGGATCCTCTTATTAGTGGCGGTGGTGGTATPTB6ACTGCTCGAGGGTACCAACATGGCAGTATCCGGTG GCAGTCTAGAGGATCCAGCGCCTCGTCTTCTCCATTTC GCAGAAGCTTGGATCCAGGGTGGGGGTGTTAGGGAG GCAGTCTAGAGGATCCAGCAGCCAACAAACAGACAAATPTB11ACTGCTCGAGGGTACCGACGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCGACGCCATCAAAAGGAAAGCC GCCGTCTAGAGGGTACCAGCAGCCAACAAACAGAACAAATPTB12TAATCTCGAGGGTACCAGCGCCATCAAAGGAAACGG GCAGTCTAGAGGATCCAGCAGCCAACAAACAGAACAAATPTB14TAATCTCGAGGGTACCAGCGCCACCACC GCAGTCTAGAGGATCCTATCCCTTCCACCCACCACC GCAGTCTAGAGGATCCTATCCCTTCCACCCACCACC GCAGTCTAGAGGATCCCACCTGAAGAGGAACGGC GCAGTCTAGAGGATCCCACCTGTACACCGAGAGGAACGGC GCAGTCTAGAGGATCCCTCTTCCATGCCGCCTGATAeCCAAGCCTTGCACACACTTTATG CCGCCAAAGAAGTACGCCACfTACCATTCCATGCGCAGGAGAACGG TACCATTCCATGCGCGCATGAAACgGCAATTGTGTGAGCTGAACG TACCATTCCATGCGCGCTTGGAGAGGAACGG GCAATTGTGTGAGCTGAACGjAGAGTAAAAGCGCGCCTACG CAGTTGGAAAACCGGTAGCC	;/;	TAATCTCGAGGGTACCGAATATCCGATGCATGCCGC
kGCCGTCTAGAGGATCCTTTCCTCGAAAAACGCACACA8ATGACTCGAGGGTACCGGGCTATGGTGTGGTATTATGC GACGTCTAGAGGATCCGCAGAAAACTCCCAACGACAATPTB3ACTGCTCGAGGGTACCAAAGAGGAGGTGAGGTCGC GCAGTCTAGAGGATCCCCCTAGGGTTCTTCGAAGCAATPTB4CTGACTCGAGGGTACCACAACATGGCAGTATCCGGTG GCAGTCTAGAGGATCCTCCTCTGGGCGTCCAATTTG GCAGTCTAGAGGGTACCCAACATGGCCAGTATCCGGTG GCAGTCTAGAGGGTACCGACGTCGTCTTCTCCATTTC GCAGAAGCTTGGATCCAGGGTACCGACGCGTCGTCTTCCCATTTC GCAGAAGCTTGGATCCAGGCGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCGACGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCGACGCCAACAAACAGACAAATPTB11ACTGCTCGAGGGTACCGACGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCGACGCCATCAAAGGAATGCC GCCGTCTAGAGGGTACCAGCGCCAACAAACAGACAAATPTB12TAATCTCGAGGGTACCAGCGCCAACAAACAGACAA GCAGTCTAGAGGATCCTTTCTCGTGCCGCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAATPTB14CAAGCCTTGCACAAACCTGAAGGCCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAeCAAGCCTTGCACAAACCTGAAGGCCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAfTTTTCTACATACCGCAGCAGT TACCATTCCATGCGCGTTGgGCAATTGTGTGAGCTGAACG TACCATTCCATGCGCGTTGgGCAATTGTGTGAGCTGAACG TACCATTCCATGCGCGTTGjAGAGTAAAAGCGGCCCTACG CAGTTGGAAAACCGGTAGCC	νj	GCCGTCTAGAGGATCCACTTCGCTCTACTGCATGCA
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GCAGTCTAGAGGATCCTATCCCTTCCACCCACCACTATPEG3TACACTCGAGGGTACCAAACCTGAAGGCCCTCACAC GCAGTCTAGAGGATCCCTCTTTCGTGCCGCCTGATAPrimers for quantification of mRNA levels by qPCReCAAGCCTTGCACACACACTTTATG CCGCAAAGAAGTACGCCACfTTTTCTACATACCGCAGCAGT TACCATTCCATGCGCGTTGgGCAATTGTGTGAGCTGAACG TACTGGCCGCATTGCATAACijAGAGTAAAAGCGCGCCTACG CAGTTGGAAAACCGGTAGCC	ATPTR14	TAATCTCGAGGGTACCGTTGAGTGAGGAGGAACGGG
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eCCGCAAAGAAGTACGCCACfTTTTCTACATACCGCAGCAGTfTACCATTCCATGCGCGTTGgGCAATTGTGTGAGCTGAACGgTACTGGCCGCATTGCATAACi/jAGAGTAAAAGCGCGCCTACGcAGTTGGAAAACCGGTAGCC	Рі	imers for quantification of mRNA levels by qPCR
Image: CCGCAAAGAAGTACGCCAC f TTTTCTACATACCGCAGCAGT f TACCATTCCATGCGCGTTG g GCAATTGTGTGAGCTGAACG j AGAGTAAAAGCGCGCCTACG i/j CAGTTGGAAAAACCGGTAGCC	0	CAAGCCTTGCACACACTTTATG
fTACCATTCCATGCGCGTTGgGCAATTGTGTGAGCTGAACGgTACTGGCCGCATTGCATAACi/jAGAGTAAAAGCGCGCCTACGi/jCAGTTGGAAAACCGGTAGCC	e	CCGCAAAGAAGTACGCCAC
TACCATTCCATGCGCGTTG g GCAATTGTGTGAGCTGAACG j TACTGGCCGCATTGCATAAC i/j AGAGTAAAAGCGCGCCTACG CAGTTGGAAAAACCGGTAGCC	£	TTTTCTACATACCGCAGCAGT
g TACTGGCCGCATTGCATAAC i/j AGAGTAAAAGCGCGCCTACG CAGTTGGAAAACCGGTAGCC	J	TACCATTCCATGCGCGTTG
<i>i/j</i> TACTGGCCGCATTGCATAAC AGAGTAAAAGCGCGCCTACG CAGTTGGAAAACCGGTAGCC	~	GCAATTGTGTGAGCTGAACG
i/j CAGTTGGAAAACCGGTAGCC	g	TACTGGCCGCATTGCATAAC
CAGTTGGAAAACCGGTAGCC	• /•	AGAGTAAAAGCGCGCCTACG
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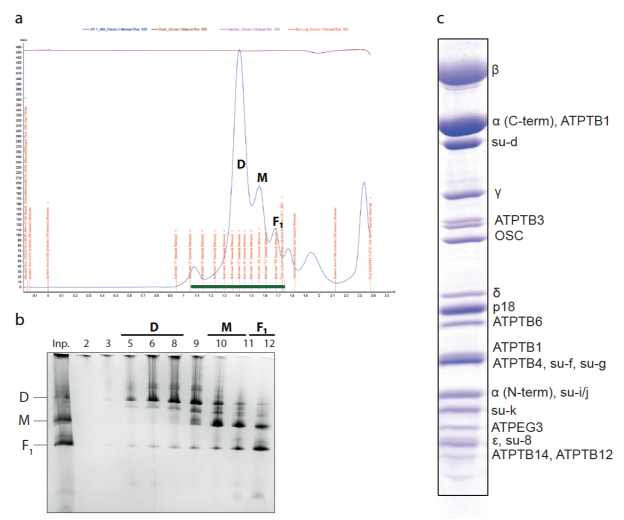
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АТРТВ6	TCGGCATAGGAGAAGTAACGA
AIFIDU	GATTCGGTTTGGAACTTGCG
АТРТВ11	CAACGGCCCCACATTCTC
AIFIDII	ACACCGCGGTCATTCATTG
ATPTB12	GCACTTCATTCTCCCGACTG
AII IDI2	ACATGATGTAACACCTCCGC
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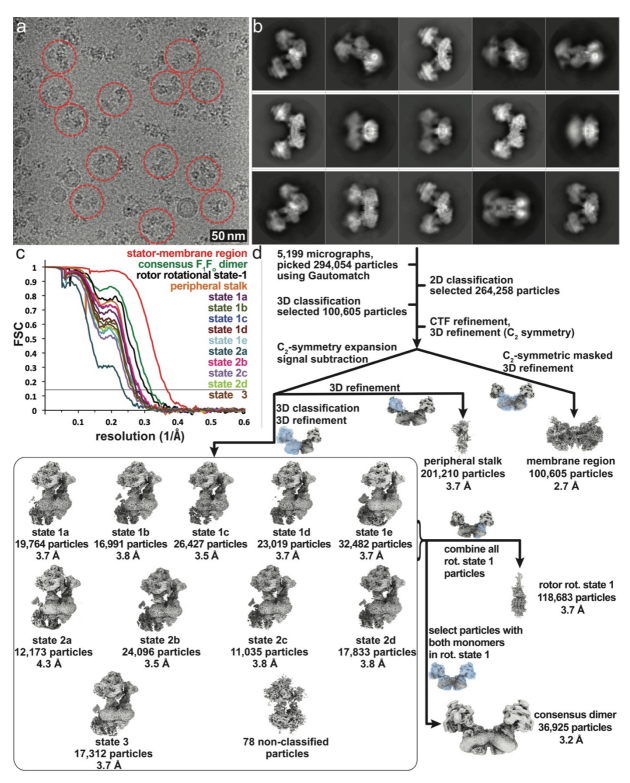
793 Supplementary Table 4. List of antibodies used in this study

Target	Туре	Reference	Dilution SDS-PAGE	Dilution BN-PAGE							
	Primary antibodies										
subunit-β	rabbit polyclonal	1	1:2000	1:2000							
p18	rabbit polyclonal	1	1:1000	-							
ATPTB1	rabbit polyclonal	1	1:1000	1:1000							
subunit-d	rabbit polyclonal	1	1:1000	1:500							
mtHsp70	mouse monoclonal	2	1:5000	-							
	Secondary antibodies										
goat anti-ra	abbit IgG HRP conjugate	BioRad 1721019	1:2000	1:2000							
goat anti-m	nouse IgG HRP conjugate	BioRad 1721011	1:2000	1:2000							

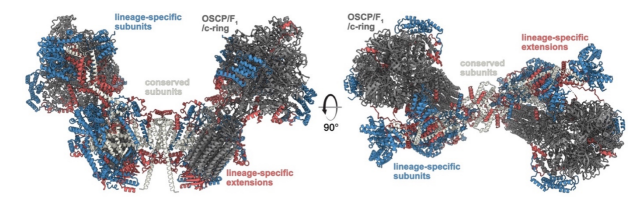
795 Supplementary Figures:



Supplementary figure 1: Purification of the *T. brucei* ATP synthase dimer. a, Size
exclusion chromatography trace with peaks enriched with ATP synthase dimers (D), monomers
(M) and F₁-ATPase (F₁) labelled. b, Fractions from size exclusion chromatography marked
with green bar in (a) resolved by native BN-PAGE. c, Dimer-enriched fraction resolved by
SDS-PAGE stained by Coomassie blue dye. Bands are annotated based on mass spectrometry
identification from excised gel pieces.

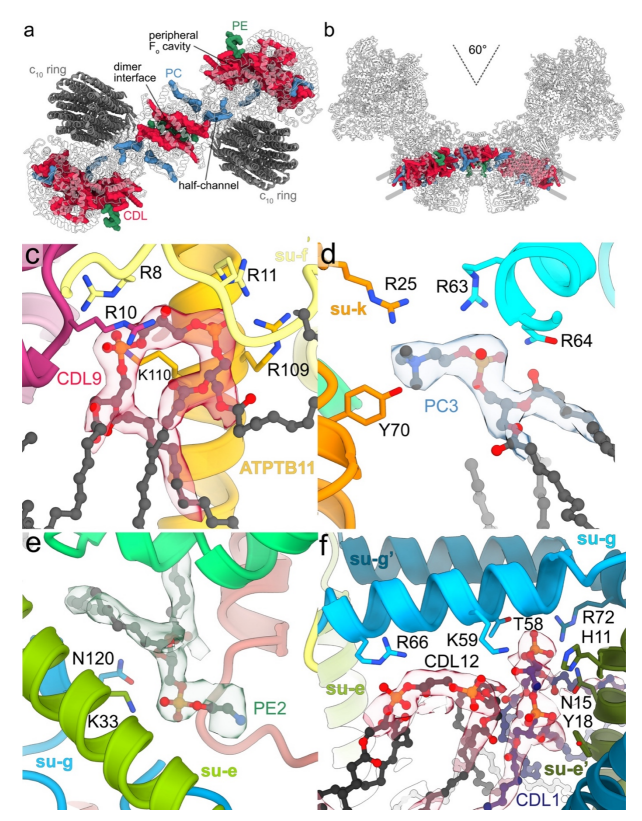


Supplementary figure 2: Cryo-EM data processing of the *T. brucei* ATP synthase dimer.
a, Representative micrograph. b, 2D class averages. c, Fourier Shell Correlation (FSC) curves
showing the estimated resolutions of ATP synthase maps according to the gold standard 0.143
criterion. d, Data processing scheme resulting in maps covering all regions of the complex, as
well as 10 rotational states.



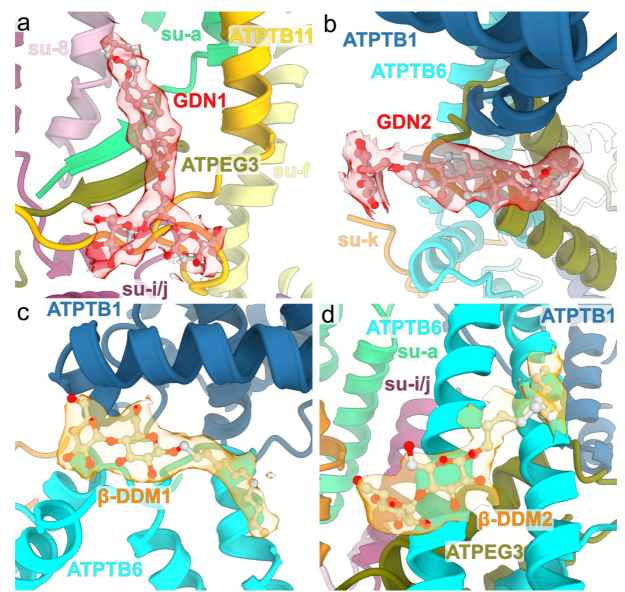
812 Supplementary figure 3: Conserved and phylum specific elements generate the *T. brucei*

- 813 ATP synthase architecture. The canonical $OSCP/F_1/c$ -ring monomers (dark grey) are tied
- $\mathbf{814}$ together by both conserved F_o subunits and extensions of lineage-specific subunits (red). The
- 815 F_o periphery and peripheral stalk attachment are composed of lineage specific subunits (blue).



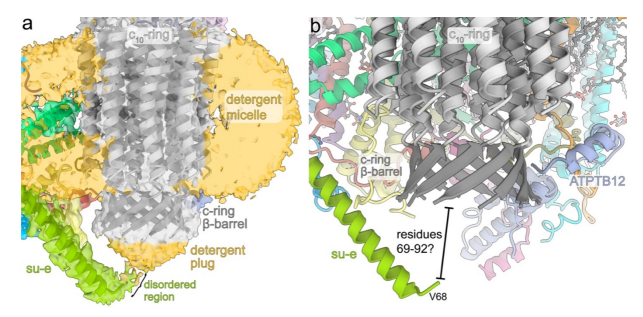
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Supplementary figure 4: The F₀ region coordinates numerous bound lipids. a, F₀ top view,
cardiolipin (CDL), phosphatidylcholine (PC) and phosphatidlyethanolamine (PE) are bound at
the dimer interface, the lumenal proton half-channel and the peripheral F₀ cavity. b, The 60°dimer angle generates a curved F₀ region with phospholipids bound in an arc-shaped bilayer.
c-f, Bound lipids with cryo-EM density and coordinating residues.



824

825 Supplementary figure 5: Bound detergents of the F_0 region. GDN (a,b) and β -DDM (c,d) 826 molecules bound in the periphery of the membrane region with cryo-EM map densities shown 827 (transparent), indicating that both glycosides are retained in the detergent micelle.



829

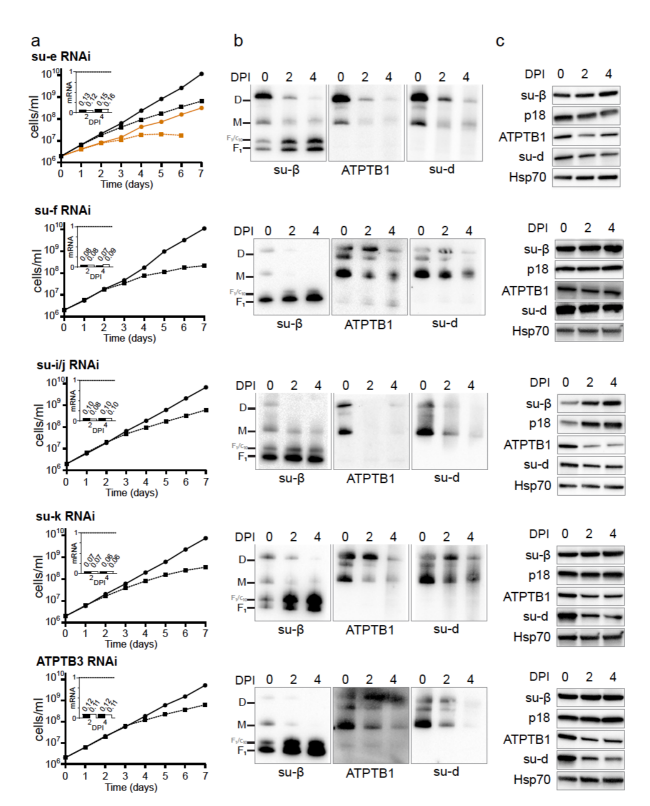
830 Supplementary figure 6. The C-terminal tail of subunit-*e* interacts with the c_{10} -ring. a,

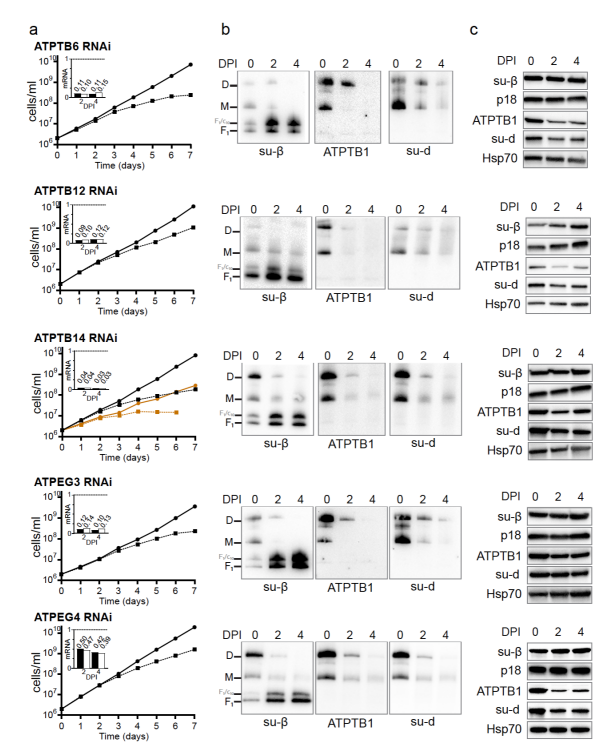
831 The cryo-EM map reveals disordered detergent density of the detergent belt surrounding the

832 membrane region as well as a detergent plug on the luminal side of the c-ring. **b**, The helical

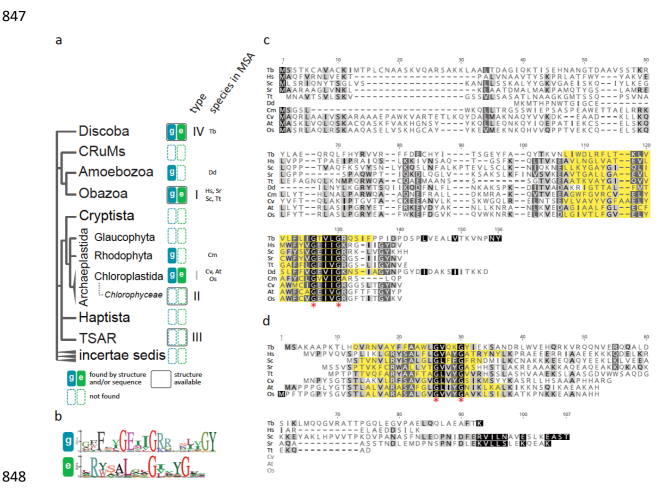
833 C-terminus of subunit-*e* extends into the lumen towards the *c*-ring. The terminal 23 residues

834 are disordered and likely interact with the β -barrel.





Supplementary figure 7. Effects of RNAi knock-down of ATP synthase subunits on 838 viability and stability and dimerization of ATP synthase. a, Growth curves of indicated 839 non-induced (solid lines) and tetracycline induced (dashed lines) RNAi cells lines in the 840 presence (black) or absence (brown) of glucose. The insets show relative levels of the 841 respective target mRNA at indicated days post induction (DPI) normalized to the levels of 18S 842 rRNA (black bars) or ß-tubulin (white bars). b, Immunoblots of mitochondrial lysates from 843 indicated RNAi cell lines resolved by BN-PAGE probed by antibodies against indicated ATP 844 synthase subunits. c, Immunoblots of whole cell lysates from indicated RNAi cell lines probed 845 846 with indicated antibodies.



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Supplementary figure 8: Phylogenetic distribution and sequence conservancy of subunits 849 e and g. a, Distribution of subunits e and g mapped on the phylogenetic tree of eukaryotes³. 850 Homologs of subunits e and g were searched in non-redundant GenBank and UniprotKB 851 protein databases by PSI-BLAST, and phmmer and hmmsearch⁴, respectively, using individual 852 853 sequences of representatives from H. sapiens and T. brucei, and in the case of hmmsearch a multiple sequence alignment (MSA) of representatives from Homo sapiens, Saccharomyces 854 cerevisiae, Arabidopsis thaliana and T. brucei, as queries. Groups, in which at least one 855 structure of ATP synthase is available, are marked. Abbreviations of species used in MSA in 856 857 panels (c) and (d) are shown. b, Sequence logo of GXXXG motifs and flanking regions of subunits e and g. Hits from hmmsearch were clustered by CD-HIT Suite⁵ to 50% sequence 858 identity and MSA of representative sequences of each cluster was generated by Clustal 859 860 Omega4⁶. The sequence logos were created from MSA in Geneious Prime (Biomatters Ltd.). c,d, MSA of sequences of subunits g (c) and e (d) from species representing major groups 861 shown in (a) generated by MUSCLE⁷ and visualized in Geneious Prime. The experimentally 862 determined or predicted transmembrane regions are highlighted in yellow. Species 863 abbreviations: Tb - T. brucei, Hs - H. sapiens, Sc - S. cerevisiae, Sr - Salpingoeca rosetta, Tt864 865 - Thecamonas trahens, Dd - Dictyostelium discoideum, Cm - Cyanidioschyzon merolae, Cv – Chlorella vulgaris, At – Arabidopsis thaliana, Os – Oryza sativa. 866 867

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