1 Cryo-electron tomography of C. elegans mitochondria reveals how the ATP

2 synthase dimer interface shapes crista membranes

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#### 17 Abstract

18 Mitochondrial ATP synthases form rows of dimers, which induce membrane curvature to give cristae their 19 characteristic lamellar or tubular morphology. The angle formed between the central stalks of ATP synthase 20 dimers varies between species. Using cryo-electron tomography and sub-tomogram averaging, we determined the structure of the ATP synthase dimer from the nematode worm C. elegans. We showed that 21 22 the angle formed by the ATP synthase dimer is different to previously determined structures. The 23 consequences of species-specific differences at the dimer interface were investigated by comparing 24 mitochondrial morphology between C. elegans and S. cerevisiae. We reveal that a larger ATP synthase 25 dimer angle in C. elegans is consequent with more lamellar (flatter) cristae compared to yeast. The 26 underlying cause of this difference was investigated by generating an atomic model of the C. elegans ATP 27 synthase dimer by homology modelling and comparing it to an existing S. cerevisiae structure. We reveal 28 extensions and rearrangements of C. elegans subunits that maintain the dimer interface. We propose that 29 increased dimer angles resulting in flatter cristae could provide an energetic advantage for species that 30 inhabit variable-oxygen environments.

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#### 32 Significance Statement

33 ATP synthase, the world's smallest rotary motor, generates the energy required for all living processes. 34 The universal formation of ATP synthase dimers in mitochondria induces membrane curvature and cristae 35 formation, generating the famously convoluted inner mitochondrial membrane that maximises ATP 36 production. Intriguingly, the angle between ATP synthase dimers varies between species. Here, we report 37 the discovery of a novel ATP synthase dimer angle in mitochondria from the nematode worm C. elegans, 38 identify a relationship between dimer angle and crista membrane morphology, and exploit AlphaFold 39 homology modelling to investigate associated structural changes. In summary, we have taken steps 40 towards understanding the importance of subunit composition at the ATP synthase dimer interface, 41 providing insights into the origins of this evolutionary divergence.

#### 42 Main Text

43

#### 44 Introduction

45 The  $F_1F_0$  ATP synthase is a molecular motor ubiquitous to all living organisms. This machine is required 46 for the essential conversion of an electrochemical gradient into the universal energy currency ATP (1). The 47 ATP synthase is composed of a catalytic  $F_1$  head connected to a membrane-embedded  $F_0$  motor by a 48 central stalk; the entire assembly is visualised as a lollipop shape when examined by electron microscopy 49 (2, 3). The central stalk transmits the torque generated by rotation of  $F_0$  to the  $F_1$  head, and a peripheral 50 stalk acts as an elastic spring, ensuring malleable coupling between  $F_1$  and  $F_0$  (4). Mitochondrial ATP 51 synthases across species share the same complement of core subunits with varying nomenclature (Table 52 S1) (5, 6). Here, we use the C. elegans nomenclature, exceptions being use of the S. cerevisiae naming 53 system for subunits missing in worms. The F<sub>1</sub> head is comprised of  $\alpha$  and  $\beta$  subunits, the central stalk of y, 54  $\delta$  and  $\epsilon$  subunits, the peripheral stalk of b d, F<sub>6</sub> and oligomycin sensitivity conferral protein (OSCP) subunits, and the F<sub>o</sub> motor contains the c-ring and subunit a. 55

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57 Mitochondrial ATP synthases are capable of assembling into dimers (7), of which there are 4 types (8): 58 Type I is present in both multicellular (9–11) and unicellular organisms (12) and types II-IV are present in 59 various unicellular organisms (13–18), reviewed in (8). When compared to type II-IV dimers, previously 60 studied type I dimers contain an additional set of subunits at the dimer interface: e, f, g, i/j, k and 8 (8). 61 Based on biochemical and imaging experiments, subunits e and g were shown to be essential for dimer 62 formation (7, 11, 19–21). Dimers of ATP synthase assemble into oligomeric rows (or ribbons) along the 63 curved ridges of crista membranes, observed by cryo-electron tomography (cryoET) (9, 11, 22). This 64 oligomerisation (formation of dimer rows) is mediated by an ancestral motif in subunits e and g (20, 21) with 65 assistance from subunit k (5, 23, 24). Formation of dimer rows is required for crista membrane curvature, 66 and thus maintenance of lamellar or tubular shaped cristae (11, 12). Deformation of cristae into balloon-67 like structures was observed in S. cerevisiae after knockdown of interface subunits e or g (11) and in ageing 68 P. anserina, when dimers disassociated into monomers (12). Moreover, molecular simulations indicated

69 that ATP synthase dimers have an innate propensity to induce membrane curvature (25). This was 70 confirmed experimentally when dimers reconstituted into liposomes spontaneously self-assembled into 71 oligomeric rows to engender this curvature, maintaining identical dimer angles to those observed in whole 72 mitochondria (26).

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74 In situ structures of type I ATP synthase dimers have been determined from native membranes (12, 25-75 28). Mammals and fungi both display an average angle between the dimer heads of  $\sim$ 86° (27). Interestingly, 76 higher-resolution single particle analysis of the purified bovine ATP synthase dimer reveals that dimer 77 angles likely vary around this average (between 76° and 95°), depending on catalytic state (29). Further 78 atomic-detail structures of purified mitochondrial type I ATP synthase dimers have also been determined 79 from mammals (Bos taurus (23) and fungi (S. cerevisiae (30) and Y. lipolytica (22)). The structure and 80 organisation of ATP synthase dimers have thus been studied across a range of different species, yet our 81 knowledge of invertebrates is lacking. The free living nematode worm C. elegans is a well-established 82 model system for the study of invertebrate cell and developmental biology (31), including the role of 83 mitochondria in metabolism, health, disease and aging (32). To complement in vivo physiological studies, 84 intact mitochondria can be stably prepared (33, 34) for biochemical and structural analyses (35). 85 Interestingly, studies have shown that nematodes lack the dimer-specific subunits i/j, k (36) and 8 (37) 86 found in mammals and fungi (Table S1). Subunit 8 is encoded by one of two overlapping ATP synthase 87 genes on the mitochondrial genome (38). Proteins encoded on the mitochondrial genome are translated 88 from essential genes (39, 40); thus it follows that subunit 8 is likely to be essential for respiration in mammals 89 and fungi. The lack of dimer-specific subunits in C. elegans provides a unique opportunity to investigate 90 how certain subunits influence ATP synthase dimer angles and mitochondrial morphology.

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In this study, we employed cryoET and sub-tomogram averaging to reveal the structure and organisation of the *C. elegans* ATP synthase, revealing a novel average dimer angle of 105°. We compared mitochondria from both *C. elegans* and *S. cerevisiae* to investigate the relationship between ATP synthase dimer angle and crista morphology. Finally, we used AlphaFold (41) to predict how protein chains in the *C. elegans* ATP

96 synthase dimer are arranged. This allowed us to analyse changes at the dimer interface and postulate the 97 cause of differences in angle. We speculate that an evolutionary divergence at the dimer interface and 98 corresponding widening of the dimer angle may be an adaptation to more variable oxygen environments.

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#### 100 Results

#### 101 The architecture of the C. elegans ATP synthase dimer

To determine the arrangement and architecture of ATP synthase dimers in C. elegans, tomograms of whole 102 103 mitochondria (Fig. 1A) and of isolated crista membranes (Fig. 1B) were analysed. ATP synthases were 104 unambiguously identified by the characteristic lollipop shape of the 10 nm diameter F1 heads positioned 105 ~10 nm away from the membrane. We confirmed the presence of oligomeric ATP synthase dimer ribbons, 106 localised at the sharp curved ridges of crista membranes, in both samples (Fig. 1A, B). Due to the obscuring 107 presence of a dense matrix in whole mitochondria, many more dimers could be visualised in crista 108 membrane samples. Therefore, 3.234 dimer pairs were extracted from the crista membrane data for sub-109 tomogram averaging. After classification, a map of the C. elegans ATP synthase dimer was determined 110 from 1,755 dimer pairs (Fig. 1C, Fig. S1, S2). Both the central and peripheral stalks were resolved clearly.

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112 Previous studies have revealed a type I dimer angle of ~86° across a range of mammalian and fungal species (12, 25-28). The architecture of the C. elegans ATP synthase dimer is unlike any other species 113 114 studied so far, with an average angle of 105° between the dimer heads (Fig. 2A). A comparison to a sub-115 tomogram average of the S. cerevisiae dimer (42), revealed that the wider dimer angle in C. elegans 116 corresponds with a sharper angle of membrane curvature (50° compared to 74°) (Fig. 2B). Accordingly, 117 opposing sides of the C. elegans crista membranes are in closer proximity than in S. cerevisiae (16.5 nm 118 compared to 20 nm). Intriguingly, the dimer interface in the C. elegans map was visually remarkably 119 different to its S. cerevisiae counterpart (Fig. 2B), and indeed all other type I dimers studied to date (12, 120 25-28). This could be due to the different complement of dimer interface subunits present in C. elegans 121 compared to S. cerevisiae (Table S1, Fig. 2C). We also analysed the inter-dimer distance and angle

between dimer heads in consecutive dimers in the oligomeric rows. This revealed an inter-dimer distance of 12.5 nm and angle between dimer heads of 20°. Despite differences in dimer angle, these values are similar to those reported previously for the type II dimer from green algae (*Polytomella* sp.) (26) (Fig. S3), suggesting that dimer angle does not influence oligomerisation of ATP synthases into rows.

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### 127 A wider dimer angle in C. elegans corresponds to flatter cristae

128 We hypothesised that the wider dimer angle associated with sharper membrane curvature in the C. elegans 129 ATP synthase dimer (Fig. 2B) would produce flatter cristae with a larger surface area to volume ratio. To 130 test this, tomographic data of whole mitochondria from C. elegans and S. cerevisiae were collected and 131 quantified. Qualitatively, C. elegans mitochondria have more lamellar shaped (or flatter) cristae, with sharp 132 curved ridges, compared to mitochondria from S. cerevisiae (Fig. 3A, 3B, Movie S1 & S2). The surface area and volume of the crista membranes were quantified, to reveal that the surface area to volume ratio 133 of the average crista membrane was significantly higher (~1.5 fold, \*\*\*\*  $p \le 0.0001$ ) in C. elegans than in 134 135 S. cerevisiae (Fig. 3C). In accordance with this, the average crista width in C. elegans (~10 nm) was half 136 the width of cristae in S. cerevisiae (~20 nm) (Fig. 3D, E and F). We speculated that narrower cristae in C. 137 elegans may allow more efficient membrane packing into the mitochondrial volume. However, no significant 138 difference was found between the total crista volume relative to mitochondrial volume between the two 139 organisms (Fig. 3G). In summary, this suggests that dimer angle exerts influence on mitochondrial 140 morphology at the level of membrane curvature, but this does not alter membrane packing.

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Mitochondria are dynamic organelles, and crista morphology can be influenced by a wide range of factors such as metabolic state (43–45). However, the average ATP synthase dimer angle remains consistent when imaged in membranes or on purification in detergent (27, 29). To ensure that morphological differences observed here in whole mitochondria were not a result of different preparation conditions, we also measured the degree of membrane curvature in isolated cristae containing *C. elegans* and *S. cerevisiae* ATP synthase dimers. Our results were consistent with the results obtained *in situ*, with a shorter

- crista width observed for the *C. elegans* dimer average (Fig. 2B). This indicates that the dimer angle remains consistent, regardless of how the sample was prepared or analysed.
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#### 151 A unique arrangement of subunits at the *C. elegans* dimer interface

152 We observed additional density at the C. elegans dimer interface (Fig. 2B), which has not been observed 153 in other type I structures determined to date (27). Nematodes are missing subunit 8 (37) (Table S1, Fig. 154 2C), which plays an key structural role in other species (22, 23, 30, 46) and is considered essential for 155 respiration (39, 40). Therefore, it is likely that considerable rearrangements at the dimer interface would 156 need to occur to account for the lack of subunit 8; this could contribute to the observed change of dimer 157 angle. To explore this possibility, we performed multisequence alignments with C. elegans, S. cerevisiae 158 and B. taurus (47-49). This revealed extensions in the 3 C. elegans subunits located at the dimer interface (e, f and g), and in 3 of the 4 subunits in the peripheral stalk (b, d and  $F_6$ ) (Fig. S4). Mass spectrometry was 159 160 used to confirm that the extensions identified by sequence in the dimer interface subunits are present in 161 the mature proteins (Fig. S5).

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163 To position the dimer interface subunits in the C. elegans ATP synthase, we used AlphaFold (41) to predict the structures of individual subunits and build a homology model. The mean pLDDT score (a per-residue 164 165 measure of local confidence on a scale from 0 - 100) of each AlphaFold prediction was > 70 (Table S2, 166 Fig. S6), and therefore defined as a "confident" prediction by the software. Structures of peripheral stalk 167 subunits b, d and F<sub>6</sub> were predicted together as a multimer (50), since individual predictions were not 168 reliable. The predicted structures were then fitted sequentially into a scaffold provided by the S. cerevisiae 169 ATP synthase dimer [PDB 6B8H] (Fig. S7, Fig. 4A). Extensions in key C. elegans dimer interface subunits 170 relative to their yeast homologues are clearly visible (Fig. S8). The C. elegans ATP synthase dimer was 171 then split into monomers and each was fitted sequentially into our sub-tomogram average dimer map (Fig. 172 S7, Fig. 4B), improving the fit considerably (Fig. S9). Our homology model correlated well to the sub-

tomogram averaging map (Fig. S10 and Table S3), allowing a comparison of *S. cerevisiae* and *C. elegans*ATP synthase dimers (Fig. 4C).

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Interestingly, the additional density identified in our sub-tomogram averaging map at the dimer interface (Fig. 2B) appears to now be filled by a rearrangement of the extended subunits f and g (Fig. 4C, D & E). In addition, extensions at the base of the peripheral stalk (subunits b, d and  $F_6$ ) also appear to contribute to the greater mass at the *C. elegans* dimer interface compared to *S. cerevisiae* (Fig. S11). Despite this, there are unfilled gaps at the dimer interface. It is plausible that specific lipids could also contribute to the mass observed, which is in line with their identification and role in stabilisation of the dimer interface (23, 51). We also cannot exclude the possibility that there are additional subunits as yet unidentified in *C. elegans*.

Finally, we fitted the *C. elegans* ATP synthase dimer model into a row of oligomeric dimer pairs along the curved edge of a crista (Fig. 4F, H). This reveals extensive inter-dimer interactions mediated by subunits e and g (Fig, 4G, I and Fig. S12), in agreement with recent work demonstrating the key role that these subunits play a role in oligomerisation / row formation (21).

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# 188 Discussion

189 Owing to the essential and universal role of the ATP synthase across eukaryotic species, it is remarkable 190 that the dimeric arrangement can be so variable (10). Until now, the arrangement of ATP synthases in invertebrates was unknown, as was the correlation between dimer angle and whole mitochondrial 191 192 morphology. In this work, a novel dimer angle for the ATP synthase from the nematode worm C. elegans 193 was discovered. By comparing worm and yeast mitochondria, we discovered that a wider ATP synthase 194 dimer angle was associated with a flatter crista membrane morphology. Since dimer row formation is known 195 to be instrumental in formation of curved ridges in crista membranes (11, 12, 26), it is consequent that 196 dimer angle influences the extent of membrane curvature.

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198 AlphaFold is able to predict protein structures with high precision (41). Our homology model of the C. 199 elegans ATP synthase dimer thus allowed us to postulate how alterations in the organisation of subunits 200 could influence dimer architecture. The ATP synthase structure is relatively well conserved across species 201 (52), but more weakly at the dimer interface and peripheral stalk (53). In accordance with this, extensions 202 in 3 C. elegans subunits (e, f and g) result in the rearrangement of subunits at the dimer interface relative 203 to S. cerevisiae. In addition, changes are detected in peripheral stalk components, with a significant N-204 terminal extension in subunit b, and a range of more subtle gaps and insertions in subunits d and F<sub>6</sub>, which 205 bulk out the width of the dimer interface. Some dimer interface subunits present in S. cerevisiae (j, k and 206 8) are absent in C. elecans. Whilst it cannot be completely excluded that an as vet unidentified subunit may 207 substitute for 8, we speculate that the absence of 8 in worms (37) highlights an interesting evolutionary 208 divergence. Subunit 8 is usually encoded on the mitochondrial genome, indicating that it is essential (39, 209 40). Additionally, 8 appears to have a key structural role in joining the dimer interface to the peripheral stalk 210 (22, 23, 30, 46). It is therefore likely that the space vacated by the missing subunit 8 is either resolved by 211 the re-arrangement of neighbouring subunits, or substituted by one of the extensions of the Fo subunits 212 close by (b, d, e, f or g).

213

214 Mitochondria have evolved their highly convoluted crista membranes to increase their surface area (54), 215 hence accommodating the maximum amount of respiratory chain complexes. This has made it possible for 216 eukaryotic organisms to deal with higher energy demands than prokaryotes (54). However, we found that 217 the flatter cristae in C. elegans relative to S. cerevisiae is not associated with more efficient cristae 218 membrane packing into the mitochondrial volume. ATP synthesis is driven by the proton-motive force (PMF) 219 and the efficiency of ATP production is directly proportional to the local proton concentration. It has been 220 suggested that cristae serve as proton concentrators that facilitate a directed flow from the source 221 (respiratory chain) to sink (ATP synthase) (9, 27). On this premise, the concentration effect exerted by the 222 cristae would be directly proportional to the number of protons pumped in a given time and the volume of 223 the crista lumen. It follows that in a flatter crista of equivalent volume to a wider one (e.g. C. elegans 224 compared to S. cerevisiae), fewer protons would need to be pumped to reach a given concentration, and

225 hence to reach the PMF required to drive the ATP synthase. In addition, protons have been proposed to 226 preferably migrate from source to sink along membrane surfaces. If this is the case, then reducing the width 227 of the crista space will reduce the solvent volume that the protons can dissipate into, facilitating the 228 efficiency of ATP synthesis. Both of these factors could allow C. elegans to maximise energy production in 229 its soil-based habitat (55), where conditions range from near hypoxia to atmospheric (56, 57). Thus, we 230 propose that a wider ATP synthase dimer angle associated with flatter cristae may be paramount for 231 capitalising on ATP production when a higher level of oxygen becomes available, and that a range of angles 232 have evolved to meet the energetic needs of different organisms. Future studies geared towards 233 investigating dimer subunit composition, angle and corresponding crista morphology in a range of species 234 inhabiting different environments will be key. The divergence in ATP synthase dimer architecture relative 235 to yeast and mammalian systems makes C. elegans an ideal model system for further investigation of the 236 role of dimer angle in mitochondrial physiology, health and disease.

237

#### 238 Conclusions

In this work, we discovered that the *C. elegans* mitochondrial ATP synthase forms a novel dimer interface with an angle between subunits of 105°. We have shown that this unusually wide angle is associated with flatter cristae compared with the yeast *S. cerevisiae*. We conclude that rearrangement of subunits at the dimer interface controls dimer angle and accordingly the extent of membrane curvature. Since an energetic advantage is likely to be proffered by an increased local proton concentration, we speculate that a range of dimer angles may have evolved to alter crista diameter and thus suit bespoke energetic needs.

# 245 Materials and Methods

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247 All standard reagents were purchased from Sigma-Aldrich (Burlington, USA).

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#### 249 C. elegans and S. cerevisiae culture

The *C. elegans* N2 Bristol strain was maintained at 20°C on 60 mm Nematode Growth Medium (NGM) plates seeded with *E. coli* OP50. For large scale preparations, a semi-synchronised population of *C. elegans* (achieved by starving so that they entered the dauer stage) (58, 59) were grown in a liquid suspension of *E. coli* NA22 in S-basal complete medium (60) at 20°C, shaking at 200 rpm for 3 days to achieve adults. For further details see (34). *S. cerevisiae* 'Bakers's yeast' S288C derivative strains YPH499 were cultured at 19 - 24°C in YPGal or YPG medium (1% w/v yeast extract, 2% w/v bactopeptone, 2% w/v galactose or 3% w/v glycerol) until OD 2-2.5 was reached. For further details see (61).

257

### 258 Mitochondrial isolation

259 C. elegans and S. cerevisiae were both harvested from liquid cultures by low speed centrifugation. C. 260 elegans preparation required an additional sucrose flotation step to remove debris. To soften the C. elegans 261 cuticle, the pellets underwent collagenase treatment (1 U/ml collagenase, 100 mM Tris-HCl pH 7.4 and 1 262 mM CaCl<sub>2</sub>), whilst S. cerevisiae pellets underwent dithiothreitol (10 mM DTT, 100 mM Tris-SO4 pH 9.4) 263 and zymolyase treatment (4.5mg/g zymolyase, 1.2 M sorbitol, 20 mM potassium phosphate, pH 7.4) to 264 disrupt the cell wall. Pellets from both species were re-suspended in homogenisation buffers. For C. 265 elegans, this was STEG/M (220 mM mannitol, 70 mM sucrose, 5 mM Tris-HCl pH 7.4 and 1 mM EGTA supplemented with 1 mM PMSF in methanol and 1% (w/v) fatty acid-free BSA). For S. cerevisiae the 266 267 homogenization buffer contained 0.6 M sorbitol, 10 mM Tris-HCl pH 7.4, 1 mM PMSF, 0.2% (w/v) BSA, 2 268 mM magnesium acetate. The re-suspended C. elegans or S. cerevisiae samples were homogenised in a 269 glass-Teflon Potter homogenisor to break open cells. Both samples were subsequently spun at low speed 270 (750 – 3000 x g for 5-15 minutes) to remove cell debris and nuclei, and then at higher speed (12,000 x g 271 for 15 minutes) to pellet mitochondria. Purified mitochondria were re-suspended in buffers that were

optimised to maintain intact mitochondria: 220 mM mannitol, 70 mM sucrose, 5 mM Tris-HCl pH 7.4 and 1
mM EGTA for *C. elegans* or 250 mM sucrose, 2 mM magnesium acetate, 10 mM Mops, pH 7.2 for *S. cerevisiae*.

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# 276 Mitochondrial crista membrane isolation

277 Crista membranes used for the sub-tomogram averaging experiments were generated by successive freeze-thaw cycles of mitochondria at -80°C. To purify mitochondrial membranes from other cellular 278 279 material, membrane extracts were incubated for 1h at 4°C with an anti-NDUFS3 primary antibody 280 (ab14711; abcam) against the matrix arm of complex I from C. elegans, followed by a 3h incubation with 281 an anti-mouse secondary conjugated to a quantum dot emitting at 625 nm (Q22085; Invitrogen). Crista 282 membranes were separated from unbound antibodies and other cellular material on an Optiprep gradient 283 with 10 layers (200 µl volume each) ranging from 0 to 27% v/v of iodixanol in STEG/M buffer, by 284 centrifugation at 80,000 × g for 30 min at 4°C using a TLS-55 rotor (Beckman Coulter Inc., Miami, FL, USA). 285 Crista membranes were identified and removed based on fluorescence under a UV lamp. Samples were 286 then diluted in STEG/M buffer to wash out the iodixanol, and spun at 20,000 x g for 15 min at 4 °C to pellet 287 the membranes. The enriched cristae were again re-suspended in STEG/M buffer.

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#### 289 Electron cryo-tomography

290 Whole mitochondria or crista membranes were mixed 1:1 with 10 nm gold fiducials (Aurion, Wageningen, 291 The Netherlands), applied to glow-discharged holey carbon EM grids (Quantifoil, Jena, Germany), and 292 blotted for 5-6 seconds, followed by plunge-freezing in liquid ethane using a Vitrobot Mark IV 293 (ThermoFisher, Massachusetts, USA) for C. elegans, or a home-made device for whole S. cerevisiae 294 mitochondria. Pre-screening of C. elegans grids was carried out using an FEI Tecnai Spirit 120kV microscope (ThermoFisher), with a Oneview CCD Camera (Gatan, Pleasanton, USA). CryoET was 295 296 performed using the same microscope for whole mitochondria, or using a 200 kV Talos Arctica 297 (ThermoFisher) for crista membranes, equipped with a K2 direct electron detector camera and a GIF

298 Quantum LS energy filter (Gatan). CryoET of whole S. cerevisiae mitochondria was performed using a 300 299 kV Titan Krios (ThermoFisher), K2 direct electron detector camera and a GIF Quantum LS energy filter 300 (Gatan). Single tilt image series' (±60, step size 1.5° - 2°) were collected at -5 to -8 µm underfocus at 301 nominal magnification of 21,000 x for whole mitochondria and 39,000 x for crista membranes, 302 corresponding to 5.4 and 3.58 Å pixel sizes respectively for C. elegans, or 26,000 x for whole mitochondria 303 from S. cerevisiae, corresponding to a 4.51 Å pixel size. The total dose per tomogram was ~120  $e^{-}/A^{2}$  for 304 whole mitochondria, and ~80 e<sup>-</sup>/Å<sup>2</sup> for isolated cristae. Tomograms were aligned using the gold fiducials in 305 IMOD (University of Colorado, United States) (62) and volumes reconstructed via weighted back-projection. 306 Contrast was enhanced by nonlinear anisotropic diffusion (NAD) filtering (63), followed by manual 307 segmentation, also in IMOD. ImageJ (64) was used to generate movies of segmentations generated in 308 IMOD.

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# 310 Subtomogram averaging

311 3,234 *C. elegans* ATP synthase dimers were picked manually in IMOD, using NAD-filtered tomograms. 312 Subvolumes containing the ATP synthase dimer were then extracted from tomograms that had not been 313 NAD filtered. These sub-volumes were CTF corrected and imported into Relion 3.1 (65) using the approach 314 and script described in (66). A reference-free initial model was generated using 3 x binned subvolumes and 315 2,481 dimers were selected by 2D classification for an unbinned refinement. Finally, 1,755 dimers were 316 selected from a 3D classification of this refined model to enter a final round of refinement and post-317 processing, resulting in a 38.6 Å resolution map. Fig. S1 details the full workflow.

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# 319 Homology model generation

AlphaFold was used to predict five structural models of each ATP synthase subunit in *C. elegans* based on their mature protein sequence (41). Mature sequences were determined using MitoFates (67) or TargetP-2.0 (68) to predict mitochondrial targeting sequences. All ATP synthase subunits known to be present in *C. elegans* were included, excepting a putative homologue of subunit j, on account of its poor alignment with

324 other homologues, and absence of any corresponding peptides in mass spectrometry analysis of the C. 325 elegans dimer. The structures of peripheral stalk subunits b, d and  $F_6$  were predicted using AlphaFold 326 multimer (50), as the individual predictions were unreliable. The models for each subunit with the highest 327 average pLDDT score were fitted sequentially to a scaffold provided by the atomic model of the S. 328 cerevisiae ATP synthase dimer [PDB 6BH8] in ChimeraX (69) using the Matchmaker tool. Where a subunit 329 had more than one isoform, the version with the highest pLDDT score was used. In the case of subunit b, 330 the isoform with the highest pLDDT score is also the only isoform expressed in somatic tissues (70). The 331 resulting structure was divided into monomers, and fitted sequentially into the sub-tomogram average of 332 the C. elegans ATP synthase dimer using the "fit in volume" tool in ChimeraX. The workflow is shown in 333 Fig. S7. The resulting homology model was converted into an MRC map using the pdb2mrc command in 334 EMAN2 (71). This map could then be fitted to the sub-tomogram average map of the C. elegans dimer for 335 comparison (Fig. S10). The yeast monomeric atomic model [PDB 6CP6] (72) was used for additional 336 analysis in Fig. S8.

337

#### 338 Mass spectrometry

The ATP synthase was purified from *C. elegans* mitochondria using a method described previously (73, 74), and analysed by Nano-LC mass spectrometry. Briefly, isolated mitochondria were solubilised and mixed with a His-tagged inhibitor protein IF<sub>1</sub>. This suspension was applied to a Nickel column to capture inhibited ATP synthase. The fraction most enriched in ATP synthase subunits was taken for mass spectrometry analysis. Further details are given in Supporting Information.

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#### 504 Data availability

505

The sub-tomogram averaging maps generated in this study have been deposited in the Electron Microscopy
Data Bank (EMDB) under accession code EMD-16216. The source image data have been deposited to the
Electron Microscopy Public Image Archive (EMPIAR) under accession number [currently unknown].

509

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511

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529

# 530 Author Contributions

- 531 EB conducted the experiments with C. elegans and analysed and quantified all tomographic data. MM
- assisted with electron microscopy and processing of tomographic data. PB prepared yeast mitochondria
- 533 for processing by VG. AB conducted bioinformatics analysis. HF purified ATP synthase from *C. elegans* for
- 534 mass spectrometry analysis. BD and PK provided resources and advice. IC and VG designed the project
- and obtained the funding. EB and VG wrote the paper. All authors commented on the manuscript.

536

537 **Competing Interest Statement:** The authors declare no competing interests.

#### 538 Figures

539





(A) Tomographic slice through a whole *C. elegans* mitochondrion (top) and corresponding segmentation (bottom; outer membrane green, inner membrane light blue, and a different colour for each crista membrane). The boxed region shows an enlarged image of a single crista membrane, with green, blue and orange arrowheads indicating the outer, inner and crista membranes respectively, and yellow arrowheads indicating ATP synthase F<sub>1</sub> heads. The crista membrane is coloured light blue in the corresponding segmentation; each ATP synthase dimer pair is coloured differently. (B) Tomographic slice through *C. elegans* isolated crista membranes (left, yellow arrowheads indicating ATP synthase F<sub>1</sub> heads) and

549 corresponding segmentation (right). The boxed region shows an enlarged image of a single crista 550 membrane, with the corresponding segmentation coloured as in panel A. Scale bars, 100 nm for 551 tomograms, and 50 nm for enlarged views of crista membranes. **(C)** Sub-tomogram average of the *C*. 552 *elegans* ATP synthase dimer at a resolution of 38.6 Å [EMD-16216]. Upper panel shows side view, lower 553 panel shows top down view.



554

#### 555 Figure 2. The C. elegans ATP synthase compared to other species.

(A) Structures depicting the range of average dimer angles observed in S. cerevisiae [EMD-7067] (30). 556 bovine heart [EMD-11436] (29), and C. elegans (this work, [EMD-16216]), using the highest resolution 557 558 structures available. (B) Direct comparison between S. cerevisiae [EMD-2161] (11) and C. elegans ATP 559 synthase sub-tomogram averages, with the angle between F1 dimer heads, the angle of crista membrane 560 curvature, and crista membrane width indicated. A bracket highlights the additional density at the C. elegans dimer interface not apparent in S. cerevisiae. Black, transparent blue and dark green mesh represent 561 562 decreasing threshold levels for the averages. (C) Cartoon detailing occurrence of ATP synthase subunits 563 in S. cerevisiae and C. elegans, each labelled with corresponding nomenclature for the species (details in 564 Table S1).



565



567 (A) Tomographic segmentations of C. elegans and S. cerevisiae mitochondria are displayed (green, outer 568 mitochondrial membrane; blue, inner mitochondrial membrane; multi-colour, crista membranes). See Movie 569 S1 (C. elegans) and Movie S2 (S. cerevisiae). (B) Enlarged examples of crista membranes (location 570 indicated by asterisks in A) to highlight the flatter cristae morphology in C. elegans mitochondria compared 571 to S. cerevisiae. (C) The mean surface area to volume ratio per crista (n = 3 mitochondria for each organism, 572 with n=47 cristae for C. elegans and n=63 cristae for S. cerevisiae) was calculated from the segmentations 573 shown in (A). (D) A single tomographic segmentation from each organism is shown with all crista coloured 574 blue. Pink dots indicate distances used to measure width at the crista tips. (E) Close up of a single crista 575 membrane from each organism (location indicated by asterisks in D, with crista tip width specified. (F) The 576 mean crista width (n= 63 crista tips for C. elegans and n= 61 for S. cerevisiae) was calculated from the 577 segmentations shown in (A). (G) The mean total crista volume relative to mitochondrial volume (n = 3

- 578 mitochondria for each organism) was calculated from the segmentations shown in (A). Error bars in C, F
- 579 and G show standard deviation of the mean and significance values were calculated using Welch's t-test
- 580 for panel C or using the Mann-Whitney U-test for panels F and G. \*\*\*\* p ≤ 0.0001. Scale bars in A & D, 200
- 581 nm; in B & E, 20nm.



584 Figure 4. AlphaFold homology model of the *C. elegans* ATP synthase dimer.

583

(A) AlphaFold predictions for *C. elegans* ATP synthase subunits (blue) overlaid with the atomic model of the yeast ATP synthase dimer ([PDB 6B8H] (30), pink), using cylinder representation. Predicted models were fitted onto 6B8H using MatchMaker in ChimeraX. (B) Two monomers from the *C. elegans* ATP synthase homology model (helical representation) fitted to the sub-tomogram average of the *C. elegans* ATP synthase dimer to generate the correct dimer angle. (C) Surface view of *S. cerevisiae* and *C. elegans* ATP synthase dimer models coloured by chain in side (top) and top-down (bottom) views. Subunits are

591 annotated and shown as  $\alpha$ , red;  $\beta$ , gold;  $\gamma$ , indigo;  $\delta$ , magenta;  $\epsilon$ , coral; c, grey; a, purple; b, blue; d, 592 turquoise; h/F<sub>6</sub>, navy; OSCP, orange; e, pale blue; f, pink; g, yellow; i/j, brown; k, dark green; 8, lime. All 593 subunits are labelled in the side views with the exception of 8 which is buried. Only the dimer interface 594 subunits are labelled in the top-down views. (D) Left, dimer interface subunits in the S. cerevisiae atomic 595 model [6B8H] (30) coloured by chain and fitted to an S. cerevisiae sub-tomogram average [EMD-2161] 596 (11). Right, dimer interface subunits in the C. elegans homology model coloured by chain fitted to the C. 597 elegans sub-tomogram average. Subunits are annotated with the same colours as panel C. (E) As per (D), 598 but with all subunits colored black, highlighting subunits missing in C. elegans relative to S. cerevisiae (j, k and 9) in red (left) and extensions in C. elegans subunits e, f and g relative to S. cerevisiae in orange (right). 599 600 (F) Top down view of the C. elegans ATP synthase dimer homology model fitted to the sub-tomogram 601 average showing sequential dimer pairs in a row. (G) As per (F), but exclusively showing dimer interface 602 subunits e, f and g coloured by chain as per panels C and D. (H) and (I) show the same interactions as in 603 (F) and (G) respectively but viewed from the side of a dimer row.

# 604 Supporting Information for

605	Cryo-electron tomography of <i>C. elegans</i> mitochondria reveals how the
606	ATP synthase dimer interface shapes crista membranes
607	
608	Emma Buzzard, Mathew McLaren, Piotr Bragoszewski, Andrea Brancaccio, Holly Ford, Bertram
609	Daum, Patricia Kuwabara, Ian Collinson & Vicki A.M. Gold.
610	
611	
612	This PDF file includes:
613	Supporting text
614	Figures S1 to S12
615	Tables S1 to S3
616	Legends for Movies S1 to S2
617	SI References
618	
619	Other supporting materials for this manuscript include the following:
620	Movies S1 to S2
621	
622	Supporting Information Text
623	Extended methods
624	

### 625 ATP synthase purification from C. elegans mitochondria

626 *C. elegans* ATP synthase was purified using a His-tagged IF<sub>1</sub> as bait, following a scaled down 627 protocol designed for purification of bovine dimers (73, 74). Residues 1-60 of the *C. elegans* F-628 ATPase inhibitor protein IF<sub>1</sub> fused to a hexa-histindine tag (cel1-60His), were overexpressed from 629 a pRSFDuet plasmid in *E. coli* BL21 (DE3), and purified by affinity chromatography on a 5 mL 630 Nickel-Sepharose column (Cytiva) attached to an ÄKTA purification system (Cytiva). Fractions 631 enriched in IF<sub>1</sub> were concentrated to ~50 mg/mL with a VivaSpin concentrator (molecular weight 632 cut-off 3 kDa; Sartorius).

633

634 C. elegans mitochondria were washed in a phosphate buffer (50 mM sodium hydrogen phosphate, 635 100 mM sucrose and 0.5 mM EDTA) and then centrifuged at 13,700 x g for 45 minutes at 4°C. This 636 wash step was repeated twice to remove endogenous C. elegans IF<sub>1</sub>. Phosphate-washed 637 mitochondria (~16 mg) were solubilised for 30 minutes at 18°C at 7.65 mg/ml with digitonin (0.92% 638 w/v) and DDM (0.76% w/v). The resulting extract was centrifuged at 24,000 x g for 20 minutes at 639 4°C, and cel1-60His was added to the supernatant at 2.7 µg per 1 mg mitochondria to form 640 ATPase:cel1-60His complexes. A solution of 200 mM ATP, 200 mM MgSO<sub>4</sub>, and 400 mM Trizma 641 (pH 8.0) was also added at 15 µl/ml before incubating for 15 minutes at 37°C, with further additions 642 of this solution being added at 5 minute intervals. Precipitate was removed by centrifugation at 643 24,000 x q for 10 minutes at 4°C. NaCl and imidazole were added to the clarified sample to reach 644 final concentrations of 150 mM and 25 mM respectively. This final extract was applied to a 1 mL 645 HisTrap FF Nickel Column (Cytiva) installed on an ÄKTA purification system (Cytiva) and 646 equilibrated in a buffer containing 20 mM Tris, pH7.4, 150 mM NaCI, 2 mM ATP, 2 mM MgSO<sub>4</sub>, 647 10% (v/v) glycerol, 0.1% (w/v) glyco-diosgenin (GDN) and a 0.1 mg/mL phospholipid mix. The 648 ATPase:cel1-60His complexes were eluted from the column by addition of a linear gradient of 649 imidazole up to 500 mM over 10 mL. 0.5mL fractions were collected and run on an SDS-PAGE gel 650 to confirm which fractions contained the ATPase:I1-60His.

651

## 652 Nano-LC Mass Spectrometry

The sample of ATP synthase was run on a 10% SDS-PAGE gel until the dye front had migrated approximately 1cm into the separating gel. The gel lane was then excised as a single slice and subjected to in-gel tryptic digestion using a DigestPro automated digestion unit (Intavis Ltd.). The resulting peptides were fractionated using an Ultimate 3000 nano-LC system in line with an Orbitrap Fusion Lumos mass spectrometer (Thermo Scientific). Spectra were acquired with Xcalibur 3.0 software (Thermo Scientific).

659

660 The raw data files were processed and quantified using Proteome Discoverer software v2.1 661 (Thermo Scientific) and searched against the UniProt Caenorhabditis elegans database 662 (downloaded October 2022; 26728 sequences) using the SEQUEST HT algorithm. Search criteria 663 included oxidation of methionine (+15.995Da), acetylation of the protein N-terminus (+42.011Da) 664 and methionine loss plus acetylation of the protein N-terminus (-89.03Da) as variable modifications 665 and carbamidomethylation of cysteine (+57.021Da) as a fixed modification. Searches were 666 performed with full tryptic digestion and a maximum of 2 missed cleavages were allowed. The 667 reverse database search option was enabled and all data was filtered to satisfy false discovery rate 668 (FDR) of 5%.



- 671 Figure S1. Flow chart of tomogram processing and sub-tomogram averaging using IMOD
- 672 and Relion.



673



675 **averaging map.** The FSC curve is an output from Relion 3.1. The corrected FSC is shown in black,

unmasked FSC in green, masked FSC in blue, and phase randomised masked FSC in red.



679 Figure S3. Inter-dimer distance and angle between consecutive dimer heads in oligomeric 680 rows of C. elegans ATP synthase dimers. (A) 2D projection showing side view of a masked map 681 of the C. elegans ATP Synthase dimer. (B) Side view shown in 3D, with distance between central 682 stalks indicated. (C) 2D projection showing top down view of an unmasked map of the C. elegans 683 ATP synthase dimer. (D) Top down view in 3D with inter-dimer distance indicated. (E) 2D projection 684 showing side view (rotated 90° compared to A) of an unmasked map of the C. elegans ATP 685 Synthase dimer. (F) Side view in 3D, with inter-dimer angle indicated. All indicated measurements 686 were made in IMOD. Scale bars, 10 nm.

### 688 Subunit e

	tr Q21732 Q21732_CAEEL sp P81449 ATPJ_YEAST sp Q00361 ATP5I_BOVIN	MSAPLKHPNAVVLQPPTVTISPLIRFGRYAALSLGVVYGFFRLRQIREYHADIREWDH-E MSTVNVLRYSALGLGLFFGFRNDMILKCNAKKKEEQAQYE MVPPVQVSPLIKLGRYSALFLGMAYGAKRYNYLKPRAEEERRLAA-E : **:** **: :* . :: *	59 40 46
	tr Q21732 Q21732_CAEEL sp P81449 ATPJ_YEAST sp Q00361 ATP5I_BOVIN	KAVAAAEEAAKKKKWLAKDEMRYLMQVVNIPFEEGVKQFGVADLYKED- EKLKLVEEAKKEYAKLHPVVTPKDVPANASFNLEDPNIDFERVILNAVESLKEAS EKKKRDEQK-RIERELAEAQEDTILK	107 95 71
689	tr Q21732 Q21732_CAEEL sp P81449 ATPJ_YEAST sp Q00361 ATP5I_BOVIN	- 107 <b>T</b> 96 - 71	

#### 690

#### 691 Subunit f

sp Q22021 ATPK_CAEEL sp Q06405 ATPK_YEAST sp Q28851 ATPK_BOVIN	MAWFRPPPPHTQLRPWVPDAIFIPISRAVERVGVFFYNRVLNKTEVGLFDKRWNKNVHGP MIFKRAVSTLIPPKVVSS	60 18 0
sp Q22021 ATPK_CAEEL sp Q06405 ATPK_YEAST sp Q28851 ATPK_BOVIN	YCHWRYYGKLDTKFMDVKLGDLPAWMARREKTPSAFYNEFMRNIWRVHNLYY KNIGSAPNAKRIANVVHFYKSLPQGPAPAIKANTRLARYKAKYF MASVVPLKEKKLLEVKLGELPSWILMRDFTPSGIAGAFQRGYYRYYNKYV ::*. * . * * *	112 62 50
sp Q22021 ATPK_CAEEL sp Q06405 ATPK_YEAST sp Q28851 ATPK_BOVIN	SGPVYNNTVKVIFRFIFAYSFLNWLVKSHRYVDFQKTMYHW153DGDNASGKPLWHFALGIIAFGYSMEYYFHLRHHKGAEEH-101NVKKGSIAGLSMVLAAYVFLNYCRSYKELKHERLRKYH-88	

# 692

#### 693

# 694 Subunit g (isoform 2)

sp Q18803 ATPL2_CAEEL sp Q12233 ATPN_YEAST sp Q28852 ATP5L_BOVIN	MAAPKLGFFEKIANLTGALYRHQHAQFPRRF-AILKAVGKHELAPPRQADWPA 52 MLSRIQNYTSGLVSKANLLSSKALYYGKVGAEISKQIYLKEGLQPPTVAQFKS 53 MAEFVRNLAEKAPALVNAAVTYSKPRL-ATFWYYAKVELVPPTPAEIPT 48 ::::::* * * * *::	
sp Q18803 ATPL2_CAEEL sp Q12233 ATPN_YEAST sp Q28852 ATP5L_BOVIN	IKADWAKV-QSFIQTGGYKNLSIREGLVYTAVTLEVVFWFFVGEMIGRRYIFGYL 10 VYSNLYKQSLNFALKPTEVLSCLKNIQKNELLKYGAYGIQLIGFYSVGEIIGRRKLVGYK 11 AIQSLKKI-INSAKTGSFKQLTVKEALLNGLVATEVWMWFYVGEIIGKRGIIGYD 10 . *	.3
sp Q18803 ATPL2_CAEEL sp Q12233 ATPN_YEAST sp Q28852 ATP5L_BOVIN	VPADYVSKSTKKTVKEQEALAALEN         131           HH         115           V         103	

695

#### 697

# 698 Subunit b (isoform 2)

sp Q19126 AT5F2_CAEEL	MSLSRCLPLGQNARVIIIPARLAHAASTQAAAATDDAPNFFQKLAHRFQGVPLKGEAHAP	60
sp P05626 ATPF_YEAST	MSMGVRGLALRSVS	16
sp P13619 AT5F1 BOVIN	MLSVLSAAAAAA	14
	*	
SD 019126 ATSE2 CAFEL		120
SP DOF626 ATDE VEAST		62
ap D12610 ATEF1 DOVIN		67
SPIPISOI9 ATSFI_BOVIN	PSLKNAALLGPGVLQAIKIFHIGQPSLAPVPPLPEHOGKVKFGLIPEEFFQFL	0/
sp Q19126 AT5F2_CAEEL	QKVTGVSGPYLFFGGLFAFLVNKELWVFEEQGHMTVGWILFYLLVTRTAGYKIDQGLYNG	180
sp P05626 ATPF_YEAST	LTKTGVLGTSAAAVIYAISNELYVINDESILLLTFLGFTGLVAKYLAPAYKD	114
sp P13619 AT5F1_BOVIN	YPKTGVTGPYVLGTGLILYLLSKEIYVITPETFSAISTIGFLVYIVKKYGASVGE	122
	*** *	
sp 019126 AT5F2 CAEEL	YOERVNEFKGLIOEDLKEAVEFKKTSAKOTESLNSIKESYPTALKESMALOL	232
SD P05626 ATPE VEAST	EADARMKKVSDVI NASRNKHVEAVKDRTDSVSOI ONVAETTKVI EDVSKETVELES	170
SP P13619 ATEE1 BOVIN		170
SPIPISOIS ATSPI_BOVIN	PADRENEQRIAQEEEVRQASIRQIQDAIDHERSQQAEVQRNHIEPDVQRNHIANAE	1/0
sp Q19126 AT5F2_CAEEL	EATYRKNVQSVATELKRRIDYLKETEESKARVEREQLLKLINSEVDKEFSDRSFKDKYLQ	292
sp P05626 ATPF_YEAST	EAFELKQKVELAHEAKAVLDSWVRYEASLRQLEQRQLAKSVISRVQSELGNPKFQEKVLQ	230
sp P13619 AT5F1_BOVIN	EVTYRERLHRVYREVKNRLDYHISVQNMMRQKEQEHMINWVEKRVVQSISAQQE-KETIA	237
	*. :. : * * :* : : * :. : : :	
sp 019126 AT5F2 CAEEL	NAIOOLKGLNVOL 305	
SP P05626 ATPF YEAST	OSISEIEOLLSKLK 244	
SD P13619 ATSE1 BOVTN		
5911 13013 [A131 1_00114		

# 699

700

# 701 Subunit d

tr Q17763 Q17763_CAEEL	MSGAAKRVATSSVNWSKLAERLVPEHAAELTRVKGVSGTFQSAVSQLPADLPKIDFAA	58
sp P30902 ATP7_YEAST	MSLAKSAANKLDWAKVISSLRITGSTATQLSSFKKRNDEARRQLLELQSQPTEVDFSH	58
sp P13620 ATP5H_BOVIN	MAGRKLALKTIDWVAFGEIIPRNQKAVANSLKSWNETLTSRLATLPEKPPAIDWAY	56
tr Q17763 Q17763_CAEEL	LKKALPAHSAVLDSLQKQYESVKIPYGEVPAEYLKEVDQWVDYNNARIK	107
sp P30902 ATP7_YEAST	YRSVLKNTSVIDKIESYVKQYKPVKIDASKQLQVIESFEKHAMTNAK	105
sp P13620 ATP5H_BOVIN	YKANV-AKAGLVDDFEKKFNALKVPIPEDKYTAQVDAEEKEDVKSCAEFLTQSKTRI-	112
	**	
tr Q17763 Q17763_CAEEL	LHEVKVADGLQEAKKVEEKWAKAPPVEHFDRQHFVEYFPAHFYDLRYQNRIPDPCNIGLN	167
sp P30902 ATP7_YEAST	ETESLVSKELKDLQSTLDNIQSARPFDELTVDDLTKIKPEIDAKVEEMVKKGKW	159
sp P13620 ATP5H_BOVIN	QEYEKELEKMRNIIPFDQMTIEDLNEVFPETKLDKKKYPYWPHRPIETL-	161
	···· ·· · *.·.· ··· * · .	
tr Q17763 Q17763_CAEEL	ETPEIENRFKDYKVLRRADKVDDH 191	
sp P30902 ATP7 YEAST	DVPGYKDRFGNLNVM 174	
sp P13620 ATP5H_BOVIN	161	

# 703 Subunit F<sub>6</sub>

tr 016517 016517_CAEEL sp Q12349 ATP14_YEAST	MFRAVQSVRS <u></u> L- <u></u> ST <u></u> TAACRQDLIQQTFVTKIREIAK MFPIASRRILLNASVLPLRLCNRNFTTTRISYNVIQDLYLRELKDTKLAPS	35 51
sp P02721 ATP5J_BOVIN	MILQRLFRLSSAVQSAISV-SWRRNIGITAVAFNKELDPVQKLFVDKIREYRTK :* . * : : : :*. :: ::::	53
tr 016517 016517_CAEEL		91
sp Q12349 ATP14_YEAST	TLQDAEGNVKPWNPPQKPNLPELELQGPEALKAYTEQNVETAHVAKESEE	101
sp P02721 ATP5J BOVIN	RQTSGGPVDAGPEYQQDLDRELFKLKQMYGKADMNTFPNFTFEDPKFEVVE	104
	** : * . : :* :.	
tr 016517 016517_CAEEL	GQTLASLLEGVKKD-HSEYVASRDAKKAEQAARNAALKQ 129	
sp Q12349 ATP14_YEAST	GESEPIEEDWLVLDDAEETKESH 124	
sp P02721 ATP5J_BOVIN	KP-QS 108	
	:	

704

# 705

#### 706 Subunit OSCP

tr P91283 P91283_CAEEL	MAQLMKRGFSTSAALAKAQLVKTPIQVHGVEGRYAAALYSAGHKQNKLDQI	51
sp P13621 ATPO_BOVIN	MAALAVSGLSQQVRCFSTSVVRPFAKLVRPPVQIYGIEGRYATALYSAASKQNKLEQV * :: :: *: *: **: **:	58
tr P91283 P91283_CAEEL	STDLNNVRSVYKDNKKFQEFVLDPTLKANKKKTAIEAI	89
sp P05626 ATPF_YEAST	DPKAKANSIINAIPGNNILTKTGVLGTSAAAVIYAISNELYVINDESI	92
sp P13621 ATPO BOVIN	EKELLRVGQILKEPKM-AASLLNPYVKRSVKVKSLSDM	95
	:* ::: . ::	
tr P91283 P91283_CAEEL	STKLGLTKETGNFLGLLAENGRLNKLESVVSSFESIMRAH	129
sp P05626 ATPF YEAST	LLLTFLGFTGLVAKYLAPAYKDFADARMKKVSDVLNASRNKHVEAVKDRIDSVSQLQNVA	152
sp P13621 ATPO_BOVIN	TAKEKFSPLTSNLINLLAENGRLTNTPAVISAFSTMMSVH	135
tr P91283 P91283_CAEEL	-RGELFVQVTS-AEELSSSNQKALSDALSKIGKSGQKLT-V-TYAVKPSIL	176
sp P05626 ATPF_YEAST	ETTKVLFDVSKETVELESEAFELKQKVELAHEAKAVLDSWVRYEASLRQLEQRQLAKSVI	212
sp P13621 ATPO_BOVIN	-RGEVPCTVTT-ASALDEATLTELKTVLKSFLSKGQVLK-L-EVKIDPSIM :: *:. : *	182
tr P91283 P91283_CAEEL	GGLVVTIGDKYVDLSIASRVKKYKDALATAI 207	
sp P05626 ATPF_YEAST	SRVQSELGNPKFQEKVLQQSIS-EIEQLLSKLK 244	
sp P13621 ATPO_BOVIN	GGMIVRIGEKYVDMSAKTKIQKLSRAMREIL 213 .: :*: * :: * .::: :	

# 707

Figure S4. Multiple sequence alignment for dimer interface and peripheral stalk subunits. Comparisons were made between *C. elegans* (O16517), *S. cerevisiae* (Baker's yeast strain ATCC 204508 / S288c; Q12349) and *B. taurus* (P02721) using Clustal Omega at EMBL-EBI (47–49). In the alignment output, an asterisk (\*) indicates a perfect alignment, a colon (:) indicates a site belonging to a group exhibiting strong similarity, and full stop (.) indicates a site belonging to a group exhibiting weak similarity. Residues are coloured according to their biophysical properties.

- 514 Small and hydrophobic residues are coloured red, acidic residues are coloured blue, basic residues
- are coloured magenta, and hydroxyl, sulfhydryl, amine and glycine residues are coloured green.
- 716 Extensions in C. elegans subunits relative to both the S. cerevisiae and B. taurus homologues are
- underlined in black, deletions are underlined in maroon. Where subunits have multiple isomers, the
- isomer used in the homology model is used for alignment.

# 719 Subunit e (Q21732)

# 720 <u>MSAPLKHPNAVVLQPPTVTISPLIR</u>FGR<u>YAALSLGVVYGFFR</u>LRQIR<u>EYHADIREWDHEKAVAAAE</u>

- 721 <u>EAAKKKKWLAKDEMRYLMQVVNIPFEEGVKQFGVADLYK</u>ED
- 722
- 723 Subunit f (Q22021)
- 724 <u>MAWFRPPPPHTQLRPWVPDAI</u>FIPISRAVERVGVFFYNRVLNKTEVGLFDKRWNKNVHGPYCH
- 725 <u>WR</u>YYGK<u>LDTKFMDVKLGDLPAWMAR</u>R<u>EKTPSAFYNEFMR</u>NIWR<u>VHNLYYSGPVYNNTVK</u>VIFR
- 726 FIFAYSFLNWLVKSHR<u>YVDFQK</u>TMYHW
- 727
- 728 Subunit g (isoform 2) (Q18803)
- 729 MAAPKLGFFEKIANLTGALYRHQHAQFPRRFAILKAVGKHELAPPRQADWPAIKADWAKVQSFIQ
- 730 <u>TGGYK</u>NLSIREGLVYTAVTLEVVFWFFVGEMIGR<u>RYIFGYLVP<mark>ADYVSK</mark>STKKTVKEQEALAALE</u>
- 731 <u>N</u>
- 732
- 733 Subunit b (isoform 2) (Q19126)
- 734 MSL<mark>SRCLPLGQNARVIIIPA</mark>RL<mark>AHAASTQAAAATDDAPNFFQKLA</mark>HR<u>FQGVPLKGEAHAPKSMFE</u>
- 735 <u>DCNKEWSAPEPLPAIPKDF</u>KEHPDR<u>DLVNYPYPARPMYPPK</u>SR<u>LLMMPDSWFTPFQK</u>VTGVSG
- 736 PYLFFGGLFAFLVNKELWVFEEQGHMTVGWILFYLLVTR<u>TAGYKID<mark>QGLYN</mark>GYQER</u>VNFFK<u>GLIQ</u>
- 737 <u>EDLKEAVE<mark>FKKT</mark>SAKQTESLNSIKESYPTALKESMALQLEATYRKNVQSVATELKRRIDYLKETEE</u>
- 738 <u>SKARVEREQLLKLINSEVDKEFSDRSFKDKYLQNAIQQLK</u>GLNVQL
- 739
- 740 Subunit d (Q17763)
- 741 MSGAAKR<u>VATSSVNWSK</u>LAER<u>LVPEHAAELTRVKGVSGTFQSAVSQLPADLPKIDFAALKKALP</u>
- 742 <u>AHSAVLDSLQKQYESVKIPYGEVPAEYLKEVDQWVDYNNAR</u>IKLHEVK<u>VADGLQEAK</u>KVEEKWA

#### 743 KAPPVEHFDRQHFVEYFPAHFYDLRYQNRIPDPCNIGLNETPEIENRFKDYKVLRRADKVDDH

744

# 745 Subunit F<sub>6</sub> (O16517)

- 746 MFRAVQSVRSLSTTAACRQDLIQQTFVTKIREIAKNAGNLANSDPAVKKALQEELNRLATKFQLA
- 747 NADVVSKLPTNFEAAK<u>VDSAVQSALE<mark>GQTLASLLE</mark>GVKK</u>DHSEYVASRDAKKAEQAARNAALKQ

748

Figure S5. Mass spectrometry data for *C. elegans* ATP synthase subunits with significant extensions. The sequence for each subunit of interest is shown and identified with a Uniprot code. The predicted mitochondrial targeting sequences are coloured red. The *C. elegans* specific extensions (revealed in sequence alignments from Fig. S4) are highlighted in yellow. Peptides identified by mass spectrometry are underlined. Where subunits have multiple isomers, the isomer used in the homology model is shown.

### Alphafold predictions gallery



## 757 Figure S6. AlphaFold predictions gallery

758 AlphaFold predictions (41) for each C. elegans ATP synthase subunit, coloured by pLDDT score 759 per residue. The pLDDT score is a per-residue measure of local confidence on a scale from 0 -760 100. The structure of subunits b d and  $F_6$  were predicted as a multimer. The confidence measure 761 for predictions made using AlphaFold multimer (50) is similar, but modified to score interactions 762 between residues of different chains. It is calculated using a weighted combination of predicted-TM 763 score (pTM) and interface predicted-TM score (ipTM), and has a scale from 0-1. The appropriate mean confidence score for each AlphaFold / multimer prediction is shown beneath each subunit 764 765 name.



768 Figure S7. C. elegans ATP synthase homology model workflow

(A) MitoFates (67) or TargetP-2.0 (68) were used to predict the mitochondrial targeting sequence

770 of individual proteins of the ATP synthase, so that the mature protein sequence could be identified. 771 The example shown is the TargetP-2.0 prediction for subunit α. (B) AlphaFold was used to predict 772 structures of all mature C. elegans ATP synthase subunits; again the example shown is the 773 prediction for subunit α. (C) Predicted models were sequentially fitted into the S. cerevisiae ATP 774 synthase model [PDB 6B8H] (30) using MatchMaker in ChimeraX. (D) The resulting homology 775 model (blue) after all subunits have been fitted to the scaffold provided by 6B8H (pink). (E) The 776 homology model without the S. cerevisiae scaffolding. (F) The C. elegans ATP synthase dimer was 777 split into separate monomers. (G) The monomers were fitted sequentially into the sub-tomogram 778 average of the C. elegans ATP synthase using matchmaker in ChimeraX in order to obtain the 779 correct dimer angle. (H) The final homology model of the C. elegans ATP synthase dimer fitted into 780 the sub-tomogram average.





- 786 sequence alignment scores output by ChimeraX when using the "fit to model" tool are shown for
- each overlay. (A) Overlays for subunits e, f, g, b and d are taken from the assembled ATP synthase
- dimers in Fig. 4A. (B) Two alternative overlays are shown for subunit F<sub>6</sub>. Left, an overlay taken from
- 789 Fig. 4A as above. However, the S. cerevisiae 6B8H (30) structure does not contain the complete
- density for subunit F<sub>6</sub>. Right, an S. cerevisiae monomeric atomic model [PDB 6CP6] (72) was used
- to display a more complete *S. cerevisiae* chain for the overlay.
- 792



793 794 Figure S9. Comparison of different models fitted to the *C. elegans* ATP synthase dimer map. 795 Different ATP synthase dimer models were fitted into the C. elegans ATP synthase in situ map. All 796 models were fitted into the map at threshold 0.0429 in ChimeraX, and the percentage of atoms 797 outside the contour is shown for each model. (A) The purified S. cerevisiae ATP synthase dimer 798 atomic model [PDB 6B8H] (30) shows a poor fit, with 14% of atoms outside the contour. (B) The 799 C. elegans ATP synthase dimer homology model following scaffolding to the S. cerevisiae model 800 also shows a poor fit, with 15% of atoms outside the contour. (C) Sequential fitting of monomers 801 from the C. elegans homology model shows an improved fit, with only 6% of atoms outside the 802 contour.



805 Figure S10. The C. elegans homology model fitted to the C. elegans ATP synthase dimer 806 sub-tomogram averaging map. Using the pdb2mrc command in EMAN2 (71), the PDB of the C. elegans homology model was converted into an MRC map at both 12 Å and 36 Å resolution. 807 808 Converted pdb2mrc maps (blue) were then fitted to the sub-tomogram averaging map of the C. 809 elegans dimer (grey) at equivalent threshold levels. Correlation scores between the homology 810 model and sub-tomogram averaging maps are displayed. (A) Maps of the 6B8H S. cerevisiae ATP 811 synthase atomic model (30) fitted to the sub-tomogram average for reference. (B) Maps of the C. 812 elegans original homology model (without adjusting for dimer angle) fitted to the sub-tomogram 813 average. (C) Maps of the dimer angle adjusted C. elegans homology model fitted to the sub-814 tomogram average.



816 Figure S11. Comparison of peripheral stalk subunit arrangement in S. cerevisiae vs C. elegans ATP synthase dimers. (A) S. cerevisiae and C. elegans peripheral stalk subunits 817 818 coloured by chain. Subunits are annotated and shown as b, blue; d, turquoise; F<sub>6</sub>, dark navy; and 819 OSCP, orange. Left, peripheral stalk subunits b, d and OSCP in the 6B8H S. cerevisiae atomic 820 model (30), and F<sub>6</sub> from the 6CP6 monomeric atomic model (72), fitted to the S. cerevisiae subtomogram average [EMD-2161] (11). The chain for F<sub>6</sub> was taken from EMD-6CP6 (see Fig. S8B) 821 822 (72). Right, C. elegans homology model fitted to the C. elegans sub-tomogram average. (B) As per 823 (A), but with all subunits colored black, highlighting extensions in C. elegans subunits b, d and  $F_6$ 824 relative to S. cerevisiae in orange.





Figure S12. Inter-dimer interactions mediated by subunits e and g in *C. elegans* ATP synthase dimer rows. Top down view (left) and side view (right) of the *C. elegans* ATP synthase homology model (grey) fitted to each dimer pair in the unmasked sub-tomogram average of the *C. elegans* dimer. Dimer interface subunits are colored (e, pale blue; f, pink; g, yellow) to highlight extensive inter-dimer interactions mediated by subunits e and g.

C. elegans	S. cerevisiae	B. taurus
F1 head		
α	α/Atp1	α
β	β/Atp2	β
F₀ head	·	
γ	γ/Atp3	γ
δ	δ/Atp16	δ
3	ε/Atp15	3
Peripheral stal	k	
b	b/Atp4	b
d	d/Atp7	d
F <sub>6</sub>	h/Atp14	F <sub>6</sub>
OSCP	OSCP/Atp5	OSCP
F <sub>o</sub> motor		
а	a/Atp6	а
С	c/Atp9	С
Type I dimer-s	pecific subunits	
е	e/Atp21	е
f	f/Atp17	f
g	g/Atp20	g
-	i/j/Atp18	6.8PL/ j
-	k/Atp19	DAPIT/ k
-	8/Atp8	A6L/ATP8

### 833 Table S1. Nomenclature for homologues of ATP Synthase subunits

Nomenclature for yeast and mammalian species are described as detailed by Song and Pfanner

835 (6).

Subunit name	<i>C. elegans</i> Uniprot Accession Number	Mean pLDDT (or weighted pTM & ipTM) <sup>1</sup>	RMSD between pruned atom pairs <sup>2</sup>	RMSD across all atom pairs	Sequence alignment score <sup>3</sup>
α	Q9XXK1	92.9066444	0.929	2.275	1869.6
β	P46561	87.5983733	1.241	2.961	1875.8
γ	Q95XJ0	92.17635904	0.914	2.058	665.4
δ	Q09544	85.82596715	0.901	2.488	224.7
ε <sup>4</sup>	O16298	65.4116052			
	P34539	71.51756013	0.541	15.704	18.9
с	Q9BKS0	95.27668763	0.74	0.74	262.1
е	Q21732	89.96603434	0.993	10.078	79.8
f	Q22021	87.12810426	1.36	14.559	50.4
g	Q18803	90.2590889	1.318	10.538	112.8
а	P24888	77.54888203	1.154	4.824	308.5
b	Q20053	84.43326886			
	Q19126	84.74422485	1.004	15.512	332.8
d	Q17763	82.60642993	1.342	9.967	116.7
F <sub>6</sub>	O16517	89.9038886	0.992	15.452	43
OSCP	P91283	76.95722866	1.035	7.553	346.3
	Q7JNG1	76.43462181			
b,d,F <sub>6</sub> multimer	Q19126, Q17763, O16517	0.712090029	1.324	11.335	327.9

#### 836 Table S2. Metrics to assess confidence and fit of AlphaFold predicted structures

<sup>1</sup> pLDDT scores are shown for subunits where structure was predicted individually, a weighted pTM and iPTM score is shown for a complex of subunits predicted using AlphaFold multimer. The pLDDT score is a perresidue measure of local confidence on a scale from 0 – 100. The predicted-TM score (pTM) and interface predicted-TM score (ipTM), and has a scale from 0-1. <sup>2</sup> RMSD (Root Mean Square Deviation) is a measure of the similarity between two superimposed atomic

<sup>2</sup> RMSD (Root Mean Square Deviation) is a measure of the similarity between two superimposed atomic coordinates, in this case for the predicted *C. elegans* subunits and the model of the *S. cerevisiae* ATP synthase dimer.

<sup>3</sup> Sequence alignment score between *C. elegans* and *S. cerevisiae*.

<sup>4</sup> Where a subunit has more than one isoform, the version with the highest pLDDT score was used to build the homology model. RMSD and sequence alignment scores are only shown for the selected protein. In the case of subunit b, the isoform with the highest pLDDT score is also the only isoform expressed in somatic tissues (70).

# 837 Table S3. Metrics to assess fit of atomic detail models to C. elegans ATP synthase dimer

## 838 sub-tomogram averaging map.

	S. cerevisiae atomic model [PDB 6B8H]	Original <sup>5</sup> <i>C. elegans</i> homology model	Adjusted <sup>6</sup> <i>C. elegans</i> homology model
PDB % atoms outside contour <sup>7</sup>	14	15	6
MRC map <sup>8</sup> correlation score	0.8757	0.8643	0.9469

839

<sup>7</sup> This value is given by Chimera when fitting a PDB model to a map using the "fit in map" command.
 <sup>8</sup> MRC map generated from PDB's using pdb2mrc command in EMAN2 (71). This metric shows level of correlation between pdb2mrc map and our sub-tomogram average at the same resolution (38.6 Å).

<sup>&</sup>lt;sup>5</sup> Homology model following scaffolding of AlphaFold predicted *C. elegans* subunits onto the *S. cerevisiae* atomic model without adjusting for dimer angle.

<sup>&</sup>lt;sup>6</sup> Homology model following fitting of dimer angle adjusted *C. elegans* ATP synthase monomers to the *C. elegans* ATP synthase sub-tomogram averaging map.

- Movie S1 (separate file). Movie showing a 360° rotation about the y-axis of a single segmented *C. elegans* mitochondrion from the upper panel of Fig. 3A. An image sequence of 100 PNG files
  was collected in IMOD, and the sequence montaged into a10fps AVI file in Image J (64).
- 843
- 844 Movie S2 (separate file). Movie showing a 360° rotation about the y-axis of a single segmented
- 845 S. cerevisiae mitochondrion from the lower panel of Fig. 3A. An image sequence of 100 PNG files
- was collected in IMOD, and the sequence montaged into a10fps AVI file in Image J (64).

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