Alternative complex III (ACIII) is a key component of the respiratory and/or photosynthetic electron transport chains of many bacteria. Like complex III (also known as the bc$_3$ complex), ACIII catalyses the oxidation of membrane-bound quinol and the reduction of cytochrome $c$ or an equivalent electron carrier. However, the two complexes have no structural similarity. Although ACIII has eluded structural characterization, several of its subunits are known to be homologous to members of the complex iron–sulfur molybdoenzyme (CISM) superfamily, including the proton pump polysulfide reductase. We isolated the ACIII from *Flavobacterium johnsoniae* with native lipids using styrene maleic acid copolymers, both as an independent enzyme and as a functional 1:1 supercomplex with an aa$_3$-type cytochrome $c$ oxidase (cyt aa$_3$). We determined the structure of ACIII to 3.4 Å resolution by cryo-electron microscopy and constructed an atomic model for its six subunits. The structure, which contains a [3Fe–4S] cluster, a [4Fe–4S] cluster and six haem $c$ units, shows that ACIII uses known elements from other electron transport complexes arranged in a previously unknown manner. Modelling of the cyt aa$_3$ component of the supercomplex revealed that it is structurally modified to facilitate association with ACIII, illustrating the importance of the supercomplex in this electron transport chain. The structure also resolves two of the subunits of ACIII that are anchored to the lipid bilayer with N-terminal triacylated cysteine residues, an important post-translational modification found in numerous prokaryotic membrane proteins that has not previously been observed structurally in a lipid bilayer.

The ACIII–cyt aa$_3$ supercomplex from *F. johnsoniae* membranes was solubilized, purified and biochemically characterized using styrene maleic acid (SMA) copolymer nanodiscs without traditional detergents (Supplementary Discussion, Extended Data Figs. 1–3). The supercomplex catalyses the two-electron oxidation of menaquinol (or ubiquinol) and the four-electron reduction of oxygen to water with a turnover number of around 21 electrons per second without the addition of exogenous cyt $c$ (Supplementary Information, Extended Data Fig. 3), indicating a functional electron transfer chain within the supercomplex. The addition of exogenous cyt $c$ did not increase the rate of electron transfer. The structure of the ACIII–cyt aa$_3$ supercomplex in SMA nanodiscs was determined by cryo-electron microscopy (cryo-EM) (Fig. 1, Extended Data Fig. 4). The supercomplex has a mass of 464 kDa (Supplementary Discussion), a transmembrane cross-section of approximately 9 nm × 13 nm (Extended Data Fig. 5), and contains 48 transmembrane $\alpha$-helices. To our knowledge, the ACIII–cyt aa$_3$ supercomplex is the largest protein complex reported to be contained within an SMA copolymer nanodisc. The SMA copolymer and lipids contribute only a thin layer of density around the supercomplex (Fig. 1a, b), which is not circular but follows the contours of the protein.

Whether this is a general feature of SMA-solubilized proteins or is due to the large size of the ACIII–cyt aa$_3$ supercomplex is not known, and will be clarified when more structures are determined using this approach. The number of loosely bound, unresolved lipid molecules is not known, nor is it known whether they are sufficient in number to form a true bilayer surrounding the protein. The SMA–supercomplex nanodiscs retain native lipids, are more stable and have 30% higher specific activity than the supercomplex isolated with detergents (for example, dodecylmaltoside) (Supplementary Discussion, Extended Data Fig. 3). Because traditional detergents are avoided in generating SMA nanodiscs, the preparative protocol is more rapid and simpler than making nanodiscs using membrane scaffold proteins.

Although the properties of the SMA nanodiscs are less well characterized than nanodiscs made with membrane scaffold proteins, our work demonstrates the utility of SMA nanodiscs for high-resolution structural studies of membrane proteins.

The resolution of the cryo-EM density map enabled construction of an atomic model for more than 90% of the sequences predicted from the ACIII gene cluster (Supplementary Discussion), including subunits ActA, ActB, ActC, ActD, ActE and ActF (Fig. 2, Extended Data Fig. 5, Extended Data Table 1). Sequence analysis shows that ACIII contains a unique combination of known modules from other respiratory complexes (Supplementary Discussion). The ACIII structure confirms this prediction and shows the structure responsible for catalysing the quinol:cyt $c$ oxidoreductase activity. The ACIII structure can be divided into three parts: a core assembly of ActC and ActB that oxidizes quinol; a haem $c$ assembly consisting of ActA and ActE that directs electrons from ActB to the terminal electron acceptor; and auxiliary transmembrane subunits ActD and ActF with unknown functions. With some key differences (Extended Data Fig. 5), the overall architecture of ActB and ActC resembles the complex consisting of the PsrA, PsrB and PsrC components of the supercomplex in SMA nanodiscs.
subunits of polysulfide reductase from *Thermus thermophilus* (PsrABC)\(^6\), a member of the CISM superfamily (Supplementary Discussion). Like PsrC, ActC contains no cofactors, but it does contain the proposed site for the oxidation of menaquinol. Residues at the menaquinol-binding site identified in PsrC\(^6\) are not conserved in ActC\(^7\). Although menaquinone is not observed in the cryo-EM map, we propose that the ActC residues His133 and Asp164 form the menaquinol-binding site in ActC near the interface with ActB (Extended Data Fig. 6). These two residues are conserved in ActC sequences and there is a crevice between transmembrane helices 3 (TM3) and 4 (TM4) of ActC that would provide access to the substrate in the membrane bilayer.

The N-terminal portion of ActB is homologous to the PsrA subunit of polysulfide reductase, which contains the molybdenum cofactor, but the molybdenum cofactor is absent in ActB\(^1\). The C-terminal domain of ActB is homologous to PsrB, and both ActB and PsrB contain iron–sulfur clusters. Like PsrB, ActB from *F. johnsoniae* is expected to contain four iron–sulfur clusters, but only two are observed in the cryo-EM map (Extended Data Fig. 7). There is one [3Fe–4S] cluster near the interface with ActC, about 10 Å from the proposed site of menaquinol oxidation, and one [4Fe–4S] cluster about 9 Å further away. There are two additional cysteine clusters present in the structure of ActB, but the cryo-EM map does not show iron–sulfur clusters at these locations. Instead, we observe disulfide bonds (Cys965–Cys938 and Cys971–Cys769) within these two cysteine clusters in ActB. The substitution of proposed [4Fe–4S] clusters by disulfide bonds may be a genuine aspect of the structure or may result from oxidation that occurred during sample preparation. However, if these two ’missing’ [4Fe–4S] clusters were present, they would form a dead-end for electron transfer from the [3Fe–4S] cluster of ActB, suggesting that their absence from the structure is not an artefact.

The [3Fe–4S] cluster in ActB is the most probable initial oxidant of menaquinol bound to ActC, and is 12.3 Å from the nearest haem c in ActA. The five haems c in ActA plus the single haem c in ActE form a probable electron transfer wire from the [3Fe–4S] cluster in ActB, with the largest edge-to-edge distance of 9.2 Å between adjacent haems (Fig. 2b). The [4Fe–4S] cluster in ActB appears to be off-pathway and its function remains to be determined.

In all *Flavobacteria*, including *F. johnsoniae*, ActA is predicted to have a monohaem domain at the N terminus in addition to the pentahaem domain at the C terminus (Supplementary Discussion). Mass spectrometry analysis shows that the N-terminal monohaem domain is present in the preparation (Extended Data Fig. 1), but no density can be assigned to this entire domain. The inability to resolve the monohaem domain may result from flexibility of the domain. Full-atom molecular dynamics simulations were performed for the entire structure of ACIII embedded in a phospholipid bilayer to determine the stability and dynamics of the structure (Extended Data Fig. 8). Notably, the pentahaem domain of ActA had the largest root-mean-square deviation (r.m.s.d.), which arises mainly from the transmembrane α-helix connected to the missing monohaem domain; this is consistent with the monohaem domain being unobservable owing to a variable position in the complex. Although ActE also had a substantial r.m.s.d., it did not appear to correlate with disorder in the cryo-EM map.

ActD and ActF are transmembrane subunits without bound cofactors, and both interact with ActC. It has not been established whether ACIII generates a proton motive force coupled to electron transport\(^8\). The absence of redox centres in ActC, ActD and ActF suggests that if ACIII contributes to the transmembrane proton gradient, it does not use the bifurcation-type Q-cycle mechanism of canonical complex III\(^9\), but instead functions as a true proton pump with a mechanism that resembles that of complex I\(^10\). ActD has two transmembrane α-helices that cross within the membrane and are adjacent to ActC. Both N and C termini are within the cytoplasm and combine to form a single globular domain that rests on the cytoplasmic surface of ActC. The ten transmembrane α-helices of ActD form a pseudo two-fold axis of symmetry with the ten transmembrane α-helices of ActC (Extended Data Fig. 5), despite the fact that ActF has less than 20% sequence identity with ActC. If ACIII is a proton pump, it is likely that conserved polar residues within the bilayer will have important roles.

The structure of ACIII reveals eleven ordered phospholipid molecules as well as triacylated cysteine residues at the N termini of ActB (Fig. 3a) and ActE (Extended Data Fig. 7). The anchoring of bacterial membrane proteins by an N-terminal triacylated cysteine is a well-characterized phenomenon\(^11\); however, to our knowledge, this is the first time the structure of a triacylated cysteine residue has been determined in the context of a membrane protein. Both lipid anchors are tilted with respect to the plane of the lipid bilayer (Fig. 2a), restricting the ability of other lipids to pack around them. This feature could alter the mechanical properties of the adjacent portion of the membrane bilayer, and also guide conformational changes in the ACIII protein. Notably, the two N-terminal lipid anchors are adjacent to each other in the membrane. These lipid anchors probably help ACIII to assemble and keep the monohaem ActE bound to the complex. The eleven lipids that are resolved adjacent to the transmembrane α-helices accommodate the rugged protein surface of the complex (Fig. 2a, Extended Data Fig. 2). The head groups of the lipids could not be identified and were all modelled as phosphatidylethanolamine. There are two ’hot spots’ for resolved lipids: the cytoplasmic interface between ActC and ActF; and the vicinity of the triacylated cysteine of ActB, which is near the proposed entry point for menaquinol into the complex. All eleven of the resolved lipids remained bound to the protein throughout 250 ns of molecular dynamics simulation (Extended Data Fig. 8), supporting the ability of SMA nanodiscs to preserve some native lipid–protein interactions and suggesting a functional role for the lipids. A large number of annular lipids, including those modelled in the structure, were observed to associate with the protein from the in silico bilayer.

Frequently, the subunits encoding ACIII are within an operon that includes subunits of an associated complex IV\(^3\) (cyt aa\(_3\) or cyt caa\(_3\)). We find that the sequences of subunit III from complex IVs that are associated with ACIII have unusual features that distinguish them from the canonical subunit III (Supplementary Discussion).
Whereas subunit III of complex IV generally contains seven transmembrane α-helices, those that are associated with ACIII lack TM1 and TM2 (Fig. 4a). Although only parts of subunit III of cyt aa₃ are resolved to better than 4 Å, the density for cyt aa₃ has sufficient resolution to identify five α-helices from the structure. A homology model of subunit III from F. johnsoniae cyt aa₃ was built on the basis of the structure of TM3 to TM7 of subunit III from Rhodobacter sphaeroides cyt aa₃, and fit into the ACIII–cyt aa₃ supercomplex density map (Extended Data Fig. 9) with high fidelity. The deletion of the first two transmembrane α-helices in subunit III of cyt aa₃ appears to be a necessary adaptation to enable formation of the supercomplex with ACIII. It is notable that the same two helices in subunit III are also absent in the cyt aa₃ obligatory cyt bcc–cyt aa₃ supercomplex found in Actinobacteria (for example, Corynebacterium glutamicum and Mycobacterium tuberculosis)²⁵.

The sequence analysis also reveals that the loop between TM5 and TM6 of subunit III in the cyt aa₃ that is part of the supercomplex is much longer in F. johnsoniae (and all Flavobacteria) than in other organisms. Typically, this loop contains eight residues, but in F. johnsoniae it contains 121 residues (Fig. 4a). Part of this long loop fits in a groove between ActB and ActD of ACIII on the periplasmic side of the membrane (Extended Data Fig. 9). The structural model reveals a π–cation interaction between Trp188 of subunit III and Arg868 of ActB (Fig. 4b), both of which are conserved among organisms containing subunit III with a long loop between TM5 and TM6 (Extended Data Fig. 9). This specific and strong interaction stabilizes the ACIII–cyt aa₃ supercomplex and appears to be a second adaptation that enables the formation of a supercomplex with ACIII. The contact between the periplasmic loop of subunit III of cyt aa₃ and ACIII is the only observed direct contact between the two complexes. The five well-resolved transmembrane α-helices of subunit III of cyt aa₃ are angled away from ACIII with only the tip of TM6 of subunit III touching ActE, forming a wedge-like space between the membrane domains of ACIII and cyt aa₃. The decrease of resolution in the portions of cyt aa₃ that are distant from the interface with ACIII suggests that there may be several conformations of the supercomplex that are all tethered by the loop in cyt aa₃. The loop could, therefore, serve as a hinge, enabling the membrane domains of ACIII and cyt aa₃ to swing into contact transiently.

Using the location of TM3 to TM7 of subunit III within the supercomplex as a guide enables a model of the entire cyt aa₃ to be placed within the density map for the supercomplex (Extended Data Fig. 9).

In the resulting model, there is a considerable distance (56 Å) between the haem c in ActE of ACIII and Cu₄ within subunit II of cyt aa₃. Electron transfer within the supercomplex does not require the addition of exogenous cyt c, which is also the case for the cyt bcc–cyt aa₃ supercomplex from C. glutamicum²⁵. It is possible, although it seems unlikely, that there is a subset of conformations in which ActE comes close enough to cyt aa₃ for direct electron transfer. It is noteworthy that the monohaem domain of ActA has substantial sequence homology (around 30% identity) with the haem c domain that is present at the C terminus of subunit II of cyt caa₃ from T. thermophilus. This observation suggests that the ActA monohaem domain within ActE of ACIII and Cu₄ within subunit II of cyt aa₃. As such, electron transfer within the supercomplex may require the monohaem domain of ACIII to swing back and forth between ACIII and cyt aa₃, allowing for electron transfer to and from interface sites on ActE from both ACIII and cyt aa₃. Moreover, the ActA monohaem domain from one complex can interact with subunit II of cyt aa₃ from the other, facilitating electron transfer between the two complexes (Fig. 4c).

Additional experimental work will be required to test this model and, indeed, to determine the physiological advantage of forming the supercomplex.

We would like to note that, contemporaneously with our studies, Sousa et al.²⁴ determined the structure of the homologous ACIII from Rhodothermus marinus by cryo-EM at 3.9 Å resolution. Aside from species-specific variations, the reported structures of the ACIIIs of F. johnsoniae and R. marinus are compatible. The observation made here that, like canonical complexes III and IV, ACIII and an adapted complex IV can also form a supercomplex hints at the importance of supercomplexes in oxidative phosphorylation. We demonstrate that high-resolution cryo-EM with SMA nanodiscs, which preserves native protein–protein and protein–lipid interactions, is ideally suited to have an important role in future studies.

**Online content**

Any Methods, including any statements of data availability and Nature Research reporting summaries, along with any additional references and Source Data files, are available in the online version of the paper at https://doi.org/10.1038/s41586-018-0061-y.
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METHODS

Bacterial strain and growth conditions. Flavobacterium johnsoniae ATCC 17061 strain UW101 was used in this study. The strain was a gift from M. McBride at the University of Wisconsin, Milwaukee. The cells were grown in casitone-extract medium at 30 °C under high aerobic conditions (500 mL cultures in 2 L flasks) [10].

Membrane preparation and protein purification. Cells grown overnight were collected by centrifugation (14,000g for 10 min). The cell pellet from 121 of culture (2.5 × 10^8 cells) was resuspended in ~200 mL of 20 mM Tris-HCl buffer, pH 8 (buffer A) with 5 mM MgSO_4, DNase I (Sigma) and a protease inhibitor cocktail (Sigma). This suspension was passed three times through a Microfluidizer at a pressure of 80,000 psi to disrupt the cells. The cell extract was centrifuged at 14,000g for 10 min to remove unbroken cells. Membranes were obtained after centrifugation at 185,500 × g for 4 h. Under the above growth conditions, the membranes contained ACIII, cyt aa_3 and cyt bd. The membrane pellet was solubilized by using either a traditional detergent or the SMA copolymer.

Purification using Triton X-100 and DDM. The membrane pellet was resuspended in buffer A (5–50 mg mL⁻¹) along with 300 μM NaCl, and solubilized by the addition of Triton X-100 (Fisher Scientific) to a final concentration of 4%. The solution was incubated at 4 °C for 2 h with mild agitation. The suspension was cleared by centrifugation at 185,500 × g for 1 h, after which the detergent was diluted fourfold by adding two volumes of buffer A to the supernatant. The diluted supernatant was then added to a chromatography column containing 10 mL of Ni–NTA resin (Qiagen) pre-equilibrated with 20 mM Tris-HCl pH 8 containing 0.5% Triton X-100 and 0.15 M NaCl (buffer B). The resin was washed with about ten column volumes of buffer B to remove any unbound sample. Detergent exchange to n-dodecyl-β-D-maltoside (DDM; Anatrace) was carried out by exchanging buffer B containing 0.05% DDM instead of Triton X-100 (buffer C). The column was further washed with five column volumes of buffer C containing 10 mM imidazole to remove the loosely bound proteins from the resin. The proteins that were well bound to the resin were eluted using 100 mM imidazole in buffer C. The eluent was concentrated to around 3 mL using Amicon Ultra-15 filters (Millipore) with a 100-KDa cutoff. The excess imidazole was removed by dialysis against buffer C. The yield of protein obtained was about 0.3 mg mL⁻¹ of ACIII and 0.16 mg mL⁻¹ of cyt aa_3 from 121 of culture. When indicated, the proteins were further purified by gel filtration chromatography using a Superdex 200 10/300 GL column (GE Healthcare-Life Sciences). The purified proteins were stored at −80 °C after adding glycerol to a final concentration of 10%.

Purification using SMA copolymer. The SMA copolymer SMA 3000HNA (styrene maleic acid copolymer, ~3:1 molar ratio of styrene: maleic acid) was a gift from T. Bricker (Louisiana State University) who used SMA copolymer made by Cray Valley USA (now Total Petrochemicals & Refining USA) successfully for the purification of ACIII–cyt aa_3 supercomplex [15]. The purified supercomplex was suspended in 4 mL of 50 mM potassium phosphate buffer pH 7.0 to a concentration of 3 μM with 25 μM each of the following redox mediators: benzyl viologen (Em = −350 mV), anthraquinone-2-sulfonate (Em = −225 mV), 2-hydroxy-1,4-naphthoquinone (Em = −220 mV), 9,10-anthraquinone-2,6-disulfonate (Em = −185 mV), duroquinone (Em = −180 mV), N-ethylphenazonium ethosulfate (Em = −65 mV), N-phenylbenzazonium methosulfate (Em = −85 mV), diaminodurene (Em = −275 mV), 2,6-dimethyl benzoquinone (Em = −180 mV), 1,2-naphthoquinone (Em = −143 mV), 1,4-naphthoquinone (Em = −36 mV) and potassium ferricyanide (Em = −435 mV) [30]. Titrations were performed with an Oxygraph+ (Oxymax, Oxford Instruments). The solution potential was adjusted by injecting aliquots of 10 mM sodium dithionite or potassium ferricyanide as reductant and oxygen, respectively. Spectra were taken at approximately 10–20-mV increments over the titration range indicated. Spectroscopic changes of the α-bands of the haems upon reduction or oxidation were monitored at the peak maxima to determine the midpoint potentials of each class of haem centre. The datasets were analysed using Origin (Origin Laboratory Corporation) to determine spectral components and fit titration curves using the Nernst equation [26].

Electron microscopy sample preparation. Holey carbon film–coated electron microscopy grids were nanostructured with regular arrays of 500–800-nm holes [17] and coated with an additional layer of gold. Cryo-EM specimens were prepared with a FEI Vitrobot grid preparation robot at 4°C and 100% humidity by applying 3 μL of sample (3 mg mL⁻¹) to glow-discharged grids, allowing the grids to equilibrate for 1 s, and blotting for 12 s before freezing in a liquid ethane-propylene mixture (1:1 v/v) [18]. grids were subsequently stored in liquid nitrogen before shipping to the New York Structural Biology Center for imaging with a FEI Titan Krios electron microscope equipped with a Gatan K2 Summit camera and automated with Leginon [19].

Electron microscopy data acquisition. Movies were acquired in electron counting mode with a pixel size of 1.1 Å, an exposure rate of 7.4 electrons per pixel per second, and a total exposure time of 10 s divided in 40 frames (418 movies) or 50 frames (1,599 movies). Frame alignment and exposure weighting were performed with Motioncor2 [40]. After screening averages from the aligned movies, 475 movies were discarded because of excessive movement, low defocus, high defocus, or overfocus. Contrast transfer function parameters were estimated from the exposure-weighted averages of movie frames with CTFIND4 [41].

Image processing. Particle images (3,044) were manually selected and subjected to 3D classification with Relion 1.4 [22]. The resulting 2D classes were used as templates for automatic selection of 899,405 particle images [3]. The number of particle images was reduced to 693,416 by further 2D classification. Subsequent image processing was carried out in cryoSPARC [44]. An initial map of ACIII–cyt aa_3 was obtained...
by ab initio 3D classification, refined to 4.1 Å resolution, and used as a reference for the multi-refine procedure in cryoSPARC producing initial maps of the ACIII and the ACIII–cyt a3. Particle images (164,239) were used to refine the ACIII–cyt a3 map to 3.4 Å resolution, but this map showed the cyt a3 portion of the complex with lower density than the ACIII part. Maps with uniform density for ACIII–cyt a3 and ACIII, both at 3.6 Å Resolution, were calculated from 81,530 and 51,547 particle images, respectively.

Model building. The 3.4 Å ACIII resolution density map was used for the de novo model building of ACIII. The density map was first segmented with UCSF Chimera45 to facilitate the identification of subunits. The connectivity of each segmented map was further examined and the result was compared with topology predictions from topcons46 and secondary structure prediction from Jpred47 to validate the subunit assignment and identify the directionality of peptide chain. With this information, model building was carried out manually in Coot48. Individual chains were first traced in Coot’s cartoon mode. Readily interpretable features from the density map, including regions rich in bulky residues, triacylated cysteines, and axial ligands of haem c, were used to register the structure to the sequence. Stretches of ~20 amino acids were built progressively around these registration points and assembled as a single chain in Coot. All six subunits of ACIII were combined and refined with phenix.real_space_refine51. For cofactors, the starting models were taken from the CCP4 ligand library directly. Cofactors were docked to the density map with Coot and merged with the apo protein structure. The complete structure was then refined with phenix.real_space_refine with geometric constraints for the protein–cofactor coordination. The final model was further examined in Coot to remove amino acid side chains with ambiguous orientations and further validated with MolProbity52. All identified lipids with CO epitopes were modelled as phosphatidylethanolamine with palmitoyl tail. The conformation of phosphatidylethanolamine was refined with interactive molecular dynamics flexible fitting (iMDFF) in the presence of the protein structure using VMD52. Lipid tails were then truncated according to the density map.

The 3.6 Å ACIII–cyt a3 density map was used for the model building of cyt a3. Part of the subunit III loop region was manually built in Coot. Homology models for individual subunits were generated with the RaptorX server53 and docked into the density map with UCSF Chimera. The model for the ACIII–cyt a3 supercomplex was assembled by fitting the ACIII structure to the ACIII–cyt a3 map and placing the cyt a3 structure from Rba. sphaeroides (PDB 1M56) into the map based on the position of cyt a3 subunit III.

Bioinformatic analysis. Homologous protein sequences were retrieved using the NCBI blastp server44. The blastp results were analysed in python 2.7 with pandas and biopython modules. Sequence hits were filtered on the basis of coverage and sequence identity. Representative sequences were selected on the basis of sequence identity to maintain the variations in sequence and aligned using the Clustal Omega server54. Figures for sequence alignment were prepared using the ESPript server55.

Simulation system preparation. The initial ACIII structure for the MD simulation was obtained from the refined structure determined by cryo-EM. Eleven α-glycerol-1,2-diol-3-phosphoethanolamine (POPE) lipids resolved in the map based on the position of cyt aa3 within the ACIII complex. The connectivity of each segmented map was further examined and the result was compared with topology predictions from topcons46 and secondary structure prediction from Jpred47 to validate the subunit assignment and identify the directionality of peptide chain. With this information, model building was carried out manually in Coot48. Individual chains were first traced in Coot’s cartoon mode. Readily interpretable features from the density map, including regions rich in bulky residues, triacylated cysteines, and axial ligands of haem c, were used to register the structure to the sequence. Stretches of ~20 amino acids were built progressively around these registration points and assembled as a single chain in Coot. All six subunits of ACIII were combined and refined with phenix.real_space_refine51. For cofactors, the starting models were taken from the CCP4 ligand library directly. Cofactors were docked to the density map with Coot and merged with the apo protein structure. The complete structure was then refined with phenix.real_space_refine with geometric constraints for the protein–cofactor coordination. The final model was further examined in Coot to remove amino acid side chains with ambiguous orientations and further validated with MolProbity52. All identified lipids with CO epitopes were modelled as phosphatidylethanolamine with palmitoyl tail. The conformation of phosphatidylethanolamine was refined with interactive molecular dynamics flexible fitting (iMDFF) in the presence of the protein structure using VMD52. Lipid tails were then truncated according to the density map.

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Extended Data Fig. 1 | Expression and spectroscopic characterization of the ACIII-cyt aa3 supercomplex. a, A schematic of the respiratory chain of F. johnsoniae. b, UV–visible spectrum and SDS–PAGE of the membranes from F. johnsoniae. Left, the difference spectrum of the membranes of F. johnsoniae, obtained from the spectrum of the air-oxidized membranes and the spectrum after reduction with dithionite. The wavelengths associated with the haem peaks are 605 nm, 560 nm, 552 nm and a broad peak at 630 nm for haems a, b, c and d, respectively. Right, the SDS–PAGE with the membranes followed by staining the gel for haems shows bands corresponding to the cytochrome subunits ActA (48 kDa) and ActE (20 kDa) of ACIII but no bands corresponding to the cytochrome subunit (around 35 kDa) from the cbb3 oxidase. c, The gene arrangement for the ACIII and the cytochrome oxidase aa3 genes in the F. johnsoniae genome. The genes for the subunits I and II from cyt aa3 oxidase are found immediately downstream of those for the act genes of the ACIII. Two different versions of subunit III are denoted as vI and vII. d, UV–visible spectra of the reduced and oxidized forms of the supercomplex in detergent and SMA nanodiscs. The dithionite reduced form of the samples is represented in red and shows the peaks for haem c at 524 nm and 552 nm and those for haem a at 443 nm and 605 nm. e, Pyridine haemochrome assay of the ACIII–cyt aa3 supercomplex in SMA nanodiscs. Plotted is the reduced-minus-oxidized difference spectrum of the pyridine haemochromes of the sample. Peaks at 520 nm and 550 nm are associated with haem c and the peak at 590 nm is associated with haem a. Quantification from the spectrum shows a ratio of 10.6:1 between haem c and haem a, which translates into a 3:2 ratio between ACIII and cyt aa3 oxidase. Data in b are representative of two independent experiments with similar results, and data in d and e are representative of six independent experiments with similar results.
Extended Data Fig. 2 | Component and size analysis of ACIII–cyt aa₃ supercomplex. a, SDS–PAGE of the detergent-solubilized preparation followed by Coomassie staining (left) and haem staining (right). b, SDS–PAGE of the SMA nanodiscs preparation followed by Coomassie staining (left) and haem staining (right). c, Mass spectrometry results for the ACIII–cyt aa₃ supercomplex preparations. d, Size-exclusion chromatography with the ACIII–cyt aa₃ supercomplex from F. johnsoniae. Top left, the chromatogram of the detergent-solubilized sample, showing traces for protein at 280 nm, haem c at 412 nm and haem a at 443 nm respectively. Top right, the chromatogram of the sample isolated using the SMA copolymer, showing traces for protein at 280 nm, haem c at 410 nm and haem a at 605 nm. I and II are the two peaks corresponding to two populations of the supercomplex. Bottom left, chromatogram of the fraction containing peak I. Bottom right, chromatogram of the fraction containing peak II. e, BN–PAGE of the ACIII–cyt aa₃ supercomplex. Left, the detergent-solubilized ACIII–cyt aa₃ supercomplex, showing a band at around 500 kDa, a smear of possible aggregates and possibly ACIII by itself. Right, the supercomplex in SMA nanodiscs, showing two different populations. f, BN–PAGE with the two different populations of ACIII–cyt aa₃ supercomplex in SMA nanodiscs purified from size-exclusion chromatography. The two chromatographic peaks correspond to the two bands observed in the BN–PAGE. Data in a, b are representative of six independent experiments and those in d–f are representative of three independent experiments with similar results.
Extended Data Fig. 3 | Functional assays of the ACIII–cyt aa3 supercomplex. 

**a**, The EPR spectrum of the air-oxidized sample showing peaks of the [3Fe–4S]$^+$ cluster from ACIII, the Cu4 from the cyt aa3 oxidase and low-spin haems with overlapping g values. Insert is a zoomed view from 3,000 G to 3,500 G to better visualize the peaks from Cu4 (black arrows) and the [3Fe–4S]$^+$ cluster. The region between 4,000 G and 5,000 G is magnified ten times to show the broad g trough of low-spin haems. The measurement condition is 10 K, 9.267 GHz, 2 mW microwave power and 20 Gauss modulation. 

**b**, The EPR spectra of the ferricyanide-oxidized sample at various temperatures. The measurement condition is 9.257 GHz, 2 mW microwave power and 5 Gauss modulation. 

**c**, The EPR spectrum of the air-oxidized sample showing peaks of iron–sulfur clusters from ACIII and low-spin haems. The measurement condition is 10 K, 9.427 GHz, 2 mW microwave power, 10 Gauss modulation. 

**d**, The EPR spectra of the air-oxidized sample at various temperatures. The measurement condition is 9.427 GHz, 2 mW microwave power, 5 Gauss modulation. 

**e**, Redox titration of the haems in the ACIII and the cyt aa3 oxidase in supercomplex in DDM. The potentiometric titration of the c haems from the ACIII (top) and the a haems from the cyt aa3 oxidase (bottom). The $E_m$ values are indicated and the solid red line represents the Nernst fitting. 

**f**, Steady-state activity of the ACIII–cyt aa3 preparations. The number of independent experiments is six for ACIII in DDM and SMA nanodiscs, and three for peak I and peak II. Data are means ± s.d. Data in **a**–**e** are representative of three independent experiments with similar results.
Extended Data Fig. 4 | Single-particle cryo-EM of the ACIII–cyt aa₃ supercomplex in SMA nanodiscs. a, Sum of an aligned movie of the ACIII–cyt aa₃ supercomplex in an SMA nanodisc. Scale bar, 20 nm. b, Two-dimensional class averages. Scale bar, 10 nm. c, Fourier shell coefficient curves between two independently refined half-maps for the ACIII–cyt aa₃ map, ACIII map and combined map. d, Surface rendering maps coloured according to local resolution. Scale bar, 5 nm. e, Euler angle distributions of particles included in the calculation of the three final maps. Data collection and structure calculation were not repeated.
Extended Data Fig. 5 | Features observed in the cryo-EM density and the de novo structure of ACIII. a. Surface representations of ACIII, cyt $aa_3$ and the ACIII–cyt $aa_3$ supercomplex. The density threshold is the same for ACIII and cyt $aa_3$. b. Different views of the ACIII density, coloured by subunit. c. Two single-span transmembrane peptides of unknown origin and sequence, denoted ActX and ActY, are present in the structure in the vicinity of ActC. These have each been modelled as a polyalanine peptide. d. $\alpha$-helices 2–10 of ActC form two four-helical up-and-down bundles, coloured in two different shades of blue. e. ActB, shown in cartoon form, has contact with ActA, ActC, ActD, ActE and ActF. Surfaces are drawn from residues that are within 4 Å of ActB and coloured according to their chain. f. The transmembrane $\alpha$-helices of ActC and ActF are arranged in a pseudo two-fold rotation symmetry. g. Side-by-side comparison of the polysulfide reductase (PDB 2VPZ) and the assembly of ActB and ActC. These two structures are aligned based on PsrB, the domain containing four iron–sulfur clusters.
Extended Data Fig. 6 | The proposed quinone pocket in ActC.

a, Sequence alignment of the ActC from *F. johnsoniae*, *R. marinus*, and *Chloroflexus aurantiacus*. The transmembrane α-helices are labelled based on the structure of ACIII from *F. johnsoniae*. The black arrows point to conserved polar residues that are within 15 Å of the [3Fe–4S] cluster in ActB.

b, Proposed quinone pocket based on the arrangement of conserved polar residues. c, Different views of the proposed quinone pocket with a docked menaquinone-1 molecule. Hydrophobic residues near the menaquinone-1 (MK1) head group are also shown. The crevice between α-helix 3 and α-helix 4 is a putative quinone entry pathway.
Extended Data Fig. 7 | Fitting of the ACIII structure to cryo-EM density. 

a, Fitting of cofactors into the cryo-EM density. The blue mesh is drawn with a higher density threshold to reveal metal centres. The numberings of nearby amino acid residues, which are shown along with these cofactors, are listed below each cofactor. 
b, Fitting of different secondary structure elements to cryo-EM density. 
c, Eleven identified lipids are modelled as phosphatidylethanolamine molecules. 
d, The triacylated cysteine at the N terminus of ActE shown along with 15 downstream amino acids. Notably, residue Tyr28 is in contact with the covalent lipid of ActE. Attachment of ActE to the membrane may also be assisted by aromatic residues Tyr30 and Phe31, which appear to be inserted into the lipid bilayer. Throughout the molecular dynamics simulation trajectory, these residues remain buried in the lipid bilayer.
Extended Data Fig. 8 | Protein stability and lipid–protein interaction analysis based on molecular dynamics simulations. a, Root-mean-square deviation (r.m.s.d.) of the protein backbone heavy atoms for the entire ACIII complex and each subunit, aligned based on ACIII backbone heavy atoms from three independent molecular dynamics simulations.  

b, Same as a, but aligned using the backbone heavy atoms of each subunit.  

c, Superposition of the initial (black) and final (coloured) conformations of each subunit after 250 ns of simulation (aligned using backbone heavy atoms).  

d, The lipid–protein contact number defined by the number of lipid atoms within 4 Å of the protein atoms calculated over the time course of the simulation. This contact number is either calculated for the eleven lipids resolved by cryo-EM (top) or all membrane lipids (bottom).  

e, The lipid–protein contact number for each of the eleven cryo-EM resolved lipids.  

f, Isosurfaces (50%) of the atom-occupancy map for the lipid anchors (orange), cryo-EM resolved lipids (red) and other membrane lipids (purple), calculated using the last 230 ns of the simulation trajectory. The stronger the lipid–protein interactions, the longer the local residence time, which leads to higher atom-occupancy values. ACIII subunits C, D, and F are shown in silver. For all plots, the raw data are shown as translucent thin lines and the block-averages are shown as dark lines.
Extended Data Fig. 9 | Structural basis for supercomplex formation between the ACIII and the cyt aa₃. a, Two contact areas between the ACIII and the cyt aa₃, the transmembrane portion of subunit III (red) and the loop from subunit III (orange). A homology model of subunit III fits the transmembrane density. The loop is modelled to the cryo-EM density. The sequence of the peptide is also shown. Trp188 and Phe189 are used to register the density to the sequence. b, Model of the ACIII–cyt aa₃ supercomplex. The cyt aa₃ structure from *Rba. sphaeroides* was positioned based on the transmembrane portion of subunit III. *α*-helices 1 and 2 of subunit III are omitted to avoid steric clashes with the ACIII structure. c, Sequence alignment of subunit III with a long loop (highlighted with the orange bar) between *α*-helix 5 and *α*-helix 6 (numbered according to subunit III from *Rba. sphaeroides*). Trp188 (red arrow) is conserved. d, Sequence alignment of ActB from organisms with a long loop in subunit III of their cyt aa₃ oxidase. Arg868 (red arrow) is largely conserved with occasional substitution to lysine.
## Extended Data Table 1 | Cryo-EM data collection, refinement and validation statistics

|                                | Combined (EMDB-7286) (PDB 6BTM) | ACIII-cyt aa₃ (EMDB-7447) | ACIII (EMDB-7448) |
|--------------------------------|---------------------------------|---------------------------|-------------------|
| **Data collection and processing** |                                 |                           |                   |
| Magnification                  | 75,000x                         | 75,000x                   | 75,000x           |
| Voltage (kV)                   | 300                             | 300                       | 300               |
| Electron exposure (e⁻/Å²)      | 61                              | 61                        | 61                |
| Defocus range (µm)             | 0.8-5.0                         | 0.8-5.0                   | 0.8-5.0           |
| Pixel size (Å)                 | 1.1                             | 1.1                       | 1.1               |
| Symmetry imposed               | C1                              | C1                        | C1                |
| Initial particle images (no.)  | 899,405                         | 899,405                   | 899,405           |
| Final particle images (no.)    | 164,239                         | 81,530                    | 51,547            |
| Map resolution (Å)             | 3.4*                            | 3.6*                      | 3.6*              |
| FSC threshold                  | 0.143                           | 0.143                     | 0.143             |
| Map resolution range (Å)       | 3.0-4.5                         | 3.0-6.0                   | 3.0-5.0           |
| **Refinement**                 |                                 |                           |                   |
| Initial model used (PDB code)  | N/A                             | N/A                       | N/A               |
| Model resolution (Å)           | 3.7†                            | N/A                       | N/A               |
| FSC threshold                  | 0.5                             |                           |                   |
| Model resolution range (Å)     | 3.7                             | N/A                       | N/A               |
| Map sharpening B factor (Å²)   | -150.1                          | -132.4                    | -129.1            |
| Model composition              |                                 |                           |                   |
| Non-hydrogen atoms             | 18,935                          |                           |                   |
| Protein residues               | 2,361                           |                           |                   |
| Ligands                        | 10                              |                           |                   |
| B factors (Å²)                 |                                 |                           |                   |
| Protein                        | 126.4†                          |                           |                   |
| Ligand                         | 119.7†                          |                           |                   |
| R.M.S. deviations              |                                 |                           |                   |
| Bond lengths (Å)               | 0.009                           |                           |                   |
| Bond angles (°)                | 1.26                            |                           |                   |
| **Validation**                 |                                 |                           |                   |
| MolProbity score               | 1.36                            |                           |                   |
| Clashscore                     | 1.58                            |                           |                   |
| Poor rotamers (%)              | 0.15                            |                           |                   |
| Ramachandran plot              |                                 |                           |                   |
| Favored (%)                    | 92.7                            |                           |                   |
| Allowed (%)                    | 7.3                             |                           |                   |
| Disallowed (%)                 | 0.0                             |                           |                   |

* Determined with cryoSPARC
† Determined with Phenix.mtriage
‡ Mean value of the B factors determined with Phenix.real_space_refine
Life Sciences Reporting Summary

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- Experimental design

1. Sample size
   - Describe how sample size was determined.
   - No statistical methods were used to predetermine sample size. The sample size was chosen following the guidelines from the instrument manufacturer.

2. Data exclusions
   - Describe any data exclusions.
   - A fraction of the acquired cryo-EM movies were discarded because of excessive movement, low defocus, high defocus, or over-focus.

3. Replication
   - Describe whether the experimental findings were reliably reproduced.
   - cyro-EM Data collection and structure calculation were not repeated.

4. Randomization
   - Describe how samples/organisms/participants were allocated into experimental groups.
   - Samples were not allocated into experimental groups because the goal of this study was not to evaluate the impact of a particular factor, but to simply determine the structure of a protein.

5. Blinding
   - Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   - Not applicable. There was no existing protein structure to refer to and the data analysis was ab initio.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
   - For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).
   - n/a Confirmed
   - □ The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
   - □ A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
   - □ A statement indicating how many times each experiment was replicated
   - □ The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
   - □ A description of any assumptions or corrections, such as an adjustment for multiple comparisons
   - □ The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
   - □ A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
   - □ Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.
7. Software

Describe the software used to analyze the data in this study.

Legion as of February 2017 used in Simons Electron Microscopy Center at New York to collect cryo-EM movies; Motioncor2 to align frames and weigh exposure; CTFFIND4 to estimate CTF parameters; Relion 1.4, cryoSPARC 0.3.2 to process images and reconstruct cryo-EM density maps; Jpred 4 server to predict protein secondary structures; Topocons server to predict membrane protein topology; RaptorX server to build homology-based protein structures; NCBI blastp server to retrieve homologous protein sequences; python 2.7.13, pandas 0.20.1, biopython 1.68 to analyze/filter protein sequences; Clustal Omega server to align protein sequences; ESPript 3.0 server to represent protein sequence alignments; UCSF Chimera 1.12, Coot 0.8.8 EL (ccp4) to examine density map and build atomic structures; MDFF, Phenix.real_space_refine (Phenix ver-2722), modelmaker (beta version) to refine atomic structures; EMringer, MolProbity and Phenix.mtriage (Phenix ver-2998) to validate atomic structures; namd 2.13 to run molecular dynamics simulations; VMD 1.9.4, Origin 9.1, matplotlib 1.5.1, Adobe Illustrator CC 22.0.1 to prepare figures.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

8. Materials and reagents

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

- **Antibodies**

  No antibodies were used.

- **Eukaryotic cell lines**

  - no eukaryotic cell lines were used
  - no eukaryotic cell lines were used
  - no eukaryotic cell lines were used
  - no eukaryotic cell lines were used

- **Animals and human research participants**

  Provide details on animals and/or animal-derived materials used in the study.

  no animals were used in the study.
12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

This study did not involve human research participants.