Structures of Respiratory Supercomplex I+III₂ Reveal Functional and Conformational Crosstalk

Highlights
- CoQ trapping within isolated respiratory supercomplex I+III₂ limits complex I turnover
- CryoEM structures of multiple 3D classes show crosstalk between complex I and III₂
- Key transmembrane helix in complex I rotates upon “closed” to “open” state transition
- CoQ density at only three complex III₂ sites indicates symmetry breaking

Authors
James A. Letts, Karol Fiedorczuk, Gianluca Degliesposti, Mark Skehel, Leonid A. Sazanov

Correspondence
sazanov@ist.ac.at

In Brief
The first chromatographic isolation of functional mammalian mitochondrial respiratory supercomplex I+III₂ (SC I+III₂) provides a new framework for studying the structure, function, and physiology of respiratory supercomplexes.

Data Resources
6Q9D 6QC5
6Q9B 6QC6
6QE9 6QC8
6QBX 6QC7
6QC3 6QC9
6QC2 6QCA
6QC4 6QCF
6QA9
Structures of Respiratory Supercomplex I+III₂
Reveal Functional and Conformational Crosstalk

James A. Letts,¹,² Karol Fiedorczuk,¹,³ Gianluca Degliesposti,⁴ Mark Skehel,⁴ and Leonid A. Sazanov¹,⁵,*

¹Institute of Science and Technology Austria, Klosterneuburg 3400, Austria
²Department of Molecular and Cellular Biology, University of California, Davis, Davis, CA 95616, USA
³Laboratory of Membrane Biophysics and Biology, The Rockefeller University, New York, NY 10065, USA
⁴MRC Laboratory of Molecular Biology, Cambridge CB2 0QH, UK
⁵Lead Contact
*Correspondence: sazanov@ist.ac.at
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SUMMARY

The mitochondrial electron transport chain complexes are organized into supercomplexes (SCs) of defined stoichiometry, which have been proposed to regulate electron flux via substrate channeling. We demonstrate that CoQ trapping in the isolated SC I+III₂ limits complex (C)I turnover, arguing against channeling. The SC structure, resolved at up to 3.8 Å in four distinct states, suggests that CoQ oxidation may be rate limiting because of unequal access of CoQ to the active sites of CIII₂. CI shows a transition between “closed” and “open” conformations, accompanied by the striking rotation of a key trans-membrane helix. Furthermore, the state of CI affects the conformational flexibility within CIII₂, demonstrating crosstalk between the enzymes. CoQ was identified at only three of the four binding sites in CIII₂, suggesting that interaction with CI disrupts CIII₂ symmetry in a functionally relevant manner. Together, these observations indicate a more nuanced functional role for the SCs.

INTRODUCTION

 Aerobic cellular respiration, the process by which cells transfer electrons from sugars, fats, and proteins to molecular oxygen (O₂), is central to the energy metabolism of all eukaryotes and many prokaryotes. The final stages of aerobic respiration are carried out in the mitochondria. The mitochondrial electron transport chain (ETC) complexes of the inner mitochondrial membrane (IMM) catalyze the terminal electron transfer reactions. The ETC is composed of four large membrane protein complexes: (1) a H+⁻pumping NADH-coenzyme Q (CoQ; ubiqui-none) oxidoreductase (complex [C]I), (2) a succinate-CoQ oxido-reductase (CII), (3) an obligatorily dimeric H⁺-pumping CoOH₂ (reduced CoQ; ubiquinol)-cytochrome c (cyt c) oxidoreductase (CIII₂; cytochrome bc₁ complex), and (4) a H⁺-pumping cyt c oxidase (CIV) responsible for O₂ reduction. We now have several atomic models for all of the isolated ETC complexes (Baradaran et al., 2013; Iwata et al., 1995, 1998; Sun et al., 2005; Tsukihara et al., 1996). Most recently, structures of mammalian mitochondrial CI, the largest and least well characterized ETC complex, have been reported from multiple sources (Agip et al., 2018; Fiedorczuk et al., 2016; Guo et al., 2017; Wu et al., 2016; Zhu et al., 2016). CI is an ~1 MDa membrane protein complex with 45 protein subunits arranged into two “arms”: a peripheral arm that extends into the mitochondrial matrix and a membrane arm that is embedded in the IMM (Hirst, 2013; Sazanov, 2015).

Although each ETC complex is capable of functioning in isolation (Hatefi et al., 1962), the individual complexes form supercomplexes (SCs) of defined stoichiometry (Schägger and Pfeiffer, 2000). In mammalian heart mitochondria, the majority of CI is found in association with CIII₂ and CIV (SC I+III₂+IV; respirasome) or in association with CIII₂ alone (SC I+III₂). Recent structural work has defined the arrangement of the individual complexes within the mammalian mitochondrial respirasome (Gu et al., 2016; Guo et al., 2017; Letts et al., 2016b; Sousa et al., 2016; Wu et al., 2016). Recent electron cryo-tomography on mitochondria from mammals, plants, and yeast revealed that it is not the respirasome that is structurally most conserved across kingdoms but SC I+III₂ (Davies et al., 2018). However, the subunits shown to be responsible for the interactions between CI and CIII₂ in mammalian mitochondria (Letts et al., 2016b) are either completely absent or largely truncated in plants and yeast (Letts and Sazanov, 2015; Subrahmanian et al., 2016), suggesting convergent evolution and, in turn, an important but still undefined physiological role for SC I+III₂ in energy metabolism (Davies et al., 2018).

The possible physiological functions of the SCs remain controversial (Letts and Sazanov, 2017; Milenkovic et al., 2017), with proposals including roles in the stability of the individual complexes (Acín-Pérez et al., 2004; Calvaruso et al., 2012; Schägger et al., 2004), CoQ substrate channeling between CI and CIII₂ (Bianchi et al., 2004; Lapuente-Brun et al., 2013; Lenaz et al., 2016), the reduction of reactive oxygen species (ROS) production (Lopez-Fabuel et al., 2016), and the prevention of non-specific protein aggregation in the IMM (Blaza et al., 2014). Evidence has been mounting against the hypothesis that substrate channeling is a significant function of the respirasome, especially against the notion that there are two pools of CoQ in the IMM, one associated with SCs and one freely diffusing (Enríquez, 2016; Lapuente-Brun et al., 2013). Recent respirasome structures (Gu et al., 2016; Guo...
et al., 2017; Letts et al., 2016b; Sousa et al., 2016; Wu et al., 2016) do not show any protein subunits blocking the free exchange of CoQ from CI or CIII2 within the membrane pool. Moreover, kinetic analyses indicate that only a single pool of CoQ exists in the membrane (Blaza et al., 2014; Fedor and Hirst, 2018; Gupte et al., 1984; Kröger and Klingenberg, 1973a, 1973b). To settle this debate, new experimental frameworks are needed to further explore the physiological roles of SCs.

Here, we report the isolation of functional SC I+III2 from ovine heart mitochondria. The preparation was highly active when isolated in amphipols, providing a soluble functional respiratory unit eminently suitable for detailed functional studies. Our results indicate that limiting amounts of CoQ-10 were co-purified with the SC and that CoQ trapping by SC particles in fact decreased rates of electron transport. Structural characterization of the SC particle by cryo-electron microscopy (cryoEM) resulted in reconstructions at 4.2–4.6 Å overall resolution of multiple structural three-dimensional (3D) classes and up to 3.8 Å resolution in focused refinements. In contrast to the isolated ovine CI (Fiedorczuk et al., 2016), subunit NDUF A11 (B14.7), the traverse helix from ND5 subunit, and a few other peripheral areas of the complex were well ordered within the SC, confirming the stabilizing role of CIII2 on CI (Letts and Sazanov, 2017). The structures here allowed us to improve the completeness and accuracy of the ovine CI model and to obtain an atomic model of ovine CIII2, with clear density for endogenous CoQ bound to three of the four possible sites. Comparison of local map resolution between SC structures revealed a CI state-dependent conformational flexibility in CIII2’s cytochrome b (MT-CYB) subunit, indicating crosstalk between the two complexes. Using a focus-revert-classify strategy to separate distinct states of CI, we resolved six open and one major closed conformation. These additional states extend over the structural states previously observed in the bovine, ovine, and mouse CI (Agip et al., 2018; Blaza et al., 2018; Fiedorczuk et al., 2016; Zhu et al., 2016) and add an intriguing possibility that a key transmembrane (TM) helix rotates during the catalytic cycle. Our approach presents a new experimental framework for structure-function analysis of the physiological roles of mitochondrial SCs. Our biochemically defined system supports a stabilizing rather than a substrate-channeling function for SCs and suggests a more subtle functionally relevant interaction between CI and CIII2.

RESULTS

Preparation of Chromatographically Pure SC I+III2

When purifying CI, we observed that the yield from the first chromatographic step (anion exchange Q-column) was lower when the membranes were extracted with the branched chain detergent lauryl maltose neopentyl glycol (LMNG) than when using the single-chain detergent dodecyl-maltoside (DDM) (Letts et al., 2016a). In LMNG, a significant proportion of CI did not elute from the column until the high salt wash (Figure S1A). By modifying the gradient, two clear peaks of CI NADH:FeCy activity could be identified (Figures S1B and 1A).
The second peak contained both CI NADH:FeCy activity and high absorption at 420 nm, indicating the presence of heme groups (Figures S1B and 1A). When this peak was concentrated and run over either another round of anion exchange (Mono Q) or a size exclusion chromatography (SEC) column, it separated into two distinct peaks, one with NADH:FeCy activity and the other with the majority of the A420 signal (Figures S1C–S1E). Mass spectrometry demonstrated that these peaks corresponded to isolated CI and CI112 (Figures S1F–S1H; Table S1). By adding the amphipathic polymer (amphipol) A8-35 directly following elution of the CI + CI112 peak on the first anion exchange step, a peak containing both CI NADH:FeCy activity and A420 signal was isolated by SEC (Figures 1B and 1C). Blue Native (BN)-PAGE indicated that this peak was SC I+III2 (Figure 1D). This was confirmed by mass spectrometry, which identified the presence of all expected CI and CI112 subunits (Figures 1E and 1H; Table S1).

In the absence of amphipol, the complexes dissociated over time and separated during subsequent purification steps (Figures S1C–S1E). Given that lipids are important to stabilize SCs (Mileykovskaya and Dowhan, 2014), de-lipidation is a possible cause for the separation of the SC I+III2 during purification in the absence of amphipols. We determined that the isolated SCs were lipid-protein particles containing on average 122 ± 8 lipid molecules per SC (Table S2).

**Amphipol-Stabilized SC I+III2 Is a Functional NADH:cyt c Oxidoreductase**

The amphipol-stabilized SC I+III2 displayed all expected enzymatic activities (see Figure 2A for a schematic of SC I+III2 substrate binding sites), that is, CI NADH:CoQ oxidoreductase activity and CI112 CoQH2:cyt c oxidoreductase activity via the Q-cycle mechanism (Cramer et al., 2011). Independent of whether NADH oxidation or cyt c reduction was monitored, clear hyperbolic concentration-activity curves for both NADH and cyt c were observed. Given the ~4:1 phospholipid/cardiolipin ratio of the IMM (Horvath and Daum, 2013), this suggests that each SC particle contains ~24 cardiolipin molecules and ~98 standard phospholipids, mainly phosphatidylcholine and phosphatidylethanolamine with minor amounts of phosphatidylinositol and phosphatidylserine.
seen in the absence of any added CoQ analogs (Figures 2B and 2C). This indicated that endogenous CoQ-10 co-purified with the SC particles. This co-purification is expected even in the absence of CoQ-10 trapping in the membrane, because of CoQ-10’s high hydrophobicity (Persson et al., 2005) and the fact that the SCs, which contain several CoQ-10 binding sites, are being extracted into an aqueous environment.

As NADH is a two-electron donor and cyt c is a single-electron acceptor, two molecules of cyt c are reduced per NADH oxidized. The observed “rate-coupling” ratio of $K_{cat}$/[cyt c] (cyt c reduction/NADH oxidation) was $1.80 \pm 0.05$ (Table S3), near the expected ratio of 2.0 for perfect rate coupling, indicating that electrons from NADH are directly transferred to cyt c without a significant buildup of reduced CoQH$_2$. Time courses of NADH oxidation and cyt c reduction showed an expected ~2-fold greater final concentration of reduced cyt c relative to NADH added (Figures 3A–3F). A possible side reaction may be the reduction of cyt c by superoxide generated from the reduced flavin of CI (Prady and Hirst, 2011), bypassing CoQ-10 and CII$_2$. To prevent this, all re-action buffers contained 50 U/mL superoxide dismutase (SOD). Together, the high rate coupling and expected final concentration of reduced cyt c indicated that very few electrons were escaping to ROS. Nevertheless, the activity of the isolated SC I+II$_2$ was low relative to what was measured in ovine mitochondrial membranes or isolated mammalian mitochondrial CI using various CoQ analogs (Hatefi and Stigglall, 1978; Letts et al., 2016a; Sharpley et al., 2006; Shinizawa-Itoh et al., 2010), suggesting that the activity of the isolated SC I+II$_2$ may be limited by insufficient levels of co-purified CoQ-10.

To address this, we tested different concentrations of the CoQ analog decyl-ubiquinone (DQ) in the reaction buffer (Figure 2D). The addition of small amounts of DQ (~10 μM) resulted in a large increase in both NADH oxidation and cyt c reduction activity; however, further addition of DQ resulted in a reduction of activity
relative to the peak at 10 μM DQ (Figure 2D). The non-hyperbolic nature of the [DQ]-activity curve has been attributed to its hydrophobic “detergent-like” properties (Estornell et al., 1993; Letts et al., 2016a). If DQ solubility dictated the shape of the [DQ]-activity curve, factors such as ionic strength should affect the profile. To test this, we exchanged the buffer from HEPES to potassium phosphate, maintaining the same pH, and re-measured the concentration activity curve (Figure S2A). This buffer change resulted in a significantly different profile at low DQ concentrations, with a broader high-activity peak (5–20 μM DQ) in phosphate compared with HEPES (tightly centered around 10 μM; Figures S2B and S2C). This is consistent with the shapes of the curves resulting from solubility effects and the low activity seen at high [DQ] being due to poor DQ solubility.

Given that maximal NADH:cyt c oxidoreductase activity was seen with 10 μM DQ, this concentration was used to evaluate NADH and cyt c activity (Figures 2E and 2F). In the case of the [NADH]-activity curve, a hyperbolic profile was seen, with a higher \( V_{\text{max}} \) (~5-fold increase) and higher apparent \( K_v \) when than when no DQ was added (Figures 2B, 2C, 2E, and 2F). However, although the activity was increased at all [cyt c] when 10 μM DQ was added, the [cyt c]-activity curves are no longer hyperbolic, instead showing a pronounced decrease in activity at high [cyt c] (Figure 2G). Given that this is not seen at higher [DQ] (see below), this reduction in activity may stem from a non-specific interaction between cyt c and DQ. The rate-coupling ratio from the [NADH] activity curves was 1.90 ± 0.06 (Table S3), again reflecting efficient electron transfer from NADH to cyt c (Figure 3B).

Although addition of DQ increased the activity of the amphipol-stabilized SC I+III2 (Figure 2D), the NADH oxidation activity of CI was still lower than that seen in mitochondrial membranes or solubilized in detergent (Letts et al., 2016a), indicating that some aspect of SC I+III2 was limiting CI turnover. The CII2 activity observed in SC I+III2 (~102 ± 2 s⁻¹; Table S3) is consistent with that measured for isolated eukaryotic CII2 (S. cerevisiae, 60–270 s⁻¹ [Covian and Trumpower, 2008; Gutierrez-Cirlos and Trumpower, 2002; Nett et al., 2000], and Bos taurus, ~600 s⁻¹ [Brandt and Okun, 1997]) and for mitochondria of living cultured cells (~250 s⁻¹ [Kim et al., 2012; Ripple et al., 2013]). However, the rate of ~100 cyt c reduced per CII2 per second is on the low end of the reported range, suggesting that the activity of CII2 may also be impeded in the amphipol-stabilized SC.

To determine whether CI and CII2 activities are limited by confinement in the SC or whether isolation by this protocol results in lower activity in general, the activities of the amphipol-stabilized SC were assayed in lipid-detergent (LD) buffer containing 0.25 mg/mL dioleoylphosphatidylcholine (DOPC)/cardiolipin (CL) 4:1, 0.1% CHAPS (w/v), and 0.1% LMNG (w/v). This LD mixture allows maximal activity for isolated ovine CI (Letts et al., 2016a). The addition of lipid and detergent to the reaction mixture resulted in a significant shift in the [DQ] activity curve with maximal activity at higher [DQ] (~100 μM) and marked increases in both NADH oxidation and cyt c reduction activity (Figure 2G). The [NADH] activity curves in these conditions were hyperbolic and displayed increased \( V_{\text{max}} \) and higher apparent \( K_v \) values than when measured in the absence of lipid and detergent (Figures 2H and 2E). Addition of lipid and detergent uncoupled NADH oxidation from cyt c reduction and resulted in robust NADH oxidation in the absence of any added cyt c (Figure 2I). However, the addition of sufficient cyt c (≥50 μM) did increase the NADH oxidation activity of CI, indicating that CII2 activity was still able to influence CI and that their association may not be wholly disrupted (Figure 2I). When cyt c reduction was monitored, the [cyt c] activity curve was hyperbolic, resulting in a 2-fold increase in \( V_{\text{max}} \) relative to the maximum activity seen in the absence of lipid and detergent (8.22 ± 0.09 U/mg versus 4.11 ± 0.09 U/mg). This rate of ~230 cyt c reduced per second per CII2 agrees more closely with fluxes measured in the mitochondria of living cells (Kim et al., 2012; Ripple et al., 2013).

Given that upon addition of the LD buffer, the CI activity increased by a larger factor than CII2 activity, the rate-coupling ratio decreased to 1.21 ± 0.02 (Table S3). When a time course of the reaction was examined, a mismatch between the amount of NADH oxidized versus cyt c reduced at each time point was observed (Figure 3F). Moreover, the reduction of cyt c became bi-phasic with a fast phase of reduction that coincided with the oxidation of NADH by CI and a slow phase of cyt c reduction after all the NADH was oxidized (Figure 3F). The two phases of cyt c reduction activity again suggest that the amphipol-stabilized SC I+III2 may not be completely disrupted in the LD buffer. The initial fast phase of cyt c reduction likely corresponds to DQ2H2 diffusing quickly within the SC between CI and CII2, and the slow phase to DQ2H2 that escapes the SC into the LD buffer and then must re-encounter CII2 to be oxidized (Figure 3F). Overall, there is a buildup of reduced DQ2H2 during the experiment as CI activity outpaces that of CII2. This is consistent with there being a kinetic advantage to keeping the CI and CII2 active sites close together, as would be expected for a diffusion-coupled process (Hakenbrock et al., 1986). Nonetheless, diffusion of CoQ has been shown not to be rate limiting during respiration (Chazotte and Hakenbrock, 1988), and decreasing the diffusion distance between CI and CII2 is not likely to be the main physiological role of SCs. Comparing the LD buffer data with those in the absence of detergent indicates that CI activity is limited in the amphipol-stabilized SC I+III2 particle.

We also note that the NADH oxidation and cyt c reduction activities of the amphipol-stabilized SC I+III2 were sensitive to known CI and CII2 inhibitors (Figure S3). We tested the CI inhibitors rotenone and piericidin A, as well as Na⁺H⁺-antiporter amiloride inhibitors. Also, the Q0-site CII2 inhibitor myxothiazol and the Qm-minus-site inhibitor antimycin A both showed strong inhibition of SC activities, with Hill coefficients close to 2, suggesting that binding to either of the Q0 or Qm sites in the dimer may be sufficient to inhibit CII2 activity (Figures S3C and S3D).

**CI Activity Is Limited by CII2 Activity When CoQ Is “Trapped” in SC I+III2**

To better understand what was limiting the activity of the SC, we measured SC activity at different DQ concentrations in the presence or absence of the non-specific hydrophobic carrier protein BSA (Figures 3G–3I). The addition of 10 μM BSA to the reaction mixture shifted the [DQ]-activity curve to higher [DQ], and the curves became more “hyperbolic-like” (Figure 3G). This indicates that higher concentrations of DQ are needed to make the substrate available for electron transport in the SC I+III2.
partitions in the presence of BSA and hence that BSA reduces the partitioning of DQ into the SC lipid-protein particles at low [DQ]. When we measured the NADH oxidation activity in the presence of BSA, we saw a slight but significant (p < 0.01) increase in V_max, whereas there was a significant decrease in the V_max of cyt c reduction (Figures 2E, 2F, and 3H). This results in a reduction of the rate-coupling ratio similar to that seen in LD buffer (Table S3), indicating that BSA provides an alternate path for DQH2 out of the SC I+III2 particle. To test whether the higher concentration of DQ was partially responsible for the differences in LD buffer and in the presence of BSA, we measured [NADH] activity and [cyt c] activity with exchange of 100 μM DQ (Figures 3J and 3K). As expected from the [DQ] activity curves (Figure 2D), the V_max measured with 100 μM DQ was lower than that using 10 μM DQ (Figures 3J and 3K). However, the decreases in NADH oxidation and cyt c reduction activities were proportional, thus maintaining a high rate-coupling ratio between CI and CIII2 (Table S3).

Whether the ability of SC-associated DQ to exchange with the bulk DQ pool is dependent on [DQ] can also be estimated by measuring NADH oxidation in the absence of any added cyt c (Figure 3L). In these conditions, reduced DQ cannot be readily oxidized by CIII2, and thus the only source of oxidized CoQ to maintain CI activity is via exchange from the bulk pool into the SC I+III2 particles. In the absence of DQ and cyt c, NADH oxidation activity was undetectable (Figures 2F and 3L). However, the decrease in NADH oxidation and cyt c reduction activities were proportional, thus maintaining a high rate-coupling ratio between CI and CIII2 (Table S3).

CryoEM Structures of Isolated SC I+III2 Reveal Multiple States

We used the amphipol-stabilized SC I+III2 to solve the atomic structure of the SC to resolutions up to 3.8 Å (in focused refinements) by cryoEM (Tables 1 and S4), using our improved protocol for model refinement (STAR Methods). CI appeared more stable within the SC than previous preparations of isolated ovine CI (Fiedorczuk et al., 2016), allowing us to improve the model in some peripheral areas that were previously disordered. Initial 3D classification identified four classes of SC I+III2 particles (Figures 4A–4D; Figure S4C). These differed mainly in the relative angles between the CI peripheral arm and membrane arm and in the angle between CI and CIII2 (Figures 4E and 4F). We concluded that the most distinct class with respect to the position of the CI peripheral arm belonged to the closed state (or active form) of CI (Agip et al., 2018; Fiedorczuk et al., 2016). The assignment of the closed state was based on clear density adjacent to the CI Q-tunnel for the NDUF52 [-1]-[-2] loop and for the ND3 TM helix 1 (TMH1)-TMH2 loop (residues Pro25–Lys54; Figure 4G). In all other classes of SC I+III2, these loops were disordered (Figure 4G), suggesting that they are distinct forms of the open state (or inactive form) of CI (Blaza et al., 2018; Fiedorczuk et al., 2016). Previously, only a single open state had been identified for CI (Agip et al., 2018; Blaza et al., 2018). We identified striking changes in TMH3 of ND6 between the open- and closed-state structures. Whereas in the closed state, ND6 TMH3 was a clear α-helix (residues Leu52–Met74), in each of the open states, the helix was interrupted by a β-bulge midway across the membrane (residues Tyr60–Met65; Figure 4H). The result of this β-to-α transition between the open and closed states of CI is a rotation of ~100° for the C-terminal half of ND6-TM3 (residues Met64–Met74; Figures 4H and 4I).

The three major changes between the closed and open states—the ordering of the NDUF52 [-1]-[-2] loop in the Q-tunnel, the ordering of the ND3 TMH1-TMH2 loop, and the π-to-α transition of ND6-TM3—occur at the interface of the CI peripheral and membrane arms and were accompanied by a significant rotation between the two arms of the complex, bringing NDUF5 and NDUF10 into close contact (Figure 4F). As shown in Figure 4I, the ordered ND3 TMH1-TMH2 loop in the closed state crosses directly above ND6 TMH3. This loop, however, was disordered in the open states. Additional large conformational changes were seen in this area: the short three-stranded β-sheet of NDUF52 that harbors the [-1]-[-2] loop was adjacent to ND6 TMH3 in the closed state but rotated away from TMH3 by as much as 10 Å in the open states (Figure 4I). This movement was accompanied by the striking rotation of the side chains of Phe68 and Tyr70 because of the π-bulge-to-α-helix rearrangement. The analogous structural differences were also observed between the active and inactive forms of CI for mouse CI (Agip et al., 2018), indicating that such large conformational transitions are conserved.

Because of the hinge-like motions between the different regions of the SC, we carried out focused refinements on distinct sub-regions to generate higher quality density maps. These focused refinements were performed on the peripheral arm of CI, on the membrane arm of CI and on CIII2, resulting in
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reconstructions at 3.8–3.9 Å resolution and improved atomic models for each region of the SC (Figure S5; Tables 1 and S5).

Focus-Revert-Classify Strategy Reveals at Least Six Distinct Open States for CI

Although we were able to separate three distinct CI open states using the standard 3D classification, we reasoned that because of the systematic worsening map quality along the peripheral arm of CI (Figure 7; Figure S7), averaging of particles with diverse angles between the membrane and peripheral arms was still occurring. Hence, we developed a strategy for the separation of different CI states on the basis of the angle between the peripheral and membrane arms. Termed “focus-revert-classify,” an initial focused refinement around the peripheral arm of CI is performed, maximizing the differences in the relative positions between the membrane arms of each CI, followed by “reverting” to a mask around the membrane arm for classification (Figures 5A and S6A).

The focus-revert-classify strategy resulted in the separation of nine distinct structural classes of CI ranging in resolution from 4.1 to 7.5 Å (Figures 5B and S6A). Of these, seven correspond to distinct open states in which the Q-site loops are disordered, one to a closed state in which the loops around the Q site are well ordered and the final class to a closed-like state at 7.0 Å, precluding the assignment of any of the active site loops. All six CI open state structures at ≤6.5 Å each had distinct angles between the peripheral and membrane arms (Figures 5B and 5C). The closed state differs from the open states as described above for the closed state of the SC (Figure 5C). When comparing the particles between the classes of the original 3D classifications of the SC, isolated CI (Figure S4C) and focus-revert-classify (Figure S6A), it is clear that the closed state particles are distinct from all other classes (Figure S6B). The majority of particles that we identified as having CI in the closed state in the original classifications were also found in the closed state class after application of the focus-revert-classify strategy. This was not the case for the three original open state classes, which were more distributed between the six focus-revert-classify open classes. These data indicate that the closed state of CI is a distinct conformation and the open state is an ensemble of nearly continuous conformations differing in the relative positions of CI’s membrane and peripheral arms.

Overall Arrangements of the Amphipol-Stabilized SC I+III2 and CoQ Active Sites

The overall arrangement of the SC I+III2 shown here is similar to that in the full respirasome (Lettis et al., 2016b) as well as in situ (Davies et al., 2018). CIll2 contacts CI at two main sites: (1) in the IMM, where CI subunit NDUF11 contacts UQCRB and UQCRQ of the adjacent CIll protomer (Figure 6A), and (2) in the mitochondrial matrix, where CI subunits NDUF4 and NDUF9 (B22) contact UQRC1 of that same CIll protomer (Figure 6B). At the interface of the mitochondrial inner membrane and matrix, only van der Waals contacts can be seen between the top of NDUF11 (Leu49) and the main chain of the adjacent UQCRB helix (between Lys11 and Trp12). However, a cluster of positively charged residues extending from both NDUF11 (Arg103 and Arg105) and UQCRQ (Arg42) suggests the possibility of
lipid-mediated bridging between CI and CIII2 (Figure 6A). On the inter-membrane space side of the membrane, several charged and polar residues are in close contact (NDUFA11: Glu16, His18, Arg19, and Gln78; UQCRQ: Tyr56, Gln64, and Lys68), which allows the formation of stabilizing hydrogen bonds or salt bridges (Figure 6A). Within the mitochondrial matrix, salt-bridging interactions can be seen between NDUFB9 and NDUFB4 of CI and UQCR1C of the adjacent CIII protomer (Figure 6B). Possible salt bridges include NDUFB9 Lys54 and UQCR1C Asp20, NDUFB9 Asp55 and UQCR1C Arg24, NDUFB4 Glu27 and UQCR1C Lys51, and NDUFB4 Arg29 and UQCR1C Glu225 (Figure 6B). Although these interacting residues are conserved in mammals, they diverge quickly in other lineages, likely reflecting different ways of forming SC I+III2 across eukaryotes (Davies et al., 2018).

Given the hydrophobic nature of CoQ-10, it would remain in the fragment of lipid bilayer that is co-purified with the SC I+III2 particles. Hence, even if only limiting amounts of CoQ-10 were
co-purified within the SC, its local concentration would be high. Accordingly, although no exogenous CoQ was added to our cryoEM sample, clear density for the CoQ-10 head group can be seen in the CIII2 focused maps at three of the four CoQ binding sites (Figures 6C–6E). Density could be seen at both QN sites, which are expected to have a higher affinity for the oxidized CoQ-10 (Figures 6C and 6D) and the Qp site distal from the CI Q-tunnel (Figure 6E). However, the QP site proximal to the CI Q-tunnel did not display any density for CoQ-10, suggesting different affinities for CoQ-10 in the QP sites of the different MT-CYB protomers.

At low contour, the cryoEM maps showed density for a disordered layer of amphipols and lipids forming a belt around the hydrophobic surfaces of the SC (Figure 6F). On the CIII2-proximal side of CI, the four-TMH-containing subunit NDUFA11, which directly interacts with CIII2, is shaped like a lipid-filled arch (Figure 6F), contacting CI with the ends (springers) and CIII2 with the top (crown). The dominance of lipid-mediated contacts between CI and NDUFA11 may explain why this subunit is so easily lost or disordered by detergent when CI is extracted. The rest of the CI-CIII2 interface within the membrane involves two large lipid-filled cavities, made discontinuous by the close interactions among NDUFA11, UQCRB, and UQCRQ. These interactions likely act as a barrier to the free diffusion of CoQH2 as it exits the CI Q-tunnel (Figure 6F).

The two Q cavities of CIII2 penetrate deep into the core of the dimer and are separated from each other by an interaction between the two MT-CYB protomers (Figure 6F). Each Q cavity contains a CoQH2-oxidizing Qp site and a CoQ-reducing Qn site; however, the Qp and Qn sites within each cavity derive from the opposite CIII protomer (Figure 6F) and, hence, the Qp and Qn sites of one MT-CYB subunit face opposite cavities. One of the Q cavities is proximal to the Q-tunnel of CI, while the other is distal on the opposite side of the dimer and much less accessible (Figure 6F). This arrangement led to the hypothesis that symmetry-breaking within CIII2 may result in the specialization of the Q cavities, with the proximal Q cavity specialized for CoQH2 oxidation and the distal cavity specialized for CoQ reduction (Lettis and Sazanov, 2017; Letts et al., 2016b). In the amphipol-stabilized SC, trapping of CoQ would limit the diffusion path from the CI Q-tunnel to CIII2 and vice versa to the lipid-amphipol belt surrounding the SC (Figure 6F). Given the overall structure of the SC (Figure 6F) and evidence of poor CoQ exchange in the amphipol-stabilized particle (Figure 3L), functional symmetry-breaking of CIII2 and specialization of the Q-cavities is highly likely.

**Local Resolution Analysis Reveals CI State-Dependent Crosstalk**

Clear differences can be seen in the local resolution of specific regions of CI and CIII2 between the different reconstructions (Figure 7; Figures S7B and S7C). In the SC I+III2 class containing...
Figure 6. Interactions, CoQ Density, and Overall Arrangement of the Amphipol-Stabilized SC I+III2 Particles

(A) Interaction between CI and CIII2 in the membrane. CI subunit NDUFA11 in cyan, CIII2 subunits UQCRB in green, and UQCRQ in light green. The putative "positive cluster" lipid-binding site is circled and labeled. Side chains of important residues are shown as sticks and colored by atom, with nitrogen blue, oxygen red, and carbon the same color as the subunit. Models shown as cartoons throughout.

(B) Interaction between CI and CIII2 in the mitochondrial matrix. CI subunits are in green and CoQ, heme blue, oxygen red, and iron orange. MT-CYB His201 are shown as sticks and colored by atom, with carbon gray, nitrogen blue, oxygen red, and iron orange.

(C) Density for CoQ-10 binding in the proximal QN-site from the CIII2 focused maps. CIII2 subunits are in green, and CoQ, heme b56, and the side chain of MT-CYB His201 are shown as sticks and colored by atom, with carbon gray, nitrogen blue, oxygen red, and iron orange.

(D) Density for CoQ-10 binding in the distal QN-site from the CIII2 focused maps. CIII2 subunits, CoQ, heme b56, and the MT-CYB side chain His201 are shown as in (C).

(E) Density for CoQ binding in the distal QP site from the CIII2 focused maps. CIII2 subunits, CoQ, and heme b56 are shown as in (C).

(F) Slice through the membrane domains of CI and CIII2 within the closed structure of the SC looking from the mitochondrial matrix. Amphipol-lipid belt is shown at low contour in red, and CI (blue) and CIII2 (green) are shown at a higher contour. Each CIII protomer in close contact to CI contains the CoQH2 oxidation site most proximal to the Q-tunnel of CI. Hence, this MT-CYB subunit is likely responsible for oxidizing the majority of CoQH2 produced by CI within the SC. The decrease in the local resolution of this subunit only in the closed state of CI strongly suggests that there is crosstalk between the two complexes and that CIII2 may be primed for CoQH2 oxidation upon the closing of CI. Nonetheless, at this resolution it remains unclear how this crosstalk between CI and CIII2 occurs.

DISCUSSION

We present here the first chromatographic isolation of functional SC I+III2 from mammalian mitochondria, followed by its detailed functional and structural characterization. Previous isolations of this SC from mitochondrial membranes displaying NADH:cyt c oxidoreductase activity have been reported (Hatefi and Stiggall, 1978; Hatefi et al., 1961). However, the purity of these preparations is unclear, and their stability is poor. The preparation presented here displays a homogeneous single peak by SEC (Figure 1C) and appears as a single major band in light green. Important side chains are shown and colored as in (A). Putative salt-bridging interactions indicated by dashed ovals.

(C) Density for CoQ-10 binding in the proximal QN-site from the CIII2 focused maps. CIII2 subunits are in green, and CoQ, heme b56, and the side chain of MT-CYB His201 are shown as sticks and colored by atom, with carbon gray, nitrogen blue, oxygen red, and iron orange.

(D) Density for CoQ-10 binding in the distal QN-site from the CIII2 focused maps. CIII2 subunits, CoQ, heme b56, and the MT-CYB side chain His201 are shown as in (C).

(E) Density for CoQ binding in the distal QP site from the CIII2 focused maps. CIII2 subunits, CoQ, and heme b56 are shown as in (C).

(F) Slice through the membrane domains of CI and CIII2 within the closed structure of the SC looking from the mitochondrial matrix. Amphipol-lipid belt is shown at low contour in red, and CI (blue) and CIII2 (green) are shown at a higher contour. Each CIII protomer is colored a different shade of green. CoQ active sites are marked: solid wedges for location on the matrix leaflet of the membrane and dashed wedges for inter-membrane space leaflet. Black and yellow ovals indicate lipid binding pockets of CI; black and yellow circle indicates lipid binding pocket of NDUFA11; black and yellow X indicates barrier to CoQ diffusion caused by contact site between CI and CIII2 in the membrane. Yellow arrow illustrates the shortest path for CoQ diffusion from the CI Q-tunnel to the proximal Qp site of CII2.
on BN-PAGE gels (Figure 1D). The characterization of amphipol-stabilized SC I+III2 NADH:cyt c oxidoreductase activity demonstrates that it is possible to study the transfer of electrons from NADH to cyt c, both in the absence or presence of added CoQ analogs (Figure 2). The rates of NADH oxidation and cyt c reduction observed when the SC was supplemented with 10 μM DQ were lower than what has been observed for CI and CIII2 turnover in membranes (Lettis et al., 2016a; Ripple et al., 2013) (Figures 2E and 2F). However, the activity of both complexes increased to native levels upon the addition of lipid and detergent (Figures 2H and 2I). With the purity and inhibitor sensitive activity achieved here (Figures 1, 2, and S3), this preparation can be used for the study of SC I+III2 by diverse biophysical methods.

This preparation is not without its limitations. Because of the lack of a sealed membrane compartment, there is no Δψ or ΔpH present in the system. This is significant, as CI normally operates near equilibrium with the proton motive force (Ripple et al., 2013), and as the redox poise of the b-hemes in CIII2 is affected by both Δψ and ΔpH (Kim et al., 2012). Also, because of the nature of the amphipol-based stabilization of the SC in solution, CoQ-10 and to a lesser extent DQ are trapped in the SC particle, and their diffusion is constrained to the lipid-amphipol belt (Figure 6F). CoQ-10 is highly hydrophobic; hence, when extracted into an aqueous solution, CoQ-10 will partition into the hydrophobic environment of the deep lipid pockets present in the SC. Together with the affinities of CI and CIII2 for CoQ-10, a concentrating effect of CoQ-10 within the extracted SC is expected. Hence, it does not follow from CoQ-10 trapping in this context that CoQ-10 is not free to diffuse and exchange with the bulk membrane pool in the hydrophobic environment of the IMM.

Nevertheless, the fact that CoQ is “trapped” by the amphipol-stabilized SC I+III2 was advantageous, as it allowed us to test the hypothesis that CoQ-10 trapping by SC I+III2 in situ would increase the rate of the ETC. This system allowed us to determine
how CoQ trapping affects CI and CIII2 activity and to place what we learned into a structural context. Given the arrangement of CI and CIII2 in the SC (Figure 6F), it is likely that only one side of CIII2 would be readily supplied with CoQH2 from CI and that the symmetry of CIII2 would be functionally broken. Nonetheless, given the Hill coefficients of 2 for both the QP- and QN-site CIII2 inhibitors (Figure S3), it is clear that the CoQ sites in the distal CIII2 Q-cavity also play an important role in turnover. Rate limitation of CI by CIII2 can be understood in the context of the SC I+III2 structure (Figure 6F). Trapping the CoQ between CI and CIII2 would limit the ability of CIII2 to oxidize CoQH2 to a single QP site. In the context of the structure, the ~2-fold increase in CIII2 activity may be due mainly to a re-establishment of two accessible QP sites caused by the free exchange of DQH2 in the LD mixture (Table S3). This indicates that CIII2 within the amphipol-stabilized SC I+III2 particles is operating at near maximal activity in the experimental context but that only one Q-cavity has access to reduced CoQH2. The rate coupling in these contexts is nearly 2.0 (Table S3), demonstrating that CI activity is limited by CIII2.

Thus, this argues against the view that trapping of CoQ-10 by SCs may increase the flux through the ETC but instead indicates that CI can function more effectively if CoQ is able to exchange with the membrane pool and is not dependent solely on the proximal CIII2 QP site for regeneration of oxidized CoQ. The native arrangement within the IMM may allow both CI and CIII2 to operate at the highest rates via exchange with the bulk pool and not via CoQ trapping. Given the structure of the SC, the only way for CoQ trapping to result in higher rates would be if the turnover of CI and the single CIII2 protomer were well matched. Because of the nature of the Q-cycle, CIII2 must oxidize two CoQH2 for each CoQ reduced by CI. Given the complexity of the Q-cycle mechanism—including electron bifurcation, conformational transitions of UQCRFS1, and three simultaneously bound substrates—this strict requirement for rate matching may be difficult to accommodate. The mismatch in rates between CI and CIII2 turnover may in part explain why there is an ~3-fold excess of CIII2 compared with CI in mitochondrial membranes (Schagger and Pfeiffer, 2001).

We demonstrate here that the CI closed-to-open state transition involves not only a rotation of the CI peripheral and membrane arms but the ordering of the NDUS2 β1-β2 and ND3 TMH1- TMH2 loops (Figure 4G) but also a notable rotation of ND6’s TMH3 by ~100° that is driven by the conversion of a π-bulge into an α-helix (Figures 4H and 4I). This ND6 TMH3 π-to-α transition was also seen in mouse mitochondrial CI between active and deactive forms (Agip et al., 2018). Although the ovine closed state is likely to be identical to the mouse active form, this is not necessarily the case for the ovine open states and the mouse deactive form. The deactive form of CI is obtained upon incubating the enzyme at 37°C without substrates and then converts back to active upon turnover (Vinogradov, 1998). No significant amount of deactive form was detected biochemically in the ovine enzyme as prepared (data not shown). This is consistent with mouse CI, which is ~90% in the active form as purified (Agip et al., 2018). Given that our preparation was largely in the active state, but many SC particles were in open states, it is likely that CI undergoes conversion between open and closed states while active as part of its catalytic cycle. In this case, the deactive form would represent one of the open-like states with likely more extensive unfolding of loops around the Q-site. A direct link between closed-to-open transitions and the catalytic cycle was also proposed in a recent cryoEM study of Y. lipolytica CI (Parey et al., 2018). Given that the ND6 TMH3 π-bulge is conserved in the structures of yeast and bacterial CI (Baradaran et al., 2013; Zickeffmann et al., 2015), this rotation is likely a conserved feature of CI turnover and may be a novel way of achieving efficient conformational coupling by the enzyme. Although the rotation was not observed in the Y. lipolytica or bacterial structures to date, this may be due to limited resolution or crystal contacts.

By careful classification of the SC particles, we show that there are state-dependent differences in local resolution (Figure 7). In the closed state of CI, conformational flexibility in the peripheral arm may be transmitted to ND4 in the membrane arm via interaction between NDUFA5 and NDUFA10 (Figure 7F), possibly facilitating coupling of CoQ reduction and H+ pumping during CI turnover. How this state-dependent conformational flexibility is further communicated to CIII2 remains unclear. However, our structures of multiple states of CI within the SC at similar resolutions suggest that the state of CI affects CIII2’s MT-CYB conformational flexibility (Figures 7D and 7E). The fact that CoQ was found at only three of the four possible binding sites within CIII2 (Figures 6C–6E) also suggests a functionally relevant effect of CI on CIII2. Overall, our data further support the hypothesis of functional symmetry breaking in the CIII2 dimer by CI.

Our pure, biochemically defined system allowed us to demonstrate that in contrast to the long-proposed substrate-channeling role for the SC, CoQ trapping in the SC limits CI turnover and would reduce the overall rate of the ETC in vivo. If CoQ trapping is inefficient, what could be the functional role of the SCs? Possible roles in the stabilization of the individual complexes, the reduction of ROS production, and the prevention of non-specific protein aggregation such that the IMM retains suitable CI/CIII2/CIV ratios remain feasible. The SCs may provide kinetic advantages by bringing the active sites into close proximity, but this advantage would exist even in the absence of CoQ trapping by the SCs, as it is only dependent on the distance between the active sites. Moreover, under high turnover, because of the structure of the mitochondrial membranes, individual CIs may outpace the capacity for CoQ pool equilibration, in which case the local CoQH2 concentration may become higher than in the bulk membrane (Budin et al., 2018). By ensuring close association of CI and CIII2 in SCs, but still allowing free exchange of CoQ with the bulk, the SC would prevent any local buildup of CoQH2, while at the same time not limiting CI’s activity by making it entirely dependent on the adjacent CIII2 for the re-oxidation of CoQH2. We show here, for the first time, functional crosstalk between CI and CIII2 within the SC. Although more work is needed to confirm and characterize the nature of this coordination, our results and our new experimental framework have significant implications for the physiological roles of respiratory SCs, suggesting more subtle functionally relevant interactions between CI and CIII2.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:
Molecular Cell 75, 1131–1146, September 19, 2019


## STAR METHODS

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Leonid A. Sazanov (sazanov@ist.ac.at)

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Ovis aries (sheep) hearts were purchased from C Humphreys & Sons (Chelmsford, UK).

METHOD DETAILS

Purification of amphipol-stabilized SC I+III2
Mitochondria were isolated and membranes prepared as previously described (Letts et al., 2016a). Complex (C) purification in LMNG was initially performed as previously described (Letts et al., 2016a), consisting of a single anion-exchange step followed by SEC. In short, 5% (w/v) LMNG was added drop-wise to the washed mitochondrial membranes to a final concentration of 1% LMNG, followed by stirring for ~30 min at 4°C and centrifugation at 48,000 g for 45 min. The supernatant was filtered (0.45 μm pore size polyethersulfone membrane) and loaded onto a pre-equilibrated 45 mL Q-Sepharose FF anion-exchange column (GE Healthcare). The Q-Sepharose buffers (A and B) contained 20 mM Tris-HCl, pH 7.4, 10% (v/v) glycerol, 1 mM EDTA, 1 mM DTT and 0.04% LMNG; additionally, buffer B contained 1 M NaCl; the Q-Sepharose column was pre-equilibrated at 5% buffer B in buffer A. After application of the mitochondrial extract, the Q-Sepharose column was washed with 50 mL of 5% buffer B, then with a 30-mL linear gradient of 5%–22%, and finally with 150 mL of 22% buffer B. CI was then eluted with a 300-mL linear gradient...
of 22%–30.5% buffer B (CI elution gradient; Figure S1A). Any remaining protein was then eluted with 100% buffer B. The Q-Sepharose gradient was run overnight at 1.0 mL/min at 4°C. CI-containing fractions were pooled based on NADH:FeCy activity and concentrated to 1.5-2.0 mL. Once it was determined that this Q-Sepharose protocol resulted in a significant proportion of CI in the high-salt wash (Figure S1A), the CI elution gradient was extended to 22%–60% buffer B over 450 mL, resulting in the isolation of two distinct peaks containing NADH:FeCy (CI) activity (Figure S1B). As a further optimization, the elution gradient was changed to 22%–50% over 250 mL. This resulted in the abundant isolation of SC I+III2 (Figure 1A). It was also essential to increase the flow rate over the column to 3.0 mL/min throughout to minimize the amount of time the protein was on the Q-Sepharose column.

Samples eluted from the Q-Sepharose column were then applied either to the same column again and run with a similar gradient to separate CI and CIi2 (Figure S1C), or to a Superose 6 prep grade XK 6/100 size exclusion chromatography column equilibrated in buffer S (20 mM HEPES, pH 7.4, 2 mM EDTA, 10% glycerol, 50 mM NaCl and either 0.05% LMNG (Figure S1D) or no LMNG for the amphipol samples (Figure 1B)), and eluted overnight at 0.35 mL/min at 4°C for each step. For the amphipol sample, amphipol A8-35 was added to a final concentration of 0.3% (w/v) immediately following elution from the Q-Sepharose gradient. This mixture was then tumbled at 4°C for ~30 min before concentrating to ~1.5-2.0 mL and loading onto the Superose 6 column. Subsequent SEC runs were done using the same column either in 0.5% LMNG (Figure S1E) or in no LMNG (Figure 1C) buffer S. After elution, the fractions containing NADH:FeCy activity were pooled and concentrated. These samples were either used directly for cryoEM grid preparation or made to 30% glycerol and stored in liquid nitrogen. Samples stored in this way maintained high activity over several months.

Blue Native PAGE
Samples diluted with 4X sample buffer (50 mM BisTris-HCl, pH 7.2, 50 mM NaCl, 10% w/v glycerol, and 0.001% Ponceau S) were loaded onto a NativePAGE 3%–12% Bis-Tris Gel (Invitrogen) and run in the cold room at 4°C. The cathode buffer was 50 mM Tricine, 50 mM BisTris-HCl, pH 6.8 plus 1X NativePAGE Cathode Buffer Additive (Invitrogen) and the anode buffer was 50mM Tricine, 50 mM BisTris-HCl, pH 6.8. Gels were run at 150 V for ~60 min and then the voltage was increased to 250 V for the remainder of the run ~90 min.

Mass Spectrometry
Mass spectrometry (MS) analysis was performed essentially as previously described (Letts et al., 2016a). Briefly, polyacrylamide gel slices (1–2 mm) containing the purified proteins (Figure S1H) were prepared for mass spectrometric analysis by manual in situ enzymatic digestion. The excised protein gel pieces were placed in wells of a 96-well microtiter plate and destained with 50% (v/v) acetonitrile and 50 mM ammonium bicarbonate, reduced with 10 mM DTT, and alkylated with 55 mM iodoacetamide. After alkylation, proteins were digested with 6 ng/μL trypsin (Promega, UK) overnight at 37°C. The resulting peptides were extracted in 2% (v/v) formic acid, 2% (v/v) acetonitrile. The digest was analyzed by nano-scale capillary LC-MS/MS using an Ultimate U3000 HPLC (ThermoScientific Dionex, San Jose, CA) to deliver a flow of ~300 nL/min. A C18 Acclaim PepMap 100 5 μm, 100 μm x 20-mm nanoflip Viper (ThermoScientific Dionex) trapped the peptides prior to separation on a C18 Acclaim PepMap100 3 μm, 75 μm x 250 mm nanoflip Viper (ThermoScientific Dionex). Peptides were eluted with a gradient of acetonitrile. The analytical column outlet was directly interfaced via a nanoflow electrospray ionization source, with a hybrid dual pressure linear ion trap mass spectrometer (Orbitrap Velos, ThermoScientific). Data-dependent analysis was carried out, using a resolution of 30,000 for the full MS spectrum, followed by 10 MS/MS spectra in the linear ion trap. MS spectra were collected over an m/z range of 300-2000. MS/MS scans were collected using the threshold energy of 35 for collision-induced dissociation. LC-MS/MS data were then searched against a protein database (mammalian subset of UniProt KB) using the Mascot search engine program (Matrix Science, UK) (Perkins et al., 1999). Database search parameters were set with a precursor tolerance of 10 ppm and a fragment ion mass tolerance of 0.8 Da. One missed enzyme cleavage was allowed, and variable modifications for oxidized methionine and carbamidomethyl cysteine were included. MS/MS data were validated using the Scaffold program (Proteome Software) (Keller et al., 2002). All data were additionally interrogated manually. See Table S1.

Organic Phosphate Assay
Measurement of organic phosphate content was performed according to the protocol of Anderson and Davis without modification (Anderson and Davis, 1982). Lipid was extracted from protein samples (~20 μL) by addition of 200 μL of 2:1 chloroform/MeOH followed by vortexing. Then, 50 μL of 125 mM NaCl was added, and the samples were vortexed again. The phases were separated by centrifugation for 3 min at ~13,000 x g, and the lower chloroform phase was desiccated and used for measurement. See Table S2.

Activity Assays
CI NADH:FeCy activity was measured by spectroscopic observation of NADH oxidation at 340 nm wavelength using a Shimadzu UV-2600 UV-VIS spectrophotometer with CPS-100 thermoelectrically temperature-controlled cell positioner and modified by Rank Brothers for continuous sample stirring. SC I+III2 NADH:cyt c oxidoreductase activity was measured at 340 nm for NADH oxidation and 550 nm for cyt c reduction using either the same Shimadzu UV-2600 UV-VIS spectrophotometer as above or a Spectramax M2e Plate-Reader (Molecular Devices). The buffer used for NADH:FeCy activity was 20 mM HEPES, pH 7.4, 50 mM NaCl, 2 mM EDTA, 0.1% DDM, 100 μM NADH and 1 mM potassium ferricyanide (KFeCy). NADH:Cyt c oxidoreductase activity of the isolated SC I+III2 were done in the standard buffer (SB): 100 mM HEPES, pH 7.4, 50 mM NaCl, 10% glycerol and 4 μM KCN, 50 mM SOD; or
lipid-detergent (LD) buffer: SB plus 0.1% CHAPS, 0.1% LMNG, 0.25 mg/mL 4:1 DOPC:CL and 10 μM BSA. Both NADH oxidation activity (340 nm, top) and cyt c reduction activity (550 nm, bottom) were monitored in separate experiments for each condition. Activity is reported as Units (1 U = 1 μmol substrate per min per mg of SC I+II2). Where Michaelis-Menten curve fitting is appropriate, the Vmax and apparent Km are shown on the plots ± standard error. When the concentration-activity curves are not hyperbolic, the maximum activity (Acmax) is shown, mean ± SEM. SC I+II2 was added to a final concentration of 1.3-10 nM depending on the experimental conditions and amount of activity. Measurements done using the Shimadzu UV-2600 UV-VIS spectrophotometer were carried out using disposable 2 mL polystyrene cuvettes (LLG Labware) with constant stirring. Measurements done using the Spectramax M2e Plate-Reader were done using 200 μL reaction buffer in 96-well plates with 10-20 s of stirring before beginning to record. All activity measurements were performed at 30 °C and initiated by addition of NADH. Concentrations of NADH stocks were determined using the Shimadzu UV-2600 UV-VIS spectrophotometer and the known extinction co-efficient of 6.22 mM⁻¹ cm⁻¹. These standards were then used to generate a standard curve on the Spectramax M2e Plate-Reader under experimental conditions in the presence of 100 μM cyt c. Standard curves for oxidized and reduced (by dithionite) cyt c were also generated in a similar fashion in situ under experimental conditions in the presence of 100 μM NADH. All protein concentrations were determined with the Pierce bicinchoninic acid (BCA) assay kit using BSA standards (Thermo Fisher, Waltham, MA). All samples for protein concentration determination were diluted at least 10-fold into 20 mM HEPES, pH 7.4, 0.1% DDM, and 50 mM NaCl buffer to reduce interference from glycerol in the sample buffers. See Figures 2, 3, and S2 and Table S3.

**CryoEM grid preparation, optimization, and data acquisition**

Initial grid preparation using the amphipol samples resulted in only broken particles. We hypothesized that this was due to instability of the SC at the large air-water interface present during blotting (Glaeser and Han, 2017; Glaeser et al., 2016) and found it necessary to add back detergent to reduce the surface tension and preserve the particles. Using the polyoxyether detergent Brij-35 generated high-quality grids with a high-density of particles in the holes, without solubilizing SCs. The final optimal blotting conditions were from application of 2.7 μL aliquots of –2 mg/mL amphipol stabilized SC I+II2 (in 250 mM NaCl, 20 mM HEPES, pH 7.7, 0.02% Brij-35) to glow-discharged holey carbon grids (Quantifoil R0.6/1 CU) followed by blotting for 30 s at 4°C, 95% humidity and flash freezing in liquid ethane using an FEI Vitrobot IV. CryoEM data acquisition was performed on a 300 kV FEI Titan Krios electron microscope with a Falcon II camera. Automated data collection was performed with the FEI EPU package. Micrographs were recorded at a nominal magnification of 100,000 X, resulting in a pixel size of 1.40 Å. Defocus values varied from −1.5 to −3.0 μm. The dose rate was ~50 electrons per pixel per second. Exposures of 2 s were dose-fractionated into 34 frames, leading to a dose of 1.5 electrons per Å² per frame and a total accumulated dose of 51 electrons per Å². A total of 1,854 micrographs were collected. See Table S4.

**Image processing and 3D reconstruction**

All processing steps were done using RELION 2.0 and 2.1 (Kimanius et al., 2016) unless otherwise stated. MotionCor2 (Zheng et al., 2017) was used for whole-image drift correction of each micrograph. Contrast transfer function (CTF) parameters of the corrected micrographs were estimated using Gctf and refined locally for each particle (Zhang, 2016). Automated particle picking in Relion resulted in ~400 k particles after manual curating. The particles were extracted using 400² pixel box and sorted by reference-free 2D classification followed by re-extraction at 512² pixel box and initial 3D classification with a regularization parameter T of 4 and a 30 Å low-pass filtered initial model generated from the PDB coordinates of CI and CII2 fit into the low resolution reconstruction of the ovine respirasome (Lettis et al., 2016b). This initial processing resulted in ~250 k particles of good quality, which separated into several classes containing SC I+II2, isolated CI or isolated CII2 (Figure S4C). As the SEC trace of the amphipol-stabilized SC I+II2 was monodisperse (Figure 1), we concluded that the isolated CI and CII2 were isolated during the grid preparation and plunge-freezing process. The isolated CI and SC I+II2 particles were separated and further classified to remove poor particles. At this stage, several distinct classes of the SC and isolated CI were identified and refined, resulting in reconstructions with nominal resolutions of 4.2 Å for three states and 4.6 Å for a fourth state according to the gold-standard FSC criteria.

Nonetheless, it was clear that the relative orientation between the two arms of CI and between CI and CII2 in SC I+II2 was variable, which resulted in lower resolution at the edges of the peripheral arm of CI and CII2 (Figure 7E; Figure S7) and likely limited the nominal resolution of the reconstructions. To improve the maps, focused refinements were independently performed around three regions of the SC: the CI peripheral arm, the CI membrane arm and CII2 (Figure S6). For the CI peripheral arm and CI membrane arm refinements, isolated CI particles were also included. After initial 3D classification, all of the CI-containing particles (i.e., isolated CI and SC I+II2) were combined and aligned via the peripheral arm of CI (Figure S5A). An additional round of 3D classification focused on the peripheral arm was performed to remove any remaining poor-quality particles. Interestingly, a small class of particles was identified that lacked the NADH-binding N-module of the peripheral arm (Figure S5A). The N-module is added to CI in the final stages of assembly (Formosa et al., 2018; Guerrero-Castillo et al., 2017) and its association may be dynamically controlled in response to ROS (Guarás et al., 2016). A class of particles lacking the N-module was also observed in the bovine respirasome structure (Sousa et al., 2016), suggesting that a sub-population of CI lacking the N-module is a common feature in mammalian mitochondria.

When we used the best class of 178,121 particles for 3D-refinement around the CI peripheral arm, we achieved a reconstruction at 3.8 Å, which allowed for atomic modeling of the CI peripheral arm (Figures S5A and S5D; Table 1). This class of 178,121 particles also defined the set of high-quality CI-containing particles that we used for all further CI 3D classifications (see below). Next, we aligned these particles via the CI membrane arm and performed an additional 3D classification to define a best-matching set of CI membrane
arm particles (Figure S5C). We further refined the best classes from this classification (174,334 particles) to generate a reconstruction at 3.9 Å resolution, again allowing for atomic modeling of the CI membrane arm (Figures SSC and SSD; Table 1). As these sets of particles contained both isolated CI and SC I+II,2, we performed the focused refinement for CIII2 with the original set of 120,596 particles identified in the original SC 3D classification (Figure SSB). After an additional round of 3D classification, we identified a set of 102,314 particles that, once refined, generated a 3.9 Å reconstruction around CII2 (Figures SSB and SSD). It was found that re-centering and re-boxing the particles around the CI peripheral arm and CII2 using the program REP (D’Imprima et al., 2017) improved map quality. This was not the case for the membrane arm which sits very near to the center of mass of the SC.

Finally, to better separate particles based on the difference between the CI membrane and peripheral arms, we developed a focus-revert-classify strategy in which the particles were first aligned by the peripheral arm of CI and then separated by 3D-classification with a mask around the membrane arm of CI without angular or translational searches (Figure 5). By pre-aligning particles by the peripheral arm, the small differences in the positions of the membrane arms should be maximized, thus facilitating efficient separation during 3D-classification without rotational or translational searches (Figure 5A). The peripheral arm was chosen for initial alignment as it contains all of the FeS cluster co-factors of CI, which generate strong peaks in the cryoEM density map. This strategy using ten classes resulted in the separation of at least seven distinct states of CI: one closed state and six open states (Figure S6A). The closed state and open states 1-3 each had significantly fewer particles than the original four SC states identified, but refined to similar or improved nominal resolutions, indicating that a more homogeneous set of particles for each state had been identified by this classification strategy. The nominal resolution of the seven classes ranged from 4.1-6.5 Å according to the gold-standard FSC criteria. Two additional classes at worse than 7.0 Å resolution were also generated but not analyzed further due to their poorer quality. Since no rotational or translational searches were performed, one class consisted of particles that did not align well with any of the other classes or each other and hence averaged out resulting in very weak and noisy density.

All local resolution maps were generated using BlockRes in Relion2.0 (Kimanius et al., 2016).

**Atomic model building and refinement**

Starting models for isolated ovine CI (Fiedorczuk et al., 2016) and bovine CIII2 (Iwata et al., 1998), corrected for the ovine sequence, were used. These models were split and fit into the highest-resolution focused refinement maps for separate atomic model building of the CI peripheral arm, CI membrane arm and CII2. Improved density in many regions relative to the previous maps of isolated CI allowed for the building of improved atomic models in COOT (Emsley et al., 2010). Real-space refinement of the model was done in PHENIX (Adams et al., 2010) and group atomic displacement parameters (ADPs) were refined in reciprocal space. Due to the unique features of cryoEM density (Coulomb potential density) relative to the electron density generated from X-ray diffraction data, such as frequent missing side chains for acidic amino acids caused by radiation damage (Baker and Rubinstein, 2010), we implemented a novel refinement protocol. First, we investigated whether the default values for group ADP refinements established for X-ray diffraction may be too stringent given the uncertainty in side chain position caused by radiation damage. Due to the moderate resolution of the reconstructions, individual (per-atom) ADP refinement was not performed, but, rather, group ADPs (one value for main chain atoms and another for side chain atoms) were assigned. ADP refinement in PHENIX is done using local sphere restraints for each atom ai with three adjustable parameters: the sphere radius, all atoms within this radius of atom ai will be used to constrain its ADP value (default value 5 Å); the distance power, this parameter determines the degree to which the distance between two atoms ai and aj within the sphere radius influences the restraint (X-ray refinement default value: 1.69); and the average power, which determines the degree to which the ADP is restrained by the average value of ADP between atom ai and aj (X-ray refinement default value: 1.03) (Afonine et al., 2005). As has been pointed out previously, the ADP can represent the convolution of two types of uncertainty: 1) the dispersion of the atom position in the molecules (which may be the result of temperature-dependent atomic vibrations or static disorder in a crystal lattice) and 2) the uncertainty of the experimenter’s knowledge about the atom position (Merritt, 2012). The second type of uncertainty can be taken care of by occupancy refinement, but that is problematic at current cryoEM data resolutions. Therefore, we choose to optimize ADP refinement specifically for cryoEM data. We found that the default values underestimated the ADPs for side chains with poor density (such as Asp and Glu) when the density for the main chain atoms was well defined. We therefore tested an array of ADP refinement parameters and found that decreasing the sphere radius to 2.5 Å and increasing the distance power to 2.3 without changing the average power resulted in more reasonable, higher group ADPs for side chains with poorly defined cryoEM density while maintaining overall consistency throughout the structure. Another complication arising from refinement of models with poor side chain density for specific residues is that, unless the ADP is calculated before coordinate refinement, the refinement will attempt to place the side chain into the nearest main chain density displacing the main chain atoms and distorting the local geometry. We therefore implemented a refinement protocol in which, after manual model building in COOT, a single macro cycle of ADP refinement was performed in PHENIX (using the new defaults above) before coordinate refinement was performed. This allowed the ADP for side chains with poor cryoEM density to rise and hence reduce their propensity to displace main chain atoms during global minimization. The single cycle of group ADP refinement was followed by three cycles of global minimization, followed by an additional cycle of group ADP refinement and finally three cycles of global minimization. This refinement protocol resulted in models with improved geometry, reasonable group ADP values and better correlation with the cryoEM maps. Additionally, the ADP values of the model agreed qualitatively more with the local resolution of the cryoEM reconstructions (Figure S7). The script to run the protocol is available from authors on request.
In previous work on isolated ovine CI, subunit NDUFA11 (B14.7), TMH4 of ND6, the C-terminal TMH16 of ND5 and the C-terminal half of the ND5 lateral helix were disordered (Fiedorczuk et al., 2016). Given that these regions were clearly resolved in the focused maps of the CI membrane arm (Figure S5C), we could now build their model at atomic resolution. Additionally, in the previous structure, subunit NDUFB8 (ASHI) was built mostly as poly-alanine (Fiedorczuk et al., 2016). In contrast, we now present it in full atomic detail here. We also made significant improvements to the models of subunits NDUFB4 (B15), NDUFB9 (B22) and NDUFB7 (B18). Besides these major improvements, we also achieved minor improvements throughout the structure due to the high quality of the focused refinements (Figure S5). Overall, the CI model presented here in the closed SC I+III2 is > 95% atomic and the remaining unbuilt residues are almost entirely found at disordered regions on the N- and C-termini of supernumerary subunits (Table S5). The overall structure of CIII2 is similar to what has been previously reported for isolated CIII2 and in the respirasome (Guo et al., 2017; Huang et al., 2005; Iwata et al., 1998; Letts et al., 2016b; Wu et al., 2016). One major difference in our structure is the lack of density for the single TM helix subunit UQCR11. UQCR11, which is located at the periphery of the Q-binding cavities on either side of the CIII2 dimer, is added late in CIII2 assembly (Smith et al., 2012), is not required for CIII2 activity and is easily lost by chromatographic de-lipidation of the complex (Huang et al., 2005; Hunte et al., 2000; Schagger et al., 1990). Although UQCR11 is considered an authentic subunit of CIII2, like many of the respiratory chain supernumerary subunits, its role is not well understood (Smith et al., 2012). UQCR11 was identified in our mass spec analysis (Table S1), which suggests that it is either sub-stoichiometric or disordered.

Due to low resolution, the focus-revert-classify reconstructions at \( R > 7.0 \) Å were not used in detailed structural comparisons, and poly-alanine models were only rigid-body fit into the reconstructions at 5.7-6.5 Å. We refined atomic models only for the four sub-4.5 Å reconstructions: the closed state and open states 1-3 (Figure 5B; Figure S6A).

QUANTIFICATION AND STATISTICAL ANALYSIS

All reported p values are from two-tailed Student’s t test for independent measures of unequal variance. Resolution estimations of cryoEM density maps are based on the 0.143 Fourier Shell Correlation (FSC) criterion (Chen et al., 2013; Rosenthal and Henderson, 2003).

DATA AND CODE AVAILABILITY

Data Resources

The accession numbers for the atomic coordinates reported in the paper are PDB: 6Q9D, 6Q9B, 6QE9, 6QBX, 6QC3, 6QC2, 6QC4, 6QA9, 6QC5, 6QC6, 6QC8, 6QC7, 6QC9, 6QCA and 6QCF.

The accession numbers for the EM density maps reported in this paper are EMDB: 4480, 4479, 4481, 4493, 4495, 4494, 4496, 4482, 4497, 4498, 4500, 4499, 4501, 4502, 4505, 4506 and 4507.