Interaction of complexes I, III, and IV within the bovine respirasome by single particle cryoelectron tomography

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The respirasome is a multisubunit supercomplex of the respiratory chain in mitochondria. Here we report the 3D reconstruction of the bovine heart respirasome, composed of dimeric complex III and single copies of complex I and IV, at about 2.2-nm resolution, determined by cryo-electron tomography and subvolume averaging. Fitting of X-ray structures of single complexes I, III2, and IV with high fidelity allows interpretation of the model at the level of secondary structures and shows how the individual complexes interact within the respirasome. Surprisingly, the distance between cytochrome c binding sites of complexes III2 and IV is about 10 nm. Modeling indicates a loose interaction between the three complexes and provides evidence that lipids are gluing them at the interfaces.

Oxidative phosphorylation (OXPHOS) in mitochondria is carried out by five multisubunit complexes (complexes I–V). They work in concert within the respiratory chain and catalyze the transfer of electrons from NADH to mitochondrial oxygen. This electron flow is coupled to proton pumping over the inner membrane, which generates a proton gradient utilized for ATP production by the ATP synthase (complex V). The five OXPHOS complexes are all catalytically active as single units, but their overall function changed after it was shown by blue-native gel electrophoresis (BN-PAGE) that they further associate into supercomplexes (1). The idea of their free occurrence within the cristae—or inner—membrane became progressively replaced by the concept of dedicated supercomplexes, which were proposed to be stable interactions of single complexes. The I + III2 and the V2 supercomplexes were the first ones to be structurally characterized (2–4), providing additional proof of their existence. The I + III2 + IV1–4 supercomplex or the respirasome is one of the most intriguing supercomplexes, because it autonomously carries out the respiration steps from electron transfer from NADH to mitochondrial oxygen. The exact reason for its presence, however, remains elusive. It was proposed that it could stabilize the single complexes, enhance the electron flow between these complexes, and prevent excess formation of oxygen radicals (5).

Single molecule averaging of EM projections of ice-embedded macromolecules can provide atomic resolution for large, water-soluble complexes (6). A common way to perform a 3D reconstruction of hydrophilic proteins is to use projections from randomly oriented particles. In amorphous ice layers, prepared on holey carbon film grids, to which the particles are adsorbed with a thinner second carbon film, to which the particles are adsorbed. An additional glassy carbon film is a suitable alternative, although this leads to preferential orientation of the complexes on the support structure, resulting in a limited range of covered orientation angles of the molecules. The random conical tilt data collection method could be used to compensate for the lack of randomness in orientations, by tilting the grids in the microscope to one specific angle (8). This method, however, requires the particles to be attached in one or several specific orientations to the support film. A powerful alternative data collection scheme that compensates for the lack of sampling over the full 3D space is electron tomography, in which the missing structural information is obtained from tilt series (9). Selected subvolumes of tomographic reconstructions (tomograms), containing specific protein complexes, can then be further averaged to enhance the signal and resolution (10, 11). In this study, tomographic subvolume averaging was applied on the I + III2 + IV supercomplex, further referred to as the respirasome, which is a highly abundant supercomplex in bovine heart mitochondria. Fitting high-resolution structures of its three components—dimeric complex III (12), monomeric complex IV (13), and monomeric complex I (14)—into a cryo-EM reconstruction at 2.2-nm resolution provides first insight into the unique interaction between these complexes within the respirasome.

Results

The bovine heart I + III2 + IV supercomplex was purified to homogeneity by BN-PAGE, followed by electrophoretion, a prerequisite to perform a cryo-electron microscopy study. Preparation of frozen hydrated cryo-EM grids by adsorption of the purified sample onto holey carbon film grids and plunge-freezing, however, resulted in inhomogeneously distributed particles in the grid holes. We therefore used holey carbon film grids that were covered with a thinner second carbon film, to which the particles could adsorb. This approach resulted in evenly distributed molecules (Fig. 1). Single particle analysis, including classification and averaging of two-dimensional projections, showed that the respirasome molecules were preferentially oriented in a few positions on carbon support film. The most abundant projection type has a triangular shape (Fig. 1, Inset), as also previously noticed in negatively stained samples (5). We performed electron tomography on these cryo-EM specimens and from 21 tilt series the 3D volumes of the samples were reconstructed. From these tomograms, 2,466 subtomograms containing single respirasomes were extracted for 3D averaging. Combination of XMIPP (X-Window-based Microscopy Image Processing Package) maximum likelihood (ML) global alignment (11) and local refinement by cross-

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Data deposition: The EM map of the respirasome has been deposited in the EMDataBank, www.emdatabank.org (accession no. EMD-5319).

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correlation with AV3 toolbox (15) was used to create an average structure. The resolution of the respirasome is anisotropic (Fig. S1) and correlates to the angular sampling (Fig. S2). The highest achievable resolution of our average structure in the direction perpendicular to the incident beam is 1.7 nm and the lowest one is 2.55 nm parallel to the beam. We have filtered the entire reconstruction to 2.2 nm, which is at the first zero of the contrast transfer function.

The reconstruction shows a respirasome with dimensions of $27 \times 21$ nm in the membrane plane (Fig. 2 B and C) and a maximal height of 23 nm (Fig. 2B), made by the giant hydrophilic domain of complex I. By comparison with known 3D structures of the individual complexes, many relevant features are recognizable, such as protruding densities at the center, related by a local twofold symmetry axis (Fig. 2, arrows), that point to the position of the core I and II subunits of dimeric complex III. Complex III$_2$ sits in the arc of the membrane arm of complex I and complex IV is attached to the tip of NADH dehydrogenase and complex III$_2$ (Fig. 2, arrowheads). To closely interpret the cryo-EM map, we superimposed the high-resolution X-ray structures of bovine complexes III$_2$ (12) and IV (13). For modeling of the complex I moiety, the medium-resolution X-ray data from the yeast *Yarrowia lipolytica* were used (14). Fitting was accomplished with correlation coefficients between 0.83–0.92 for the bovine structures, indicating a unique, close fit with high confidence of components in the final hybrid structure (Fig. 3 A, B, and D, and Fig. S3). No substantial parts of the single complexes extend the 3D cryo-EM reconstruction, indicating that the respirasome is basically the sum of its components. Based on the fitting, our 3D structure presents a complete respirasome, including the peripheral arm domain of complex I (Fig. 2 A and B, arrowheads), which sometimes is partly lacking in negatively stained samples (16). Some minor differences exist between the EM reconstruction and the X-ray maps (Fig. 3 A, B, and D, orange arrowheads), which are most likely detergent molecules surrounding the respirasome. Iterative multireference alignment and classification (17) showed the presence of the peripheral arm in all five classes with a slight amount of variation (Fig. S4). Subtraction of two reconstructions representing two different classes revealed a density (Fig. S4, red asterisk), which can be accounted for a possible conformational change in the hydrophilic domain of complex I. To prove it, additional experiments are required. The full complex I moiety of bovine and *Yarrowia* are equal; the peripheral matrix arms make the same angle (Fig. 3 A, yellow) and the bend of the membrane arm is closely the same (Fig. 3 D, double arrowhead). The superposition of the smaller prokaryotic complex I from *Thermus thermophilus* indicates a similar overall shape (Fig. S5) and, in addition, clearly shows the locations of the

Fig. 1. Cryoelectron micrograph of isolated bovine respirasomes. (Scale bar: 100 nm.) (Inset) Two-dimensional projection map of the respirasome in a top view position from single particle image analysis. (Scale bar: 10 nm.)

Fig. 2. Cryo-EM map of I + III$_2$ + IV supercomplex: (A and B) seen from aside, (C) seen from the matrix side; arrow points to complex III$_2$, arrowhead to complex IV, and double arrowhead to the curved membrane arm of complex I, respectively. (D) Seen from aside from the tip. Contour level is 0.1. Horizontal lines on A and D indicate the position of the membrane. (Scale bar: 10 nm.)
accessory subunits of the eukaryotic enzymes are located. These accessory proteins appear to be arranged mostly in clusters.

Discussion

Single particle reconstruction can be performed in several ways. The most straightforward way is to reconstruct 3D structures from randomly oriented particles in a nontilted thin layer of vitreous ice (7). The respirasome, however, showed a strong preferential orientation on the carbon film. Therefore electron tomography reconstruction and subvolume averaging was applied. This method was previously used to visualize a number of membrane complexes in situ, like the nuclear pore complex in its native membrane (10) or protein–protein interactions in intact organelles (18, 19). Recent advances in automation of electron tomography data collection allowed more efficient subtomogram averaging, as applied for the study of capsids of herpes virions (20). Lately, the effectiveness of the maximum-likelihood processing approach of subtomograms was demonstrated on chaperonin GroEL/GroES complexes, showing a more reliable convergence at lower signal-to-noise levels (11). We performed an ML analysis to computationally extracted single respirasome subvolumes, which minimizes the reference bias for the reconstruction (21). Analysis of 2,466 subtomograms resulted in a 3D structure of the respirasome at an anisotropic resolution of 1.7–2.55 nm. However, the first zero in the contrast transfer function (CTF) curve at 2.2 nm$^{-1}$, the details with spatial frequencies of 2.2 to 1.7 nm$^{-1}$ have a reversed phase sign and are removed from the final reconstruction. This finding indicates that, in principle, the applied electron tomography and extensive 3D averaging could even be pushed to higher resolution, if reliable methods to perform CTF correction on the highly noisy images of the original tilt series would become available. From the 3D reconstructions, the position toward the support film of the single particles could be retrieved. About 70% of the respirasomes were found in one specific orientation, resulting in a triangular projection view (Fig. 1), and another 12% of the particles were in a side position with respect to the carbon support film (Fig. S2B). There was angular variation around both preferential orientations, and 10% of the particles were found in different, random orientations. This additional variation plus the tomographic tilting of the grids lead to a good sampling of angles over the Fourier space, demonstrating the advantages of subtomographic averaging for membrane proteins, when particles preferentially attach to a carbon film.

Although the 2.2-nm resolution cryo-EM model does not provide direct information about the secondary structure, the fitting allowed an interpretation at the level of α-helices, reflecting that the fitting procedure corresponds to a resolution gain by a factor of approximately 5 (22). From the obtained model, we can conclude that the interaction sites of complexes III$_2$ and IV are different from the negatively stained data of the bovine respirasome (16) and the III$_2$ + IV$_2$ supercomplex from *Saccharomyces cerevisiae* (23). Complex IV contacts to complex III$_2$ with the side opposite to the complex IV dimer interface in the X-ray structure (13) (Fig. 3 A, B, D, and F). Cytochrome $b$, the Rieske protein, and subunit 11 of one particular monomer of complex III are in close proximity to subunits III, VI$_a$, and VII$_a$ of complex IV. The cytochrome $c_1$ subunit is far away from the complex III$_2$–IV interface (Fig. 3 C, and F, blue helices), in contrast to what was concluded from the negative stain map of the bovine respirasome. The previously available respirasome EM map had resulted from negatively stained samples and did not show any distinct complex IV features that could have been of help in the proper positional assignment of complex IV (16). In contrast, in the current cryo-EM map, we can clearly distinguish subunits protruding outside the membrane. Subunit II of complex IV does not seem to be involved in the interaction between complex III and IV as it was proposed before (16). Subunits II, VI$_a$, and VB form a concave surface on top of subunit I, which could form an interaction site for a cytochrome $c$ molecule (2.5 nm in diameter; ref. 13).

Interestingly, the single complexes I, III$_2$, and IV appear after fitting to be at some distance in the model, suggesting that there is little close contact (Fig. 3 C, E, and F). The section through the model on the level of the membrane demonstrates the gap between complexes III$_2$ and IV within the membrane (Fig. 3C). The complexes appear to contact each other in the membrane close to the matrix (Fig. 3F). It is possible that interactions are enhanced by lipid molecules such as cardiolipids, which were proposed to be essential in supercomplex formation (24, 25). In the structure of bovine complex IV, one phosphatidylethanolamine was found on subunit VI$_a$ and two phosphatidylethanolamines plus one
phosphatidylglycerol molecule were revealed in the larger cleft of subunit III, which may serve as a lipid pool (13). The involvement of lipids may be special for maintaining the structure of membrane supercomplexes because in another large membrane protein complex, dimeric Photosystem II, 12 lipid molecules were at the interface between the monomers, of which four sulfonovosyldiaclyglycerols might promote dimerization (26).

Complexes I and III* II are physiologically linked by the movement of quinones. The ubiquinone binding site at complex I in the matrix arm of the interface with the membrane arm (27) and cytochrome b of complex III binding hydroquinone (12) are facing each other in our cryo-EM map and are at a distance of 13 nm (Fig. 4A), allowing efficient ubiquinone transport between binding sites of complexes I and III. The distance that cytochrome c has to diffuse between the sites of complexes III and IV is 10 nm in the respirasome (Fig. 4B), but less than 4 nm in the III* + IV* supercomplex (23). Nevertheless, the formation of the respirasome brings these sites closer to each other than in a situation of randomly distributed complexes III and IV molecules in the membrane, which is probably beneficial, because in free random diffusion the time is proportional to the square of the distance. But although diffusion coefficients have been determined for cytochrome c (28), the rate at which cytochrome c diffuses is not directly measurable, because it depends on the local free water volume. Hence, we cannot quantify the effect of the (variable) distance on cytochrome c diffusion time. However, it is questionable if distance between electron transport components is the main reason for the supercomplex formation. Moreover, flux control experiments have not demonstrated substrate channeling between complexes III and IV via cytochrome c, although substrate channeling was found between bovine complexes I and III via ubiquinone (29). Likely, there are other reasons for the respirasome formation, such as overall regulation of oxidative phosphorylation. Another one is the assembly of the components.

It was shown earlier that complex III can stabilize complex I in mammalian mitochondria (30). Another may be on an even higher level of organization, which is the respiratory string, a linear association of respirasomes. Our model is not in conflict with previously proposed models of respiratory strings (31, 32). According to Wittig et al. (31), the ratio between complex I, III, and IV in bovine mitochondria is equal to 1:3:6. Based on this ratio, they proposed the string in which every respirasome is connected via complex IV to another respirasome and III* + IV* supercomplex. Although the interfaces of complexes III*–IV are different than in yeast supercomplex, the site of complex IV, which is involved in dimerization, as it is revealed by X-ray crystallography, is available for the attachment of the next respiratory unit via the complex IV–complex IV interface. Finally, in mammalian mitochondria, the respirasome may prevent superoxide formation, a very harmful, naturally occurring process in mitochondria. Most of the mitochondrial superoxide originates from complex I and complex III (33).

**Materials and Methods**

Beef heart mitochondria were solubilized in 30 mM Hepes buffer with 10% digitonin, 0.15 M potassium acetate, 10% glycerol, pH 7.4 at 4°C at a ratio of 25 g detergent per 1 g of protein. The solubilized fraction was separated with BN-PAGE as described (2). The band with respirasomes was excised from an unstained gel, supplemented with 0.01% digitonin and electroeluted in a Bio-Rad electrophoresis (34). For tomography, the sample was concentrated by centrifugation in Microcon tubes; aliquots of 2.5 μl were mixed with 10-nm gold particles as fiducial markers and applied on glow-discharged 200 mesh Quantifoil support grids (Quantifoil Micro Tools, GmbH) with an additional carbon support film. Grids were blotted at 100% humidity for 4–6 s at a blot offset (the longitudinal grid positioning) setting of ~3.5, using a Vitrobot Mk3 (FEI). Data collection was performed with a 300-kV G2 Polara electron microscope (FEI) equipped with a Gatan energy filter. Images were recorded at 200 kV accelerating voltage for tomography or 300 kV for 2D projections processing with a 2,000 × 2,000 CCD camera (Gatan) at 2 μm underfocus (first CTF zero at 2.2 and 2 nm, respectively) and 78,000 of magnification, resulting in a sampling with 0.38-nm pixels. Tilt series were acquired with tilt angle increments covering a range of ±65°, with a total dose of 8,000 electrons/nm². Two-dimensional image processing of projections was done with the Groningen Image Processing software package (2).

A total of 21 tomograms were reconstructed by weighted back-projection using IMOD software (35). No CTF correction was performed. In total, 2,466 subvolumes of respirasomes with a box size of 128 × 128 × 128 pixels were selected from the tomograms by cross-correlation to an artificial globular reference followed by manual selection of particle volumes. An average structure was generated using the XMiPP ML for tomography software (11) on volumes downsampled to a pixel size of 0.76 nm. Averaging was initiated with a sum of randomly oriented particles and after 20 iterations with 2° global angular search the structure reliably converged. Further decreasing the angular search to 10° and 5° yielded an average at resolution of 3.2 nm. XMiPP ML calculations were run on 16 or 32 cores on the Maia cluster of the University of Basel. The alignment transforms were imported to the “AV3” routine for Matlab (15) and local 6D refinements on the volumes with initial voxel sizes of 0.38 nm were performed reaching the final average structure at resolution 2.2 nm. About one-half of the particles (1,174) with the best correlation coefficients were used to produce the final average. Resolution estimation was performed by the 0.5 information threshold criterion (36). Resolution criterion yielded a value that is below the first zero of the CTF, thus the resolution was assumed to be 2.2 nm. Maximum likelihood classification (17) into five classes was performed starting with the average structure at 3.2 nm with eight iterations allowing local angular search of 10°.

**Fig. 4.** Scheme of electron flow within the respirasome mediated by cytochrome c and ubiquinol. (A) Side view of the I + III* II + IV supercomplex. Black arrows point how the ubiquinol and the cytochrome c move between complexes I, III* II, and IV. (B) Top view of the respirasome fragment consisting of complexes III* II and IV, seen from the intermembrane space. The distance between cytochrome c binding sites of complexes III* II and IV is 10 nm. In yellow, the density map of complex I from Yarrowia lipolytica, marked as I; in gray, X-ray structures of bovine complexes III* II and IV, marked as III* II and IV correspondingly. Q, ubiquinol; cyt c, cytochrome c. Colors: red, cytochrome c; dark green, subunit II of complex IV; blue, cytochrome c; green, subunit of complex III* II; green, cytochrome b of complex III* II; magenta, “Rieske” protein of complex III* II. Horizontal lines on A indicate the position of the membrane.
For the anisotropy test, we measured Fourier shell correlation in cones of Fourier space for different orientation of the cone according to unit sphere \( \varphi-\theta \). The cone had a semiaperture angle of 22.5°, angles were spaced every 18°. The resolution in the direction of each cone was determined by the 0.5 threshold. This routine is a feature of a yet unreleased Dynamo software package for subtomogram alignment.

X-ray structures of the bovine dimeric cytochrome bc, complex [Protein Data Bank (PDB) ID 1BGY; ref. 12], monomeric cytochrome c oxidase (PDB ID 1OCC; ref. 13), Yarrowia lipolytica NADH:ubiquinone oxidoreductase (14), and NADH:ubiquinone oxidoreductase from Thermus thermophilus (PDB ID 3M9S; ref. 27) were used for modeling of the electron density map of the respirasome at a voxel size 3.8 Å. The monomeric complex I (PDB ID 3M9S; ref. 27) were used for modeling of the electron density map of the respirasome at a voxel size 3.8 Å. The monomeric complex I was generated from the density map by manually erasing of the densities assigned to the neighboring molecule in the crystal unit by using the Chimera program from University of California, San Francisco (UCSF) (37). The maps were first prealigned manually and then automatically by cross-correlation using the Fit in Map tool in the UCSF Chimera program (37).

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