Molecular characterization of a complex of apoptosis-inducing factor 1 with cytochrome c oxidase of the mitochondrial respiratory chain

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Combining mass spectrometry–based chemical cross-linking and complexome profiling, we analyzed the interactome of heart mitochondria. We focused on complexes of oxidative phosphorylation and found that dimeric apoptosis-inducing factor 1 (AIFM1) forms a defined complex with ~10% of monomeric cytochrome c oxidase (COX) but hardly interacts with respiratory chain supercomplexes. Multiple AIFM1 intercross-links engaging six different COX subunits provided structural restraints to build a detailed atomic model of the COX-AIFM12 complex (PDBDEV_00000092). An application of two complementary proteomic approaches thus provided unexpected insight into the macromolecular organization of the mitochondrial complexome. Our structural model excludes direct electron transfer between AIFM1 and COX. Notably, however, the binding site of cytochrome c remains accessible, allowing formation of a ternary complex. The discovery of the previously overlooked COX-AIFM12 complex and clues provided by the structural model hint at potential roles of AIFM1 in oxidative phosphorylation biogenesis and in programmed cell death.

In the present study, we explored the molecular interactions of AIFM1 with the multiprotein complexes of the OXPHOS system in heart mitochondria using our recently established complementary experimental approach (11) that combines cross-linking mass spectrometry (XL-MS) and complexome profiling (Fig. 1). To increase the depth and confidence of the study, bovine heart mitochondrial membranes (BHM) were treated with three different chemical cross-linkers: DSSO (12), PhoX (13) and DMTMM (14). While DSSO and PhoX predominantly generate lysine–lysine residue cross-links, DMTMM acts as a condensation reagent of acidic side chains of aspartic or glutamic acids with lysine side chains, resulting in the formation of a stable bond between those residues. We found that a significant fraction of AIFM1 in its dimeric form is specifically bound to monomeric COX, an interaction that has been overlooked so far. By using the identified cross-links as structural restraints, we generated a structural model of dimeric AIFM1 docked to monomeric COX.

Mitochondria are considered the powerhouse of aerobic eukaryotic cells, as they contain the major pathways of oxidative energy metabolism and produce the bulk of ATP by oxidative phosphorylation (OXPHOS) necessary for cellular homeostasis. Only at the end of the last century it became evident that mitochondria also are key players in apoptosis and that this process is tightly linked to OXPHOS components (1). Apoptosis-inducing factor 1 (AIFM1) was one of the proteins found to be released from the mitochondrial intermembrane space during programmed cell death and to have the capacity to induce chromatin condensation and DNA fragmentation in a caspase-independent fashion (2). A mutation found in AIFM1 has been associated with Cowchock syndrome [OMIM 310490] (3). Early on, it was also reported that ablation of AIFM1 leads to OXPHOS deficiency (4) in line with findings that AIFM1 mutations cause combined oxidative phosphorylation deficiency 6, a severe mitochondrial encephalomyopathy [OMIM 300816] (5). More recently, it has been proposed that AIFM1 is involved in the disulﬁde relay of the mitochondrial intermembrane space by serving as an import receptor of CHCHD4/MIA40 (6–8). However, the speciﬁc mechanisms and molecular interactions by which these different functions of AIFM1 are connected in health and disease are not well resolved. For example, AIFM1 deficiency affects OXPHOS predominantly by lowering the amount of respiratory chain complex I (4). Other components were found to be affected in a tissue-specific manner. In AIFM1-deficient patients (5), ablation of AIFM1 in skeletal and heart muscle affected cytochrome c oxidase (COX) in addition to complex I, whereas in liver, deficiency of complexes I and V was observed (9, 10).

Significance

Apoptosis-inducing factor 1 (AIFM1) resides within the intermembrane space of mitochondria and upon programmed cell death was found to induce chromatin condensation and DNA fragmentation. While the apoptosis-related role of AIFM1 is well understood, recent findings pointed to additional, not well-characterized functional roles of AIFM1 in oxidative phosphorylation. Using cross-linking mass spectrometry and complexome profiling, we uncover that a substantial amount of dimeric AIFM1 is engaged with ~10% of monomeric cytochrome c oxidase (COX). Further structural modeling and restraint-driven docking structurally characterize a COX-AIFM12 complex, not only highlighting how AIFM1 might be N-terminally inserted into the inner mitochondrial membrane but also providing clues on potential functional implications including an involvement in promoting apoptosis.

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Results and Discussion

We analyzed the organization and interaction landscape of protein complexes in BHM by combining XL-MS and complexome profiling (11, 15), thereby adding new information on native state multiprotein complexes of interest and expanding previous work that explored the interactome of mitochondria from different organisms, tissues, and cells by XL-MS (16–20).

Exploring Mitochondrial Complexes by Combined Cross-linking and Complexome Profiling. To increase the depth of the protein interaction map of BHM, we applied multiple cross-linkers (DSSO, PhoX, and DMTMM). Throughout the manuscript, the term cross-link is used to describe a link between two residues coming from two unique peptides, with an intra-cross-link describing a linked residue pair within a protein and an inter-cross-link describing a linked residue pair between two different proteins. Additionally, identified cross-links were filtered so that only cross-links corresponding to protein–protein interactions that were reported for at least two cross-linkers and with at least two cross-link spectrum matches (CSMs) were kept. Covering 215 proteins listed in MitoCarta 3.0 (21), we obtained a total of 4,413 unique cross-links (3,261 intra- and 1,152 interprotein cross-links; SI Appendix, Fig. S1A and Dataset S1). In accordance with previously published studies (22, 23), the abundance of detected cross-linked proteins was higher than the median of all identified proteins in the BHM sample [8.8 versus 6.9 log_{10} intensity-based absolute quantification (iBAQ); SI Appendix, Fig. S1B]. Reflecting the large number and high abundance of membrane integral multiprotein complexes and the very high protein density, especially within the inner mitochondrial membrane, ~75% of the cross-links identified involved membrane proteins (SI Appendix, Fig. S1C). For the same reasons and corroborating previous studies using mouse and human mitochondria (16, 17, 19, 21), the largest number of cross-links reflected interactions between the many subunits of OXPHOS complexes and their association to supercomplexes of respiratory chain complexes I, III, and IV (1,431 out of 4,131 cross-links; SI Appendix, Fig. S1D), also called respirasomes (24). Furthermore, interdomain cross-links for complex I-III and COX were very good agreement with previously published structural models but providing no indications for homodimerization (complex I and II and COX) or multimerization (complex III) (Dataset S1). In contrast, a significant portion of interdomain cross-links for complex V showed substantially more apparent restraint violations (Dataset S1). Most likely, this reflected the formation of higher order ATPase assemblies involved in shaping tightly curved cristae ridges (25–27), which can also be observed in the complexome profiling data (SI Appendix, Fig. S1E).

Complexome profiling analysis of untreated (i.e., non-cross-linked) BHM yielded very similar results as those obtained previously with rat heart mitochondria using the same approach (15) showing a very similar migration pattern of the OXPHOS complexes and respirasomes (SI Appendix, Fig. S1E and Dataset S2). When the samples were cross-linked with PhoX and DMTMM before subjecting them to complexome profiling, the overall abundance of detected proteins was not affected substantially. However, it was evident from the migration profiles of OXPHOS complexes that cross-linking to some extent prevented dissociation of complex V (CV) dimers and other fragile higher order respiratory supercomplexes during native electrophoresis (SI Appendix, Fig. S1E). Importantly, in most cases, the apparent molecular masses of the bulk of the OXPHOS complexes were not markedly affected by cross-linking. A notable exception was complex III (CHII) in the DMTMM-treated sample, where the obligatory dimer did not migrate predominantly at the predicted apparent mass of ~500 kDa as in all other conditions but appeared at ~650 kDa and showed multiple peaks at higher masses. The shift of the CHII dimer to higher masses suggested that, possibly through the large hydrophilic domains of its two core subunits, this OXPHOS complex cross-linked to a much larger extent to other mitochondrial proteins than the others. The ~800 kDa peak corresponds to a previously described supercomplex between one CHII dimer and one complex IV (COX) monomer (28). The latter was also found in untreated and PhoX cross-linked samples but was much more pronounced after cross-linking with DMTMM. The peaks at ~1,100 and ~1,300 kDa can be interpreted as dimers of CHII dimers without and with one monomer of COX, respectively.

Taken together, these results establish that classical XL-MS analysis alone and in combination with complexome profiling delivered consistent results. Separating native complexes prior to mass spectrometric analysis provided additional key information on their apparent molecular masses and multimeric state. Cross-linking them beforehand allowed for more reliable detection of...
more fragile assemblies that otherwise partially or completely dissociate during solubilization and/or electrophoresis.

**A Specific Complex between AIFM1 Dimers and COX Revealed by XL-MS.** When we performed an in-depth analysis of all detected cross-links involving OXPHOS complexes in addition to engaging their canonical components themselves, one specific protein stood out: in all our XL-MS datasets combined, AIFM1 had intercross-links with no less than six subunits of COX, with 82% of them involving COX6B1 and COX6C (Fig. 2A and Dataset S3). Cross-links with COX subunits accounted for 86% of the total inter cross-links with AIFM1. Adenylate kinase 2 (AK2) and adenine nucleotide carrier isoform 1 (SLC25A4) were the only other two proteins featuring multiple interprotein cross-links with AIFM1.

The association of AIFM1 with this OXPHOS complex is remarkable in particular since COX from a bovine heart is undoubtedly the longest and best studied version of COX (29). Therefore, we interrogated an earlier cross-linking dataset of mouse heart mitochondria for this interaction (16). Corroborating our findings, the majority of AIFM1 cross-links identified in this study engaged three different COX subunits, with COX6C being the most prominent by far (SI Appendix, Fig. S2A). Of note, Liu and coworkers (16) detected multiple cross-links between AIFM1 and AK2 as well in mouse heart mitochondria. Moreover, charting large affinity purifications MS (AP-MS) depositories, we found that they contained multiple instances of COX subunits interacting with AIFM1 (30–32) (Dataset S2). Yet, buried in datasets generated by large-scale analyses of the mitochondrial interactome,

**Fig. 2.** Dimeric AIFM1 forms a defined complex with monomeric COX. (A) Interaction network of AIFM1 in cross-linked BHM. Bold numbers indicate the observed cross-links for each interaction, and the thickness of lines indicate the cumulative evidence (CSMs) for each interaction (number in parentheses). Orange lines indicate cross-links involving AIFM1, while cross-links between AIFM1 interactors are presented as gray lines. (B) Xi-net plot of the COX-AIFM1 interaction. Purple colored links indicate intracross-links. Green colored links indicate intercross-links. Respective sequence and cross-link features are indicated accordingly. (C) Migration profiles of AIFM1 (orange) and averaged COX (green) from non–cross-linked (untreated) and cross-linked (PhoX or DMTMM) mitochondria separated by BN-PAGE (4 to 16%). Peaks are annotated based on the apparent molecular mass of AIFM1 (≈62 kDa) and monomeric COX (≈220 kDa). In all samples, peaks corresponding to monomeric AIFM1 and COX as well as a peak corresponding to a COX-AIFM1 complex are observed. Although already present in the non–cross-linked sample, treatment with DMTMM seems to somewhat stabilize the COX-AIFM12 complex.
these indications for AIFM1 binding to COX seem to have gone unnoticed so far.

Detailed evaluation of the observed cross-links between AIFM1 and COX (Fig. 2B and SI Appendix, Fig. S2B) revealed that they were predominantly within the pyridine nucleotide-disulfide oxidoreductase domain (Pfam: PF07992; residues 136 to 460) of AIFM1 comprising one FAD- and one NADH-binding domain. Suggesting that AIFM1 had not been cleaved to its truncated proapoptogenic form (33), additional intra- and interprotein cross-links were observed at the N-terminal end of the proppeptide (residues 55 to 101) of AIFM1 that is predicted to cross the inner mitochondrial membrane reaching to the matrix side. Notably, these cross-links were the only ones to the matrix-facing subunit COX5A, while all other cross-links engaged domains of COX subunits facing the intermembrane space.

Our three independent cross-linking analyses strongly suggested that AIFM1 and COX formed a specific complex but provided no information on the multimeric state of the interaction partners and how much of this unexpected complex was present in BHM. Therefore, we applied complexome profiling to analyze complexes containing AIFM1 and COX using the same samples as in the XL-MS analysis (Fig. 2C and Dataset S2). In all cases, COX was predominantly present as a monomer (~220 kDa), and a prominent fraction of AIFM1 was found to migrate at an apparent mass consistent with its monomeric state (62 kDa). Substantial amounts of AIFM1 dimers were only observed in untreated BHM, indicating that they may be destabilized by the cross-linking protocol. This was possibly due to partial oxidation of NADH known to be required for AIFM1 dimerization (8). Importantly, however, a significant amount of AIFM1 consistently showed up as a peak at an apparent mass of ~350 kDa in untreated mitochondria as well as after cross-linking with PhoX or DMTMM. This peak coincided with a shoulder next to the prominent peak at ~220 kDa of monomeric COX in all samples analyzed, thus suggesting the presence of an ~350 kDa complex containing a dimer of AIFM1 (~124 kDa) bound to monomeric COX (~220 kDa). Notably, a shoulder on the higher mass side of the COX monomer can also be observed in complexome profiling data of human cells published earlier, but its significance was not evident at the time (34). Label-free quantification revealed that hardly any of the other respiratory chain subunits COX6B1, COX6C, MT-CO2, and NDUFA8 are too far apart from each other to be consistent with simultaneous binding of the two OXPHOS complexes to AIFM1 within the respirasome. This is fully in line with our cross-link data and proposed model of the complex.

Creation of a Cross-link–Guided Structural Model of the COX-AIFM12 Complex. Next, we aimed at building a structural model for the COX-AIFM12 complex guided by the distance restraints obtained by cross-linking, also including those involving the N-terminal sequence of AIFM1 comprising its proppeptide sequence (residues 55 to 101). We first derived a de novo model of this so far structurally unresolved region using trRosetta (39) to complement a homology model of the bovine AIFM1 dimer that we derived from the human structure [PDB: 4BUR (40)] using Robetta (41). The structural model obtained for the N-terminal domain of AIFM1 agreed well with secondary structure predictions and featured three alpha-helices (residues 67 to 88; 94 to 98; 105 to 112), of which the first is predicted to be a transmembrane segment (SI Appendix, Fig. S3A). We then used restraints derived from our cross-linking data to dock this model of the bovine AIFM1 dimer to the 1.8 Å structure of COX (PDB: 1V54) (42). Unfortunately, this COX structure does not contain the more loosely attached NDUFA4 subunit. Therefore, we used Robetta (41) to complement it with a homology model derived from the human NDUFA4 structure (PDB: 5Z62 chain N) (43).

Mapping the cross-links onto the structural models of AIFM12 and COX, respectively, revealed that the majority of cross-links were below 30 Å, with a combined mean distance of 19.1 Å for DSSO/PhoX and 20.8 Å for DMTMM cross-links (SI Appendix, Fig. S3 B and C and Dataset S3). The mean distance for DSSO and PhoX cross-links was well within the theoretical maximal range of ~30 and ~25 Å, respectively. DMTMM cross-links averaged somewhat above the theoretical maximum of ~15 Å, in line with previous observations (14). Note, that eight cross-links for AIFM1 and 10 cross-links for COX were not included in these calculations because they involved intracross-links from AIFM1 (residues 128 to 613) to its de novo modeled N terminus (residues 55 to 124) or regions not resolved in the structural models (AIFM1 residues 517 to 550; COX4L1 residues 23 to 25; SI Appendix, Fig. S3C).

Based on solvent accessibility and distance restraints obtained from both structures, accessible interaction interfaces between COX and the AIFM1 dimer as well as COX and the N-terminal region of AIFM1 were calculated using DisVis (44). While this analysis suggested that the AIFM1 dimer attaches to the intermembrane space side of COX, the predicted interaction space for the N-terminal region of one AIFM1 protomer covers the transmembrane domain at the matrix side of COX making contacts to subunits COX6B1, COX6C, MT-CO2, and NDUFA4 (Fig. 3A). Scoring the interface models using the restraints imposed by the cross-linking data suggests that the COX-AIFM12 interface is mostly occupied by just one AIFM1 protomer. In agreement with this notion, cross-links suggested that only one N-terminal region, not both of the AIFM1 dimers, interacted directly with COX.
Therefore, the final modeling of the COX-AIFM1\textsubscript{2} complex was performed by docking just one AIFM1 protomer and one N-terminal region of AIFM1 to the COX monomer. Docking was restrained by 46 of 59 unique cross-links detected for the COX-AIFM1 interaction. Haddock (45) generated six acceptable clusters (with negative Haddock score). One cluster (cluster 7) showed better scores (overall lowest Haddock score and interface root-mean-square deviation [RMSD]) compared to the other clusters (SI Appendix, Fig. S4A). An investigation of the individual structures of produced clusters revealed that only for structures of cluster 7, the N-terminal domain of AIFM1 was positioned in accordance with its transmembrane domain. Furthermore, structural alignment of models from cluster 7 showed high cluster precision and only differed marginally from each other (RMSD = 0.942 Å; SI Appendix, Fig. S4B). Lastly, clusters were validated by mapping the used cross-link restraints on the highest scoring model of each cluster, with the structure of cluster 7 satisfying them best (Dataset S4). Based on overall best cluster scoring and high cluster precision, the highest scoring structure of cluster 7 was chosen as a representative model for the COX-AIFM1\textsubscript{2} complex.

In the final structural model of the COX-AIFM1\textsubscript{2} complex, the AIFM1 dimer “sits” on COX, facing the intermembrane space side, and makes contact through one AIFM1 protomer covering parts of COX6\textsubscript{B1}, COX6\textsubscript{C1}, MT-CO\textsubscript{2}, and NDUFA4 (Fig. 3B). The second AIFM1 protomer points away from COX, making just one very limited contact to COX through its C-terminal loop. At the opposite side of COX, the N-terminal region of the interacting AIFM1 moiety is highlighted in red. Membrane boundaries of the inner mitochondrial membrane are sketched as gray spheres. The final complex consists of monomeric COX, dimeric AIFM1 (residues 128 to 516, 551 to 613), and the N-terminal region of one AIFM1 protomer (residues 55 to 127).

![Fig. 3. Cross-link–derived structural model of the COX-AIFM1\textsubscript{2} complex. (A) Visualization of the cross-link–driven accessible interaction space models for a COX-AIFM1\textsubscript{2} complex. COX is represented in green, while the bright orange volume represents the center-of-mass position of the AIFM1 dimer, and the dark orange volume represents the center-of-mass position of the model of the AIFM1 N terminus (residues 55 to 124). The cross-linking data are consistent with the interaction space available for docking dimeric AIFM1 and the N-terminal region of one AIFM1 protomer to monomeric COX. (B) Cross-link–derived structural model of the COX-AIFM1\textsubscript{2} complex. COX is represented in green, and AIFM1 protomers (residues 128 to 516; 551 to 613) with and without N-terminal region (residues 55 to 127) are represented in orange and yellow, respectively. The transmembrane residues (67 to 85) of the N terminus of the interacting AIFM1 moiety is highlighted in red. Membrane boundaries of the inner mitochondrial membrane are sketched as gray spheres. The final complex consists of monomeric COX, dimeric AIFM1 (residues 128 to 516, 551 to 613), and the N-terminal region of one AIFM1 protomer (residues 55 to 127).](https://doi.org/10.1073/pnas.2106950118)
good agreement with the final structural model of COX-AIFM12 (SI Appendix, Fig. S4 C and D). The combined mean distance for DSSO/PhoX cross-links used for docking was slightly lower (26.5 Å, 20 cross-links) than for DMTMM (28.2 Å, 26 cross-links) (Dataset S4). Most of the obtained over-length cross-links (>33 Å, 15 cross-links) involve the N-terminal domain of AIFM1 (12 out of 15 cross-links). Interactions to the N-terminal domain are predominately mapped with DMTMM (20 out of 24 detected cross-links), explaining the slightly higher mean distance. The over-length cross-links involving the AIFM1 N terminus were obtained for cross-links to a flexible segment of COX6C (six cross-links) and a defined residue stretch (221 to 244) of AIFM1 (six cross-links) located closely to the AIFM1 N terminus. These over-length cross-links predominantly involving specific domains could indicate that cross-links are derived for several assemblies rather than one assembly, which is a challenge of in-solution XL-MS described previously (11). In our case, cross-links observed between AIFM1 and the AIFM1 N terminus might well result also

![COX - AIFM1 interface](image)

**Fig. 4.** Deciphering interaction interfaces in the COX-AIFM1 structural model. (A) Three distinct interfaces between COX subunits and respective AIFM1 protomers were found. Subunits (COX) and protein domains (AIFM1) with residues in respective interfaces are colored in gray. Active COX residues are shown as green-colored sticks and active AIFM1 residues as orange-colored sticks. (B) Analysis of the number of residue contacts between respective COX subunits and AIFM1 domains. Colored circles indicate residue contacts between single subunits (COX) and domains (AIFM1) with the size of each circle corresponding to the number of residue-residue interactions. (C) COX-AIFM1 complex with CytC (purple) bound to its COX-binding site. The structural model presented here was merged with a previously published model of CytC docked to COX from bovine heart (54). COX subunits are colored green, while AIFM1 protomers are colored orange and yellow. The transmembrane (TM) domain of the N-terminal domain of AIFM1 is highlighted in red. Boundaries of the inner mitochondrial membrane are indicated as gray spheres.
from monomeric or dimeric AIFM1 assemblies (Fig. 2C), potentially featuring different domain orientations as compared to AIFM1 in complex with COX. Such averaging likely explains the observed over-length cross-links. Notwithstanding these considerations, the COX-AIFM1 model is in good agreement with the cross-linking data.

Next, we performed an interaction interface analysis of the docking model that predicted three distinct interfaces between COX and AIFM1 (Fig. 4A and Dataset S5). The first, extensive interface is defined by the N-terminal residues of AIFM1, which interact with neighboring residues of the COX subunits MT-CO2, NDUF4A, and COX6B1. Secondly, residues of the pyridine nucleotide-disulfide oxidoreductase domain of AIFM1 comprising the NADH- and FAD-binding domains intimately interact with MT-CO2, COX6B1, and COX6C. The third, rather small interaction interface is defined by residues of the C-terminal region of the second AIFM1 protomer and residues of MT-CO2, COX4I1, and COX7B1 (Fig. 4B). The interface between the hydrophilic parts of the N-terminal region of AIFM1 facing the intermembrane space is mostly driven by contacts to COX6B1 and NDUF4A, whereas its transmembrane domain predominantly interacts with one of the transmembrane segments of MT-CO2. Notably, the very N-terminal residues of AIFM1 facing the matrix side reside within a ∼25 Å distance from residues 44 to 54 of COX5A, consistent with the observed cross-links to this COX subunit (Fig. 2B).

Potential Functional Implications of a COX-AIFM1 Complex. While the predominant consequence of AIFM1 deficiency is impaired complex I assembly (4), additional COX deficiency has been reported in skeletal muscle and heart (9, 10) as well as in Drosophila melanogaster (48) and Caenorhabditis elegans (49). Conversely, AIFM1 expression was found to be significantly increased along with several COX assembly factors in human COX-negative muscle fibers (50). However, it seems unlikely that the COX-AIFM12 complex described here contributes to the assembly or stabilization of COX because it accounted only for 10% or less of the total amount of this OXPHOS complex. Moreover, no apparent commutation between AIFM1 and any of the individual COX subunits or subassemblies at apparent masses lower than ∼350 kDa was observed, suggesting that the association of AIFM1 occurred only with fully assembled COX. Conditional involvement of AIFM1 in the maturation of COX assembly factors that are substrates of the disulfide relay of the intermembrane space (6–8) appears as a more likely explanation for the link between AIFM1 and COX deficiency in some tissues.

It has been reported that AIFM1 is a member of the NDH-2 family of proteins (51) and thus exhibits NADH:ubiquinone oxidoreductase activity (52). For this reason, we examined the possibility of direct electron transfer between the FAD and CuA of COX within the COX-AIFM12 complex. The minimal distance between the isoalloxazine moieties of the FADs and the CuA center was ∼50 and ∼55 Å (SI Appendix, Fig. S5), which is more than three times larger than the 14 Å considered as the maximum distance for efficient electron tunneling in a protein matrix (53). It would be conceivable that this distance is bridged by cytochrome c (CytC) serving as an electron shuttle between AIFM1 and COX. Therefore, we explored whether CytC could still bind to its substrate-binding site in the COX-AIFM1 complex. Merging our structural model with a previously obtained model of CytC bound to COX from bovine heart (54) suggested that the AIFM1 dimer does not hamper CytC from binding to COX (Fig. 4C). However, distances of ∼45 and ∼47 Å between the heme moiety of CytC and the isoalloxazine rings of FAD in both AIFM1 protomers excluded direct electron transfer also in the presence of the additional heme. Yet, a substrate channeling mechanism could still be imaginable, implying movement of CytC forth and back between AIFM1 and COX without leaving a crevice. A crevice in the second AIFM1 protomer facing COX could potentially reduce the distance between the redox centers indeed to about 14 Å. However, this would require CytC to turn within the pocket formed by AIFM1 and COX in order to bring its heme as close as possible to the isoalloxazine ring. Thus, while such a substrate channeling mechanism cannot be excluded, it does not seem very likely. Moreover, oxidizing NADH would transiently destabilize dimerization of AIFM1 (6) and thus the entire complex, arguing further against any oxidoreductase activity of the COX-AIFM12 complex.

Since our structural model excludes electron transfer from AIFM1 to COX, it seems unlikely that the complex between them serves to drain electrons from the disulfide relay of the intermembrane space by regenerating CHCHD4/MAIA40 (6). Therefore, it remains to be established whether there is any functional link between the COX-AIFM12 complex described here and the import machinery for proteins of the mitochondrial intermembrane space containing disulfide bonds.

If COX-AIFM12 is not a catalytic complex, it is still tempting to speculate that a ternary interaction of COX, AIFM12, and CytC could play a role in mitochondrial proapoptotic mechanisms. Apart from directly promoting programmed cell death (5, 55, 56), AIFM1 could play an indirect role in apoptosis by modulating the release of CytC (1) through its binding to the COX-AIFM12 complex. For this, it is important to note that CytC makes direct contact to the first AIFM1 protomer in the ternary complex (Fig. 4C). It is important to note that the N-terminal propeptide with its transmembrane helix provides a significant portion of the AIFM1/COX interface, the complex is expected to destabilize upon cleavage of AIFM1, thereby activating its proapoptotic function. In addition to cleaved AIFM1, any previously bound CytC would be released, concomitantly further promoting apoptosis, potentially providing a synergistic boost to the cell death program already underway.

Conclusions

We show that ∼10% of monomeric COX in BHM are engaged in a defined complex with dimeric AIFM1. Using structural restraints provided by cross-linking data, available high-resolution structures, and structural modeling, we could derive a model of the COX-AIFM12 complex with and without bound CytC. Combining chemical cross-linking and complexome profiling provided useful complementary information and represents proof of concept for our experimental approach, demonstrating that it can be used to define and characterize multiprotein assemblies in detail that may have been overlooked by other means.

While our structural model excludes direct electron transfer between AIFM1 and COX, it provides clues on potential functional implications of the formation of the COX-AIFM12 complex including a possible involvement in promoting apoptosis. The structural insights into this unexpected mitochondrial complex will
stimulate and guide further studies on the role of AIFM1 in OXPHOS biogenesis and apoptosis.

Materials and Methods

Isolation and Purification of BHm. Mitochondrial membranes from bovine heart were isolated and preserved as described in ref. 11. In order to increase the purity of the preparation and for Tris buffer removal, frozen crude mitochondria (4 × 15 mL aliquots; 60 mg protein/mL) were thawed on ice, diluted (1:4) with ice-cold SEH buffer (250 mM sucrose, 1 mM ethylenediamine tetraacetic acid [EDTA], 20 mM Heps, pH 7.4 adjusted with NaOH) and centrifuged at 1,000 × g (10 min; 4 °C). The supernatants were recovered and centrifuged at 40,000 × g (20 min; 4 °C), and each resulting pellet was suspended in 5% DMSO. Afterward, mitochondria were loaded onto a 50-cm two-layer sucrose gradient (1 M sucrose, 20 mM Heps, pH 7.4 /1.5 M sucrose, 20 mM Heps, pH 7.4) and centrifuged at 60,000 × g (20 min; 4 °C). The pure mitochondrial fractions accumulated at the interphase were carefully recovered and pooled into one tube. After resuspension in 20 mL ice-cold SEH buffer, pure mitochondria were centrifuged at 10,000 × g (20 min; 4 °C) and finally suspended in 5 mL ice-cold SEH buffer supplemented with protease inhibitor mixture (GIMAFAST). Protein concentration was determined by the DC protein assay (Bio-Rad), and aliquots of pure mitochondria were shock frozen in liquid nitrogen and stored at −80 °C until use.

Cross-linking of BHm Sample with DSSO, PhoX, and DMTMM. Purified BHm membranes were buffer exchanged into cross-linking buffer (10 mM Heps pH 7.8, 1 mM EDTA, 1 mM ethylene glycol tetraacetic acid [EGTA], 10 mM NaCl, 150 mM KCl, protease inhibitor). After optimization of the cross-link reaction, −2 mg of BHm were either incubated with DSSO (0.5 mM freshly resuspended in anhydrous dimethyl sulfoxide [DMSO]; Thermo Fisher Scientific), PhoX (1 mM freshly resuspended in anhydrous DMSO; made in-house), or DMTMM (10 mM freshly resuspended in cross-linking buffer; Sigma-Aldrich) in 2 mL cross-linking buffer at room temperature (RT). The cross-link reaction was quenched after 30 min by the addition of 50 mM Tris (1 M Tris buffer, pH 8.5) for an additional 30 min at RT.

Sample Preparation for X-MS Analysis of Cross-linked BHm. Cross-linked mitochondria were solubilized with digitonin (9 g/g protein) for 30 to 60 min on ice. Proteins were denatured and purified as described previously (61). Briefly, denatured proteins were resuspended and digested overnight at 37 °C with Lys-C followed by trypsin. The final peptide mixtures were desalted with solid-phase extraction C18 columns (Sep-Pak, Waters). Samples cross-linked with DSSO and DMTMM were fractionated with an Agilent 1200 HPLC pump system (Agilent) coupled online with an Agilent 1200 Infinity UHPLC system (Agilent) on a 50-cm analytical column packed with C18 beads (Dr. Maish Reprosl C18, 3 μm) coupled online to an Orbitrap Fusion Lumos (Thermo Fisher Scientific). We used the following liquid chromatography–tandem mass spectrometry (LC-MS/MS) parameters: after 5 min of loading with 100% buffer A (water with 0.1% formic acid), peptides were eluted at 300 nL/min with a 97 min gradient from 4 to 39% of buffer B (80% acetonitrile and 20% water with 0.1% formic acid). For MS acquisition, we used an MS1 Orbitrap scan at 120,000 resolution from 310 to 1,600, an automatic gain control (AGC) target of 2×10⁶ ions, and a maximum injection time of 50 ms. The ions with a charge from +3 to +8 were fragmented with collision-induced dissociation (CID) (normal-collision energy [NCE] of 30%) and analyzed with an MS2 Orbitrap at 30,000 resolution, an AGC target of 2×10⁶ ions, and a maximum injection time of 54 ms for the detection of DSSO signature peaks (difference in mass of 4,200) with a collision energy of 80%. The resulting MS2 spectra were searched against human protein database using X!Tandem (http://x!tandem.hudsonalpha.org) with the following parameters: static carbamidomethyl and dynamic oxidation. The cross-linked peptides were searched with an X!Tandem node for analysis of cross-linked peptides as reported by Kljukov et al. (64). Data were searched against a FASTA file containing the −4,200 most abundant proteins, which were previously determined following a classical bottom-up workflow. Where applicable, mitochondrial target peptides were removed from respective protein sequences. For an XlinkX search, we selected fully tryptic digestion with three maximum missed cleavages, 10 ppm error for MS1, 20 ppm for MS2, and 0.5 Da for MS3 in Ion Trap. For modifications, we used static carbamidomethyl and dynamic oxidation. The cross-linked peptides were accepted with a minimum score of 40, minimum score difference of 4, and maximum false discovery rate (FDR) (controlled at peptide-spectrum match [PSM] level for cross-linked spectrum matches) rate set to 5%. Both noncleavable cross-linkers were analyzed with pLink2 (65) and the same FASTA used for DSSO. For PhoX, we manually added the cross-linker to the list (alphabetical sites “X”, “K”, linker composition (C)H2N(Ο)S(Ρ)I (mass of 209.971 Da), and for both cross-linkers, the same parameter settings as described for XlinkX was used for all gel pieces excepting: no minimum score option, and the FDR was calculated separately for intra- and intercross-links. Finally, cross-links were additionally filtered: only cross-links corresponding to protein–protein interactions that were reported for at least two cross-linkers and with at least two CSMs were kept for the final interaction analysis and structural modeling.

Complexome Profiling Analysis. Aliquots of untreated and PhoX and DMTMM cross-linked mitochondrial membranes (see Cross-linking of BHm Sample with DSSO, PhoX and DMTMM for details) were thawed on ice, solubilized with digitonin (9 g/g protein) in 50 mM NaCl, 50 mM imidazole-HCl, 2 mM 6-aminohexanoic acid, 1 mM EDTA, pH 7, and kept on ice for 20 min. Samples were further centrifuged at 22,000 × g (20 min; 4 °C), and the supernatants were transferred into clean tubes and supplemented with Coomassie blue (final concentration of 0.1% v/v), 100 μg protein/mL. In order to increase the sensitivity of the subsequent gel electrophoresis run, the gel was fixed overnight in 50% methanol, 10% acetic acid, and 100 mM ammonium acetate following staining with 0.025% Coomassie blue G-250 (Serva G) in 10% acetic acid for 30 min, de-stained twice in 10% acetic acid (1 h each), and kept in deionized water overnight. The next day, the gel was color scanned using a flatbed Image Scanner III (GE) to use it as a template for the cutting procedure.

Proteins were identified by LC-MS/MS after in-gel tryptic digestion following the protocol described in Heide, et al. (15) with some modifications. In short, each gel lane was cut into 60 even slices starting at the bottom of the gel. The slices were cubed and transferred into 96-well filter plates (Millipore, MABVNN1250) adapted manually to 96-well plates (MaxiSorp Nunc) as waste collection. Gel pieces were equilibrated with 50% methanol and 50 mM amonium hydrogen carbonate (AHC) under moderate shaking; the solution was refreshed until the blue dye was removed completely. The removal of excess solution was done by centrifugation (1,000 × g, 15 s). In the next step, gel pieces were reduced with 10 mM dithiothreitol in 50 mM AHC for 1 h. After removing excess solution, 30 mM chloroacetic acid in 50 mM AHC was added to each well, incubated in the dark for 45 min, and removed. A short incubation step with 50% methanol and 50 mM AHC was performed for gel pieces dehydration (∼15 min). The latter solution was removed, and gel pieces were dried for ∼30 min at RT. Later, 20 μL 5 mg · μL⁻¹ trypsin (sequencing grade, Promega) in 50 mM AHC plus 1 mM CaCl₂ were added to each well and incubated for 20 min at 4 °C. Gel pieces were covered by adding 50 μL 50 mM AHC followed by an overnight incubation at 37 °C for protein digestion. The next day, the peptide-containing supernatants were collected by centrifugation (1,000 × g, 30 s) into clean 96-well PCR plates (Axygen). The gel pieces were finally incubated with 50 μL 30% acetonitrile (ACN) and 3% formic acid (FA) for ∼30 min prior to elution of the remaining peptides on the previous eluates by centrifugation. The peptides were dried in a SpeedVac Concentrator Plus (Eppendorf) for 2.5 to 3 h, resuspended in 20 μL 5% ACN and 0.5% FA, and stored at −20 °C until MS analysis.

After thawing the frozen resuspended peptides and a 30 min gentle sonication of each individual sample fraction, the dried and separated by reverse-phase LC and analyzed by MS/MS in a Q-Exactive Orbitrap Mass Spectrometer equipped with a nano-flow-ultra-HPLC system (Easy nLC1000, Thermo Fisher Scientific). In brief, peptides were separated using 100 μm inner diameter
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1. M. Saraste, Oxidative phosphorylation at the fin de siècle. Science 283, 1488–1493 (1999).
2. S. A. Susin et al., Molecular characterization of mitochondrial apoptosis-inducing factor. Nature 397, 441–446 (1999).
3. C. Rinaldi et al., Cofactor chondrodiac syndrome is associated with a mutation in apoptosis-inducing factor. Am. J. Hum. Genet. 91, 1095–1102 (2012).
4. N. Vathsen et al., AIF deficiency compromises oxidative phosphorylation. EMBO J. 23, 4679–4689 (2004).
5. D. Ghezzi et al., Severe X-linked mitochondrial encephalomyopathy associated with a mutation in apoptosis-inducing factor. Am. J. Hum. Genet. 86, 639–649 (2010).
6. C. Petrungho et al., The Ca(2+)-dependent release of the Mia40-induced MUC1-MUC2 dimer from MUC regulates mitochondrial Ca(2+) uptake. Cell Metab. 22, 721–733 (2015).
7. K. Meyer et al., Loss of apoptosis-inducing factor critically affects MIA40 function. Cell Death Dis. 6, e1814 (2015).
8. E. Hanget et al., Interaction between AIF and CHCHD4 regulates respiratory chain branching. Mol. Cell 60, 1001–1014 (2015).
9. J. A. Pospisilik et al., Targeted deletion of AIF decreases mitochondrial oxidative phosphorylation and protects from obesity and diabetes. Cell 131, 476–491 (2007).
10. N. Joza et al., Muscle-specific loss of apoptosis-inducing factor leads to mitochondrial dysfunction, skeletal muscle atrophy, and dilated cardiomyopathy. Mol. Cell. Biol. 25, 10261–10272 (2005).
11. J. F. Heveler et al., Selective cross-linking of coining protein assemblies by in-gel cross-linking mass spectrometry. EMBO J. 40, e106174 (2021).
12. A. Kao et al., Development of a novel cross-linking strategy for fast and accurate identification of cross-linked peptides of protein complexes. Mol. Cell Proteomics 10, M110 002212 (2011).
13. B. Steigenberger, R. J. Pieters, A. J. R. Heck, R. A. Scheltema, X. PhoX: An IMAC-enrichable cross-linking reagent. ACS Cent. Sci. 5, 1514–1522 (2019).
14. A. Leitner et al., Chemical cross-linking/mass spectrometry targeting acidic residues in mitochondria by cross-linking mass spectrometry provides evidence for coexisting respiratory supercomplexes. Mol. Cell Proteomics 17, 216–232 (2018).
15. D. K. Schwepe et al., Mitochondrial protein interaction elucidated by chemical cross-linking mass spectrometry. Proc. Natl. Acad. Sci. U.S.A. 111, 9455–9460 (2014).
16. N. Heide et al., Complexome profiling identifies TIM126B as a component of the mitochondrial complex I assembly complex. Cell Metab. 16, 538–549 (2012).
17. F. Liu, P. Lassé, B. M. Robbins, R. S. Balaan, A. J. R. Heck, The interaction of intact mitochondria by cross-linking mass spectrometry for coexisting respiratory supercomplexes. Mol. Cell Proteomics 17, 216–232 (2018).
18. A. Linden et al., A cross-linking mass spectrometry approach defines protein interactions in yeast mitochondria. Mol. Cell Proteomics 19, 1161–1178 (2020).
19. P. J. S. Ryl et al., In situ structural restraints from cross-linking mass spectrometry in human mitochondria. J. Proteome Res. 19, 327–336 (2020).
20. J. D. Chavez et al., Mitochondrial protein interaction landscape of 55-31. Proc. Natl. Acad. Sci. U.S.A. 117, 13583–13537 (2020).
21. S. Rath et al., Mitocarta AIIF1 and mitochondrial proteome now with suborganellar localization and pathway annotations. Nucleic Acids Res. 49, D1541–D1547 (2021).
22. M. A. Gonzalez-Lozano et al., Stiching the synapse: Cross-linking mass spectrometry into resolving synaptic protein interactions. Sci. Adv. 6, eaax783 (2020).
23. J. Fürsch, K. M. Kammer, S. G. Kreft, M. Beck, F. Stengel, Proteome-wide structural probing of low-abundant protein interactions by cross-linking mass spectrometry. Anal. Chem. 92, 4016–4022 (2020).
24. H. Schägger, K. Pfeffer, Supercomplexes in the respiratory chains of yeast and mammalian mitochondria. EMBO J. 19, 1777–1783 (2000).

25. T. B. Blum, A. Hahn, T. Meier, K. M. Davies, W. Kühlbrandt, Dimers of mitochondrial ATP synthase induce membrane curvature and self-assemble into rows. Proc. Natl. Acad. Sci. U.S.A. 116, 4250–4255 (2019).

26. D. E. Kim, D. Chivian, D. Baker, Protein structure prediction and analysis using the deep neural network model. Bioinformatics Acta Mol. Basis Dis. 1864, 3650–3658 (2018).

27. T. E. Spikes, M. G. Montgomery, J. E. Walker, Structure of the dimeric ATP synthase from bovine mitochondria. Proc. Natl. Acad. Sci. U.S.A. 117, 23519–23526 (2020).

28. A. M. Bonvin, S. Arnold, The power of life

29. J. Gu

30. H. Antonicka

31. G. C. van Zundert, A. M. Bonvin, DisVis: Quantifying and visualizing accessible inter-

32. C. D. Go

33. I. F. Sevrioukova, Apoptosis-inducing factor: Structure, function, and redox regulation. Antioxid. Redox Signal. 14, 2545–2579 (2011).

34. S. Guerrero-Castillo

35. J. Hirst, J. Carroll, I. M. Fearnley, R. J. Shannon, J. E. Walker, The nuclear encoded inhibitory protein IF1. EMBO J. 150 (2003).

36. A. Abdrakhmanova, K. Dobrynin, K. Zwicker, S. Kerscher, U. Brandt, Functional sul-

37. E. Balsa

38. T. Tsukihara

39. J. Yang

40. P. Ferreia et al., Structural insights into the coenzyme mediated monomer-dimer transition of the pro-apoptotic apoptosis inducing factor. Biochemistry 53, 4204–4215 (2014).

41. D. E. Kim, D. Chivian, D. Baker, Protein structure prediction and analysis using the Robetta server. Nucleic Acids Res. 32, W526–W531 (2004).

42. T. Suzuki et al., The low-spin heme of cytochrome c oxidase as the driving element of the proton-pumping process. Proc. Natl. Acad. Sci. U.S.A. 100, 15304–15309 (2003).

43. S. Zong et al., Structure of the intact 14-subunit human cytochrome c oxidase. Cell Rep. 28, 1026–1034 (2018).

44. G. C. van Zundert, A. M. Bonvin, DisVis: Quantifying and visualizing accessible inter-

45. G. C. van Zundert et al., The HADDOCK2.2 web server: User-friendly integrative modeling of biomolecular complexes. J. Mol. Biol. 428, 720–725 (2016).

46. Z. Yang et al., UCSF Chimera, MODELLER, and IMP: An integrated modeling system. J. Struct. Biol. 179, 269–278 (2012).

47. E. F. Petterson et al., UCSF Chimera-A visualization system for exploratory research and analysis. J. Comput. Chem. 25, 1605–1612 (2004).

48. N. Joza et al., The molecular archaeology of a mitochondrial death effector: AIF in Drosophila. Cell Death Differ. 15, 1009–1018 (2008).

49. K. Troulis et al., WAH–TIAF regulates mitochondrial oxidative phosphorylation in the nematode Caenorhabditis elegans. Cell Death Discov. 4, 2 (2018).

50. M. Murgia et al., Proteomics of cytochrome c oxidase-negative versus -positive muscle fiber sections in mitochondrial myopathy. Cell Rep. 29, 3825–3834.e4 (2019).

51. S. J. Kerscher, Diversity and origin of alternative NADH-ubiquinone oxidoreductases. Biochim. Biophys. Acta 1459, 274–283 (2000).

52. M. M. Elguindy, E. Nakamura-Ogiso, Apoptosis-inducing factor (AIF) and its family member protein, AMID, are rotenone-sensitive NADH:ubiquinone oxidoreductases (NDH-2). J. Biol. Chem. 290, 20815–20826 (2015).

53. C. C. Moser, J. L. Anderson, P. L. Dutton, Guidelines for tunneling in enzymes. Bio-

54. H. Schägger, K. Pfeffer, Supercomplexes in the respiratory chains of yeast and mammalian mitochondria. EMBO J. 19, 1777–1783 (2000).