

Mini Review

Cytochrome *c*: Surfing Off of the Mitochondrial Membrane on the Tops of Complexes III and IV

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ABSTRACT

The proper arrangement of protein components within the respiratory electron transport chain is nowadays a matter of intense debate, since altering it leads to cell aging and other related pathologies. Here, we discuss three current views—the so-called *solid*, *fluid* and *plasticity* models—which describe the organization of the main membrane-embedded mitochondrial protein complexes and the key elements that regulate and/or facilitate supercomplex assembly. The soluble electron carrier cytochrome *c* has recently emerged as an essential factor in the assembly and function of respiratory supercomplexes. In fact, a 'restricted diffusion pathway' mechanism for electron transfer between complexes III and IV has been proposed based on the secondary, *distal* binding sites for cytochrome *c* at its two membrane partners recently discovered. This channeling pathway facilitates the surfing of cytochrome *c* on both respiratory complexes, thereby tuning the efficiency of oxidative phosphorylation and diminishing the production of reactive oxygen species. The well-documented post-translational modifications of cytochrome *c* could further contribute to the rapid adjustment of electron flow in response to changing cellular conditions.

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1. Introduction

Mitochondria are the powerhouses of the cell, generating the greatest amount of chemical energy in the form of ATP during cellular respiration. During respiration electrons from reduced nicotinamide adenine dinucleotide (NADH) and reduced flavin adenine dinucleotide

(FADH₂) are transferred via the electron transport chain (ETC) to molecular oxygen (O₂). Electron flow through the ETC is coupled with the generation of an electrochemical proton gradient across the inner mitochondrial membrane that in turn yields ATP, a process known as oxidative phosphorylation (OxPhos) [1].

The ETC has four transmembrane protein complexes: complex I (CI; NADH-ubiquinone oxidoreductase), complex II (CII; succinate-ubiquinone reductase), complex III (CIII; cytochrome *bc*₁ complex) and complex IV (CIV; cytochrome *c* oxidase). In addition to the transmembrane complexes, two mobile carriers participate in the OxPhos

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process: lipophilic ubiquinone (Q) and hydrophilic cytochrome *c* (Cc), which transfer electrons from CI/CII to CIII and from CIII to CIV, respectively. The electrochemical proton gradient generated by the ETC complexes (except CII which does not pump H^+) drives the synthesis of ATP through complex V (CV; ATP synthase). Beyond the ETC components abovementioned, several mitochondrial enzymes—namely, electron-transferring flavoprotein Q oxidoreductase, dihydroorotate dehydrogenase or glycerol-3-phosphate dehydrogenase—may act as auxiliary elements of the respiratory chain [2].

The organization and dynamics of protein components within the ETC is currently a matter of debate. Three different models have been proposed to explain their organization (Fig. 1). On one hand, the *fluid* model states that the membrane components diffuse freely and independently [3] (Fig. 1A, right panel). While on the other hand, the *solid* model proposes that respiratory complexes are stably associated forming supercomplexes, thus facilitating the flow of electrons [4,5] (Fig. 1A, left panel). However, the idea that these two scenarios are not exclusive gave rise to the *plasticity* model; this model consists of a balanced distribution between free respiratory complexes and supercomplexes, adapting the efficiency of the ETC by means of partitioned Q and Cc pools [6] (Fig. 1B).

Supercomplex formation occurs in mammals, plants, yeast and some bacteria [7–9]. Among these supercomplexes, one stands out; the respirasome, which contains CI, CIII and CIV and is capable of NADH: O_2 oxidoreduction *in vitro* [10]. The degree of association and the

stoichiometries of respiratory complexes in supercomplexes depend on cell type, tissue-specific isoforms and cellular conditions [11–14]. The mitochondrial intermembrane lipid content also affects supercomplex assembly [15–18].

The diverse composition and stoichiometry of supercomplexes has been evidenced by blue native polyacrylamide gel electrophoresis (BN-PAGE), cryo-electron tomography and single-particle electron microscopy studies of mitochondrial fractions solubilized with mild detergents. Fig. 1B represents the different supercomplexes described in the literature. In brief, CI mainly interacts with CIII and CIV to form the respirasome (CI + CIII₂ + CIV_{1–4}) [10,19–26]. Alternatively, CIII binds to CI (CI + CIII₂) or CIV as a monomer (CIII₂ + CIV_{1–2}) [7,22,23,25,27,28]. Generally speaking, CV assembles oligomers and CII acts as a stand-alone complex. However, their associations with complexes I, III, and IV have also been described [25,27].

2. Key Elements for Respiratory Supercomplex Assembly

The mechanisms which regulate the assembly of mitochondrial supercomplexes remain unsolved. However, several biomolecules have been recently described to regulate/facilitate the formation of certain supercomplexes.

As mentioned above, phospholipids mediate protein–protein interactions within the inner mitochondrial membrane. Cardiolipin—an anionic phospholipid found primarily in the inner mitochondrial

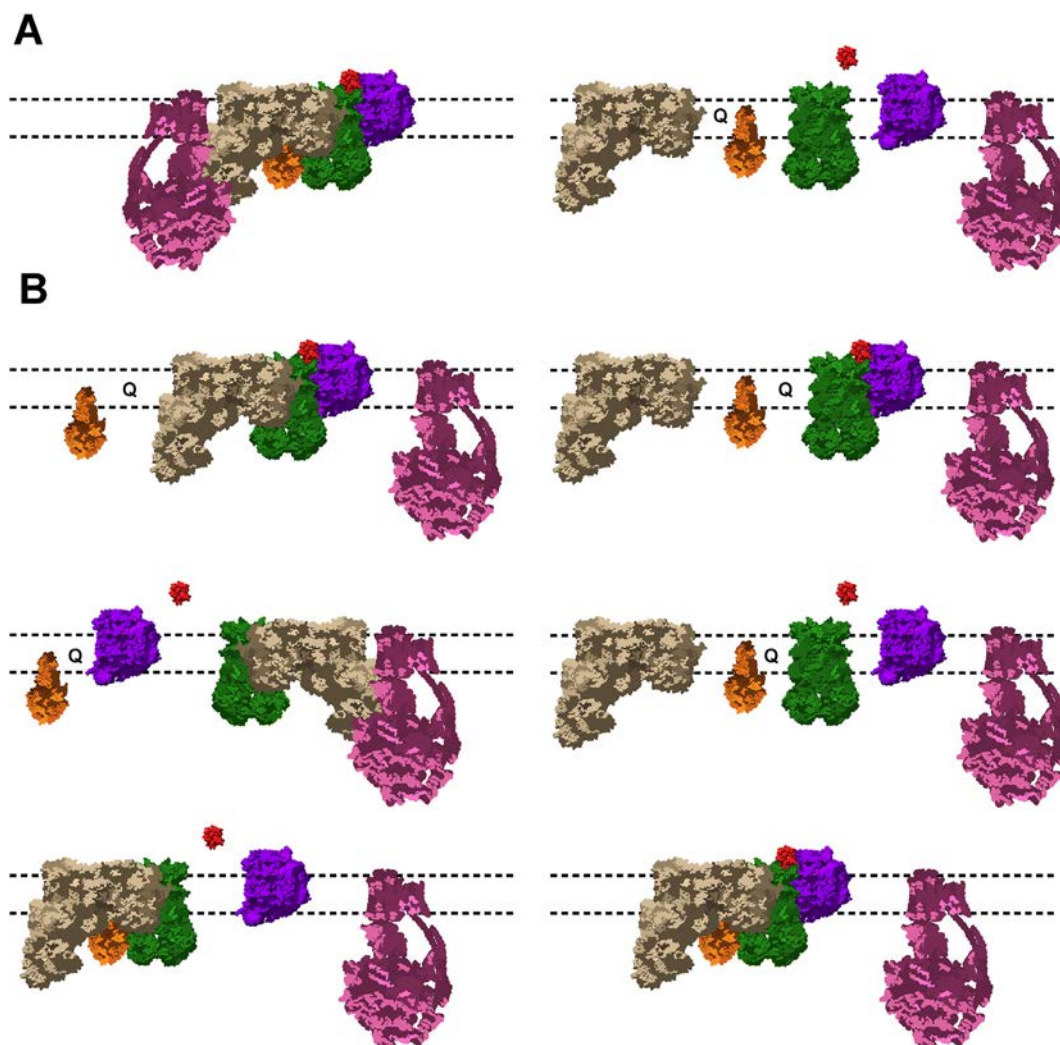


Fig. 1. Schematic representation of the organization of respiratory complexes. A) *Solid* (left) and *fluid* (right) models. B) *Plasticity* model. All supercomplexes described in the literature are displayed. The stoichiometry of each respiratory complex is not represented. CI is in tan, CII in orange, CIII in green, CIV in purple, CV in pink, Cc in red and ubiquinone as Q.

membrane—represents 20% of all membrane lipids [29]. Lack of cardiolipin has been linked with metabolic disorders, such as Barth's Syndrome, and affects the stability of supercomplexes and the activity of CIV [30]. Phosphatidylethanolamine is a cationic phospholipid present in all cellular membranes which when depleted favours the formation of large supercomplexes [17]. Therefore, the distribution of phospholipids may play an important role in the reorganization of respiratory complexes into supercomplexes.

Several proteins have been postulated to function as chaperones or assembly factors for supercomplexes. For instance, the depletion of 28 different NADH:ubiquinone oxidoreductase supernumerary subunits (NDUF) out of 30, which are necessary for CI assembly and stability, has also been described to reduce supercomplexes containing CI + CIII₂ and CI + CIII₂ + CIV₁ in human cells [31]. Although the transmembrane NDUF4A subunit has initially been defined as a component of CI, new reports suggest that it belongs to CIV [32,33]. Interestingly enough, NDUF4A has recently been postulated as a factor required for assembly of supercomplexes in growing cells and cancer tissues [34]. The MJC/DnaJC15 protein, a mitochondrial repressor of the ETC that interacts with CI, has been described as a regulator of supercomplex formation [35].

Recently, the CIV subunit COX7A2L (also known as SCAFI) has been proposed as a CIII binding protein which stabilizes the CIII₂ + CIV₁ supercomplex, independently of respirasome formation [36,37]. In yeast, two respiratory complex factors (Rcf1 and Rcf2), members of the conserved hypoxia induced gene 1 (Hig1) protein family, regulate the formation and stabilization of the CIII₂ + CIV₁₋₂ supercomplex [38–40]. Rcf1 has two eukaryotic orthologues, namely HIG1 hypoxia inducible domain family 1A and 2A (HIGD1A and HIGD2A, respectively). Indeed, HIGD2A knockdown causes depletion of the CIII₂ + CIV₁ supercomplex [41]. Further, the ADP/ATP carrier protein has been reported to exist in association with CIII₂ + CIV₂ and mediates such supercomplex assembly in yeast [42]. Prohibitin has been hypothesized to act as a chaperone in aiding the assembly of respiratory complexes—in particular CI [43]. Indeed, recent data indicates that the CIII₂ + CIV supercomplex content decreases in the absence of prohibitin [44].

Finally, the ETC mobile carriers Q and Cc co-migrate with supercomplexes in BN-PAGE analyses [27]. Strikingly, the absence of Cc destabilizes CI and CIV [45].

3. Cytochrome c: An Efficient Electron Carrier between CIII and CIV

Cc is an essential metalloprotein for mitochondrial metabolism and homeostasis, playing a dual-purpose role in cell life and death [46–48]. Under homeostasis, Cc acts as electron carrier within the respiratory chain in the mitochondria. Voltammetry assays and current crystal structures of CIII and CIV complexed with Cc resolve only one Cc molecule per monomer [49–51]. However, earlier reports had suggested the existence of additional binding sites in both CIII and CIV [52–56]. Multiphasic kinetics observed in early assays for the oxidation of reduced Cc by CIV can be fitted to a model with just one catalytic site. Such a simple model includes alternative binding conformations of the transient complex, with some of them being unable to transfer electrons but affecting the ET rate at the catalytic site [53,57]. Direct Cc-CcO binding studies performed by gel filtration evidenced a 2:1 stoichiometry, in which Cc bound at a first site with a low dissociation constant and at a second site with less affinity [49]. Non-ET conformations within the complex between CIII and Cc were also evidenced by steady state kinetic analysis, and so a binding model with more than one molecule of Cc per molecule of CIII was proposed to explain the observed multiphasic kinetics [58]. More recently, *in solution* nuclear magnetic resonance (NMR) and isothermal titration calorimetry (ITC) experiments have confirmed the presence of a secondary binding site on cytochrome c₁ (Cc₁) for Cc, with an affinity 10-fold lower than that of the principal binding site [59,60]. The two Cc-binding sites of Cc₁ in CIII were named *proximal* and *distal*, according to Brownian Dynamics (BD) calculations and NMR-driven docking computations (Fig. 2). In the *proximal* site, the heme groups of Cc and Cc₁ lay near each other, allowing electron transfer. The interaction surface on Cc involves a hydrophobic patch around the heme cleft surrounded by positively charged residues. Such binding surface configuration determines the correct orientation of Cc towards the heme group of Cc₁ and is common to other ET complexes [61,62]. However, the surface used by Cc to bind the *distal* site of Cc₁ is less-well defined when compared with the *proximal* one, in agreement with its lower affinity. Furthermore, the *distal* site lies far from the heme pocket of Cc₁, meaning there is low compatibility at the secondary site for efficient electron transfer.

Secondary binding sites may play multiple roles in ETCs, including photosynthetic ones. For instance, in eukaryotic Photosystem I [63], secondary binding sites assure fast replacement of electron donor

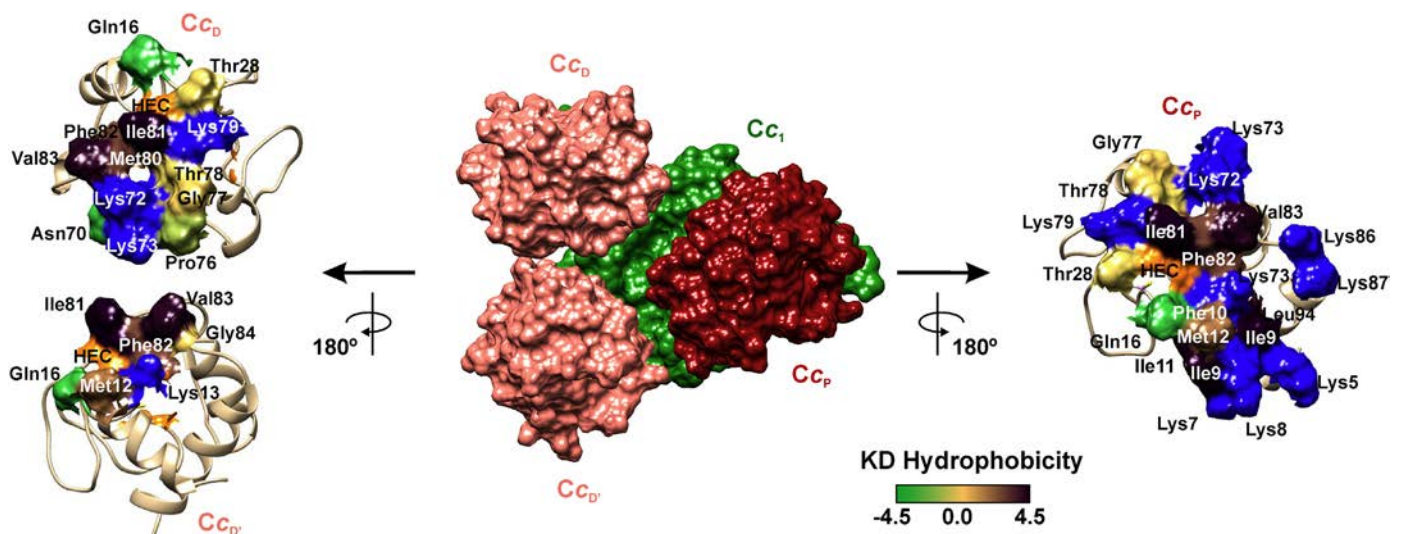


Fig. 2. Binding surfaces of cytochrome c on cytochrome c₁. Brownian Dynamics calculations and NMR-driven docking computations accommodate three Cc molecules in the soluble domain of cytochrome c₁. In the middle panel, two Cc molecules are located at the *distal* sites of cytochrome c₁ (Cc_D and Cc_{D'}), whereas one Cc molecule is at the *proximal* site of cytochrome c₁ (Cc_P). On *left* and *right* panels, Cc molecules are rotated 180° to expose interaction areas, with the residues colored according to the Kyte and Doolittle (KD) hydrophobicity scale; basic residues are in blue and the heme group (HEC) is in orange. Pictures were created with the UCFS Chimera software using the PDB structures for CIV (5XTH) and Cc (2N9I) [88].

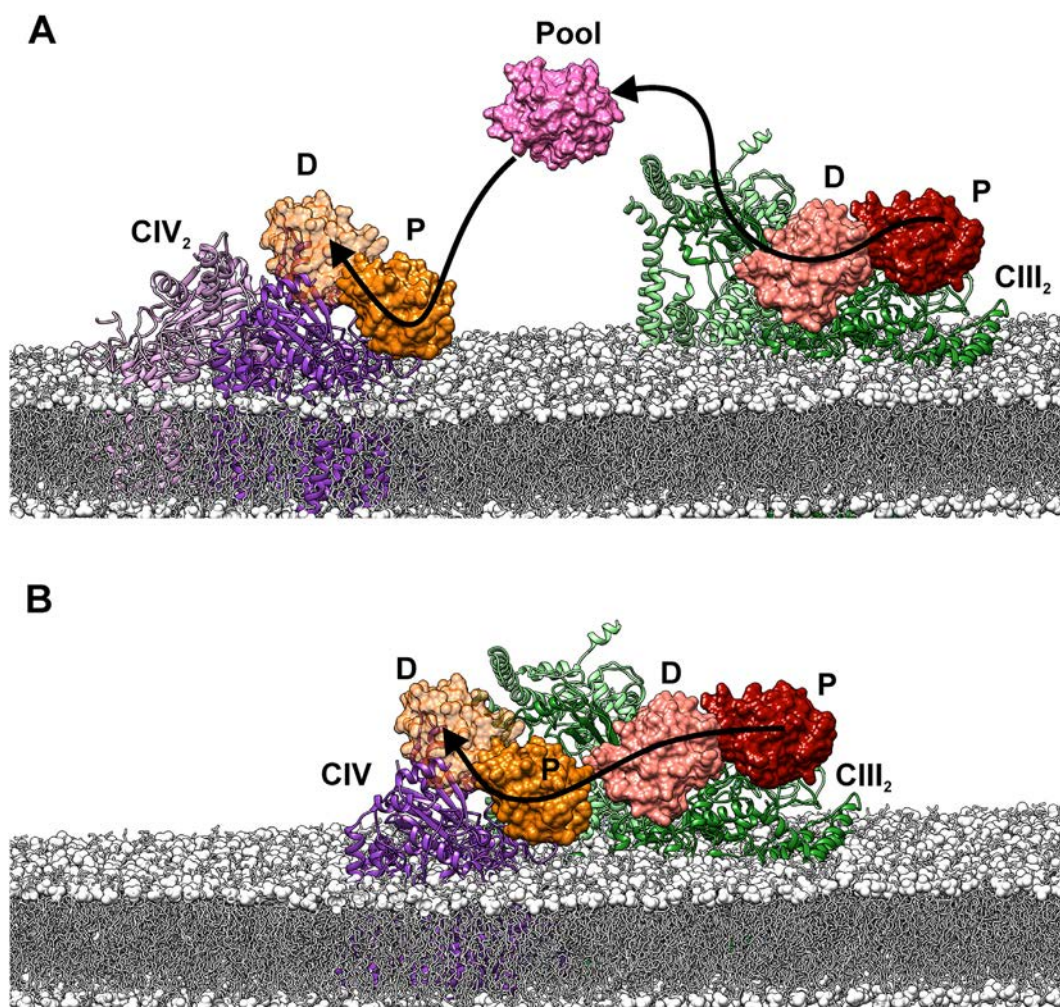


Fig. 3. Cytochrome *c* molecules surfing between respiratory complexes. A) Fluid model of dimeric CIII (green) and dimeric CIV (purple) embedded in the mitochondrial membrane. Isolated CIV is classically a dimer, as inferred from X-ray crystal structures [87]. B) Solid model of dimeric CIII and monomeric CIV, as shown in the recent Cryo-EM structure for human respiratory supercomplex complex [26]. P and D respectively stand for *distal* and *proximal* Cc binding sites on each complex. “Pool” denotes Cc population within the bulk intermembrane space. Pictures were created with the UCFS Chimera software using the PDB structures for CIII, CIV (5XTH) and cytochrome *c* (2N9I) [88].

molecules once an ET reaction takes place, and disabling them is lethal. In the context of supercomplexes, these additional *distal* sites at CIII and CIV could facilitate the channeling of electron carriers by building a ‘restricted diffusion pathway’ for Cc molecules across the surface of the two mitochondrial respiratory complexes instead of carrying electrons by random diffusion across the intermembrane mitochondrial space (Fig. 3). This fractal diffusion pathway substantially increases the efficiency of the OxPhos process in respect to the bulk 3D search, and decreases the production of reactive oxygen species (ROS) [64,65]. Indeed, restraining diffusion by weak interactions with membranes, cytoskeletal structures and DNA substantially lowers the time a molecule needs to reach its target [66]. Both effects—increasing turnover and restraining diffusion of mobile carriers between two membrane complexes—are clear advantages of secondary sites in ETCs.

4. Functional Implications of Post-Translational Modifications of Cytochrome *c*

The ETC is tightly regulated by post-translational modifications (PTMs), widening the function of its components under homeostatic and stress conditions [67,68]. Notably, the structure and activity of Cc are controlled *in vivo* by phosphorylation of threonine 28, serine 47 and tyrosine 48 and 97 [69–79].

Because of the low yield of most phosphorylated Cc species obtained from cell extracts and the specific Cc kinase being unknown, phosphomimetic Cc variants are the best alternative approach to analyze the effect of Cc phosphorylation [76,80,81]. Actually, an extensive study of the functional effect of phosphorylation is unfeasible with purified phosphorylated Cc species, but can be performed in an equivalent way with phosphomimetic mutants.

Traditionally, tyrosine phosphorylation has been mimicked by mutating tyrosine to glutamate/aspartate. However, it has recently been established that the substitution of tyrosine with the non-canonical amino acid *p*-carboxymethyl-L-phenylalanine (*p*CMF) better mirrors tyrosine phosphorylation. Notably, all mutants maintain the 2:1 stoichiometry (Cc:Cc₁ and Cc:CIV) found for the wild-type (WT) species. Nevertheless, whereas all tested phosphomimetic Cc species show similar affinities for the *proximal* site at Cc₁ (Table 1), binding at the *distal* site is substantially hindered by the additional negative charge of Cc [75,77]. The affinity of Cc towards bovine CIV changes depending on the residue of Cc that is phosphorylated (Table 1) [65,77].

As noted in the Introduction section, the formation of supercomplexes has been proposed to increase ET efficiency during OxPhos, thereby minimizing the generation of ROS. ROS are generally generated in complexes I and III, and can damage key components of cells [82,83]. The OxPhos process has been determined by the ability

Table 1
Effect of post-translational modifications on cytochrome *c* binding and activity assays.

| Cc species | Affinity towards CIII ^a | | Affinity towards CIV | | CcO activity | | |
|--------------|------------------------------------|-------------|----------------------|-------------|--------------------|--------------------|--------------------|
| | Proximal site | Distal site | Proximal site | Distal site | Fluid ^b | Fluid ^c | Solid ^d |
| T28D [75] | ~ | ↓ | n.d. | n.d. | ↑ | | n.d. |
| T28E [76] | n.d. | n.d. | n.d. | n.d. | | ↓↓ | |
| pT28 [76] | n.d. | n.d. | n.d. | n.d. | | ↓↓ | n.d. |
| S47D [75] | ~ | ~ | n.d. | n.d. | ↑ | | n.d. |
| Y48pCMF [77] | ~ | ↓ | ↓ | ↓↓↓ | ↑ | | ↓ |
| pY48 [78] | n.d. | n.d. | n.d. | n.d. | | ↓ | n.d. |
| Y97pCMF [65] | n.d. | n.d. | ↑↑↑ | ↑↑↑ | ↑ | | ↑↑ |
| pY97 [70] | n.d. | n.d. | n.d. | n.d. | | ↓↓ | n.d. |

n.d.: not determined.

'Fluid' and 'Solid' refer to the two proposed models in which ETC are organised in the inner mitochondrial membrane.

^a Affinity determined with the soluble N-terminal domain of *Arabidopsis thaliana* cytochrome *c*₁.

^b Activity measured as Cc oxidation by CIV isolated from bovine or equine heart.

^c Activity measured as O₂ consumption by CIV isolated from bovine tissues under conditions preserving the enzyme phosphorylation status.

^d Activity measured as Cc oxidation in isolated yeast mitochondria.

of Cc to shuttle electrons to CIV. When Cc performs a three-dimensional search visiting the bulk of the intermembrane mitochondrial space, every well-described phosphorylation in Cc enhances the cytochrome *c* oxidase (CcO) activity *in vitro*, in Cc oxidation assays using commercial CIV from bovine or equine heart (Table 1) [65,75,77]. Nevertheless, CIV isolated from bovine tissues under conditions preserving the phosphorylation status of CcO shows increased *K_M* values for phosphorylated Cc species and T28E mutant in O₂ consumption assays [70,76,78]. Thus, simultaneous phosphorylation of both heme-containing proteins, Cc and CcO, could act synergistically in inhibiting CcO activity. This apparent discrepancy could mainly be ascribed to PTMs of CcO.

When CcO activity is measured in mitochondria from cells grown in medium promoting the CIII-CIV supercomplex assembly (as part of the *solid* model), the Y48pCMF mutation decreases the electron donor ability of Cc towards CcO. In contrast, *in vitro* observations mimicking the *fluid* model enhance Cc oxidation (Table 1). In the context of the *fluid* model, the *proximal sites* for Cc on CIII and CIV govern ET reactions. However, in the *solid* model, the *distal sites*, which are non-ET functional sites, become important in limiting the diffusion of Cc molecules on the respiratory CIII-CIV supercomplex.

The balance between relative affinities towards the *proximal* and *distal* sites of CIII and CIV determines Cc exchange rates, alike in photosynthetic membrane complexes [63]. Indeed, the higher affinity of reduced Y48pCMF Cc towards the *proximal* site at CIII impairs its replacement by a new, oxidized Cc molecule. Further, the channeling of Cc molecules at the CIII-CIV ensemble would be disrupted by its weaker binding to the *distal* sites, altogether affecting steady-state, CcO-dependent oxidation rate of Cc (Table 1) [77]. On the other hand, in the assays with isolated CIV, a decrease in the binding affinities may inhibit CcO activity [78], although an increased turnover could yield the opposite under steady-state conditions [77] (Table 1).

In addition, the higher affinity of Y97pCMF Cc towards both the *proximal* and *distal* sites of CIV could facilitate the electron Cc-CcO flow and the pathway of Cc molecules on CIII-CIV supercomplexes [65]. Although Tyr97 in WT Cc is far from the adduct interface [51,84], its phosphorylation may affect long-range interactions, Cc dynamics and/or its orientation within the complex.

To sum up, phosphorylation of Cc regulates the physico-chemical properties—such as alkaline transition, redox potential, stability, affinity towards partners, among others—that are essential for shuttling electrons. This would enable the rapid adjustment of Cc function to changing cellular conditions.

5. Concluding Remarks

According to the data herein reviewed, Cc plays a major regulatory role in OxPhos owing to its highly dynamic interactions with redox targets, and displays a fast turnover during the electron exchange cycle. Multiple binding sites in Cc partners are essential to facilitate Cc turnover and to define the diffusion pathway of the electron carrier. The effectiveness of such a constraint relies on the ability of cells to modulate both the intricate arrangements of ETC complexes into higher-order assemblies and the ample set of PTMs on Cc and other ETC components. Whereas PTMs of CI, CIII and CIV could regulate their organization into supercomplexes, PTMs of Cc could control the binding mode and affinity between the heme protein and its partners (CIII and CIV). In this context, the 'restricted diffusion pathway' of Cc molecules between CIII and CIV could generate a large encounter ensemble of Cc on the surface of ETC complexes previously assembled in supercomplexes, in which the heme protein would adopt several orientations in equilibrium with stereospecific positioning. In addition, the extra regulation layer of OxPhos activity by post-translational phosphorylation of Cc could be essential to control oxygen consumption and the generation of harmful reactive species during homeostasis and pathology. Cc—either phosphorylated or not—might play additional roles in aiding the assembly of ETC membrane complexes or even supercomplexes. Actually, the lack of Cc disrupts the stability/assembly of CIV in yeasts, mammals and plants [45,85,86] and the heme protein migrates with CIII and CIV in BN-PAGE assays [27].

Conflict of Interests

All authors declare no conflict of interest.

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