

Structural basis of mitochondrial dysfunction in response to cytochrome *c* phosphorylation at tyrosine 48

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Regulation of mitochondrial activity allows cells to adapt to changing conditions and to control oxidative stress, and its dysfunction can lead to hypoxia-dependent pathologies such as ischemia and cancer. Although cytochrome *c* phosphorylation—in particular, at tyrosine 48—is a key modulator of mitochondrial signaling, its action and molecular basis remain unknown. Here we mimic phosphorylation of cytochrome *c* by replacing tyrosine 48 with *p*-carboxy-methyl-L-phenylalanine (pCMF). The NMR structure of the resulting mutant reveals significant conformational shifts and enhanced dynamics around pCMF that could explain changes observed in its functionality: The phosphomimetic mutation impairs cytochrome *c* diffusion between respiratory complexes, enhances hemeprotein peroxidase and reactive oxygen species scavenging activities, and hinders caspase-dependent apoptosis. Our findings provide a framework to further investigate the modulation of mitochondrial activity by phosphorylated cytochrome *c* and to develop novel therapeutic approaches based on its prosurvival effects.

cytochrome *c* | mitochondrial dysfunction | nuclear magnetic resonance | phosphorylation | respiratory supercomplexes

Oxidative phosphorylation (OxPhos) relies on the electron transport chain (ETC) to generate the membrane potential that drives ATP synthesis (1). Components of the ETC oxidize substrates to reduce molecular oxygen, thereby producing water. Nevertheless, around 2% of the electrons flowing across the ETC yield reactive oxygen species (ROS) (2). ROS are a source of oxidative stress and act as signaling molecules at low concentrations (3–5). Hindering redox reactions within the distinct ETC membrane complexes (complexes I to V) leads to enhanced ROS production (6). The activity of the ETC is tightly regulated by posttranslational modifications of its components, isoform swapping, and modulation of the equilibria for the association of the membrane protein complexes into supercomplexes (3, 7). Such associations allow substrate channeling while modulating ROS production (3, 7).

Oxidative stress response involves redox signals modulating protein phosphorylation (8). In the ETC, the major phosphorylation targets are NADH:UQ oxidoreductase, cytochrome *c* (Cc), and cytochrome *c* oxidase (CcO) (9). Besides being essential for oxidative respiration (Fig. 1*A*), Cc acts as a redox regulatory protein within the mitochondrial intermembrane space (10). In addition, Cc aids in controlling ROS levels by oxidizing superoxide anions (11) and exhibiting peroxidase activity (12); the latter, however, also yields lipid peroxidation (13, 14). Furthermore, Cc plays a crucial role in programmed cell death, a process that is only partially understood (15–21). In this context, a fraction of Cc binds and oxidizes cardiolipin (CL) at the internal mitochondrial mem-

brane, thereby facilitating the release of unbound Cc to the cytoplasm (16, 17). In mammalian cells, extramitochondrial Cc interacts with apoptosis activating factor 1 (Apaf-1) in the cytoplasm to spark the caspase proteolytic cascade (15). It has recently been shown that Cc can interact with several other proteins outside the mitochondria in humans and plants (18–21). The similarities between the Cc signaling networks in both organisms suggest that key programmed cell-death pathways are conserved throughout evolution (22).

Cc phosphorylation is an alleged modulator of the mitochondrial cell-death pathway (23, 24). Its deregulation is believed to be related to the onset of neurological disorders and cancer (24). Phosphorylation of Cc is easily reversed by phosphatases, hampering the isolation of the modified protein from tissues (25). Tyr-to-Glu substitutions of Cc, designed to emulate Tyr48 phosphorylation,

Significance

Cell response to physiological changes and oxidative stress involves the modulation of mitochondrial metabolism. Its dysfunction favors the development of hypoxia-dependent pathologies, including ischemia and cancer. A key modulator of mitochondrial activity is cytochrome *c*, whose cell function is regulated by tyrosine phosphorylation. However, how such modification affects cytochrome *c* structure and function is barely known. Here we report that a phosphomimetic mutant of cytochrome *c* exhibits enhanced dynamics, which could be responsible for the observed differences in cytochrome *c* functionality in oxidative stress and cell death. Thus, phosphorylation of cytochrome *c* becomes a target for further development of robust therapeutic approaches.

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Data deposition: The NMR assignments and coordinates for Y48pCMF cytochrome *c* reported in this paper have been deposited in the Biological Magnetic Resonance Data Bank (BMRB entry code 25660) and Protein Data Bank (PDB ID code 2N3Y) databases, respectively.

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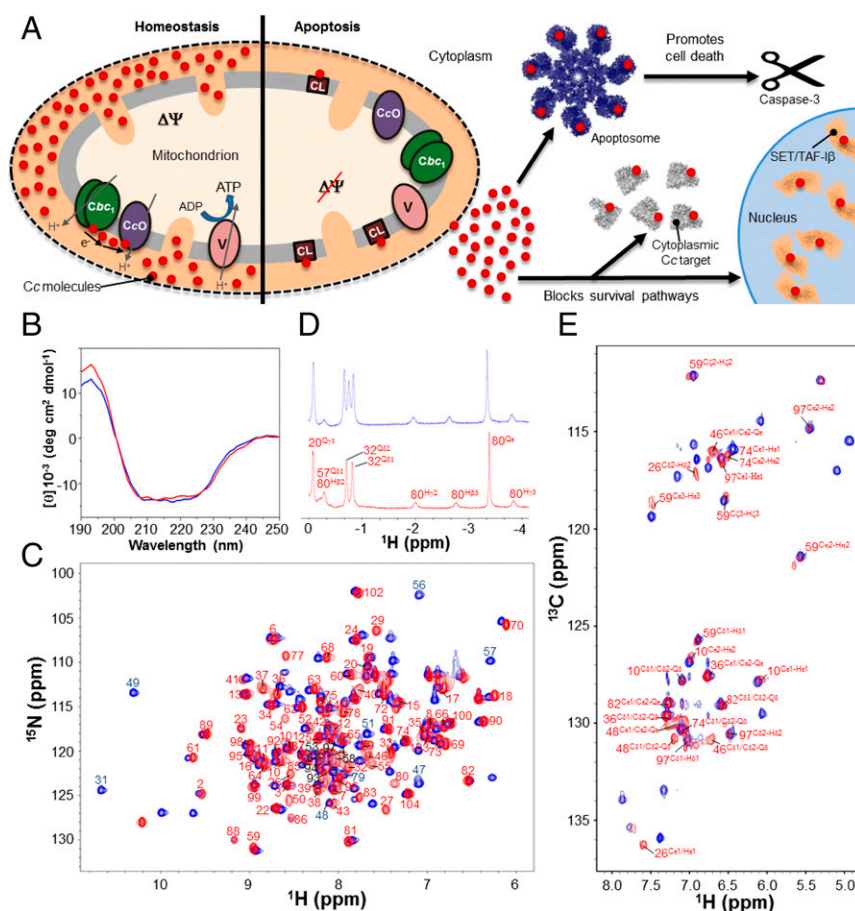


Fig. 1. Control of human cell fate by the Cc-based signalosome, and biophysical and structural characterization of the Y48pCMF variant of Cc. (A) Diagram of the role of Cc in homeostasis and apoptosis. (A, Left) Under homeostatic conditions, Cc (red circles) transfers electrons from the cytochrome *bc*₁ complex (*Cbc*₁) to the cytochrome c oxidase complex (*CcO*). (A, Right) Upon apoptotic stimuli, Cc is released to the cytosol to induce apoptosome formation and block prosurvival pathways. A portion of Cc remains bound to cardiolipin. (B) Far-UV CD spectra of the reduced forms of WT (blue) and Y48pCMF Cc (red). The same color code is maintained in the following panels. (C) Superimposition of the ^1H - ^{15}N HSQC spectra of uniformly ^{15}N -labeled forms of WT and Y48pCMF Cc. Backbone amide resonances of Y48pCMF Cc are labeled in red and black. Particular amide resonances of WT Cc are labeled in blue. (D) Detailed view of the ^1H NMR spectra of WT and Y48pCMF Cc at negative ppm values. Resonances for Met80 side-chain protons are shown for both Cc species. Assigned signals of all residues within this region are displayed for Y48pCMF Cc. The extra signal of WT Cc corresponds to the Qδ₁ protons of Ile53. (E) Superimposition of the aromatic region of the ^1H - ^{13}C HSQC spectra of WT and Y48pCMF Cc acquired in ^{13}C natural abundance. Assigned aromatic resonances of Y48pCMF Cc are displayed in red.

impair both the electron donation to CcO and the activation of caspase-9 mediated by Apaf-1 in vitro (26, 27). Notably, nitration of this residue also hinders the ability of Cc to activate Apaf-1 (28–30). Thus, these posttranslational modifications alter both mitochondrial and cytoplasmic functions of Cc. Tyr48 phosphorylation also affects the physical-chemical properties of Cc, such as its alkaline transition and midpoint reduction potential (E_0) (27, 31). Understanding the origin of these effects and how they affect Cc activities requires solving the 3D conformation of the phosphorylated species. However, deciphering the effects of Tyr48 phosphorylation on the structure and dynamics of Cc is highly challenging because of its reactivity toward phosphatases. Indeed, no atomic-resolution structure has been reported for either phosphorylated Cc or any reliable mimic mutant.

Here we have elucidated the structure of a phosphomimetic Cc variant in which Tyr48 is replaced by the synthetic, noncanonical amino acid *p*-carboxy-methyl-L-phenylalanine (*p*CMF). We show that such replacement induces local perturbations of the Cc structure and enhanced internal dynamics of the mutation's surroundings. We used biochemical assays to show that the Y48pCMF mutation impairs Cc channeling between cytochrome *bc*₁ (*Cbc*₁) and CcO, enhances peroxidase activity, and induces an antiapoptotic function of Cc.

Results

Phosphorylation of Tyr48 Induces Local Structural Changes in Cytochrome c. To understand how phosphorylation affects the structure of human Cc, we tackled the challenge of fully characterizing the phosphomimetic mutant Y48pCMF Cc in its reduced form, which is the redox state of Cc donating electrons to CcO in homeostasis and is essential for its apoptotic activity, because Cc becomes highly

reduced upon its release from the mitochondria to the cytosol (32). Y48pCMF Cc maintains the overall secondary structure and global fold of wild-type (WT) Cc, as inferred from circular dichroism (CD) (Fig. 1B) and ^1H - ^{15}N heteronuclear single-quantum correlation (HSQC) NMR spectra (Fig. 1C), respectively. Also, the native heme axial coordination was preserved, as indicated by 1D ^1H NMR data (Fig. 1D). The NMR spectra of reduced Y48pCMF Cc were extensively assigned: Triple-resonance experiments (*SI Appendix, Table S1*) allowed us to detect and assign 91 backbone amide signals and the sequential connectivities for most residues of the protein. Aliphatic side-chain signals were assigned using 3D HBHA(CO)NH and 3D HCCH-total correlation spectroscopy (TOCSY) experiments, leading to the assignment of most (96.4%) of the side-chain proton resonances. The ^{15}N -H resonance of *p*CMF48 was undetectable because the residue was not ^{15}N -enriched. Four proline residues (Pro30, Pro44, Pro71, and Pro76) interrupt the sequential HN–HN connectivity. Additionally, in contrast to WT Cc, we could not detect the amide protons of the Thr49, Ala51, Gly56, and Ile57 residues. However, the WT spectra only had four amide protons that could not be detected (Gly1, Glu21, Thr28, and Gly84) under similar experimental conditions. Further, in contrast to WT Cc, the signals from Asn31, Gly45, and Ser47 in *p*CMF48 Cc were significantly weaker than the rest. The largest chemical-shift perturbations for backbone amides induced by the modified residue at position 48 are confined to nearby residues (*SI Appendix, Fig. S1*).

Assigning ^1H resonances to the heme substituents and ^1H (and ^{13}C) signals to the aromatic side chains was required for analyzing additional 2D maps (i.e., COSY and ^1H - ^{15}N NOESY) (*SI Appendix, Table S1*). Assignment of the Cδ₁, Cδ₂, Cε₁, and Cε₂ aromatic signals from the *p*CMF48 side chain required the acquisition of an aromatic ^1H - ^{13}C HSQC spectrum recorded in a

natural abundance of ^{13}C (Fig. 1E). The assignment of the *p*CMF48 side-chain signals involved the complete assignment of the 2D ^1H – ^{15}N NOESY.

Numerous residues (Val20 to Asn31, Thr40 to Trp59, and Ile75 to Glu90) displayed signals attributable to a second, minor protein conformation, which had a 1:10 ratio in intensity. Hereafter, only the major form was considered for structure calculations.

We assigned 96% of all the ^1H signals for the major form. Structural information derived from 2D and 3D NOESY maps supported the presence of five α -helical regions (labeled α_1 to α_5) with typically strong HN–HN (*i*, *i*+1), medium-range H α –HN (*i*, *i*+3), and H α –HN (*i*, *i*+4) interresidual NOEs. These regions spanned the sequence stretches Val3 to Lys13, Ala50 to Asn54, Glu61 to Glu69, Pro71 to Tyr74, and Lys88 to Thr102, resembling those in the NMR structure of reduced WT Cc (33). In total, we observed and assigned 2,176 meaningful NOEs, corresponding to 20.8 relevant restraints per residue on average (SI Appendix, Fig. S2). The 71 ϕ and 71 ψ dihedral-angle constraints were derived from ^{15}N , $^{13}\text{C}'$, $^{13}\text{C}\alpha$, $^{13}\text{C}\beta$, and H α chemical shifts, using TALOS+ (34). The heme moiety was included in the calculations as previously reported for WT cytochromes (35, 36), assuming an intact heme iron coordination as supported by X-ray absorption spectroscopy (XAS) data (next section).

Two hundred structures were calculated by CYANA (37), and the 20 structures with the lowest target-function (TF) value were selected to form a representative family. The range of TF values was 0.38 to 0.98 $\text{\AA}^2$, highlighting the high accuracy between calculated and experimental distances. Further refinement of the 20 lowest TF structures involved restrained energy minimization and restrained molecular dynamics (RMD) computations. A final restrained energy minimization was carried out on the structure with the lowest root-mean-square deviation (rmsd) from the average for each of the 20 trajectories. The overall quality of the 20 lowest TF ensembles was good, according to the PROCHECK G factor (38), MolProbity clashscores (39), and other structural quality indicators (SI Appendix, Table S2). Most residues were in the Ramachandran plot favored regions, whereas Cys14, Cys17, His18, Val20, Lys37, and Asn70 were in the generously allowed ones, as also observed for the solution structure of WT Cc (33, 40). Cys14, Cys17, and His18 are covalently bonded to the heme moiety, thereby straining their backbone conformation, as already described for *c*-type cytochromes (35, 36). As an additional control, we performed 20-ns unrestrained molecular dynamics (MD) simulations of the final minimized conformers without any geometrical restraint. The rmsd for the main-chain atoms was about 1.55 $\text{\AA}$ at the plateau, hardly drifting ($0.122 \text{ pm}\cdot\text{ns}^{-1}$), as expected for a stable structure (SI Appendix, Fig. S3).

The overall fold of Y48pCMF Cc is very similar to that of the WT species (SI Appendix, Fig. S4), with rmsd values for the backbone nuclei of $1.67 \pm 1.01 \text{ \AA}$ (Fig. 2A and B). However, the two structures differ in the mutation-containing loop Ω_{NY} (residues 40 to 57; Fig. 2B). This is consistent with the chemical-shift differences (SI Appendix, Fig. S1) and the rmsd between the NMR solution structure of WT Cc [Protein Data Bank (PDB) ID code 1J3S] (33) and the refined lowest-TF Y48pCMF Cc (SI Appendix, Fig. S5A). An increased dynamics in the Ω_{NY} -loop has recently been observed in the G41S variant of Cc (42). The Ω_{NY} -loop (also known as foldon V) and helix α_2 constitute the less-stable folding unit of Cc (43). Unlike Tyr48 in the WT species, the *p*CMF48 residue presents a very low number of ^1H – ^1H NOEs (SI Appendix, Fig. S2) and a high rmsd value within the family (Fig. 2C and SI Appendix, Fig. S5B). A decrease in the number of detectable NOEs can generally be attributed to either a partial assignment of the residue or an increased internal mobility. In our case, the highly dynamic behavior of the *p*CMF48 residue was confirmed from further analysis of internal motions within the nanosecond-to-picosecond timescale (see below).

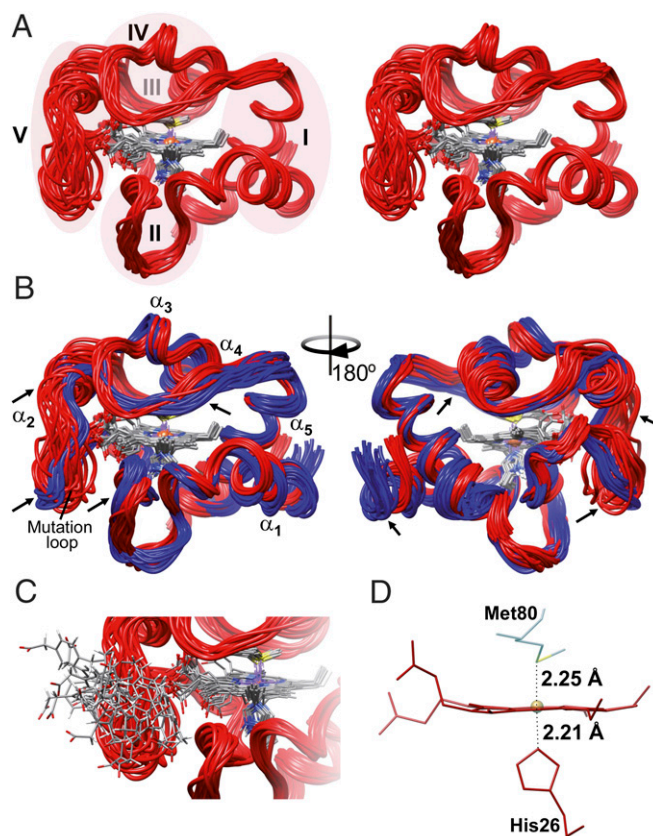


Fig. 2. NMR solution structure of the Y48pCMF variant of Cc. (A) Stereoview ribbon representation of the 20 best conformers of Y48pCMF Cc. Heme group atoms are displayed for all conformers. Ribbons are colored in red, whereas atoms from the heme group are colored following the CPK (Robert Corey, Linus Pauling, and Walter Koltun) color scheme. Foldons of Y48pCMF Cc are shadowed and marked with roman numerals, except for foldon III, which is located behind foldon IV. (B) Comparison between the NMR solution structures of WT Cc (PDB ID code 1J3S) (33) and Y48pCMF Cc (this work). The ribbon for WT Cc is in blue. The five α -helices of both Cc species, as well as the mutation-containing loop of Y48pCMF Cc, are marked. Arrows point to the regions on the Y48pCMF Cc ribbon with substantial structural changes compared with the WT form. (C) Detailed view of the loop harboring the *p*CMF48 residue. *p*CMF48 atoms follow the CPK color scheme. Protein structures are presented by UCSF Chimera software (41). (D) Detail of the heme group and axial ligands. Labels display iron-to-axial ligand distances for the Y48pCMF mutant obtained from the EXAFS analysis (SI Appendix, Fig. S5).

Other regions of the Y48pCMF Cc structure that differed from those of WT Cc belong to the 19 to 36 Ω_{G} -loop (part of foldon II) and the 71 to 85 Ω_{R} -loop (foldon IV) comprising Met80 and helix α_4 . The rmsd values for backbone atoms of residues from Val20 to Gly29 of Y48pCMF Cc are higher ($2.75 \pm 1.50 \text{ \AA}$) than those for the same residues of WT Cc (SI Appendix, Fig. S5A). In the second loop, the observed differences are mainly restricted to side chains. In fact, the rmsd values for backbone and heavy atoms are 1.18 ± 0.37 and $1.77 \pm 0.71 \text{ \AA}$, respectively. Residues included in the Ω_{R} -loop indeed displayed a double conformation, suggesting the presence of conformational equilibria (SI Appendix, Fig. S5A).

The ensemble of structures for Y48pCMF Cc is very precise, except for residues surrounding the mutation. The backbone rmsd from the mean is $0.89 \pm 0.01 \text{ \AA}$ for the whole protein and drops to $0.53 \pm 0.12 \text{ \AA}$ when the mutation surroundings are excluded (residues 40 to 57) (SI Appendix, Fig. S5B). As expected, the highest rmsd values correspond to the 40 to 57 Ω_{NY} -loop and the nearby residues Val20 to Gly29 in loop Ω_{G} , whose global backbone

rmsd values were 1.79 ± 0.53 and 0.78 ± 0.23 Å, respectively. These segments also exhibited a larger conformational variability in their secondary structure elements along the MD trajectory (SI Appendix, Fig. S3 B and C). High rmsd values along the Val20-to-Gly29 stretch are typical in Cc homologs (35, 36). They are also consistent with NMR signals revealing secondary conformations for the His26-to-Pro30 stretch. Moreover, the nearby Asn31 amide signal is weak, suggesting high mobility. Notably, all these residues contact the Ω_{NY} -loop, comprising the Y48pCMF mutation.

Further, the highest rmsd values mapped to the Ω_{NY} -loop. Consistently, signals from this region undergo a drastic reduction in their ^1H - ^1H NOE cross-peaks. The intensities of the amide signals of Gly45 and Ser47 in Y48pCMF Cc were severely decreased, whereas those from Thr49, Ala51, Gly56, and Ile57 were undetectable. The Ω_{R} -loop also contains some residues with high rmsd values (SI Appendix, Fig. S5B). The global rmsd value for backbone atoms of the 71 to 85 Ω_{R} -loop is 0.76 ± 0.20 Å. Notably, the end of the Ω_{R} -loop shows high rmsd values in WT cytochromes (35, 36).

Heme Iron Coordination Is Insensitive to Tyr48 Phosphorylation. We tested the effects of the mutation on the heme iron coordination environment and the axial coordination restraints used in our structure computations by using X-ray absorption spectroscopy, studying both WT and Y48pCMF species (Fig. 2D and SI Appendix, Fig. S5 C–F). The absorption spectra of the two proteins are almost identical. The X-ray absorption near-edge structure regions of the absorption spectra for both proteins are superimposed (SI Appendix, Fig. S5C). The absence of any shift in the energy position of the absorption edge indicates that the mutation did not affect the electron density at the Fe center.

Likewise, the extracted extended X-ray absorption fine structure (EXAFS) signals of the WT and Y48pCMF Cc species were highly similar. However, small differences in high wave vector (k) values can be observed (SI Appendix, Fig. S5D), as well as in the slightly lower amplitude of the Y48pCMF protein. Whereas the corresponding Fourier transforms are also very similar (SI Appendix, Fig. S5E), the amplitude of the first peak is lower and broader in the Y48pCMF Cc species, suggesting a larger degree of disorder. The dynamic disorder for the two proteins should be similar, as the measurements were performed at cryogenic temperatures in both cases. Hence, the differences in disorder are due to a larger static disorder in Y48pCMF Cc. The main scattering paths contributing to this peak originate from the four nitrogen atoms of the porphyrin ring, although contributions of the nitrogen and sulfur axial ligands have also been included. The fit to the data requires the addition of the paths involving the eight porphyrin carbons closest to the iron atom, beyond the first coordination sphere. In addition, the multiple scattering paths involving these atoms were also included. Fits were performed in R space but also reproduced the spectra well in q space (SI Appendix, Fig. S5F). The parameters obtained from the best fit to the data revealed that the distances between the iron atom and its first coordinating ligands are insensitive to the Y48pCMF mutation (SI Appendix, Table S3). Specifically, the distances from the iron center to the axial S ligand are 2.26 ± 0.001 and 2.25 ± 0.001 Å in the WT and phosphomimetic mutant species, respectively (Fig. 2D). Data analyses also showed that the value for the Debye-Waller factor corresponding to the path involving the four porphyrin nitrogen atoms increased from 0.0012 ± 0.0006 Å² in the WT species to 0.003 ± 0.001 Å² in the Y48pCMF mutant. These data are consistent with the preserved chemical-shift pattern of the iron axial ligands and heme substituents observed in 1D ^1H NMR (Fig. 1D), which have been reported to be sensitive indicators of the heme iron electronic structures (35, 36). The pattern of the observed NOEs for the heme substituents also supports an overall intact heme pocket, with the exception of the mutation site. Still,

the ensemble of structures shows a small change in the orientation of Phe82 with respect to the porphyrin ring (SI Appendix, Fig. S6).

Phosphorylation of Tyr48 Enhances Internal Mobility in Cytochrome c. NMR relaxation measurements were performed to evaluate the dynamics of WT and Y48pCMF Cc. The Y48pCMF substitution slightly affected both relaxation rate (R_1 and R_2) parameters (SI Appendix, Table S4). The rotational correlation time of the phosphomimetic mutant (6.96 ± 0.02 ns) was higher than that of the WT form (6.33 ± 0.02 ns), in agreement with the small increase (p_{null} values of $\sim 10^{-251}$, where p_{null} is the probability for the null hypothesis being the right one) in the average gyration radius, from 12.95 ± 0.07 to 13.04 ± 0.06 Å, as calculated by MD. Indeed, phosphorylation can alter protein dynamics at different timescales and cause conformational rearrangements, such as the formation of secondary conformations (44, 45).

Comparing R_1 , R_2 , and heteronuclear NOE (HetNOE) relaxation measurements recorded on the two proteins revealed that the Ω_{NY} -loop of Y48pCMF Cc exhibits a high mobility in the

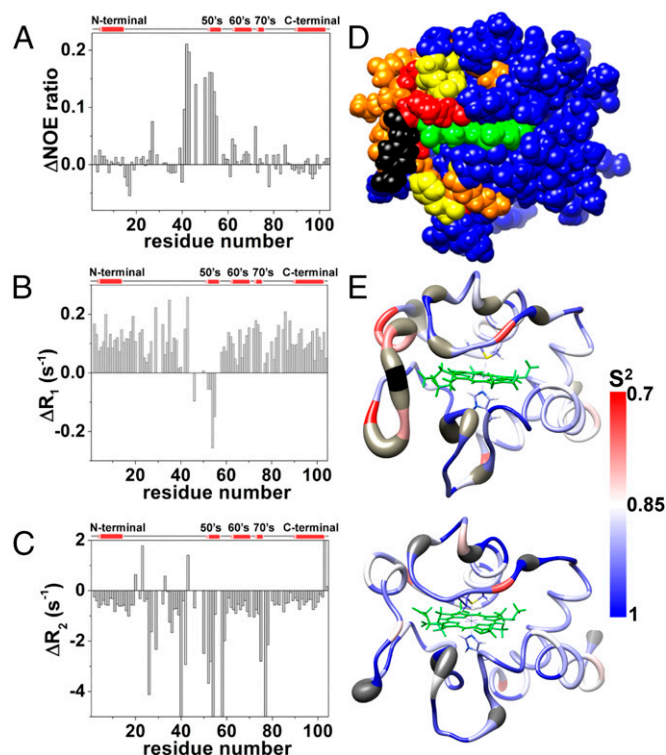


Fig. 3. Relaxation NMR measurements and dynamic properties of WT and Y48pCMF Cc. (A–C) Differences in heteronuclear NOE (A), relaxation rate R_1 (B), and relaxation rate R_2 (C) between the experimental values for the reduced forms of WT and Y48pCMF Cc, plotted as a function of the residue number. Gaps in data result from overlapping resonances, broadened resonances beyond the detection limit, and unassigned resonances. A scheme of the secondary structure elements is included (Top). (D) Map of the Y48pCMF Cc residues colored according to their dynamic properties. Affected residues in the heteronuclear NOE and relaxation rate R_2 parameters are colored in yellow and orange, respectively. Residues with backbone amide resonances that are undetectable in the ^1H - ^{15}N HSQC spectrum of Y48pCMF Cc but detectable in the ^1H - ^{15}N HSQC spectrum of WT Cc are colored in red. pCMF48 is shown in black, and the heme group is in green. Unaffected, unassigned, and proline residues are in blue. (E) Internal mobility comparison between Y48pCMF Cc (Upper) and WT Cc (Lower) Cc are represented on the respective NMR ribbon structures using a blue–red scale. Undetectable backbone resonances are in gray. Heme atoms are in green, with the axial ligands depicted as sticks.

picosecond-to-nanosecond timescale (Fig. 3). Indeed, the Gly41-to-Lys55 segment showed a drastic drop of HetNOE values in the mutant species (Fig. 3A). Further, the amide R_1 rates for the sequence stretch of Tyr46 to Lys55 in Y48pCMF Cc differ from those in WT Cc (Fig. 3B). In addition, R_2 analysis reveals three regions undergoing conformational exchange in the microsecond-to-millisecond timescale: His26 to Thr28, Thr40 to Trp59, and Ile75 to Thr78 (Fig. 3C). This behavior agrees with the reduced intensity or lack of detection of the amide signals belonging to these stretches, compared with the WT form. Furthermore, some regions within the protein displayed signals corresponding to double conformations in the 2D ^1H NOESY spectra—namely Val20 to Asn31, Thr40 to Trp59, and Ile75 to Glu90—indicating the presence of conformational equilibria between two different structures occurring on a slow timescale with respect to the NMR chemical shift. This dynamics involves residues located in a defined region surrounding the noncanonical amino acid (Fig. 3D). Altogether, the NMR relaxation measurements of Y48pCMF Cc agree with the per-residue S^2 -order parameter values computed with TENSOR (46) (Fig. 3E).

Hydrogen–deuterium exchange experiments showed that a common core region is protected from solvent amide–hydrogen exchange in both Cc species. Nevertheless, a substantial number of amides become unprotected in pCMF48 Cc (SI Appendix, Fig. S7). The newly accessible amide protons in the phosphomimetic mutant (Gly29, Gly37, Arg38, Thr40, Gln42, and Trp59) are located in the surroundings of pCMF48 and the nearby Ω_{G} -loop, in agreement with their high mobility in the microsecond-to-millisecond timescale.

Tyr48 Phosphorylation Modulates the Interaction of Cytochrome *c* with Its Mitochondrial Partners in the Electron Transfer Chain. Cc carries electrons from *Cbc*₁ to CcO within the ETC. To elucidate more details about the process, we analyzed the molecular recognition between the soluble N-terminal domain of plant cyto-

chrome *c*₁ (Cc₁) or bovine CcO and the phosphomimetic human Y48pCMF Cc. The human and plant N-terminal domains of Cc₁ have a 62% overall sequence identity and similar charge distributions on their molecular surfaces (47), whereas the bovine and human CcO are evolutionarily related proteins, with 91 and 96% sequence identity and homology, respectively.

Two binding sites for human and plant Cc on plant Cc₁ have been recently reported (47, 48). The proximal site is located near the heme moiety and is compatible with electron transfer, whereas the distal site lies far from the heme group and probably constitutes a local energy minimum of the encounter ensemble.

To test how Tyr48 phosphorylation affects the conformation of the Cc₁–Cc complex, we recorded ^1H – ^{15}N HSQC spectra upon titration of ^{15}N -labeled reduced Cc with unlabeled reduced Cc₁. Several amide signals exhibited significant chemical-shift perturbations (CSPs), thus indicating a fast exchange rate within the NMR timescale (Fig. 4A and B). Average amide CSPs ($\Delta\delta_{\text{avg}}$) were larger than 0.075 ppm for 11 residues: Gln16, Lys27, Gly29, Ala50, Lys55, Ile58, Lys72, Gly77, Met80, Ile81, and Val83 (Fig. 4B). The fits of CSPs along the titration suggested that Cc₁ binds two Y48pCMF Cc molecules, as previously found with the WT Cc species (47, 48) (Fig. 4C). Using NMR spectroscopy, the dissociation constant (K_D) values for the Cc₁–Y48pCMF Cc complex were estimated to be 0.6 and 102 μM for the proximal and distal binding sites, respectively. All of the perturbed residues, except Lys55 and Ile58, surround the heme crevice, as previously described for the interaction between WT Cc and Cc₁ (47, 48) (Fig. 4D). In fact, this region constitutes a very well conserved interaction surface in *c*-type cytochromes (49–52). Lys55 and Ile58, in turn, are located in the Ω_{NY} -loop, which undergoes a conformational exchange in free Y48pCMF Cc (Fig. 4D). In addition, significant CSPs ($\Delta\delta_{\text{avg}} \geq 0.05$ ppm) were detected for Lys7, Lys13, His26, Lys39, Ala43, Thr78, Lys86, Lys88, and Glu89. Interestingly, Lys8, Lys13, Lys27,

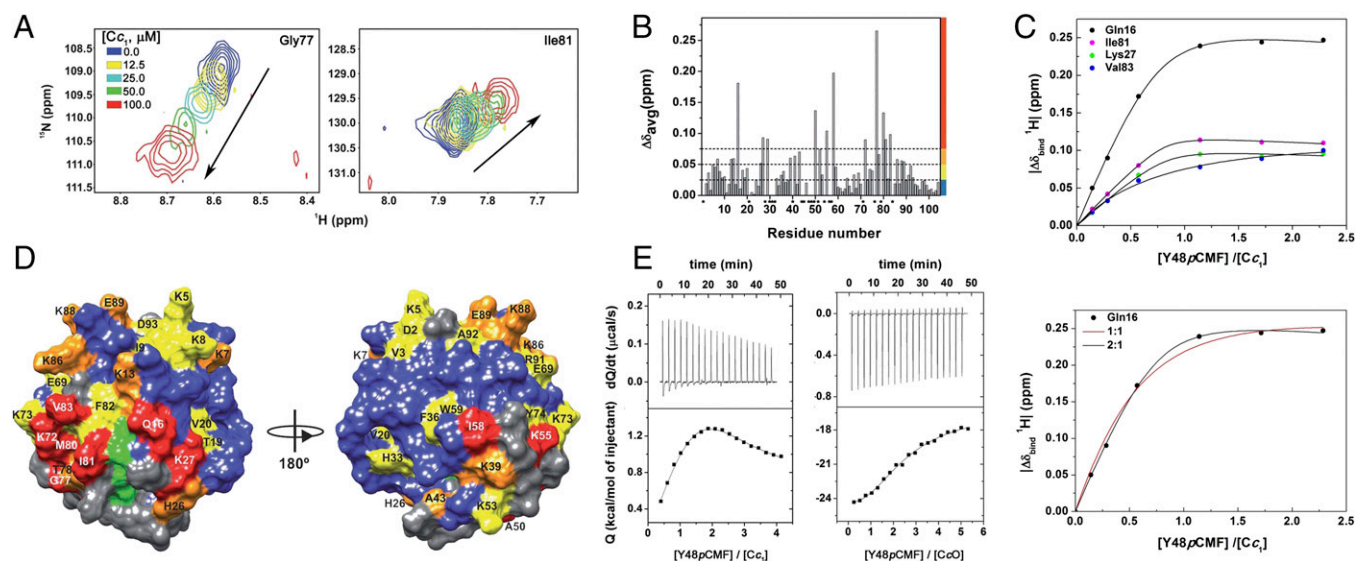


Fig. 4. Binding assays between Y48pCMF Cc and its respiratory partners. (A) Overlay of selected residues of ^1H – ^{15}N HSQC spectra of ^{15}N -labeled Y48pCMF Cc along with titration with Cc₁. Signals corresponding to different titration steps are colored according to the code indicated. (B) Plot of chemical-shift perturbations of ^{15}N -labeled Y48pCMF Cc as a function of residue number. Proline and nonassigned residues are marked by asterisks. Color bars stand for the $\Delta\delta_{\text{avg}}$ categories: insignificant $\Delta\delta_{\text{avg}} < 0.025$ ppm, blue; small $0.025 \leq \Delta\delta_{\text{avg}} < 0.050$ ppm, yellow; medium $0.050 \leq \Delta\delta_{\text{avg}} < 0.075$ ppm, orange; and large ≥ 0.075 ppm, red. (C, Upper) Curves representing the best global fit of several amide signals in the ^1H dimension to a 2:1 ratio for the Y48pCMF Cc–Cc₁ binding model with two different global K_D values. (C, Lower) Binding curves of Gln16. Lines represent the best fit to 1:1 (red) and 2:1 (black) binding models. (D) CSP map of reduced Y48pCMF Cc upon addition of reduced Cc₁ at a 1:1 ratio. Residues are colored according to $\Delta\delta_{\text{avg}}$ categories, as indicated in B. Proline and nonassigned residues are in gray. (E) ITC measurements of the Y48pCMF Cc–Cc₁ and Y48pCMF Cc–CcO complexes in their reduced states. Experimental data were fitted to a 2:1 binding model. Thermograms (Upper); binding isotherms (Lower).

Lys72, and Lys86 also experienced large CSPs, as previously reported for the interaction between WT Cc and Cc₁ (47, 48).

To obtain further data on the binding affinity and stoichiometry of the interaction between the two redox proteins, isothermal titration calorimetry (ITC) experiments were performed on both redox states. The isotherms obtained by titrating reduced Y48pCMF Cc with reduced Cc₁ are displayed (Fig. 4E, Left). For the interaction between Cc and Cc₁, the isotherm clearly shows two components with opposite signs in their respective enthalpy terms, thereby indicating the presence of two binding sites, in agreement with the NMR data herein presented and previous reports on the WT species (47, 48). Indeed, all data fit a model with two independent binding sites in the corresponding Cc partner (SI Appendix, Table S5). The interaction between Y48pCMF Cc and Cc₁ was entropy-driven. At pH 7.4, the K_D value at the proximal site was half that observed for WT Cc, whereas that at the distal site was four times higher (SI Appendix, Table S5). This could be ascribed to the extra negative charge at position 48, which alters the surface electrostatic potential of the hemeprotein (SI Appendix, Fig. S8). Both Cc₁–Y48pCMF Cc and Cc₁–Cc complexes in their oxidized states showed similar thermodynamic and equilibrium parameters at both acidic and basic pH values. The only exception was the K_D value for the oxidized Cc₁–Y48pCMF Cc complex, at the proximal site of Cc₁, which was approximately four times lower than that for the oxidized Cc₁–WT Cc adduct at pH 8.5. ITC measurements of the oxidized Cc₁–Y48pCMF Cc complex were not run at pH 7.4 because the pK_a for the alkaline transition, which is specific to the ferric state (53), shifts to neutral pH upon Tyr48 phosphorylation (31). Hence, the WT and phosphomimetic forms of oxidized Cc can differ in their sixth axial ligand at pH 7.4 (SI Appendix, Table S5).

In addition, we analyzed the binding affinity of the cross-complex between the reduced species of bovine CcO and human Y48pCMF Cc by ITC. The isotherm inflection point lies at a 2:1 Y48pCMF-Cc:CcO ratio, thereby indicating the presence of two binding sites on CcO, as previously found for WT Cc (47, 54, 55). The best fit was achieved when the model used distinct K_D values for the two independent binding sites (SI Appendix, Fig. S9). The resulting isotherms likewise fit to such a model, yielding different micromolar-ranging K_D values (Fig. 4E, Right and SI Appendix, Table S6) and enthalpy-driven interactions. Both the CcO proximal and distal sites had lower affinities for Y48pCMF Cc than the WT species, as previously reported (47) (SI Appendix, Table S6). Altogether, these data agree with previous steady-state kinetic analysis that demonstrated a binding model with more than one molecule of Cc per molecule of Cbc₁ (56) and with kinetic-based models that proposed the existence of alternative, nonproductive binding sites for horse Cc in bovine CcO (55). In addition, direct binding studies performed by gel filtration between several mammalian Ccs and bovine CcOs evidenced a 2:1 stoichiometry, in which Cc bound to a first site with a dissociation constant within the nanomolar range and to a second site with less affinity (54).

To assess the functional ability of Y48pCMF Cc to reduce respiratory complex IV, we tested the CcO activity of isolated complex IV or Cc-free mitochondria (ΔCc) from yeast cells grown with glucose as a carbon source (Fig. 5A and SI Appendix, Fig. S10A). In both cases, the Cc oxidation rate was at least twofold higher with Y48pCMF Cc than with WT Cc, thereby suggesting that Tyr48 phosphorylation enhances the ability of Cc to donate electrons to complex IV. Direct measurements of O₂ consumption were consistent with such data (SI Appendix, Fig. S10B). Interestingly, the CcO activity was positively regulated by the human membrane proteins hypoxia-inducible domain family members 1A and 2A (HIGD1A and HIGD2A), which promote cell survival under hypoxia. HIGD2A was successfully expressed in cell-free expression systems combined with *n*-dodecyl- β -D-maltoside, as previously reported for HIGD1A (57) (SI Appendix, Fig. S11A–C). HIGD1A significantly increased the rate of CcO-catalyzed

oxidation of Y48pCMF Cc (Fig. 5B), as reported for WT Cc (58). Strikingly, HIGD2A induced an even stronger positive effect than HIGD1A (Fig. 5B). Nevertheless, the HIGD-dependent increase in CcO activity was slightly lower with Y48pCMF Cc (Fig. 5B). This may be due to HIGD-dependent changes in either the complex IV affinity toward Cc or the restraints of Cc diffusion (channeling) from Cc₁ to complex IV. A direct Cc–HIGD interaction can also not be excluded.

To confirm the HIGD-mediated regulation of the CcO activity in a cellular context, we isolated mitochondria from different yeast strains grown either with glucose (YPD medium), which supports fermentation and respiration, or with the non-fermentable carbon sources lactate and galactose (YP-Gal) (59). Under both metabolic conditions, the isoformic respiratory supercomplex factors 1 (Rcf1, formerly Aim31) and 2 (Rcf2, formerly Aim38) are constitutively expressed (Fig. 5C, Inset), in agreement with their role in CcO activity and supercomplex stability (59–61). Rcf1 is a yeast ortholog of the human HIGD1A and HIGD2A proteins, whereas Rcf2 is specific to yeast. The external membranes of isolated mitochondria were then permeabilized to allow the entry of exogenous WT or Y48pCMF Cc. Under these conditions, mitochondria isolated from a yeast strain deficient in both Rcf1 and Rcf2 ($\Delta Rcf1/2$), as verified by Western blot, displayed an endogenous CcO activity lower than that isolated from WT yeast (WT_{Rcf}), no matter which type of exogenous hemeprotein—WT or Y48pCMF Cc—was used for supplementation (Fig. 5C). This indicates that Rcf1 and Rcf2 act as positive modulators of CcO activity, similar to human HIGD proteins. In mitochondria isolated from yeast grown with non-fermentable carbon sources, the Rcf-mediated increase in CcO activity was less prominent when Y48pCMF Cc rather than WT Cc was added as an exogenous electron donor, in agreement with the in vitro behavior of isolated proteins (Fig. 5B). This suggests that Tyr48 phosphorylation makes Cc less sensitive to the enhancer mechanism of the Rcf proteins and/or to the ability of Rcfs to stabilize the Cbc₁–CcO supercomplexes (Fig. 5C). In fact, Rcf1 and Rcf2 promote the supercomplex assembly, preferably in mitochondria from yeasts grown in a respiratory-based medium (Fig. 5D and SI Appendix, Fig. S11D). This is especially remarkable when comparing the band intensities in an anti-COX-II immunoblot (Fig. 5D). Note that such OxPhos supercomplexes could show a certain degree of heterogeneity in their composition, because the gene transcription of COX-Va and COX-Vb—which encode for two CcO isoforms—is repressed and active, respectively, in the aerobic-to-anaerobic metabolic transition (62). Moreover, the faint band pattern of $\Delta Rcf1/2$ strains grown in YPD medium on blue native (BN)/PAGE corresponds to supercomplexes, thus suggesting that other factors may contribute to their assembly (Fig. 5D and SI Appendix, Fig. S11D).

Altogether, our data suggest that phosphorylation of Cc at Tyr48 modulates the mitochondrial ETC (Fig. 5E). Such a post-translational modification allows a fast adaptation of the hemeprotein function to changing cell conditions. The population of Cbc₁–CcO supercomplexes is less prominent in the presence than in the absence of glucose, although Rcf proteins are still expressed. In any case, Cc channeling from Cbc₁ toward CcO is impaired. At a physiological pH, Tyr48 phosphorylation thus favors binding of Cc to the proximal rather than to the distal site of Cc₁. Under respiration-based growth, the Rcf proteins preferably associate directly with the OxPhos supercomplex, bridging Cbc₁ and CcO. However, the weaker Y48pCMF Cc–CcO binding, along with the loss of the distal site on Cc₁ due to phosphorylated Cc, impairs the channeling of Cc molecules that functionally connects Cbc₁ with CcO, which has been proposed to occur with WT Cc (47, 48). As a consequence, Y48pCMF Cc is less efficient than WT Cc as an electron carrier toward CcO in the context of OxPhos supercomplexes.

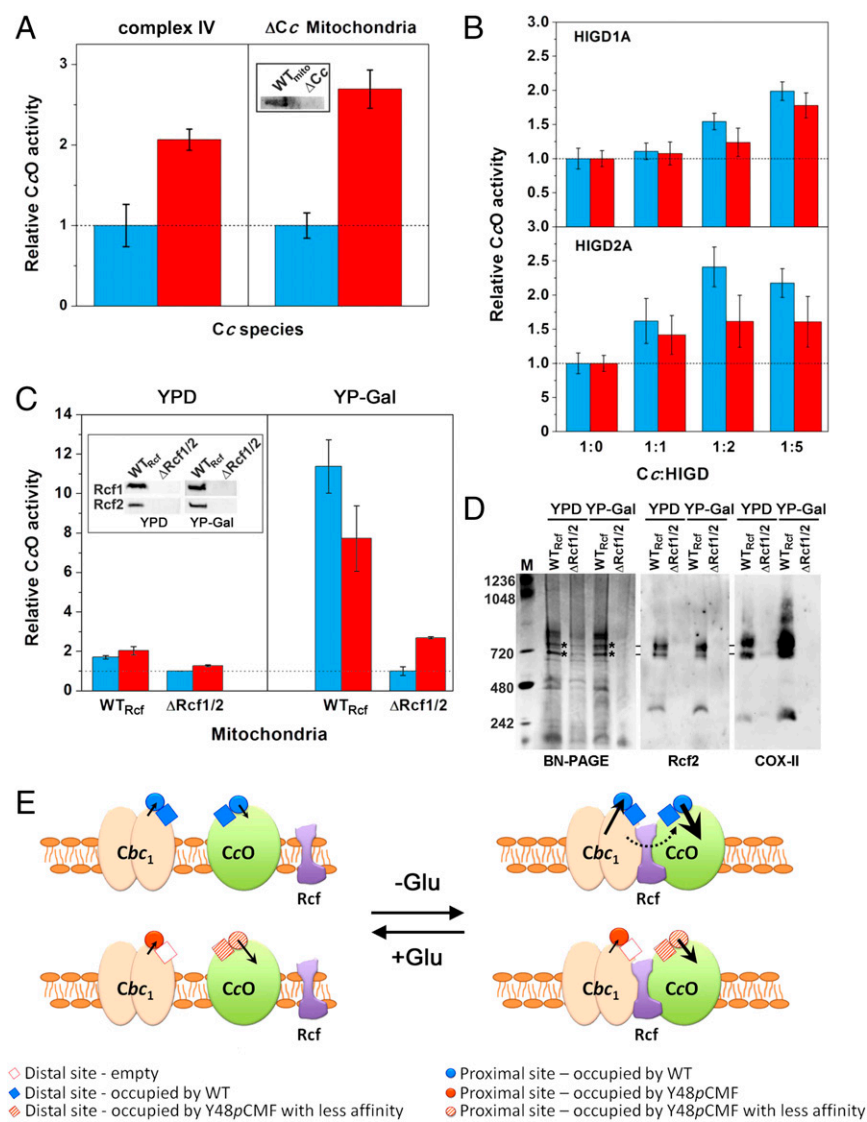


Fig. 5. CcO activity with WT or Y48pCMF Cc as the electron donor. (A) CcO activity of isolated complex IV and of mitochondria lacking Cc (ΔCc) upon addition of exogenous WT (blue bars) or Y48pCMF Cc (red bars). Western blot results confirmed the lack of endogenous Cc in ΔCc mitochondria (Inset). (B) In vitro modulation of CcO activity by HIGD1A and HIGD2A. WT Cc (blue bars) or Y48pCMF Cc (red bars), along with HIGD1A or HIGD2A at the indicated ratios, was added to isolated complex IV. (C) Effect of the modulators Rcf1 and Rcf2 on the CcO activity of mitochondria isolated from yeasts grown in YPD or YP-Gal media with either WT Cc (blue bars) or Y48pCMF Cc (red bars). All data represent the mean $\pm$ SD of three independent experiments. In all cases, CcO activity was detected only upon addition of exogenous Cc but not with endogenous Cc. (C, Inset) Western blots of WT_{Rcf} mitochondria (lane 1) and mitochondria lacking Rcf1 and Rcf2 ($\Delta Rcf1/2$) (lane 2). (D) BN/PAGE and Western blots of mitochondria from WT_{Rcf} and $\Delta Rcf1/2$ strains, using antibodies against Rcf2 and COX-II. Bands submitted to tryptic digestion (SI Appendix, Fig. S11D) are highlighted by asterisks. (E) Scheme of the interactions within the electron transport chain involving Cbc₁, CcO, Rcf proteins, and WT or Y48pCMF Cc, as a function of glucose (Glu) availability. The Rcf proteins facilitate the interaction between Cbc₁ and CcO to form OxPhos supercomplexes, mainly under glucose deprivation (Right). The thickness of solid arrows refers to the electron transfer rate at the Cc-binding proximal sites of Cbc₁ and CcO by WT or Y48pCMF Cc—the longer and thicker the arrow, the more efficient the electron transfer. The dashed line highlights the channeling of WT Cc molecules.

Tyr48-Phosphorylated Cytochrome c Acts as an Improved Peroxidase-Like Enzyme, in Particular When Bound to Cardiolipin-Containing Liposomes.

Assembly of the mitochondrial protein membrane complexes Cbc₁ and CcO into OxPhos supercomplexes enables more efficient electron flow and decreases ROS levels generated by the ETC (63). The phospholipid CL, located in the inner mitochondrial membrane, also stabilizes the resulting supercomplexes (64) in a HIGD/Rcf-independent manner (59). In addition, CL interacts with Cc in a two-step binding reaction (65): In the first step, the so-called A site at the Cc surface makes transient electrostatic contacts with the membrane; in the second step, hydrophobic forces drive the formation of a tight and stable Cc–CL complex, with one of the CL acyl chains entering the hydrophobic groove of Cc (termed the C site) (66–68). C-site binding then triggers Cc-regulated CL peroxidation under oxidative stress and induces early apoptosis (40). Interestingly, several residues of Cc (Lys22, Lys27, and His33) become protonated at low pH to form an extra binding site—the so-called L site—that facilitates the electrostatic interaction of Cc with mitochondrial membranes (69).

Within this framework, we analyzed how tyrosine phosphorylation can fine-tune the affinity of Cc toward CL-containing liposomes, analyzing binding of Cc species to liposomes in Hepes buffer at a physiological pH value by electrophoretic mobility shift

assays (EMSAs) in native agarose gels (Fig. 6 A and B). To compare the binding properties of Cc species to the CL-containing liposomes of 4:1 DOPC:TOCL (1,2-dioleoyl-*sn*-glycero-3-phosphocholine:1,1',2,2'-tetraoleoylcardiolipin) or to liposomes of DOPC alone, we measured the mobility profiles of Cc at different Cc:lipid ratios. Although both WT and Y48pCMF Cc bound to DOPC:TOCL and DOPC vesicles, their binding affinity for the CL-free DOPC liposomes seemed to be lower (Fig. 6 A and B). Notably, the presence of free hemeprotein at high lipid concentrations suggested that Y48pCMF Cc has a lower affinity than WT toward DOPC:TOCL liposomes (Fig. 6B), as recently observed for the phosphomimetic mutant S47D Cc (70). Another interesting finding is the unspecific interaction of Cc—either the WT or Y48pCMF species—with DOPC vesicles (Fig. 6 A and B).

ITC measurements corroborated these EMSA data. ITC experiments were performed under aerobiosis to mimic physiological conditions, despite using degassed samples. This may result in partial oxidation of Cc upon binding to liposomes, as previously reported (71). ITC analysis yielded the apparent K_D values for the first binding event of 427 μ M (WT Cc) and 780 μ M (Y48pCMF Cc) (Fig. 6C). Data for the interaction of WT Cc with liposomes are consistent with the K_D values reported before (68, 72–74), but they diverge from the lower K_D values reported in the literature

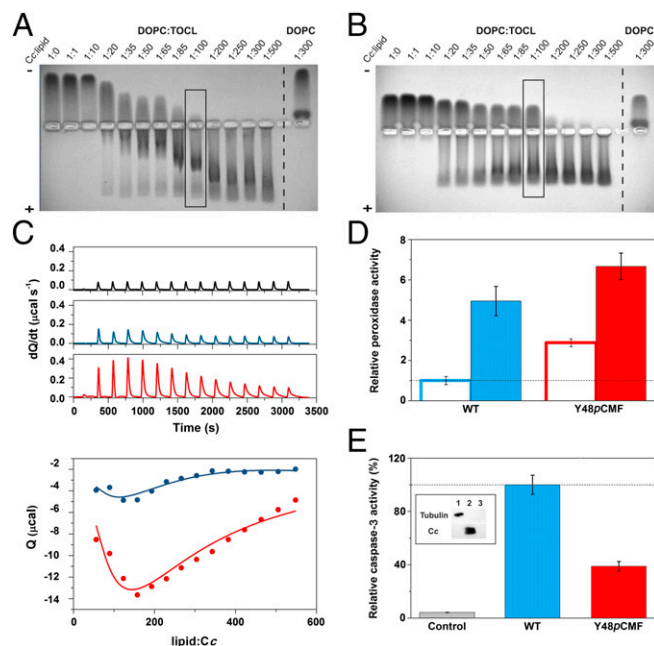


Fig. 6. Liposome-binding assays with caspase-3 activity induced by WT and Y48pCMF Cc. (A and B) EMSA of Cc in the presence of increasing concentrations of lipids. DOPC:TOCL (4:1) or DOPC liposomes were incubated with WT (A) or Y48pCMF (B) Cc. Note that free Cc species moved to the cathode, whereas liposome-bound Cc migrated to the anode. Lanes marked by rectangles correspond to the Cc:lipid ratio at which the peroxidase activity was determined (see below). (C) Calorimetric assays for lipid binding to Cc. (C, Upper) ITC thermograms, corresponding to titrations of DOPC:TOCL 4:1 liposomes (black), WT Cc (blue), or Y48pCMF Cc (red). (C, Lower) Binding isotherms with WT Cc (blue dots) or Y48pCMF Cc (red dots). Continuous lines represent the best fits to a sequential binding, as computed with NanoAnalyze software (TA Instruments) with a stoichiometry of 30 molecules of lipid per molecule of Cc. All data represent the mean $\pm$ SD of three independent experiments. (D) Relative peroxidase activities of WT Cc (blue) or Y48pCMF Cc (red) in the presence of liposomes containing DOPC (empty bars) or DOPC:TOCL (4:1) (filled bars). (E) Relative caspase-3 activity in HEK293 cell extracts devoid of endogenous Cc upon addition of exogenous WT Cc (blue) or Y48pCMF Cc (red). A lack of caspase autoactivation was verified in a run without the addition of Cc (gray). Western blots confirmed the lack of endogenous Cc in cytoplasmic cell extracts after immunoblotting with anti- α -tubulin (cytosolic marker) and anti-Cc antibodies (inset). Lane 1, cytoplasmic cell extracts; lane 2, Cc; lane 3, BSA as a negative control. All data represent the mean $\pm$ SD of three independent experiments.

(75) and the A/C two-site binding model described by Kinnunen and coworkers (66, 67). Such discrepancies are still unresolved, and further research is necessary to harmonize data from all these different binding assays in a unified and single model. The interaction with CL resulted in apparent ΔH values equal to 2.75 kcal $\cdot$ mol $^{-1}$ for WT Cc and 38.97 kcal $\cdot$ mol $^{-1}$ for Y48pCMF Cc. The differences in K_D indicate a small change in binding energies (of $\sim$ 6 kcal $\cdot$ mol $^{-1}$). Hence, the changes in ΔH are indicative of enthalpy-entropy compensation effects, which are compatible with the electrostatic change resulting from the extra carboxylate group of the Y48pCMF species. Further, an exothermic process can be observed, with longer peak equilibrium times upon successive lipid additions. In this case, the apparent ΔH values with the WT and Y48pCMF Cc species were equal to -6.97 and -47.84 kcal $\cdot$ mol $^{-1}$, respectively, but the apparent K_D value (of $\sim$ 1 mM) was practically the same with the two Cc species. Altogether, our EMSA and ITC assays indicate that Y48pCMF Cc binds to CL-containing liposomes with a slightly lower affinity than WT Cc.

WT Cc undergoes CL-dependent conformational changes that allow H_2O_2 to access the heme crevice (74). Hence, we addressed whether the affinity differences between WT and Y48pCMF Cc

for CL could affect their peroxidase activity. In the absence of CL-containing liposomes, Y48pCMF Cc exhibited a threefold higher peroxidase activity than WT Cc (Fig. 6D). However, the presence of DOPC:TOCL vesicles, at a 1:100 Cc:lipid ratio, increased the enzymatic activity of both WT and Y48pCMF Cc, similar to that observed for other phosphomimetic Cc mutants (70). Note that the slightly lower peroxidase activity increment observed for Y48pCMF Cc is likely due to its higher population of free protein compared with WT Cc (Fig. 6B and D).

Tyr48 Phosphorylation Is an Antiapoptotic Posttranslational Modification.

Phosphorylated Cc may be more easily released from mitochondria because of its lower affinity toward CL, as inferred from EMSA and ITC assays. Translocation to the cytosol could thus enable phosphorylated Cc to interact with Apaf-1 and to assemble the caspase-activating apoptosome. However, cytosolic caspase-3 activation was decreased by about 60% in the presence of the Y48pCMF Cc mutant (Fig. 6E). This is in agreement with the behavior previously reported for the Y48E Cc mutant, which exhibits a lower ability than WT Cc to activate not only caspase-3 (26) but also procaspase-9 by nonfunctional apoptosome assembly (27). These results could thus be indicative of an antiapoptotic function of Cc when phosphorylated at position 48.

Discussion

Here we tackle the structural and functional characterization of the Y48pCMF variant of human Cc. This mutation mimics protein phosphorylation at Tyr48 by adding a negative charge and slightly increasing the side-chain size while keeping the aromatic ring. A recent spectroscopic analysis of Y48pCMF Cc showed a singular shift of the typical alkaline transition pK_a to physiological pH values (31), as is the case with the Y48E Cc mutant (26, 27). Here, we report NMR-based structure computations that indicate that Tyr48 phosphorylation maintains the core fold of Cc but increases internal motions in the loops Ω_{NY} , Ω_R , and Ω_G . Specifically, the Ω_{NY} -loop, the most unstable folding unit of the hemeprotein, becomes looser and reaches conformational equilibria in Y48pCMF Cc. Enhanced motions at the Ω_G - and Ω_R -loops could be associated with the shift in the alkaline transition pK_a . Indeed, these two loops hold the residues that provide the iron axial ligands, which are observed to at least partially lose their metal coordination in the alkaline form (76). This is in contrast to XAS data that suggest that axial coordination remains untouched. However, the cryogenic temperatures at which the XAS spectra were recorded may

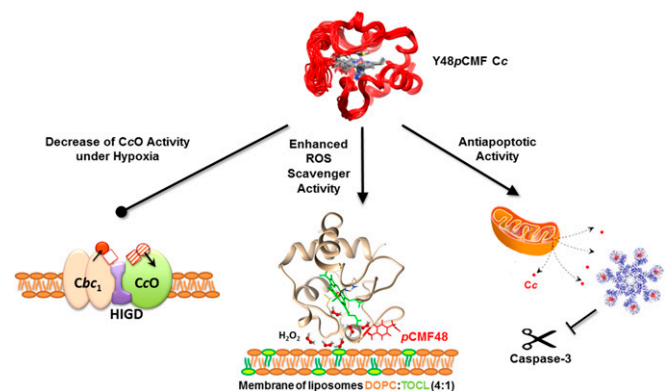


Fig. 7. Schematic diagram illustrating the changes induced in cell function by phosphomimetic Y48pCMF Cc. The negative charge at position 48 decreases CcO activity by disrupting Cc channeling in OxPhos supercomplexes (Left), enhances ROS scavenger activity by increasing the peroxidase activity of CL-bound Cc (Middle), and promotes the antiapoptotic function of Y48pCMF Cc by inhibiting its ability to activate the caspase-3 cascade (Right).

amplify the most stable structures and reduce contributions from minority species with a high disorder, which are undetectable.

The local but substantial changes in conformation and dynamics of the regions surrounding the Y48pCMF mutation increased the solvent accessibility of the porphyrin ring, thereby enhancing cytochrome peroxidase activity. Thus, Y48pCMF Cc proficiently scavenges ROS and avoids damage of cellular components (Fig. 7). Our findings also indicate that Y48pCMF Cc binds to mitochondrial CL with a lower affinity than WT Cc but that Y48pCMF Cc shows a higher peroxidase activity upon binding to liposomes. According to the literature, both events may facilitate Cc translocation to the cytoplasm during apoptosis (77). However, phosphorylation at Tyr48 hinders the proapoptotic activity of extramitochondrial Cc, similar to the function of HIGD1A protein in ischemia and tumorigenesis (78). Indeed, both HIGD1A and Y48pCMF Cc act as prosurvival proteins by preventing apoptotic caspase activation (Fig. 7) (79).

As expected, the Y48pCMF mutation affects the binding mode of Cc to its well-known respiratory partners *Cbc*₁ and CcO. Not surprisingly, the enhanced internal motions of the loops Ω_{NY} and Ω_R and the electrostatic change at the interaction patch resulting from the additional carboxyl group affect the interaction of Y48pCMF Cc with the two membrane complexes. The mutation-induced changes in electrostatics and in solvation at the interface also concur for the interaction of reduced Y48pCMF Cc with the distal sites of CcO (and Cc₁), as inferred from the high decrease in dissociation enthalpies and the strong (or moderate) enthalpy-entropy compensation herein reported. Indeed, the binding of Y48pCMF Cc on the Cc₁ surface is weak and functionally irrelevant at the distal site but is favored at the proximal site (Fig. 5E). When the turnover of the redox carrier at the main site is considered, the high affinity of reduced Y48pCMF Cc inhibits its own replacement by oxidized Cc molecules, thereby impairing the electron transport.

The Cc-CcO complex keeps its interface solvated, according to the recent X-ray diffraction structure (80), which displays a protruding lysine side chain in CcO pointing to Tyr48 in Cc. A salt bridge involving phosphorylated Tyr48 would restrain the Ω_{NY} -loop. The increase in the K_D of the complex between Y48pCMF Cc and CcO (this work) and the decrease in the Michaelis constant (K_M) of phosphorylated Cc with CcO (81) indicate that K_M is mainly governed by the catalytic step. This agrees with the larger CcO-driven oxidation rate of Y48pCMF Cc compared with that of WT Cc at the limiting protein concentrations in the enzymatic activity assays used here. In this context, the ca. 60-mV decrease in the E_0 of Y48pCMF Cc (31) may facilitate the electron flow from Cc to CcO, although it may impair Cc reduction by *Cbc*₁. In other words, Y48pCMF Cc exhibits a reduced ability as oxidizing agent (it is less capable than WT Cc of receiving electrons from Cc₁) but displays an increased capacity as reducing agent (it is a better electron donor to CcO than WT Cc). It is well-known that the catalytic step can be tuned by conformational changes of Cc upon

binding to CcO (52). Strikingly, these changes resemble the prominent internal dynamics of Y48pCMF Cc, which could also explain not only the change in E_0 but also the shift in the alkaline transition pK_a of oxidized Y48pCMF Cc (31). Therefore, the transition from reduced to oxidized Y48pCMF Cc, which is accompanied by a change in the iron axial ligand at physiological pH, is expected to have a high-energy barrier (82). Hence, such conformational changes might be the limiting step for Y48pCMF Cc oxidation inside the complex.

Under nutrient and oxygen depletion, a dysfunctional mitochondrial ETC and OxPhos can lead to many human diseases, including pathologies such as ischemia and cancer (24). The nexus in both disorders could be the control of mitochondrial ET rate by Cc phosphorylation (23), which has indeed been reported to efficiently fine-tune respiratory rates (81). Even though HIGD-mediated assembly of *Cbc*₁ and CcO into OxPhos supercomplexes preferably occurs under low-glucose conditions, the decrease in both the CcO activity and binding affinity toward Cc₁ and CcO resulting from the enhanced dynamics of Y48pCMF Cc could explain the disruption of Cc channeling inside the *Cbc*₁-CcO supercomplex, thus slowing down the ETC flow (Fig. 7).

Intriguingly, not only respiratory supercomplex formation (60, 61)—whose structure has recently been solved (83, 84)—but also Cc phosphorylation, and the resulting decrease in the ETC rate, could help to keep ROS levels low and guarantee cell survival.

Tyr48-phosphorylated Cc could be targeted as a biomarker of mitochondrial dysfunction for associated pathological states such as ischemia/reperfusion and cancer. Deciphering the details of the phosphorylated Cc-controlled complex network requires accurate structural and dynamic analyses to eventually develop robust therapeutic approaches to foster or silence—as required—the prosurvival action of phosphorylated Cc reported here.

Methods

Methods and associated references are available in *SI Appendix, Methods*. All biological samples were obtained from bacteria, yeast, or human cell extracts, in full compliance with University of Seville Ethical Committee bylaws.

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