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The Charge Distribution on a Protein Surface Determines Whether Productive or Futile Encounter Complexes Are Formed

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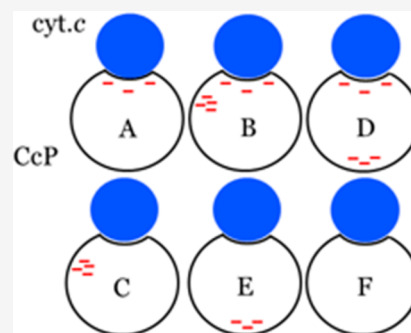
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ABSTRACT: Protein complex formation depends strongly on electrostatic interactions. The distribution of charges on the surface of redox proteins is often optimized by evolution to guide recognition and binding. To test the degree to which the electrostatic interactions between cytochrome *c* peroxidase (CcP) and cytochrome *c* (Cc) are optimized, we produced five CcP variants, each with a different charge distribution on the surface. Monte Carlo simulations show that the addition of negative charges attracts Cc to the new patches, and the neutralization of the charges in the regular, stereospecific binding site for Cc abolishes the electrostatic interactions in that region entirely. For CcP variants with the charges in the regular binding site intact, additional negative patches slightly enhance productive complex formation, despite disrupting the optimized charge distribution. Removal of the charges in the regular binding site results in a dramatic decrease in the complex formation rate, even in the presence of highly negative patches elsewhere on the surface. We conclude that additional charge patches can result in either productive or futile encounter complexes, depending on whether negative residues are located also in the regular binding site.

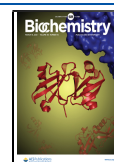


Protein complex formation proceeds through several steps. Freely diffusing proteins can collide and either move away again or form an encounter complex, which may result in the formation of the stereospecific and active complex. The encounter complex is an ensemble of configurations in which the proteins sample each other's surface.¹ Formation is initially driven by electrostatic interactions, but as the proteins get close, hydrophobic interactions can also come into play.^{2–5} If the encounter results in the stereospecific complex, it is called productive; otherwise, it is futile.^{10,11} The rate of formation of the active complex defines the association rate constant (k_a). If complex formation is not optimized, k_a can be 4–5 orders of magnitude lower than the number of collisions, because in most cases the proteins do not collide with binding sides oriented toward each other. Highly optimized protein complexes have k_a values close to the diffusion limit, due to strong electrostatic preorientation.¹² Harel et al.¹⁰ studied complex formation of TEM1- β -lactamase (TEM1) and its inhibitor, β -lactamase inhibitor protein (BLIP). They showed that charge mutations can enhance productive encounter complex formation, thus increasing the association rate without affecting the dissociation rate. However, other mutations modify the encounter complex without affecting the association rate or increase k_a without altering the encounter complex. The fractions of productive and futile encounter complexes can be altered with charge mutations on the proteins surface, even if they are far from the stereospecific binding site. By enhancing or disrupting a “pathway” from the encounter site to the binding site in the stereospecific complex, they can either facilitate the formation of the active complex or

promote the early dissociation of the proteins.¹³ The relation between productive versus futile encounters and such electrostatic pathways on the protein surface was also suggested for the complex of cytochrome P450cam and putidaredoxin.¹⁴ Despite their name, futile encounter complexes can have a role in protein functionality. Recently, it was shown that futile encounter complexes can regulate the activity of a protein complex through the formation of competitive encounters with a third protein.¹⁵

Electron transfer (ET) protein complexes often are highly optimized in complex formation and usually have a high fraction of the encounter state. We selected the complex formed by cytochrome *c* (Cc) (CYC1, UniProtKB P00044) and cytochrome *c* peroxidase (CcP) (CCP1, UniProtKB P00431) to study the importance of the charge distribution on the protein surface in relation to the rate of formation of the stereospecific complex. The protein complex formed by Cc and CcP has been extensively studied and has become a paradigm for biological ET. CcP catalyzes the reduction of H₂O₂ in yeast mitochondria through a cycle of reactions. The crystallographic structure of the complex with Cc published in 1992 represents the stereospecific complex,¹⁶ and a second low-affinity binding

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site is present at ionic strength values below ~ 100 mM.^{17–20} Electrostatic interactions are very important in the association reaction.^{18,20–23} Erman and co-workers^{24–26} measured the rate of association between Cc and several charge-reversal mutants of CcP, showing that, interestingly, not only residues close to the stereospecific binding site but also more distant ones have a role in the association. In a study from our groups, the charge distribution on the surface of CcP was changed strongly by addition of a negative patch on one side of CcP.²⁷ Monte Carlo simulations and paramagnetic relaxation enhancement (PRE) experiments showed that Cc interacts with the added patch, enlarging the surface visited in the encounter complex. The added charges also slightly enhance k_a . To investigate further the degree to which the distribution of charges on the surface of CcP affects the association with Cc, here we analyze four additional mutants of CcP with radically different charge distributions. Monte Carlo simulations show that the mutations alter the encounter complex, attracting Cc to the added negative patches. The association rates were measured through stopped flow experiments for a wide range of ionic strength values. Surprisingly, the added patches turn out to yield either productive or futile encounters, depending on whether the stereospecific site has its charges intact.

MATERIALS AND METHODS

Mutagenesis. *Saccharomyces cerevisiae* CcP (CCP1, UniProtKB P00431) considered the wild type (WT) in this study (CcP_A) contains the C128A mutation to avoid dimerization and the additional sequence MKT at the N-terminus for enhanced production levels.^{28,29} The gene was subcloned into a pET28a(+) vector.³⁰ CcP_B additionally has the K21E, K29E, K90E, and K97E mutations.²⁷ CcP_C contains the same mutations as CcP_B and D35N, E36Q, E119Q, E202Q, E210Q, E215Q, D225N, and E291Q that neutralize the charges of the WT binding site. CcP_D has mutations K75E, K76Q, N79D, Q87E, K91Q, K98Q, and N142D in comparison to CcP_A. CcP_E has the same mutations as mutant CcP_D and the mutations that neutralize the charges of the WT binding site as in CcP_C. CcP_F has only the neutralization mutations for the WT binding site (see Table S1). The DNA constructs were produced by a commercial company and verified by DNA sequencing.

Protein Production. CcP was expressed in *Escherichia coli* BL21 Star (DE3)pLysS (Life Technologies Europe BV, Bleiswijk, The Netherlands). The production and purification were performed as previously described^{28,30,31} with some differences according to the CcP variant purified. Buffers with different ionic strength values were used to build the gradient for elution during anion-exchange chromatography: 50 mM potassium phosphate (pH 5) and 500 mM potassium phosphate (pH 5) for CcP_A and CcP_C and CcP_E; 500 mM KCl was added to the second buffer to elute CcP_B and CcP_D. The purification of CcP_F comprised a precipitation step with 40% (w/w) ammonium sulfate at 25 °C, followed by dialysis against 20 mM potassium phosphate (pH 5) overnight. Finally, the sample was loaded on an SP column; the column was washed with 20 mM potassium phosphate (pH 5) and eluted with 20 column volumes of a gradient to 20 mM potassium phosphate and 1 M NaCl (pH 5). *S. cerevisiae* Cc (CYC1, UniProtKB P00044) was expressed using the iso-1-cytochrome *c* gene in a pUC19-based plasmid and purified following the protocols previously described.^{30,32}

Monte Carlo Simulations. The structures were prepared, and the Monte Carlo docking simulations were performed as described in ref 27. The structures of CcP and Cc from Protein Data Bank (PDB) entry 2PCC¹⁶ were used as a basis for the simulations. The hydrogens of the proteins were added with HBUILD³³ in CHARMM³⁴ and minimized with the CHARMM force field,³⁵ while the heavy atoms were fixed. The hemes within the protein were considered to be in the oxidized state. For each version of CcP (CcP_A–CcP_F), the mutations, described in Mutagenesis, were introduced using Pymol.³⁶ The electrostatic potential was calculated with APBS³⁷ for each CcP mutant separately. The potential was calculated in the presence of 120 mM NaCl at a temperature of 303 K. The Monte Carlo docking simulations were performed with the program suite MCMMap.³⁸ Separate simulations were performed for each CcP mutant considering its respective electrostatic potential. In the simulation, Cc (ligand) moves randomly in the electrostatic potential of CcP (receptor). From each simulation, 5000 randomly chosen encounters were used for further analysis. A Cc:CcP protein pair was considered as an encounter complex if the distance between at least two atoms of each protein was < 3 Å.

Kinetic Measurements. The results of the kinetics measured for the Cc:CcP_A and Cc:CcP_B complexes at ionic strengths from 752 to 122 mM were presented in our previous work.²⁷ The same protocols were followed to measure the reactions between Cc and the other mutants of CcP. The kinetic measurements were performed at ionic strengths 752, 622, 502, 392, 292, 192, 122, and 44 mM on a SX20 stopped flow spectrometer (AppliedPhotophysics, Leatherhead, Surrey, U.K.) with a 1.2 ms dead time at room temperature following the absorbance at 416 nm, which indicates oxidation of Cc.³⁹ Equimolar solutions of Cc and CcP were mixed to a final concentration of 0.5 μ M or 1.0 μ M of both Cc and CcP.

Kinetic Data Analysis. The stopped flow data were fitted to eq 1 as previously described:²⁷

$$\Delta A_{416} = B_0 \Delta \epsilon_{416} \left(1 - \frac{1}{1 + k_{\text{obs}} B_0 t} \right) + C \quad (1)$$

where B_0 is the starting concentration of Cc(Fe^{2+}) and CpdI after mixing, $\Delta \epsilon_{416}$ is the difference in extinction coefficient at 416 nm for oxidized and reduced Cc, $-40 \text{ mM}^{-1} \text{ cm}^{-1}$,^{40–42} k_{obs} (inverse seconds) is the observed rate constant of reduction, t is the time (seconds), and C corrects for the baseline voltage of the spectrometer.

An independent analysis of the stopped flow data was performed through simulations. Kinetic simulations using GNU Octave⁴³ were performed to simulate the experimental measurement of the kinetics obtained from a stopped flow measurement. The script is included in the Supporting Information (Script S1). To do so, reactions 2–4 (see Results) that influence the absorbance at 416 nm during the catalytic cycle were considered. The following differential equations were used to describe the change in the concentration of each species in solution during the cycle:

$$\frac{d[\text{Cc}(\text{Fe}^{2+})]}{dt} = -k(\text{II})[\text{Cc}(\text{Fe}^{2+})][\text{CpdI}] - k(\text{III})[\text{Cc}(\text{Fe}^{2+})][\text{CpdII}] \quad (2)$$

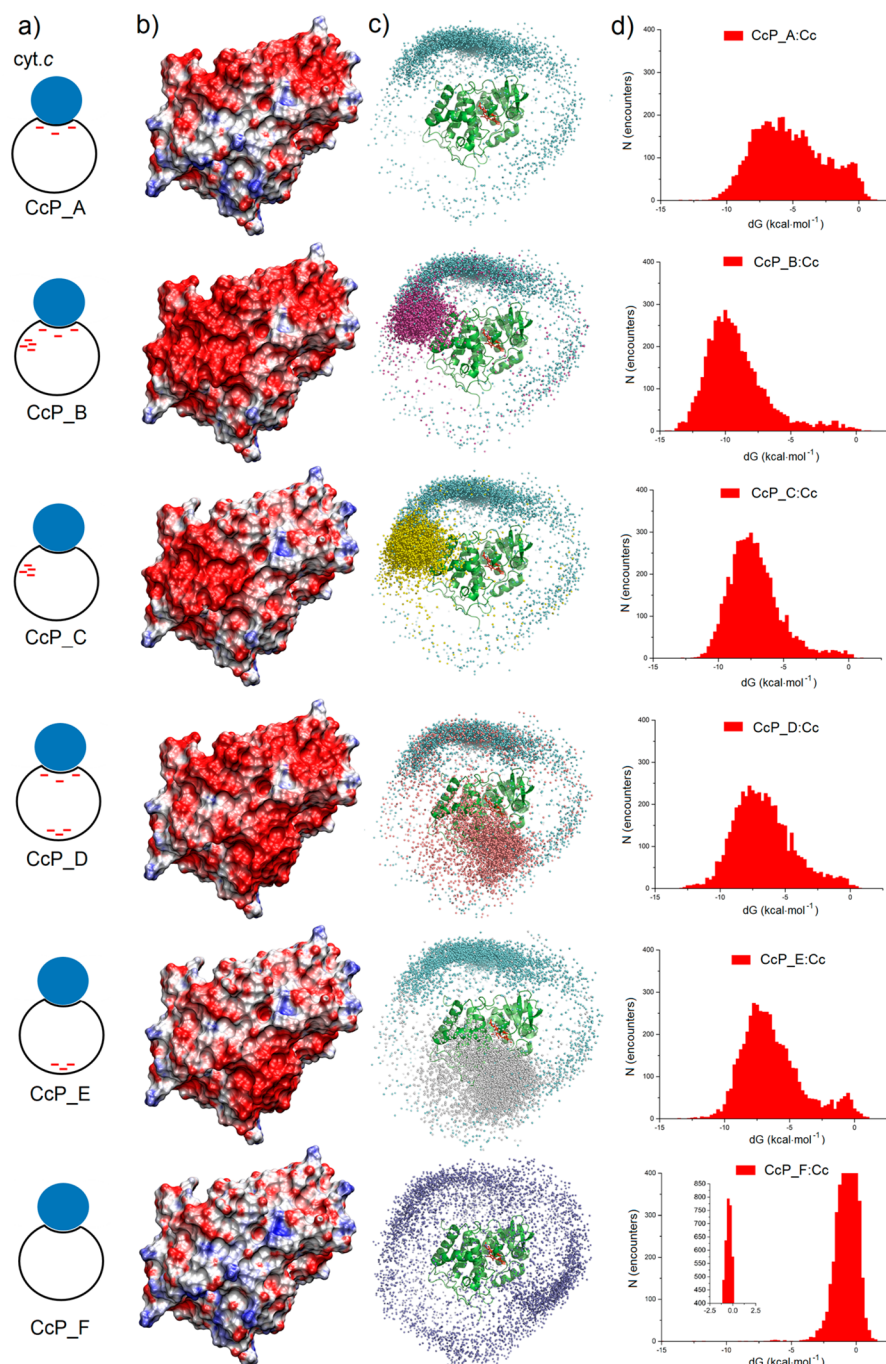


Figure 1. CcP variants. (a) Schematic representation of the complexes formed by Cc and the CcP variants. Cc is represented as a blue circle bound at the stereospecific binding site, and the CcP variants are represented as empty circles with the negative charges on the surface indicated as red dashes. (b) Electrostatic potential plotted on the surface of the CcP variants ranging from -5 (red) to 5 kcal/e⁰ (blue) at an ionic strength of 120 mM. (c) Structure of CcP (green ribbon, red heme) surrounded by the centers of the mass of Cc in the ensemble of encounters of the Cc:CcP_A (cyan), Cc:CcP_B (magenta), Cc:CcP_C (yellow), Cc:CcP_D (salmon), Cc:CcP_E (gray), and Cc:CcP_F (purple) complexes as obtained from rigid body Monte Carlo simulations. (d) Energy distribution of the encounter complexes between Cc and the CcP variants as obtained from rigid body Monte Carlo simulations. The inset shows the entire distribution for CcP_F using a different vertical scale.

$$\frac{d[\text{CpdI}]}{dt} = -k(\text{II})[\text{Cc}(\text{Fe}^{2+})][\text{CpdI}] + k(\text{IV})[\text{CpdII}]^2 \quad (3)$$

$$\frac{d[\text{CpdII}]}{dt} = k(\text{II})[\text{Cc}(\text{Fe}^{2+})][\text{CpdI}] - k(\text{III})[\text{Cc}(\text{Fe}^{2+})][\text{CpdII}] - k(\text{IV})[\text{CpdII}]^2 \quad (5)$$

$$\frac{d[\text{Cc}(\text{Fe}^{3+})]}{dt} = k(\text{II})[\text{Cc}(\text{Fe}^{2+})][\text{CpdI}] + k(\text{III})[\text{Cc}(\text{Fe}^{2+})][\text{CpdII}] \quad (4)$$

$$\frac{d[\text{CcP}]}{dt} = k(\text{IV})[\text{CpdII}]^2 + k(\text{III})[\text{Cc}(\text{Fe}^{2+})][\text{CpdII}] \quad (6)$$

From the simulated concentrations, the stopped flow trace is re-created by summing the contributions of each component to the absorbance at 416 nm following the Beer–Lambert law:

$$A_{416} = \epsilon_{\text{Cc(Fe}^{2+})}[\text{Cc(Fe}^{2+})] + \epsilon_{\text{CpdI}}[\text{CpdI}] + \epsilon_{\text{Cc(Fe}^{3+})}[\text{Cc(Fe}^{3+})] + \epsilon_{\text{CpdII}}[\text{CpdII}] + \epsilon_{\text{CcP}}[\text{CcP}] + C \quad (7)$$

The extinction coefficients at 416 nm are 129.1 mM⁻¹ cm⁻¹ for Cc(Fe²⁺),⁴⁴ 88.8 mM⁻¹ cm⁻¹ for Cc(Fe³⁺),⁴⁴ and the values in Table S2 for CcP and CpdI. The extinction coefficient for CpdII was considered to be the same as for the corresponding CpdI. The baseline correction term C was optimized manually (variable prod in the script in the Supporting Information). The values of *k*(II), *k*(III), and *k*(IV) were independently changed until the simulation was a good fit as judged by visual inspection. The main error for the stopped flow measurement is caused by the error in the concentration of Cc and CpdI. Repeated measurement of the concentration of the samples resulted in a maximum deviation of 20% from the theoretical one. To take this into account in the simulations, every kinetic measurement, consisting of at least three replicates per salt concentration, was simulated at the expected experimental concentration, and at a concentration of CcP that is 20% lower. The rates of the reactions at a particular salt concentration were calculated as the average of the values obtained from the single simulations. The error of the rates at each salt concentration was calculated as the standard deviation between the rates at different protein concentrations. A *t* test was performed to verify whether differences between the association rates of the Cc:CcP complexes are statistically significant. The Bonferroni correction with *n* = 5 was applied when comparing the data of a CcP variant with the other five variants (Tables S5 and S6). The analysis of the data suggests that dismutation of CpdII (reaction 4) is much slower than the other reactions. Nonetheless, the occurrence of this reaction explains the increase in the absorbance at 416 nm visible in kinetic traces obtained at low ionic strengths.

RESULTS

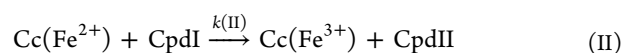
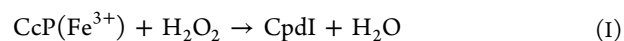
Rearranging the CcP Charge Distribution Creates New Interaction Sites. Electrostatic interactions play an important role in the complex of Cc and CcP and appear to guide Cc to the binding site for fast electron transfer.^{45–48} To test the degree to which the electrostatic interactions are optimized for the association between Cc and CcP, we produced five CcP variants with a different charge distribution on their surface (Figure 1a,b). The CcP mutants were designed with three strategies, creation of an additional negative patch, neutralization of the negative charges in the wild type binding site for Cc, or a combination of both. Wild type CcP is named here CcP_A. In variants CcP_B and CcP_D, an additional negative patch was introduced (change in charge of –8) on the side and the back of CcP, respectively. Variants CcP_C and CcP_E carry the same additional charges as CcP_B and CcP_D and have a +8 charge change around the stereospecific binding site as defined by the structures of the wild type complex determined by crystallography and nuclear magnetic resonance (NMR),^{16,29} resulting in a net charge change of zero. Finally, CcP_F has the mutations only in the stereospecific binding site, resulting a charge change of +8 and rendering CcP close to neutral. Some data on CcP_A and

CcP_B were reported in a previous paper but are included here for the sake of completeness.²⁷

Monte Carlo simulations in which Cc was docked to CcP a large number of times based solely on electrostatic interactions and steric constraints showed that the negative charges introduced on the surface of CcP attract Cc to the new sites (Figure 1c). The new patches strongly attract Cc, in both CcP_B and CcP_D, and this effect is further enhanced in CcP_C and CcP_E, suggesting that in all four mutants the encounter states would have Cc mostly or entirely at the new patches. Note that in the stereospecific complex next to electrostatic interactions, other favorable interactions are present in the WT type complex that are not modeled in these calculations, so it is expected that the shift in the interactions for the entire complex could be smaller than that shown in Figure 1c. For CcP_F, the calculations show a more even distribution of Cc around CcP, as expected for a nearly neutral partner.

The energy distribution of the encounter complexes formed by the WT complex (Figure 1d) ranges from 0 to approximately –10 kcal/mol. The energy of most populated encounters is between approximately –4 and –7 kcal/mol; nevertheless, a significant fraction of the encounters, ~25%, has an energy between –3 and 1 kcal/mol. The encounter complexes formed by Cc in the presence of CcP_B–CcP_E show that the encounters are distributed in more narrow peaks shifted to more negative energy compared to the WT encounter complexes, suggesting that they have a more favorable interaction than in the Cc:CcP_A complex. In particular, the most favorable interactions are formed between Cc and CcP_B, with a peak at –10 kcal/mol. As expected, the least favorable encounters are formed by the Cc:CcP_F complex, which presents a very narrow peak centered on –0.5 kcal/mol.

CcP Turnover Involves Two Reduction Steps. Cc and CcP have high association (*k*_a) and dissociation (*k*_{-a}) rate constants: *k*_a = 10⁸–10⁹ M⁻¹ s⁻¹ at an ionic strength of 200 mM,³⁹ and *k*_{-a} ≥ 6400 s⁻¹.²⁷ The encounter complex represents 30% of the total complex;^{29,47} the affinity is in the micromolar range (*K*_D = 5 μM),⁴⁹ and ET is fast (>50000 s⁻¹).⁵⁰ To establish the effect of charge redistribution on the formation of the stereospecific, electron transfer active complex, the rate of oxidation of reduced Cc, Cc(Fe²⁺), was measured for the different CcP variants. Stopped flow experiments were performed following the work by Miller et al.³⁹ Cc(Fe²⁺) oxidation was followed by measuring the change in absorbance at 416 nm over time in the cycle of reactions that allows CcP to reduce H₂O₂ to water. First, resting state CcP, CcP(Fe³⁺), is reacted with hydrogen peroxide to create compound I (CpdI, reaction I).^{51–54} Then, after mixing with Cc(Fe²⁺), a first molecule of Cc is oxidized to yield compound II (CpdII) and Cc(Fe³⁺) (reaction II). A second Cc(Fe²⁺) molecule can then reduce CpdII to resting state CcP (reaction III).⁵⁵ Reactions II and III both contribute to the change in absorbance at 416 nm. Furthermore, a slow regain of absorbance was observed, which we attributed to dismutation of CpdII into CpdI and resting state CcP (reaction IV).



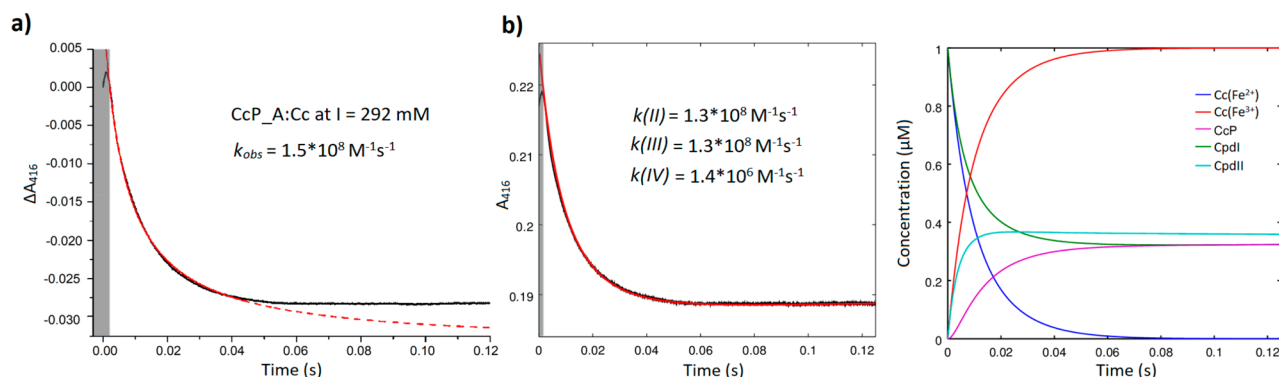
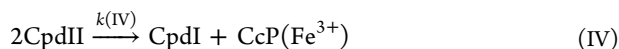
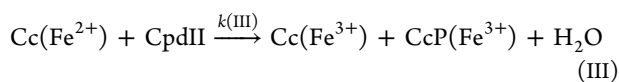
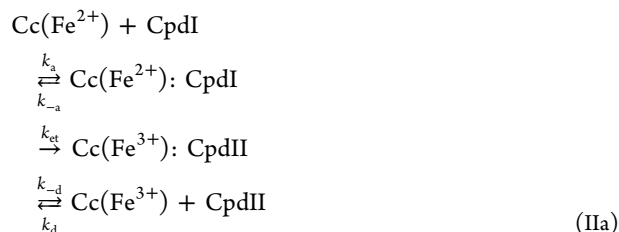


Figure 2. Comparison between fitting and simulation of the reaction performed by Cc and CcP_A at an ionic strength of 292 mM measured at the stopped flow. A gray area covers the first part of the measured data affected by a stopped flow artifact and thus excluded from the analysis. (a) Fitting of the fast decay to eq 1 (see Materials and Methods). The data are shown as a solid black line, and the fit is represented by the red line. The extrapolation of the fit is shown as a dashed line. (b) Simulation (red line) of reactions II–IV to fit the same data (black) as in panel a (left). Concentrations over time of all of the species involved in the Cc:CcP cycle as derived from the simulation (right).



Immediately after mixing, reaction II dominates because the concentration of CpdI is maximal and that of CpdII still zero. Reaction II can be broken up into several steps (reaction IIa):



The observed color change occurs after the electron transfer step (k_{et}), and thus, the observed rate constant in the stopped flow trace, k_{obs} , describes the first two parts of reaction IIa. It can readily be shown that under pseudo-steady state conditions $\left[\frac{\partial \text{Cc}(\text{Fe}^{2+}) : \text{CpdI}}{\partial t} = \frac{\partial \text{Cc}(\text{Fe}^{3+}) : \text{CpdII}}{\partial t} = 0 \right]$ eq 8 holds, giving the relation between the simulated rate $k(\text{II})$ and the association rate constant k_a :

$$k(\text{II}) = \frac{k_a k_{et}}{k_{-a} + k_{et}} \quad (8)$$

To test whether the pseudo-steady state assumption is applicable, the k_{obs} was obtained by fitting the initial part of the curve and the complete curve was simulated using differential equations that describe reactions II–IV, yielding estimates of $k(\text{II})$, $k(\text{III})$, and $k(\text{IV})$. The two methods are illustrated in Figure 2 and Figure S1. Note that the fitting method using the analytical solution given in eq 1 can be applied only at equal concentrations of Cc and CcP and neglects reactions other than reaction II, so it fits only the initial part of the decay (Figure 2). The remaining part is strongly affected by the other reactions. The simulation method covers the entire curve and has the advantage of not being limited to a specific ratio of concentrations between the proteins, enabling us to evaluate the variation in the obtained rate due to concentration errors. This approach results in a

more reliable error estimation for the rate constants. Excellent simulations could be obtained, but it should be noted that under some conditions there is correlation between the obtained rate constants; therefore, the three variables were not completely independent in all data sets (Table S3). Because of this correlation between parameters, we used simulations rather than fittings of the differential equations to avoid ending up in unrealistic local minima or with extreme values. For the analysis of the mutants, the primary interest is in $k(\text{II})$. The values for this rate constant obtained by simulation matched the fitted k_{obs} values (Table S4), indicating that eq 8 is approximated well and $k(\text{II})$ can be used as a lower limit for association rate constant k_a . For WT Cc and CcP, the electron transfer rate (k_{et}) is much larger than the dissociation rate constant (k_{-a}), so the observed rate constant approximates the association rate constant, $k(\text{II}) \approx k_a$.^{12,56} In the remainder of the work, simulations were used for data analysis.

Cc Association Rates Vary Strongly for CcP Variants.

Due to the favorable electrostatic interactions between Cc and CcP, the association rate constant for the WT proteins is high at low ionic strengths and strongly decreases with an increasing salt concentration (Figure 3).^{18,20–23} It is estimated that at high salt concentrations, at an ionic strength of 1.25 M, all of the mutants converge to the same k_a . At a low salt concentration, the complexes formed by Cc and CcP variants A, B, and D have a high k_a , likely to be very close to the diffusion limit (Figure 3). At ionic strength values between 292 and 622 mM (range of 17–25 mM^{1/2} on the horizontal scale in Figure 3), the association rate constant for binding of Cc to CcP variant B is slightly higher than with CcP_A (Table S5). Similarly, CcP_D is slightly faster than CcP_A in the ionic strength range of 392–622 mM (Figure 3 and Table S5). Only at ionic strength values of <122 mM, the association with CcP_D is slightly slower than with wild type CcP. The association rate constant for binding of Cc to CcP variants C, E, and F is strongly reduced compared to that for CcP_A. At ionic strength values of >170 mM (14 mM^{1/2}), the complexes formed by Cc and the mutants of CcP C and E show the same basal association rate as the complex formed with CcP_F. Only a significant difference is present at an ionic strength of 44 mM (7 mM^{1/2}) (Table S5). Under this condition, mutant CcP_C binds faster to Cc than CcP_E and CcP_F.

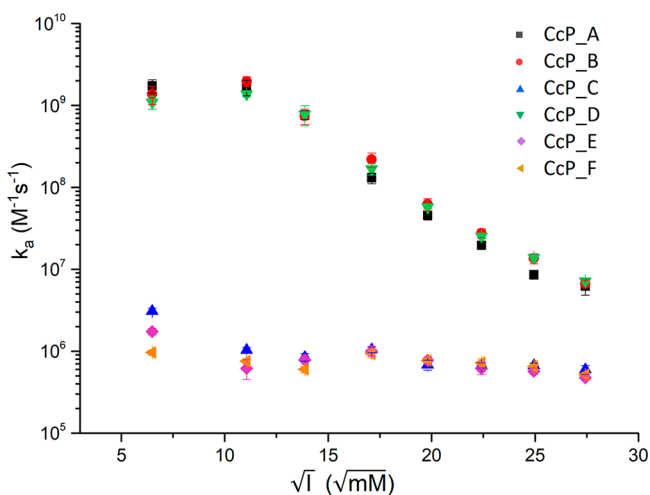


Figure 3. Rates of association (k_a) between Cc and the CcP variants. The k_a values, plotted as a function of the square root of the ionic strength, were obtained from the simulations of the stopped flow kinetics. Errors were calculated as the standard deviation between replicates and simulations performed at different CcP concentrations (see [Materials and Methods](#) for details).

DISCUSSION

To test the importance of the charge distribution on the surface of CcP for the interactions with Cc, five CcP mutants were created with different electrostatic potential and charge distribution on their surfaces (Figure 1a,b). Monte Carlo simulations were performed for the wild type and mutated Cc:CcP complexes. The energy distributions for the Cc:CcP complexes (Figure 1d) show that the charge interactions between Cc and the CcP mutants, apart from the more neutral CcP_F, are more favorable than for the WT complex, in line with the increased overall negative charge on mutants CcP_B and CcP_D. Interestingly, also mutants CcP_C and CcP_E, which have the same net charge as CcP_A, yield somewhat more favorable interactions. The reason could be that in the mutants an artificially highly negative patch is constructed. Often, charges on protein surfaces are compensated by nearby residues to ensure better protein stability. The simulations also demonstrate that these new patches strongly attract Cc, an effect that is enhanced further for the mutants in which the charges in the normal binding site are removed [CcP_C and CcP_E (Figure 1c)]. Thus, it can be expected that the encounter complex of Cc and CcP is shifted toward binding at the new patches, which we demonstrated before for CcP_B using paramagnetic NMR spectroscopy.²⁷ In the Monte Carlo simulations, it is assumed that all residues in the negative patches are charged but the possibility that mutual repulsion leads to increased pK_a values cannot be excluded, rendering the patches with a lower net charge at the experimental pH value of 6.0. Furthermore, the simulations consider only electrostatic interactions and therefore neglect other favorable interactions that could play an important role at the stereospecific binding site. To determine whether productive complex formation, leading to ET from Cc to CpdI, can occur in the mutant complexes, stopped flow experiments were performed and association rate constants determined. The six CcP variants analyzed can be divided in two groups: CcP variants with the wild type charges in the binding site intact (CcP_A, CcP_B, and CcP_D) and variants in which these charges were neutralized (CcP_C, CcP_E, and CcP_F). The rate constants

of association of the first group of CcP variants are similar and strongly dependent on the ionic strength due to the favorable electrostatic interactions between Cc and CcP (Figure 3).^{18,20–23} The second group instead presents a much lower association rate constant that is nearly independent of ionic strength. Clearly, the charges around the stereospecific binding site for Cc are therefore fundamental for the formation of a productive encounter complex and efficient ET, in agreement with previous literature.^{24–26} All variants of CcP bind the heme group, yielding ultraviolet–visible spectra similar to those of WT CcP, and they can all form CpdI and be reduced by Cc. Thus, the mutations do not affect the integrity of CcP. We cannot exclude, however, the possibility that the mutations around the stereospecific binding site subtly affect the protein surface, thus contributing to a reduced level of formation of the ET active complex.

It can be expected that at a high salt concentration, at an ionic strength of ~ 1.25 M, all variants converge to the same basal association rate constant of $\sim 5 \times 10^5$ $M^{-1} s^{-1}$, which is independent of ionic strength. The highest value observed ($k_a = 2 \times 10^9$ $M^{-1} s^{-1}$) shows that the electrostatic interactions can increase the rate 4000-fold compared to that of ET by random collision. The obtained rate constant $k(II)$ is a lower-limit estimate of k_a (eq 8), so the actual k_a could be even larger; however, it seems to approach the diffusion limit, which is suggested by the leveling off of the k_a at a very low ionic strength (Figure 3). Another explanation for such leveling off relates to the balance of monopole–dipole effects on the interactions as a function of ionic strength.^{57–59}

The results indicate that the charges added on the surface of CcP_B and CcP_D do not negatively affect the formation of the stereospecific complex. We previously reported that the enlarged encounter region on CcP_B in fact slightly increases the association rate constant.²⁷ Our hypothesis is that the added charges enhance the electrostatic interactions between Cc and CcP, outweighing the negative effect of interacting at a site more distant from the stereospecific binding site. In other words, the additional charges enhance the probability of productive complex formation and binding at the new patch therefore, by definition, represents a productive encounter. Surprisingly, a similar result is observed with CcP_D. The association rate constants between Cc and CcP_D at ionic strength values between 392 and 622 mM (from 20 to 25 $mM^{1/2}$) are significantly higher than the association measured for the wild type complex (Table S5). The differences are small for both CcP_B and CcP_D but are consistently higher than for CcP_A in that ionic strength range (Tables S5 and S6). The rigid body Monte Carlo simulations show that Cc is strongly attracted by the new negative patch of CcP_D, located on the opposite side of the protein with respect to the stereospecific binding site (Figure 1), and it still results in a slight increase in the rate of productive complex formation at moderate to high ionic strength values. Thus, analogous to CcP_B, we conclude that the positive effect of the extra charges outweighs the remote binding relative to the reaction site, yielding more productive encounters for this variant. At ionic strength values of <122 mM (11 $mM^{1/2}$), the association rate constants between Cc and CcP_D are significantly lower than for the WT complex. This suggests that at low salt concentrations the interaction between Cc and the new negative patch on CcP_D reduces the probability of Cc finding the stereospecific binding site. Similar biphasic behavior with

reduced ET rates at low ionic strengths was reported before, for example, for plastocyanin and plant cytochrome *f*.⁶⁰

Perhaps the most surprising result is that the new patches cannot compensate at all for the loss of the charge interactions at the stereospecific binding site. At ionic strength values of >192 mM (14 mM^{1/2}), the complexes formed by Cc and CcP_C and CcP_E, which have a net charge identical to that of WT CcP, show the same, basal association rate constant as the complex formed with CcP_F, which has a charge change of +8 relative to WT CcP and is overall close to neutral. Therefore, whereas the added charged patches appear to enhance the productive encounter formation in the presence of the charges in the stereospecific binding site, they have no effect on productive complex formation whatsoever if the binding site charges have been neutralized, at least at moderate and high ionic strengths. An effect is observed only at an ionic strength of 44 mM (7 mM^{1/2}), under which condition the association rate constants decrease in the following order: CcP_C > CcP_E > CcP_F. Apparently, at this salt concentration, the electrostatic interactions are strong enough to result in slightly more productive encounters, with the largest effect for the charged patch that is closest to the stereospecific binding site (CcP_C). Similar results were obtained by Harel et al.,¹⁰ who studied the complex formed by TEM1 and BLIP. Creating different mutants of BLIP and using stopped flow association rate measurements in combination with Brownian dynamics simulations, they showed that increasing the number of charge interactions that led to successful binding (fruitful trajectories) increases the rate of association between the two proteins.

CONCLUSIONS

Our results indicate that in the presence of favorable charges in the binding site, the efficiency of active protein complex formation depends more on the overall strength of the electrostatic interactions than on the distribution of charges on the protein surface. Nonetheless, the position of the charges on the protein surface can slightly affect productive complex formation, depending on the ionic strength conditions. At moderate and high ionic strengths, additional charges enhance it, whereas at low ionic strengths, it can be reduced, as seen for CcP_D. The neutralization of the charges at the stereospecific binding site for Cc strongly affects the association of the two proteins, abolishing the electrostatic interactions entirely, as judged from the ionic strength independence of complex formation with CcP_F. Addition of highly charged patches distant from the stereospecific binding site only slightly enhances complex formation at low ionic strengths and as no effect at moderate or high ionic strengths. In this case, although the Monte Carlo simulations clearly show that electrostatic interactions are strong for CcP_C and CcP_E, apparently no trajectories are present for Cc to diffuse from the charged patch toward the stereospecific binding site, rather than to dissociate from CcP.^{10,14} Therefore, in the absence of the negative charges in the stereospecific binding site, the encounters at the new patches have become futile ones.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.biochem.1c00021>.

Example of kinetics at low ionic strengths, list of mutations in the CcP variants, extinction coefficients of the proteins, association rate constants, and script of the kinetic simulations (PDF)

Accession Codes

CYC1, P00044; CCP1, P00431.

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A.D.S., J.M.F., and M.U. performed research and analyzed data. A.D.S., J.M.F., G.M.U., and M.U. wrote the paper.

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Notes

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