ARTICLE

Long distance electron transfer through the aqueous solution between redox partner proteins

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Despite the importance of electron transfer between redox proteins in photosynthesis and respiration, the inter-protein electron transfer rate between redox partner proteins has never been measured as a function of their separation in aqueous solution. Here, we use electrochemical tunneling spectroscopy to show that the current between two protein partners decays along more than 10 nm in the solution. Molecular dynamics simulations reveal a reduced ionic density and extended electric field in the volume confined between the proteins. The distance-decay factor and the calculated local barrier for electron transfer are regulated by the electrochemical potential applied to the proteins. Redox partners could use electrochemically gated, long distance electron transfer through the solution in order to conciliate high specificity with weak binding, thus keeping high turnover rates in the crowded environment of cells.
Electrons flow through biological membranes, yet they are not transported by passive conductive wire structures. Instead, they are carried individually by redox proteins, whose diversity of interactions and electrochemical properties can meet multiple biochemical functions and regulatory requirements. As a consequence, the demands on the electron transfer (ET) capabilities of these proteins are conflicting: their binding must be tight in order to keep ET rates high, but binding should be sufficiently weak to allow a high turnover rate and overall ET efficiency. These demands can be traded off by a stepwise association process between the protein partners: as the distance between redox partners is reduced, an initial encounter complex is formed that leads to a final active complex in which ET occurs between the redox-active sites located within nanometer-scale proximity. The structure of some active complexes between redox protein partners has been revealed by X-ray crystallography. However, substantial differences can be observed in the distance between the active sites of several redox protein partner complexes, which raises the question of whether ET between proteins already occurs while the protein is approaching its partner site. The distance-dependence of inter-protein ET has been studied in various protein partners and mutants that alter the geometry of the wild-type complex, which leads to different distances between active sites, and coupling mechanisms. However, these studies portray the final bound state(s) of the partner proteins, rather than the dynamic events leading to complex formation.

ET studies in proteins have mostly focused on intramolecular transfer because of its relative simplicity. In this case, ET has been shown to proceed by tunneling with a distance decay factor (β) of 11 nm, and is limited to a distance between donor and acceptor of 2.5 nm if a single step is involved. Interprotein ET between diffusing proteins in solution is more complex to interpret than intramolecular ET because of the wide range of geometries that the partners may access. The β value of ET between a protein active site and a metallic electrode has been measured by changing the thickness of molecular monolayers on a metal surface, and more directly by varying the aqueous gap separation between the protein and the metallic probe of an electrochemical scanning tunneling microscope (ECSTM).

Here, we aim to obtain, for a single redox protein pair, a functional readout of the ET rate during the steps involved in the transition from the uncoupled partner proteins to binding, complex formation, and ET. Specifically, the objective is using electrochemical tunneling spectroscopy (ECTS) to measure the current between two ET partner proteins and its dependence with their separation in an aqueous electrolyte.

We observe that current decays along more than 10 nm in the solution, corresponding to distance decay factors that are tenfold lower than those reported for protein-mediated tunneling or for tunneling through organic or water glasses. Molecular dynamics simulations reveal a reduced ionic density and extended electric field in the volume confined between the proteins. In addition, the distance decay factor is regulated by the electrochemical potential applied to the proteins, which highlights the physiological relevance of these results.

**Results**

**Long-distance electron transfer between protein partners.** We chose the ET process between cytochrome c (Cc) and the mitochondrial complex III (CIII), or cytochrome bc1, as a well-known and representative step of the mitochondrial respiratory chain. Cc forms a natural complex with the cytochrome c1 (Cc1) subunit of cytochrome bc1. Although human Cc has not yet been obtained in soluble form, this interaction can be conveniently studied within the well-characterized cross complex between human cytochrome c (hCc) and the soluble domain of plant cytochrome Cc1 (pCc1), which were respectively bound to the ECTS probe and sample electrodes. In particular, the E104C hCc mutant (hereinafter hCc) was chosen because this cysteine is located opposite to the redox-active heme group. In this way, upon covalent binding to the probe electrode, the orientation of the hCc active site is directed toward the solution, and the rotational freedom of the thiol bond facilitates the interaction with its partner. In addition, E104C hCc displays high conductivity in films. The attachment of the soluble domain of pCc1 by means of its N-terminal Cys10, also exposes its active site to the solution and provides rotational freedom. Therefore, the hydrophobic pockets surrounding the heme groups of hCc and pCc1 can face each other and interact during the experiments (Supplementary Figure 1).

ECTS experiments were initially performed at electrochemical potentials, in which pCc1 on the sample electrode is reduced (Up = -200 mV) and hCc on the ECTS probe is oxidized (Up = 600 mV), resulting in a constant positive bias Ubias = Up - Uc = 800 mV. Under these conditions, electrons are transferred from pCc1 to hCc in analogy to physiological ET (Supplementary Figure 1). To obtain each I-z plot, the current between the probe and sample was initially set to 0.4 nA. Upon disconnection of the feedback loop, the piezoelectric scanner holding the probe was retracted at 12 nm/s while the current was recorded. The feedback was then restored and the probe reapproached, allowing subsequent recordings at different locations on the sample. The I-z plots obtained with bare gold sample and probe electrodes show that the current is exponentially reduced below 10 pA in less than 1 nm (gray plots in Fig. 1a; note that a distance-independent faradaic leakage current is measured below a few pA depending on the probe insulation). The corresponding distance decay factors are distributed around β = 9 ± 2 nm⁻¹ (black histogram in Fig. 1b), in agreement with previous studies. Strikingly, I-z plots for sample-bound pCc1 and probe-bound hCc span several nanometers (red plots in Fig. 1a) with a nearly exponential dependence that allows inferring a distance decay factor, which is distributed around 1.5 ± 0.8 nm⁻¹ (red histogram in Fig. 1b).

These β values are directly measured from the current at constant electrochemical potential, during retraction of an accurately calibrated piezoelectric positioner. Thus, they are not subject to models or approximations. In order to rule out protein-related artifacts, we verified that such long-distance currents are specific between pCc1 and hCc. First, we added wild-type human Cc (WT hCc) without C-terminal cysteine to the medium at a concentration that saturates the pCc1 immobilized on the gold sample, in order to inhibit the pCc1-hCc interaction. The decay of I-z curves between pCc1 and hCc (red plots in Fig. 1c) became more pronounced upon addition of WT hCc (blue plots in Fig. 1c), doubling the β value to 3 ± 1 nm⁻¹ and broadening the distribution (blue histogram in Fig. 1d). Similarly, the β value of the non-specific interaction between hCc-funcionalized probe and sample (self-ET) nearly doubled the β of the specific pCc1-hCc interaction (Supplementary Figure 2). The interaction between each protein and a bare gold electrode produced even higher β values (Supplementary Figure 3). Finally, since subsequent I-z retraction curves might be subject to post-interaction effects, such as direct protein contact and protein deformation upon probe retraction, we also recorded I-z curves by approaching the electrodes. Prior to these measurements, the probe was retracted 100 nm from the sample surface (initial set point at 0.4 nA), which is a distance longer than the extended protein length. Although the risk of probe crashing in these I-z recordings was higher, the current increase was again nearly
exponential (Fig. 1e) with a distribution of $\beta$ values similar to that obtained in retraction curves (Fig. 1f). Thus, the observations of Fig. 1 (red plots) correspond to interprotein ET. Note that the distance-dependence of the measured current is assumed to arise mainly from ET between pC$\_1$ and hC, but electron transport between the probe and substrate also involves ET between probe-hC and substrate-pC$\_1$.

Such a gradual current decay is not compatible with tunneling at room temperature, and ET is more likely supported in this case by hopping or some process other than electron tunneling. ET spanning lengths up to 7 nm (corresponding to $\beta$ values below 1 nm$^{-1}$) have been observed in peptides, in ferritin complexes, and in polymers with redox groups, which are believed to act as "bridge" mediators of superexchange tunneling and hopping. In the absence of such redox mediators in the aqueous space between the proteins, the low decay factor (Fig. 1) cannot be accounted for in calculations of pathways including only through-bond and through-space segments, as it is usually assumed from crystal structures of protein complexes.

**Role of the aqueous solution in long-distance electron transfer.**

To account for these results, we hypothesized an ET pathway through the solution where water molecules are tunneling mediators in the constrained aqueous interface between protein partners. In this local region, the dipole resonance of oriented water layers leads to multiple intermediate states or "channels" through which electrons can switch. Similar mechanisms were observed in simulations of ET between metallic electrodes and of cytochrome b$_5$ self-ET. Experimentally, ET enhancement has been linked to a lower energy barrier caused by discrete water layers in the gap between metallic electrodes, and was also reported between a semiconductor and multiheme C$_2$. In the latter study, ET distances increased upon decreasing ionic strength, leading to $\beta = 5$ nm$^{-1}$ at 1 mM, and we observe a
similar trend with even lower values in pCc1–hCc (red plot in Supplementary Figure 4). The influence of the ionic concentration on I–z profiles between the proteins suggests that the conduction process depends on the electrical double layer at the surface. In a low ionic concentration solution, the electric potential extends along several nanometers near the surface of an electronic conductor, within the diffuse ionic layer (Gouy–Chapman layer)\(^22\). Thus, the situation could be similar in the aqueous gap between proteins due to ion depletion near the active sites.

In order to study the distribution of ions at the pCc1–hCc interface, we performed molecular dynamics (MD) simulations of pCc1 and hCc, whose redox-active sites were facing each other at 3 nm separation, in a solvent with explicit water molecules and 50 mM ionic concentration (see Methods section for details). This configuration, anions bind to basic residues in the charged ring of lysines of hCc (K13, K27, K72, K73, K79, K86, and K87) but are otherwise homogeneously distributed between the active sites of the proteins (indicated by the heme groups in Supplementary Figure 5). However, the cationic concentration (Fig. 2a) is significantly reduced at the interface and nearly zero in the proximity of the hCc active site (Fig. 2b–d). The computed electrostatic potential within a 50 mM ionic concentration shows equipotential lines that are extended between the two proteins forming a conduit (Fig. 2d–f).

The concentration of water molecules is homogeneous between the proteins except near the surface of hydrophilic residues (e.g., Q16 and K13 of hCc in Supplementary Figure 6) where it is significantly higher. Bound water molecules are also observed in the crystal structure of many redox protein active sites and act as mediators of their ET pathways\(^23–25\). The importance of water dynamics around key residues in these regions has also been highlighted in MD simulation studies\(^20,26\).

In view of the simulation results, we asked if low ionic concentrations are sufficient to observe long-distance ET between the electrodes, regardless of the presence of the proteins. I–z curves obtained in bare gold at 50 mM phosphate buffer (Fig. 1a and Fig. 1b) show that long-distance ET is not sustained at this buffer concentration. However, I–z measurements of bare gold in pure water (Supplementary Figure 7a) yield decay factors similar to those obtained for pCc1–hCc (Fig. 1a), yet more broadly distributed (Supplementary Figure 7b). The local barrier height (\(\Phi_{\text{local}}\)) can be calculated directly from I–z plots (see Supplementary Information)\(^27\). In bare gold at high ionic concentration (50 mM), barrier values oscillate in the range 0.5–1.2 eV with a spatial periodicity of 0.35 nm corresponding to the diameter of water molecules (Supplementary Figure 8a), as previously reported\(^27\). In contrast, the local barrier along the gap is reduced below 50 meV both between bare gold electrodes in pure water (Supplementary Figure 8b) and between pCc1 and hCc in the 50 mM solution (Supplementary Figure 9), in agreement with studies in metals\(^9\) and proteins\(^21\).

Electrochemical gating of long-distance electron transfer. To further investigate the specificity and physiological relevance of long-distance ET between redox partner proteins, we asked if the local barrier between pCc1 and hCc is altered around their redox potentials at moderate bias, near biological membrane potentials. Strikingly, I–z plots at different probe and sample potentials display a marked potential dependence, with \(\beta\) and \(\Phi_{\text{local}}\) reaching minima near the redox potentials of pCc1 and hCc in stark contrast with the potential-independent \(\beta\) and \(\Phi_{\text{local}}\) between metallic electrodes (Fig. 3 and Supplementary Figure 10). These electrochemical gating experiments\(^28\) were performed at different probe and sample potentials that keep a constant moderate bias of 200 mV and a higher bias of 500 mV (see the Supplementary Information for experimental details and the definition of the electrochemical gate potential, EC gate). Average \(\beta\) values obtained as a function of the EC gate display a minimum of 0.5 ± 0.3 nm\(^{-1}\) at −0.25 V EC gate potential for pCc1–hCc and allow long-distance ET beyond 10 nm (red plot in Fig. 3a and red dots in Fig. 3b). This potential value corresponds to \(U_b\) close to the midpoint redox potentials of pCc1 (0.28 V vs. SSC) and hCc (0.35 V vs. SSC) (Supplementary Figure 10). Thus, the minimum of \(\beta\) and the maximum spatial extent of long-distance ET occur when the Fermi levels of the electrodes and the redox levels of pCc1 and hCc are approximately aligned and electrons can be transferred in quasi-isoelectronic conditions (Fig. 3c). No minimum was observed for bare gold in the same potential range (black dots in Fig. 3b), which highlights the contribution of the redox centers in ET between pCc1 and hCc. No changes in \(\beta\) were observed in gating experiments at more positive probe potentials, far from the hCc redox midpoint (500 mV bias, gray dots in Fig. 3b). As a
The on/off ratio of \( \beta \) in these conditions is around 7 (red and blue dots in Fig. 3b). Thus, the redox state of the proteins strongly regulates the decay rate at moderate probe-sample bias (200 mV), which shows that ET is electrochemically gated in a narrow potential and spatial range. The control of ET rate by small changes in the redox potential suggests a plausible biological role for long-distance ET through aqueous solution.

**Discussion**

The atomic structure of the yeast CIII–Cc complex revealed the “core interface” of Cc, i.e., the minimal site that ensures a geometry suited for ET and a central feature of the interaction between Cc and its functionally diverse redox partners. It was proposed that protein–protein recognition is mediated by the charged rings around the hydrophobic patches and that the function of the latter is to expel the solvent, leaving a layer of water molecules entrapped at the interface in the final interaction state. The dynamic intermediate steps leading to the final complex formation have been difficult to address experimentally and are the subject of debate. In these processes, knowledge on the molecular structure of the solution between the proteins largely relies on models and MD simulations, and new methods are required to elucidate it. On hydrophilic materials surfaces, water is densely packed, strongly oriented, and weakly hydrogen-bonded with its neighbors, whereas it presents low density, slow orientation dynamics, and bulk-like hydrogen bonding next to hydrophobic surfaces. Although water structuring does not extend beyond a few layers in the vicinity of hydrated solutes and open surfaces, hydrogen-bonded water chains and networks have been observed in confined volumes of water channels, enzymes, and ET proteins. Since water has an active role in ET at solid/liquid interfaces, it must be taken into account to calculate the overall ET rate between redox sites. However, considering that charge screening is high in physiological solutions, the molecular structure of the solution between the proteins is similar (red plot in Supplementary Figure 9). The fact that the spatial extent of the current and of the potential profile in I–z curves extends along several nanometers (Fig. 2d), following an exponential dependence as a result of Maxwell–Boltzmann statistics, is around 1 nm at 100 mM, we wondered how the electrical driving force is maintained beyond a few water layers from the open protein surface and leads to the observed long-distance currents (Fig. 1). The MD simulations of Fig. 2 and Supplementary Figure 6 show that the ring of lysines in hCc accumulates anions, and that cations are depleted from the hydrophobic patch in a region that extends between the proteins. Such ionic distribution supports a reduced electrical screening, long \( \kappa^{-1} \), and explains that the potential profile in I–z curves extends along several nanometers (Fig. 2d), in contrast to the abrupt potential drop of the Debye–Hückel length, \( \kappa^{-1} \). We consider how the electrical driving force is maintained beyond a few water layers from the open protein surface and leads to the observed long-distance currents (Fig. 1). The MD simulations of Fig. 2 and Supplementary Figure 6 show that the ring of lysines in hCc accumulates anions, and that cations are depleted from the hydrophobic patch in a region that extends between the proteins. Such ionic distribution supports a reduced electrical screening, long \( \kappa^{-1} \), and explains that the potential profile in I–z curves extends along several nanometers (Fig. 2d), following an exponential dependence as a result of Maxwell–Boltzmann statistics (Gouy–Chapman layer). This behavior is in contrast to the abrupt potential drop of the Helmholtz layer in protein surface regions exposed to high ionic concentration (Fig. 2d).

Although the electronic and physicochemical properties of metal and protein surfaces are very different, I–z recordings between gold electrodes in pure water (Supplementary Figure 7) yield low \( \beta \) and \( \Phi_{\text{local}} \) as observed in experiments with pCc and hCc in a 50 mM electrolyte (Fig. 1, Supplementary Figures 8 and 9). The fact that the spatial extent of the current and of the potential between proteins is similar (red plot in Supplementary Figure 4) indicates that the observed long-distance ET current is driven by the diffuse double layer. In order to account for our experimental observations and MD simulations, we interpret that when pCc1 and hCc approach, a region of reduced ionic concentration is formed in the confined volume between their hydrophobic patches. The weak electrical screening allows the potential profile to extend along several nanometers between the proteins, and to give rise to an electric field all the way (\( E = dU/dz \)). Thus, the Gouy–Chapman layer can act as a charge conduit for long-distance ET through the aqueous
solution. The identity of charge carriers and the role of water in the ET reaction remain to be elucidated. One possibility is that the enhanced electronic conduction is due to ET active small molecules in the vicinity of the proteins, as farther “leaks” throughout the solution would reduce the specificity of ET and cause biological damage.

The direct experimental observation of electrochemically gated, long-distance ET between pCc1 and hCc through the aqueous solution is relevant to understand interprotein ET and its regulation. By molding the ionic distribution between their active sites, these redox protein partners could use long-distance ET to conciliate high speciation. In this regard, several insights could be derived from the experiments described in this work and from other recent measurements highlighting that electrons must be shuttled between the active sites of CIII and CIV. This process could be efficiently bridged by Cc proteins located near the active sites in two sequential long-distance ET steps without requiring slow diffusion and binding–unbinding events.

Methods

Sample preparation. E. coli BL21 (DE3) cells were transformed with pETR1–E104C plasmid to express Homo sapiens E104C hCc in a recombinant way. Protein expression was performed as described in ref.11, for WT hCc. The purification protocol of the E104C hCc mutant was previously described35. The soluble domain of Arabidopsis thaliana pCc1–containing a Cys residue at position 10 that facilitates the protein binding to an electrode—was expressed and purified similarly as in ref. 36, with minor modifications. E. coli BL21 (DE3) cells co-transformed with the plasmids pET-pCc1 and pCc6 were used to produce the soluble form of pCc1. Both E104C hCc and pCc1 samples were dialyzed against 10 mM sodium phosphate, pH between 6.5 and 7.0. Atomically flat Au(111) single-crystal disks of 10 mm diameter and 1 mm thickness (MaTeK) were flame annealed and electrochemically polished prior to use.27 Au(111) electrodes were incubated with a 56.8 M solution of pCc1. ApiezonTM insulated gold (Au wire 0.25 mm in diameter; GoodFellow) ECSTM probes were incubated with a 12.5 M solution of pCc1 and pCc6 used to produce the soluble form of pCc1. Both E104C hCc and pCc1 samples were dialyzed against 10 mM sodium phosphate, pH between 6.5 and 7.0. Atomically flat Au(111) single-crystal disks of 10 mm diameter and 1 mm thickness (MaTeK) were flame annealed and electrochemically polished prior to use.27 Au(111) electrodes were incubated with a 56.8 M solution of pCc1. ApiezonTM insulated gold (Au wire 0.25 mm in diameter; GoodFellow) ECSTM probes were incubated with a 12.5 M solution of pCc1. Protein solutions were used in sodium phosphate buffer 10 mM at pH = 6.5. Incubations were performed at 4 °C overnight.

ECSTM measurements. Distance-tunneling characteristics (I–z) were measured with a PicospFM microscope head and a Picostat potentiostat (Molecular Imaging) controlled by Dulcinea electronics (Nanotec Electrónica). This setup allows adjusting U0 and Uc independently. A homemade electrochemical cell was used in four-electrode configuration, using a 0.25 mm diameter Pt80/Ir20 wire (Advent) as a counter electrode and a miniaturized ultralow leakage membrane Ag/AgCl (SSC) reference electrode filled with 3 M KCl (World Precision Instruments). Electrode potentials were referenced to the reversible hydrogen electrode. Prior to measurements, the electrochemical cell was cleaned with piranha solution (3:1 v/v solution of H2SO4 and H2O2). Caution: piranha solution is a strong oxidizer and a strong acid. It should be handled with extreme care, as it reacts violently with most organic materials. For protein experiments, the aqueous electrolyte solution was sodium phosphate buffer 50 mM (pH 6.5). In the inhibition experiments, after recording the first set of I–z curves for pCc1–hCc interaction, the electrolyte was replaced by wild-type hCc25, 12.5 M in the sodium phosphate buffer. Measurements in pure water (18 MΩ cm−1; 4 ppb TOC, Milli-Q, Millipore) were performed under potential control. All solvents were degassed with nitrogen and filtered with a 0.2 μm diameter sterile filter (Anotop) prior to use. The I–z curves were acquired by setting an initial current set point of 0.4 nA, turning the feedback loop off, and recording the probe current at 12 nm/s and constant bias. Retrace I–z curves were obtained along 15 nm of piezo retraction. A total of 150 curves (20 samples per point, 1024 points) were recorded for each independent experiment. Approach I–z curves were acquired by retracting the piezo 100 nm from the initial current set point and recording the probe current during the approach until saturation was reached (probe current = 10 nA). A total of 30 curves were acquired for each independent experiment (35 samples per point, 4096 points). At least three independent experiments were conducted in each case.

Data processing. Data were acquired using WSxm 4.0 software, treated using a custom-generated MATLAB code (The MATHWORKS, Inc.; code available in the Supplementary Information), and analyzed with OriginPro 8.5 SR1 (OriginLab Corp.). Errors are indicated as standard deviation of the mean.

Molecular dynamics simulations. The initial model of the hCc and pCc1 proteins was built using the PDB structure 3CSX3, using the Multisq program36,37. The two proteins were separated along one axis by 3 nm. The MD simulation was set up employing the program LLeap included in the Amber suite of programs38 and the ff99SB protein force field. All titratable residues were modeled in their corresponding protonation state at pH 6.5, which was further checked by analysis of their intermolecular interactions. The hCc parameters were taken from the AMBER parameters database (http://research.bmh.man.ac.uk/bryce/amber), considering RESP charges for the reduced and oxidized cases from Artiñeh et al.39. The system was solvated with explicit TIP3P water molecules40, and it was neutralized with one sodium atom, in addition to the 33 NaCl molecules to reproduce a concentration of 50 mM. The final system contains 103610 atoms. Molecular dynamics simulations were performed using Amber1440. A restraint on the residues 1–15 (pCc1) and 97–104 (hCc) was applied to model the experimental conditions where Cys10 and Cys104 are attached to the gold probes. A thermal equilibration to 300 K was done prior to the equilibration dynamics in the NPT ensemble for 36 ns. The SHAKE algorithm, with an integration time step of 2 fs, was used. APBS electrostatics calculations41 were performed in the postprocessing stage to obtain the equipotential surfaces. The Volmap Plugin 1.1 was used to obtain average concentration maps. Figures were drawn with Gnuplot42 and VMD43.

Data availability

Additional data that support the findings of this study are available from the corresponding author upon request.

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Author contributions

P.G. and A.L. designed the ECSTM experiments and A.L. performed the ECSTM experiments and data analyses; I.D.-M. and M.A. de la R. discussed the experiments and designed protein production; A.G.-C. produced and purified the proteins; P.G. and C.R. designed the MD experiments and A.N.-H. performed the simulations. P.G. and A.L. wrote the paper with contributions from I.D.-M., M.A. de la R., A.N.-H., C.R., and J.S.

Additional information

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