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PAPER

Fast electron transfer through a single molecule natively structured redox protein†

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The electron transfer properties of proteins are normally measured as molecularly averaged ensembles. Through these and related measurements, proteins are widely regarded as macroscopically insulating materials. Using scanning tunnelling microscopy (STM), we present new measurements of the conductance through single-molecules of the electron transfer protein cytochrome b_{562} in its native conformation, under pseudo-physiological conditions. This is achieved by thiol (SH) linker pairs at opposite ends of the molecule through protein engineering, resulting in defined covalent contact between a gold surface and a platinum–iridium STM tip. Two different orientations of the linkers were examined: a long-axis configuration (SH-LA) and a short-axis configuration (SH-SA). In each case, the molecular conductance could be ‘gated’ through electrochemical control of the heme redox state. Reproducible and remarkably high conductance was observed in this relatively complex electron transfer system, with single-molecule conductance values peaking around 18 nS and 12 nS for the SH-SA and SH-LA cytochrome b_{562} molecules near zero electrochemical overpotential. This strongly points to the important role of the heme co-factor bound to the natively structured protein. We suggest that the two-step model of protein electron transfer in the STM geometry requires a multi-electron transfer to explain such a high conductance. The model also yields a low value for the reorganisation energy, implying that solvent reorganisation is largely absent.

1 Introduction

Long range electron transfer is central to processes essential to life, including photosynthesis, respiration and many enzymatic catalytic events.^{1,2} Electron transfer (ET) proteins are the components responsible for these processes and perform their function through binding, organising and tuning redox active metals and co-factors. To date, most experimental studies on ET through proteins have been achieved by a combination of chemical modification and spectroscopic measurement.^{3,4} These have largely been interpreted in terms of transport through a relatively isotropic dielectric medium, well-approximated by widely applicable parameters,⁵ or a through-bond model in

which the detailed protein structure influences electron transport to a significant extent.^{3,4}

High resolution tools, such as mechanically controlled break junctions, scanning tunnelling microscopy (STM)-based methods and conducting atomic force microscopy (cAFM), have greatly advanced our ability to investigate single molecule conduction and have been widely applied to a range of small organic and inorganic single molecules.^{6–15} However, to apply these tools to measure reliably electronic conductance of structurally complex, and mechanically fragile, single protein molecules, several conditions must be fulfilled. These include (a) effective organisation of protein molecules on a solid surface with sufficient stability against desorbing, (b) good electronic coupling between a protein molecule and external electrodes (*i.e.* electrical contacts to form molecular junctions), (c) control of molecular orientation and (d) conformation relevant to their biological structure and function under the experimental conditions required for measurements. These conditions have only been partially fulfilled in previous attempts.^{16–26} Conductance measurements on proteins by scanning probe techniques have nearly all involved a tunnelling barrier between tip and molecule,^{20–26} or the application of mechanical force to improve the electrical contact between tip and molecule.²³ But recently, we have been able to measure STM conductance of protein

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molecules in air,^{27,28} contacted to *two* metallic electrodes *via* chemical anchoring groups. In the present study we show the first single-molecule electrochemical-STM conductance measurements of a heme protein, cytochrome *b*₅₆₂ (cyt *b*₅₆₂), of controlled orientation, and contacted directly *via* two thiol anchoring groups. By operating in solution under electrochemical control, the potential of the tip and substrate can be independently adjusted with respect to a pseudo-reference electrode in the solution. With fixed tip–substrate potential difference, the molecular conductance was found to depend strongly on the substrate potential, with the conductance of the equilibrium redox state being higher than any other reported molecule of comparable dimensions. The origin of this high conductance for the protein is discussed and related to its electron transfer behaviour.

2 Results and discussion

2.1 Controlled protein orientation, conformation and electrode binding

Cyt *b*₅₆₂ is an ET protein found in the periplasm of the bacteria *Escherichia coli*. The protein binds non-covalently the redox-active iron protoporphyrin IX (heme) *via* two axial ligands. The central iron ion can switch between the Fe²⁺ and Fe³⁺ state, with the redox potential defined by heme–protein interactions.²⁹ Heme binding is also critical to the stability of the protein's native conformation. In the absence of heme (apo-protein) the protein is partially folded and highly dynamic.^{26,29} Using rational protein engineering, pairs of cysteine residues were introduced that allow cyt *b*₅₆₂ to bind through metal–thiol interactions in defined orientations with respect to the electrode surface and an STM tip.²⁷ Two sets of cysteine pairs were introduced to sample the two orientations of the approximately prolate spheroidal cyt *b*₅₆₂: (a) the Cys21–Cys50 pair located at opposing ends of protein's long axis (denoted as SH-LA cyt *b*₅₆₂), leading to a tip-to-surface distance of ~ 5.2 nm (Fig. 1(a), left) and (b) the Cys5–Cys104 pair binding across the short axis of the protein (SH-SA cyt *b*₅₆₂) leading to a tip-to-surface distance of ~ 2.4 nm (Fig. 1(a), right). Cyt *b*₅₆₂ with only the Cys50 mutation (D50C) was constructed as a control to confirm the effects of direct linkage of the cysteine pair protein variants to both tip and substrate.²⁶ All the cysteine mutants bound heme in the same manner as the original wild-type protein, as verified by UV-Visible spectroscopy and hemin titration experiments.²⁷

Protein together with the reducing agent 1,4-dithiothreitol (DTT), which prevents formation of disulphide bonds, was absorbed on a gold surface to form a mixed self-assembled monolayer.²⁶ All the measurements were performed in a pseudo-biological buffer (sodium phosphate, pH 6.2) at room temperature (23 ± 2 °C). These combined experimental conditions create a well-defined environment compatible with the protein retaining its native structure and ET function for single-molecule measurements.²⁶

Cyt *b*₅₆₂ deposition on Au(111) substrates produced species with defined molecular orientations in line with positioning of the cysteine residues and with dimensions consistent with retention of their native structure. STM images (Fig. 1(b)) obtained at different surface locations show well-dispersed protein

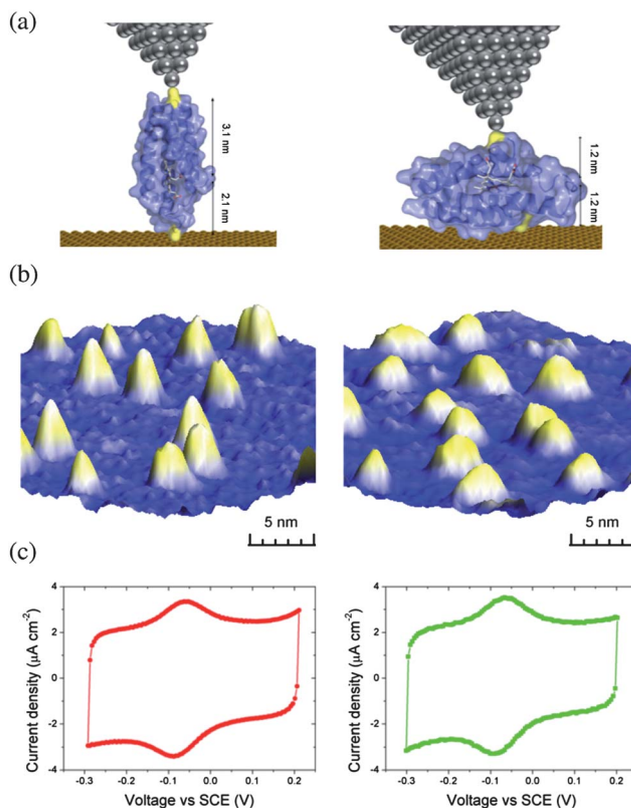


Fig. 1 Self-assembly, STM imaging and long-range protein electron transfer. (a) Schematic illustrations of self-assembly of SH-LA (left) and SH-SA (right) cyt *b*₅₆₂ molecules on a Au(111) surface in the two distinct orientations; (b) 3D view of STM images of cyt *b*₅₆₂ SH-LA (left) and SH-SA (right) recorded in sodium phosphate buffer (10 mM, pH 6.2). $I_t = 35$ pA, $V_b = -0.4$ V, $E_w = -0.10$ V (vs. SCE), z -range = 0.5 nm; (c) cyclic voltammograms of cyt *b*₅₆₂ SH-LA (left) and SH-SA (right) recorded in the same conditions as for STM imaging with a scan rate of 0.5 V s⁻¹.

molecules. The observed lateral dimensions of the imaged single molecules of either cyt *b*₅₆₂ SH-LA (circular, with diameter 3.2 ± 0.5 nm, Fig. 1(b), left) or SH-SA (laterally elongated, with a long axis of 4.5 ± 0.7 nm, Fig. 1(b), right) are in good agreement with crystallographic and NMR data for the holo-protein.^{29,30} (ESI, Fig. S1† presents a more detailed analysis of molecular dimensions.) In contrast, the apo-protein did not retain its native structure on adsorption onto the gold surface (ESI, Fig. S2†), in keeping with the protein being only partially folded and highly dynamic in solution.²⁶ Cyclic voltammograms in Fig. 1(c) demonstrate reversible ET between the protein and the electrode surface, which is essential for precise control of the protein redox state by electrochemical potentials applied to a working electrode or a STM substrate. No Faradaic response was observed in any of the cyclic voltammetry control measurements performed with apo-SH-SA, apo-SH-LA or holo-wild-type cyt *b*₅₆₂ (ESI, Fig. S3 to S6†). The measured equilibrium potential E^0 of -68 ± 2 mV vs. SCE for both cyt *b*₅₆₂ SH-LA and cyt *b*₅₆₂ SH-SA variants is very close to those reported for the wild-type cyt *b*₅₆₂ adsorbed onto hydrophobically modified silver and on the Au(111) surface modified with a peptide linker monolayer.³¹ The electrochemical measurements together with the measured protein dimensions confirm that the protein retains its functional conformation

when adsorbed onto the Au(111) surface *via* an introduced cysteine residue.

2.2 Electrochemical gating of single cyt *b*₅₆₂ protein molecules

The apparent height of individual molecules, obtained from direct STM imaging, was found to depend on redox overpotential. Fig. 2(a) shows a series of STM images of cyt *b*₅₆₂ SH-SA molecules recorded with the bias voltage $V_b = V_t - V_s$ between the tip (voltage V_t) and the substrate (voltage V_s) kept constant while adjusting the substrate potential relative to the reference electrode to control the redox state of the proteins. Apparent height *versus* overpotential $\eta = V_s - E^0$ in Fig. 2(b) clearly shows, as the redox state of the protein molecules is shifted relative to the Fermi energy of the metal substrate, a peak in the apparent height for both the SH-LA and SH-SA variants. The maximum in apparent height is, as observed in related experiments,^{16,17,26} close to the protein equilibrium redox potential $\eta = 0$. No change in lateral dimension was seen. STM-derived heights in proteins are known to be lower than the true height due to a relatively large separation between tip and surface during STM imaging at low set-point current, with only a slight lifting of the tip as it passes over the less conducting molecule.²¹ Variation in protein conductance rather than any significant protein conformational change is the likely dominant factor affecting the apparent height^{9,21} as also confirmed by tunnelling current–distance results discussed below. These results can be described quantitatively with a protein conductance model of ‘two-step’ ET between tip and substrate through the redox level of the protein.^{15–17,26} In Fig. 3 we schematically illustrate the energy alignment of redox level, substrate and tip in the two-step redox processes. The current I_t can be expressed:

$$I_t = 2en_{el} \frac{k_{sm}k_{mt}}{k_{sm} + k_{mt}} \quad (1)$$

where e is the electronic charge, k_{sm} and k_{mt} are the rate constants for ET between redox molecule and substrate (*i.e.* working electrode) and between STM tip and redox molecule, respectively, n_{el} is the number of electrons coherently transferred whilst the redox level relaxes through the energy window eV_b between the Fermi levels of the tip and substrate.³² The term n_{el} describes the fact that, in contrast to the normally encountered case of redox ET between a metal electrode and a molecular electronic level, in the STM junction there is a strong coupling to two electrodes (tip and substrate). A large number of electrons is thus transferred *via* the molecular level between the electrodes before the reduced molecular level has time to reach its relaxed state, which may be below the Fermi level of the other electrode. (We picture just the initially oxidised state for ease of description, but this applies similarly for an initially reduced state.) This transfer can occur because within the energy window eV_b there is a quasi-continuum of available empty states in one metal which can accept an electron, and full states in the other metal which can provide an electron. The rate at which the molecule can be oxidised/reduced by an electrode (which is limited by nuclear reorganisation processes) therefore does not limit the STM current for the strong coupling case, although it does influence the overall current. Using standard expression for rate constants,¹⁶ eqn (1) takes the following form at small overpotentials and bias voltages ($|e\xi\eta|, |e\gamma V_b| \leq \lambda$):

$$I_t = I_0 n_{el} \left\{ \exp \left[\frac{(\lambda + e\xi\eta + e\gamma|V_b|)^2}{4\lambda k_B T} \right] + \exp \left[\frac{(\lambda - e\xi\eta - e\gamma|V_b| + e|V_b|)^2}{4\lambda k_B T} \right] \right\}^{-1} \quad (2)$$

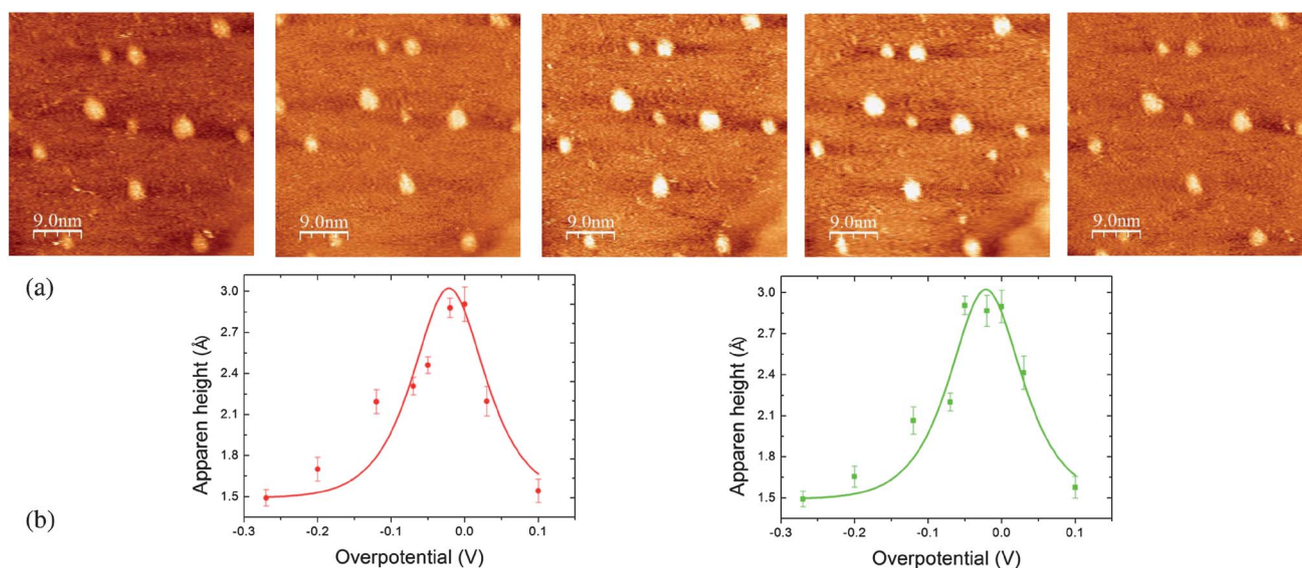


Fig. 2 *In situ* single-molecule mapping of redox state-dependent conductance. (a) Series of STM images for SH-SA molecules obtained at different overpotentials ($I_t = 35$ pA, $V_b = -0.4$ V, $\eta = E_s - E^0 = -210, -100, -10, +80, +130$ mV from left to right, scan area 45×45 nm, z -range 0–1.5 nm). (b) Dependence of apparent contrasts of SH-LA (left) and SH-SA (right) on the overpotentials applied to the working substrate electrode. The solid lines indicate theoretical fitting according to eqn (2) described in the text. Fitted parameters $\xi = 1$, $\gamma = 0.5$, $\lambda = 0.10 \pm 0.03$ eV and $\lambda = 0.15 \pm 0.03$ eV for SH-SA and SH-LA respectively. The measured vertical height of the protein in STM scans does not represent the true height of the protein²¹ due to the low conductance of biological molecules compared with the substrate, and the properties of the surrounding medium.^{22–25}

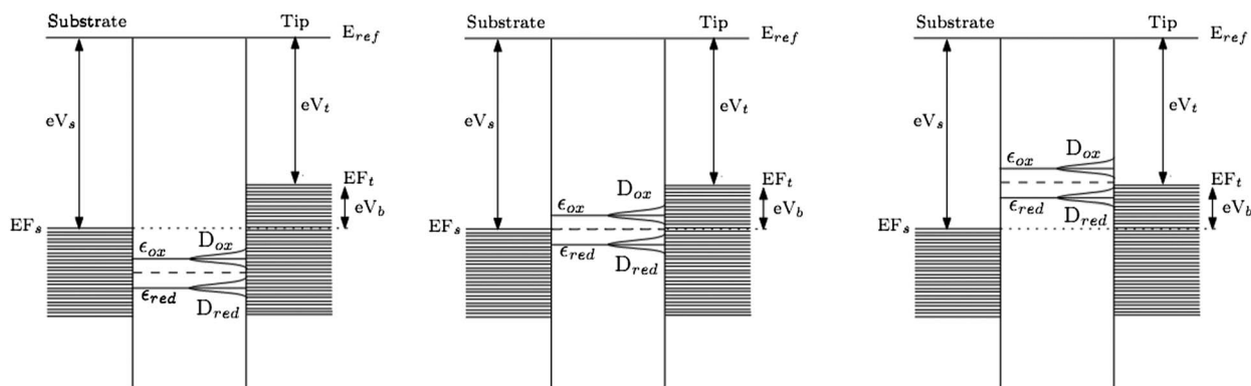


Fig. 3 Energy diagrams describing the sequential two step electron transport mechanism. The tip and the substrate Fermi levels (EF_t and EF_s) are separated by energy equal to eV_b which is kept constant through the experiment. The two Fermi energies EF_t and EF_s can be shifted in parallel with respect to the reference Fermi level tuning the substrate electrode potential. The dotted line, centred at the cyt b_{562} equilibrium potential, indicates the protein redox state. Depending on the applied bias, electrons tunnel from one electrode to the other *via* the redox molecular level. If the protein equilibrium energy is aligned to the substrate Fermi level ($\eta = 0$, middle diagram), electrons transfer first from the tip to a molecular vacant state and, whilst the level relaxes vibrationally below the tip Fermi energy, to the substrate. The probability of resonant tunnelling from one electrode to the other decreases when the molecular equilibrium level is brought below (left diagram) or above (right diagram) the metallic Fermi level by applying a negative and a positive overpotential to the system, respectively. The model shows that, by changing the working electrode potential, the molecular redox states can be modulated in and out of resonance, resulting in increase and decrease of molecular conductance.

where λ is the total nuclear (protein and solvent) reorganisation free energy, η the substrate overpotential, k_B Boltzmann's constant and T the temperature. The terms ξ and γ represent formally the fraction of the substrate potential and bias voltage drops at the site of the heme group redox centre.^{16,33} I_0 is a constant, the most important part being the effective nuclear vibrational frequency of all the nuclear modes reorganised. The number n_{el} (which is proportional to eV_b) is large when the electronic coupling between the molecule and the enclosing electrodes is strong and the adiabatic limit of interfacial ET prevails, as in the present case.

Eqn (2) formally represents well the data from the STM contrast (solid lines in Fig. 2(b)). ξ and γ are not known exactly and are correlated in the confined space of the *in situ* STM tunnelling gap.¹⁶ As an illustration, if the redox center experiences all the substrate potential applied electrochemically ($\xi = 1$) and half of the bias voltage ($\gamma = 0.5$), the reorganisation free energy parameter λ can be obtained from fitting, giving $\lambda = 0.10 \pm 0.03$ eV and $\lambda = 0.15 \pm 0.03$ eV for SH-SA and SH-LA, respectively. Even with correlations or non-ideal values of ξ (< 1), these fitted values of λ represent *upper* limits on the reorganisation energy. Contributions to λ from heme and other porphyrins have been determined experimentally and theoretically to be in the range 0.05–0.15 eV.^{34–36} The low limiting value of λ indicates that reorganisation of the heme occurs during ET but much of the reorganisation of the solvent and of the protein scaffold ($\lambda \approx 0.7$ –1.1 eV)³⁷ is decoupled in the molecular scale solvent confinement of the STM tunnelling gap. We also note that transient spectroscopy measurements inevitably involve solvent reorganisation for surface-bound donors or acceptors.³⁸ In contrast, in the STM measurement electrons are transferred between two free-electron reservoirs (metal substrate and tip, which essentially play the role of donor and acceptor of electrons during electron transfer) and their charge state is fixed by the applied bias. Thus reorganisation of the solvent surrounding the electrodes is not induced during electron transfer.

2.3 High single molecule conductance of cyt b_{562}

STM imaging gives an indirect measure of the electrochemical gating of the current through the protein *via* changes in image contrast. A more powerful approach is to quantify the current through the molecule *versus* applied bias or tip–surface distance. The latter, usually referred to as the I_{t-z} technique, is commonly used for small molecules and allows unambiguous identification of data where a junction is clearly formed between the molecule and both tip and surface.^{9–14} Recently, the I_{t-z} measurements have been applied to extract valuable information on the ET process for the protein azurin²² bound to the surface through a *single* cysteine residue present in the protein, but without a defined interaction with the tip. In our present study we take advantage of the free thiol group to make I_{t-z} measurements which provides a defined covalent tip–molecule interaction and avoids a tunnelling barrier between tip and molecule, in the same manner that most small-molecule studies are performed. Furthermore, we ensure that only a single protein is measured at a time by imaging before and after the I_{t-z} scans.

Molecules were first identified by imaging a small scanned area of 10 or 20 nm², (ESI, Fig. S7†) and the STM tip was then positioned above a protein molecule with small set point tunnelling current (*e.g.* 35 pA at -0.1 V). After disengaging the STM feedback system, the change in I_t was recorded as a function of the tip displacement while the tip was repeatedly moved towards and away from the target molecule. As the tip is withdrawn, a plateau in the measured current may be observed in the I_{t-z} curve just prior to losing contact with the molecule, at which point the current decreases rapidly (Fig. 4(a), left and middle). No current plateaus were observed for D50C proteins having just one cysteine residue, bound to the Au surface (Fig. 4(a), right) or for I_{t-z} measurements away from proteins where the surface is only populated with 1,4-dithiothreitol molecules. Conductance histograms were constructed, without selection, from I_{t-z} data and are shown in Fig. 4(b). Approximately 20% of the I_{t-z} traces

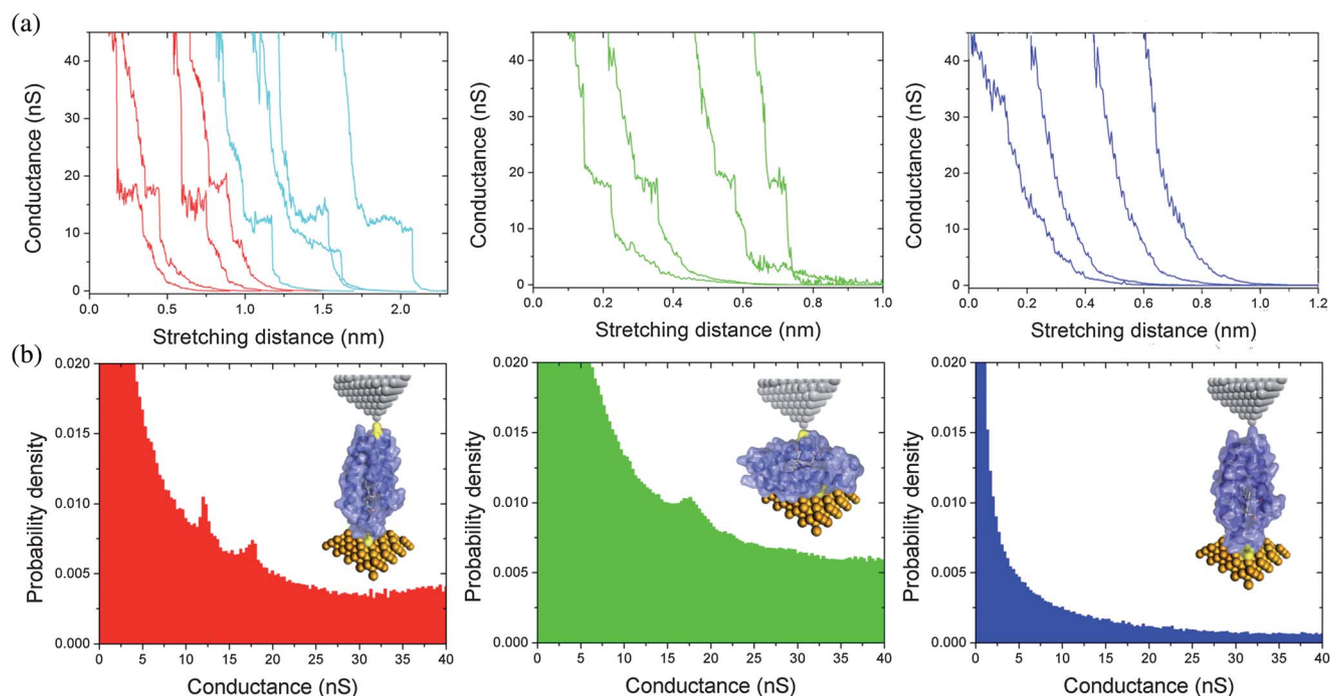


Fig. 4 Direct I_t - z measurements of protein conductance based on single-molecule junctions. (a) Representative I_t - z curves for SH-LA (left), SH-SA (middle), D50C molecules (right). (b) All-data histograms of molecular conductance. Each histogram was constructed from I_t - z curves on 12 SH-SA molecules (200 on each with a total of 2400 curves) and 11 SH-LA molecules (2200 curves) and 10 D50C molecules (2000 curves). No data selection was employed, but each I_t - z trace was weakly pre-smoothed as described in the ESI,[†] where histograms based on un-smoothed traces are also presented (ESI, Fig. S9[†]). None of the D50C curves showed any evidence of plateaus. The measurements were performed in 10 mM phosphate buffer (pH 6.2) at a fixed tip-substrate bias voltage of -0.1 V, and at the equilibrium redox potential of the protein molecules (*i.e.* $\eta = 0$ mV).

were non-exponential and exhibited plateaus with length ranging between 0.05 and 0.3 nm. These current plateaus are assigned to the cleavage of individual protein-tip junctions. Around 70% of traces failed to form a molecular junction and showed a monotonic exponential decay, which represents direct electron tunnelling from the STM tip. The remaining 10% of traces were too noisy to classify. The step lengths observed in the I_t - z traces are small (0.1–0.2 nm) compared with the molecular length, as is expected for these protein molecules since they bind to the surface in defined orientations, rather than lying more nearly flat on the surface as for measurements on conventional larger organic single molecules. In the latter case, there can be a larger range of distance over which the tip binds to one end of the molecule (as the tip is withdrawn from the surface and the angle between molecule and surface is altered) and so lead to a long plateau. One consequence of small step length is that peaks in the histograms are less pronounced.

The conductance histogram for cyt b_{562} SH-LA shows a main peak at 12.0 nS accompanied by a smaller peak at 17.8 nS, while only a single distinct peak at 17.8 nS is observed for cyt b_{562} SH-SA. The observation of two conductance values for cyt b_{562} SH-LA is unlikely to correspond to one and two protein molecules, since the higher conductance value is not simply double that of the lower and imaging of well-separated protein molecules was achieved. Similar features have been observed in several small organic molecular systems^{39–41} where it is known that the contact configuration is a factor in determining conductance.¹⁴ Another possible interpretation²⁷ is that the two conductance values

correspond to the two possible orientations of the SH-LA molecule (Fig. 1(a), left) on the surface, since the iron redox centre of the heme is asymmetrically positioned along the protein's long axis. In contrast, the two possible orientations for cyt b_{562} SH-SA have similar distances³⁰ between the heme centre and either thiol group, see Fig. 1(a). Further experimental and theoretical work is required to test these interpretations.

The I_t - z measurements were also performed at various substrate potentials to observe redox-state dependent conductance. Fig. 5 shows the conductance histograms for cyt b_{562} SH-SA in (a) the fully oxidised form, (b) at the equilibrium redox potential, and (c) in the fully reduced form. It is clear that the protein in its equilibrium state is significantly more conducting than in its fully oxidised and fully reduced form. This strong overpotential dependence of the conductance in I_t - z measurements confirms that the apparent height dependence on overpotential cannot purely be attributed to conformational changes. A comparison of our results with those for selected molecules of similar length is summarised in Table 1. Combined with the results for SH-LA, this suggests that the placement of the iron moiety in the context of the protein plays a crucial role in ET. It is immediately noticeable that cyt b_{562} is highly conductive, even in its oxidised and reduced states. Its conductance is large compared to that for a dsDNA fragment or a conjugated polymer wire of similar length. In particular, the conductance of cyt b_{562} in its equilibrium state approaches 10 times higher than that of a single zinc porphyrin monomer with two thiol contact groups (*ca.* 2.2 nS).⁴³ The conductance values of SH-LA and

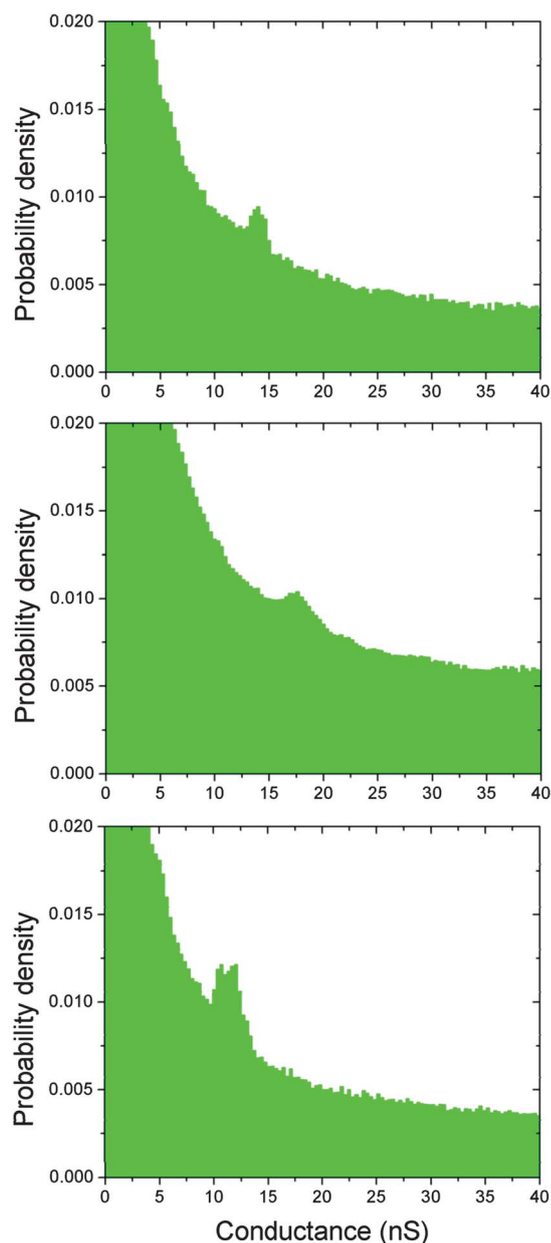


Fig. 5 Single-molecule conductance modulation of cyt b_{562} SH-SA by its redox state. The redox state of the protein was controlled by the working substrate potentials applied: (top) the oxidised form ($\eta = +300$ mV), (middle) the equilibrium state ($\eta = 0$ mV), and (bottom) the reduced form ($\eta = -300$ mV). The measurements were performed in 10 mM phosphate buffer (pH 6.2) at a fixed bias voltage of -0.1 V.

SH-SA cyt b_{562} are around one order of magnitude greater than that measured for azurin, another ET protein,^{18,22,50} having a single cysteine linker to the gold electrode. This is consistent with the observation that adding a second thiol contact between a measured molecule and the STM tip increases the measured conductance by an order of magnitude or more.^{6,51} The somewhat lower cyt b_{562} conductance values measured in air²⁷ of 1.5 and 2.7 nS for the SH-LA and SH-SA molecules respectively might arise from a number of factors. For example, solvent has been reported to change the Au effective work function⁵² and increase single molecule conductance. The protein charge

distribution and reorganisation energy, whose influence on the measured conductance is difficult to estimate, may also be affected. Strong solvent effects have been observed in some molecules.⁴²

Cyclic voltammetry measurements on azurin give ET rates^{53,54} around 10^2 to 10^3 s⁻¹, which is much lower than might be expected on the basis of the STM measurements. However, our EC-STM measurements involve strong coupling to two electrodes, whereas the measured transfer rate for voltammetry is limited by the lowest value of the ET rate between molecule and electrode, between molecule and solution (necessarily involving solvent reorganisation) and the inverse charging time constant of the double layer. Meaningful comparisons require ultra-fast voltammetry.⁵⁵

The nature of this high conductance for cyt b_{562} can be appreciated by the following considerations. Thiol linkers vastly increase conductivity of metal–molecule–metal junctions, and thiolated molecules are now ubiquitous in non-protein single-molecule electronics. The mechanism of the improved conductivity is the stronger quantum mechanical coupling to both tip and substrate of the chemisorbed molecular levels,^{56,57} resulting in a significant broadening ΔE in their energy. Our results suggest that this is the case for the heme level of the proteins, although this requires verification by detailed electronic structure calculations since ΔE depends sensitively on the molecular length, details of the contact and nature of the molecule. From the Heisenberg uncertainty relation $\Delta E \Delta t \sim \hbar$ the lifetime of this state Δt , which we denote as t_{el} , governs the intrinsic ET rate $1/t_{\text{el}}$, as outlined by Datta.⁵⁸ A broadening of around 1 meV, as found for example in DFT calculations⁵⁹ of the porphyrin monomer of Table 1, corresponds to a rate of 10^{12} s⁻¹. A crude estimate based on the β factor of the $I_{\text{t-z}}$ curves⁶⁰ is also consistent with a broadening of 1 and 10 meV, for the LA and SA molecules. There are uncertainties in such estimates, and this value might easily be exceeded.⁹

In terms of the two-step model above, when the relevant redox state lies between the Fermi energies of tip and substrate, electrons are transferred at this rate between molecule and tip/substrate. The number of electrons that are transferred in a single vibrational relaxation passage of the molecular redox state through the energy window between the substrate and tip Fermi levels depends on the nuclear environment. This is characterised by a vibrational relaxation time t_{rel} that includes all the classical vibrational modes of the solvent that are dominated by low frequency relaxation,¹⁶ giving altogether $t_{\text{rel}} \sim 10^{-12}$ s. This value includes a spectrum of values and could be larger if slow protein conformational modes and solvent confinement in the tunnelling gap are also included. If the redox level relaxes slowly enough (large t_{rel}) and the broadening is strong enough, then the ratio $t_{\text{rel}}/t_{\text{el}}$ can become large and many electrons are transferred whilst the level relaxes. The diabatic limit of relatively weak interaction corresponds to $\Delta E \sim 10^{-4}$ to 10^{-3} eV or $t_{\text{el}} \sim (10 - 1) \times 10^{-12}$ s. Larger values take us into the adiabatic range:³² in the adiabatic limit we could take $\Delta E \sim 10^{-2}$ to 10^{-1} eV giving $t_{\text{el}} \sim (10 - 1) \times 10^{-14}$ s. Even for moderately adiabatic transitions the ratio $t_{\text{rel}}/t_{\text{el}}$ is thus likely to assume significant values (orders of magnitude), allowing for the coherent transfer of a large number of electrons in a single *in situ* STM event.

Table 1 Comparison of single-molecular conductance of cyt b_{562} with selected systems^a

Molecule	Length/nm	Conductance/nS	Conditions	Reference
Cyt b_{562} SH-LA (redox protein)	5.2	12.0 (equilibrium state), 5.6 (oxidised), N/A (reduced)	Buffer solution and EC control	This work
Cyt b_{562} SH-SA (redox protein)	2.4	17.8 (equilibrium state), 11.2 (oxidised), 14.3 (reduced)	Buffer solution and EC control	This work
6PTTF6 (redox molecule)	2.5	2.5 (on form), 0.5 (off form)	Buffer solution, EC control	Leary <i>et al.</i> ⁴²
Porphyrin monomer	1.8	2.15	Air	Sedghi <i>et al.</i> ⁴³
Poly-Zn-porphyrin ($N = 4$)	4.5	0.61	Air	Sedghi <i>et al.</i> ⁴³
Short α -helix polypeptide	2.2	1.7	Buffer solution	Scullion <i>et al.</i> ⁴⁴
His(GluLeu)5Cys				
Short polypeptide Cys-Gly-Cys	1.5	0.41	Buffer solution	Xiao <i>et al.</i> ⁴⁵
Cu-binding polypeptide	1.2	1.77	Buffer solution	Xiao <i>et al.</i> ⁴⁵
Cu(Cys-Gly-Cys)				
dsDNA fragments	3.8	10	Salt solution	Hihath <i>et al.</i> ⁴⁶
5'GGAGCCCGAGG3' (11 bp)				
(dA) ₁ 5-(dT) ₁ 5 (15 bp)	5.0	0.51	Air	van Zalinge <i>et al.</i> ⁴⁷
(dG) ₁ 5-(dC) ₁ 5 (15 bp)	5.0	1.2	Air	Ashwell <i>et al.</i> ⁴⁸
Polyethylene-acryl molecular wire	7.0	1.02	Air	Ashwell <i>et al.</i> ⁴⁸
Polyconjugated molecular wires ($N = 1-4$)	3.1 (1-mer) 5.2 (2-mer) 7.3 (3-mer) 9.4 (4-mer)	0.76 0.01 0.01 0.004	Organic solvent	Hines <i>et al.</i> ⁴⁹
Azurin (redox protein)	3.5	1.6 (equilibrium state)	Buffer solution and EC control	Artés <i>et al.</i> ⁵⁰

^a The molecules were selected for their biological relevance or for the dimensions similar to the cyt b_{562} and covalent contact to the electrodes. All the reported conductance values were evaluated by I_t-z experiments. Note that comparison of molecular conductance for various systems poses some difficulties, because (i) many factors (some of which are even unknown) affect molecular conductance even for a given system, and (ii) the measurements of single-molecule conductance for various systems have so far been carried out under diverse experimental conditions such as in air, organic solvents, vacuum or in an aqueous solution, and with or without potential control.

3 Conclusions

In summary, we present the first single-molecule conduction measurement of a protein chemically coupled, with controlled protein orientation, to two enclosing electrodes under pseudo-physiological conditions with electrochemical control. Measured electrochemical equilibrium potentials (-68 ± 2 mV vs. SCE) are consistent with the native cyt b_{562} conformation. Electrochemical gating of the molecular conductance was demonstrated both indirectly from apparent height contrast and directly from the I_t-z curves. The latter measurements showed that cyt b_{562} has notably higher conductance for two controlled molecular orientations than those of comparable synthetic molecules of similar dimensions. The maximum conductance at the cyt b_{562} electrochemical potential is consistent with its suggested role as an electron shuttle in the inter-membrane region, in which it can be oxidised or reduced according to the electrochemical potential of its interacting molecule. Two possible origins of the high conductance are (i) the solvent contribution to the reorganisation free energy is notably reduced, the main contribution being attributed to the heme, (ii) the strong electronic coupling due to the double metal-sulfur bond gives strongly adiabatic ET at both contacts and a multi-electron transfer *via* the heme redox level. The observation of ET in which solvent reorganisation is significantly reduced calls for a new systematic study of reorganisation energy in redox proteins, complementing traditional photoexcitation experiments of a surface-bound donor group. The methods developed here hold promise for effective new ways for single-molecule protein ET studies broadly, with prospects for incorporation of single ET protein molecules in its native

conformation in an external electronic circuit for future nano-scale bioelectronic devices.

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