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1 **Single-molecule study of redox control involved in establishing the** 2 **spinach plastocyanin-cytochrome *b₆f* electron transfer complex**

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6

7 **Abstract**

8 Small diffusible redox proteins play a ubiquitous role in bioenergetic systems, facilitating electron
9 transfer (ET) between membrane bound complexes. Sustaining high ET turnover rates requires that
10 the association between extrinsic and membrane-bound partners is highly specific, yet also sufficiently
11 weak to promote rapid post-ET separation. In oxygenic photosynthesis the small soluble electron
12 carrier protein plastocyanin (Pc) shuttles electrons between the membrane integral cytochrome *b₆f*
13 (*cytb₆f*) and photosystem I (PSI) complexes. Here we use peak-force quantitative nanomechanical
14 mapping (PF-QNM) atomic force microscopy (AFM) to quantify the dynamic forces involved in
15 transient interactions between cognate ET partners. An AFM probe functionalised with Pc molecules
16 is brought into contact with *cytb₆f* complexes, immobilised on a planar silicon surface. PF-QNM
17 interrogates the unbinding force of the *cytb₆f*-Pc interactions at the single molecule level with
18 picoNewton force resolution and on a time scale comparable to the ET time *in vivo* (ca. 120 μs). Using
19 this approach, we show that although the unbinding force remains unchanged the interaction
20 frequency increases over five-fold when Pc and *cytb₆f* are in opposite redox states, so complementary
21 charges on the *cytb₆f* and Pc cofactors likely contribute to the electrostatic forces that initiate
22 formation of the ET complex. These results suggest that formation of the docking interface is under
23 redox state control, which lowers the probability of unproductive encounters between Pc and *cytb₆f*

24 molecules in the same redox state, ensuring the efficiency and directionality of this central reaction in
25 the 'Z-scheme' of photosynthetic ET.

26 **Introduction**

27 Photosynthetic and respiratory electron transfer (ET) reactions sustain virtually all life on Earth. Small,
28 water-soluble redox proteins play a ubiquitous role in biological ET by shuttling electrons between
29 generally larger membrane integral redox complexes (Moser et al., 1995). The interactions between
30 the soluble protein and its membrane-bound partner must balance conflicting requirements: they
31 must be highly specific to facilitate efficient ET, yet they must also be sufficiently weak to allow rapid
32 post-ET separation in order to sustain a high turnover rate (reviewed in Hope 2000; Ubbink, 2009).
33 Somehow therefore, the forces that brought the ET complex together initially must be reversed
34 following the sub-millisecond ET event to ensure their rapid dissociation, yet how this is achieved
35 remains poorly understood (Ubbink 2009). One particularly well studied small diffusible redox protein
36 is plastocyanin (Pc), found in oxygenic photosynthetic systems in cyanobacteria, plant and algal
37 chloroplasts, which transfers electrons from cytochrome *b₆f* (*cytb₆f*) to photosystem I (PSI) (reviewed
38 in Gross 1993; Hope 2000). The redox active co-factor in Pc is a copper atom, which cycles between
39 +1 and +2 oxidation states (Haehnel et al., 1980). Pc accepts an electron from the *c*-type haem of
40 cytochrome *f* and delivers it to the P700 special-pair chlorophylls of PSI. To achieve this transfer,
41 oxidised Pc must diffuse over distances up to 250 nm between these two membrane integral
42 complexes, through the narrow (ca. 10 nm wide) protein-crowded thylakoid lumen (Haehnel et al,
43 1984; Kirchhoff et al., 2011).

44 Most current models suggest that soluble redox proteins and their membrane integral partners
45 associate in a stepwise manner, with the first step involving an initial encounter complex steered by
46 long-range electrostatic interactions. The encounter complex is generally thought to be the prelude
47 to the formation of the productive or active ET complex in which short-range hydrophobic interactions
48 between aromatic and non-polar residues surrounding the redox co-factor binding sites are important

49 (Gong et al., 2000; Hart et al., 2003). The effect of the encounter complex is to bring the ET partners
50 together in an orientated state that minimises the number and scale of rotational and translational
51 movements of the partners required to find the optimum conformation for productive ET (Ubbink
52 2009). Factors that sustain rapid ET have been investigated by structural and mutational studies, which
53 have highlighted the importance of complementary electrostatic interactions between the acidic
54 patch of residues on Pc (D42, E43, D44, E45, E59, E60, D61) and a basic patch of residues on *cyt f* (K58,
55 K65, K66 and K187), as well as hydrophobic patches surrounding both the haem and copper cofactors
56 on *cyt f* and Pc, respectively (Modi et al., 1992b; Lee et al., 1995; Soriano et al., 1996; Sigfridsson,
57 1998; Hope, 2000; Illerhaus et al., 2000; Ueda et al., 2012). The importance of the encounter complex
58 for ET rates is readily demonstrated by their dependence on the ionic strength of the aqueous medium
59 (e.g. Qin and Kostic 1993; Meyer et al., 1993; Illerhaus et al., 2000). At high ionic strength the ET rate
60 declines since the attractive electrostatic interactions are screened. Nuclear magnetic resonance
61 (NMR) studies of biological ET complexes show that the encounter complexes are characterised by
62 very small chemical shift perturbations spread out over relatively large areas of the proteins; thus, the
63 interactions that establish ET complexes are highly dynamic and lack a single well-defined
64 organisation, with the electrostatic interactions remaining fully-solvated and any salt-bridges being
65 mediated by intervening water molecules (Ubbink and Bendall 1997; Crowley et al., 2002; Worrall et
66 al., 2002; Volkov et al., 2005; Ueda et al., 2012). Molecular dynamics simulations of the cytochrome
67 *c₂* (*cyt_{c2}*) - cytochrome *bc₁* (*cyt_{bc1}*) interaction similarly suggest a ‘fuzzy’ encounter complex with
68 multiple conformational states that bring the redox co-factors into close proximity (Singharoy et al.,
69 2016). Moreover, this study suggested that the majority of the stored redox energy is expended on
70 partner recognition and binding rather than on driving electron transfer (Singharoy et al., 2016).

71 While the ensemble studies described above have been useful in defining the general characteristics
72 of ET complexes, the averaging involved obscures the heterogeneity inherent within the system.
73 Ideally it would be possible to quantify the forces involved in the putative encounter and productive
74 complexes at the single-molecule level and to understand their temporal, ionic strength and redox

75 dependence. Recently, we used the pico-Newton force sensitivity and nanometre spatial resolution of
76 the AFM, specifically PF-QNM, to quantify the unbinding force between the soluble redox protein cyt
77 c_2 attached to the scanning probe and its ET partner, the reaction centre (RC) - light harvesting 1 (LH1)
78 complex from *Rhodobacter (Rba.) sphaeroides* immobilised on a gold substrate (Vasilev et al., 2013).
79 In another such PF-QNM study we mapped the organisation of the $cytb_{6f}$ complex in spinach
80 thylakoids using its interaction with Pc attached to the AFM probe as a spatial marker (Johnson et al.,
81 2014). However, the statistics of this latter study were necessarily limited by the relatively small
82 number of membrane-embedded $cytb_{6f}$ molecules that could be measured. In the present study we
83 use purified, surface-immobilised $cytb_{6f}$ complexes, which allowed a more detailed investigation of
84 the redox and salt dependence of the Pc- $cytb_{6f}$ interaction at the single-molecule level.

85

86 **Results**

87 **Single molecule force spectroscopy of the interaction between cyt b_{6f} and Pc.**

88 Pc and dimeric $cytb_{6f}$ complexes were purified from spinach (see Supplementary Figs. 1-3). For these
89 experiments the $cytb_{6f}$ was pre-reduced using sodium ascorbate and Pc was pre-oxidized using
90 potassium ferricyanide. The dimeric $cytb_{6f}$ complexes were immobilised on a SiOx substrate
91 functionalised with (3-mercaptopropyl)trimethoxysilane (MPTMS) and attached via the short linker
92 sulfosuccinimidyl 4-(N-maleimidomethyl)cyclohexane-1-carboxylate (SMCC) (Supplementary Fig. 4A).
93 Purified Pc was attached to an AFM probe covered in a MPTMS monolayer, using a heterobifunctional
94 10 nm-long polymer linker SM(PEG)₂₄, terminated with *N*-hydroxysuccinimide ester (NHS) for reaction
95 with primary amines (lysines) on the surface of Pc (Supplementary Fig. 4B). The crystal structure of
96 spinach Pc (Xue et al., 1997) reveals that 6 lysine residues (K30, K54, K71, K77, K81 and K95) are
97 available to form covalent NHS-ester linkages with the functional monolayer on the AFM probe, all of
98 which are distal to the Cu-containing active site; thus, coupling of Pc to the AFM probe is unlikely to
99 prevent binding to $cytb_{6f}$ (see Supplementary Fig. 5). In the PF-QNM experiment the Pc-functionalised

100 probe is brought into contact with the *cytb_{6f}* molecules on the surface, whereupon a specific
101 interaction can occur; subsequently the upward movement of the probe fully extends the flexible
102 linker before rupturing the interaction (Fig. 1A). The distribution of immobilised *cytb_{6f}* complexes on
103 the SiO_x substrate is shown in the AFM topology image in Figure 1B, as imaged in 10 mM HEPES pH
104 7.4, 5 mM NaCl and 0.05% GDN. The average 8.0 nm height (Fig. 1C) is consistent with the expected
105 size of the *cytb_{6f}* complex (Stroebel et al., 2003); the lateral dimensions are larger than expected based
106 on the structure due to geometrical tip-convolution effects. The interactions between the *cytb_{6f}*
107 complexes on the substrate and Pc on the probe are visualised in the simultaneously recorded
108 adhesion image (Fig. 1D), which shows the locations of unbinding forces required to rupture the
109 probe-surface interactions. The density of *cytb_{6f}* on the surface is ca. 430 per 500 x 500nm scan
110 window and the pixel density was 128 x 128; given that each pixel covers ca. 4 nm and the ~11 x 8 nm
111 dimensions of *cytb_{6f}* (Figure S3C), each complex would be contacted roughly 9-12 times during the
112 imaging process). The force-curve repetition rate was 0.5 KHz for these experiments, which
113 corresponds to a dwell-time of the AFM probe on the surface of ca. 500 μs, slightly longer than the
114 *cytb_{6f}*-Pc ET time of 70-130 μs (Haehnel et al., 1980; Delosme et al., 1991). Overlaying the topographic
115 (Fig. 1B) and adhesion images (Fig. 1D) reveals a good correlation between the points on the image
116 where high unbinding forces are recorded (pink) and the 8 nm topographic features that correspond
117 to immobilised *cytb_{6f}* complexes (Fig. 1E). The slight offset of the high adhesion force events from the
118 centre positions of the *cytb_{6f}* complexes most likely result from interaction with Pc molecules attached
119 with an offset (i.e. not directly at the apex) to the AFM probe, together with a scan direction artefact
120 during the image acquisition. We were able to quantify these interactions by extracting and analysing
121 the actual force-distance data obtained for each pixel of the adhesion image (see Materials & Methods
122 section). Fig. 1F shows examples of force-distance curves extracted from Fig. 1E; the top two curves
123 (green) are representative of the vast majority of pixels in the image where we observe low unbinding
124 forces comparable to the noise level of the measurement. The middle two force-distance curves in Fig
125 1F (shown in pink) are recorded for some of the pink pixels in Fig. 1E. These curves show a large

126 increase in the unbinding force (250-300pN) centered on a probe-sample separation distance of 10
127 nm, which indicates that the flexible linker attaching Pc to the probe is fully extended before
128 interaction is ruptured, indicating a specific Pc- *cytb_{6f}* unbinding event. In contrast, the bottom two
129 curves in Fig. 1F are recorded on the SiOx surface that lies in between *cytb_{6f}* complexes. They show a
130 similarly high unbinding force but importantly at a probe-sample separation distance close to 0 nm,
131 indicating a non-specific interaction between the bulk surface of the probe and the SiOx surface. This
132 type of analysis allowed us to use the probe-sample separation distance as a selection criterion and
133 to pool only the data specifically related to the interaction between *cytb_{6f}* complexes and Pc on the
134 probe (Johnson et al., 2014). Each dataset was statistically analysed to evaluate the frequency of the
135 occurring interaction events (interaction frequency) as well as the most probable unbinding force (see
136 Materials and methods).

137
138 Fig. 2A presents a detailed statistical analysis of the distribution of the unbinding forces for over 6000
139 separate events, centered on a 10 nm probe sample-separation distance. To further verify that these
140 represent specific Pc-*cytb_{6f}* unbinding events we tested the effect of saturating the available binding
141 sites on the surface immobilised *cytb_{6f}* complexes by injecting free oxidised Pc into the AFM liquid cell
142 at a final concentration of 90 μ M, several times higher than the reported K_D of 25 μ M (Meyer et al.,
143 1993). After approximately 5 min incubation with free Pc new data were recorded with the same
144 imaging parameters. Analysis of the data obtained before and after the blocking of the docking site
145 revealed a nearly 4-fold lower frequency for interaction between the *cytb_{6f}* complex and the Pc
146 attached to the AFM probe (Fig. 2B), but that the distribution of unbinding forces was unaffected (Fig.
147 2A). Given the 25 μ M K_D and 90 μ M concentration of added Pc, the blocking control still allows residual
148 binding events. . A similarly lowered interaction frequency was found previously for the excess Pc
149 control used for the PF-QNM study of *cytb_{6f}* in grana membranes (Johnson et al., 2014).

150 **Changes in binding dynamics between different redox states.**

151 Having established that the unbinding events centered on a 10 nm probe-sample separation distance
152 correspond to *bone fide* Pc-cytb_{6f} interactions, we next assessed the effect of the redox state of the
153 partners by prior incubation with either sodium ascorbate (to reduce) or potassium ferricyanide (to
154 oxidise). To ensure differences in probe preparation did not affect the interaction frequency (e.g. a
155 probe with more Pc attached giving a higher interaction frequency), the initial experiment of reduced
156 cytb_{6f} – oxidised Pc (cytb_{6f}[red] – Pc[ox]) was performed for 5 images, followed by a change in redox
157 state to the given condition. The adjusted frequency was thus a comparison of the new condition to
158 the cytb_{6f}[red] – Pc[ox]. Four different experiments were conducted where cytb_{6f} – Pc were either in
159 complementary redox states; reduced cytb_{6f} – oxidised Pc (cytb_{6f}[red] – Pc[ox]) and cytb_{6f}[ox] –
160 Pc[red], or in the same redox state; cytb_{6f}[ox]–Pc[ox] and cytb_{6f}[red] –Pc[red]. Fig. 3A compares the
161 cumulative binding frequencies for all four redox pairs. It is evident that when the cytb_{6f} and the Pc
162 are in the same redox state, the interaction frequency is substantially lower (by a factor of ca. 5)
163 compared to the cytb_{6f}[red] – Pc[ox] pair, in qualitative agreement with our previous findings on
164 membrane-bound cytb_{6f} (Johnson et al., 2014). The results for the ‘post ET’ state, cytb_{6f}[ox] – Pc[red],
165 showed an equally high (within experimental error) interaction frequency. Due to the low driving force
166 (30mV, Haehnel et al., 1980) for electron transfer in this complex the similar interaction frequencies
167 for cytb_{6f}[ox] – Pc[red] and cytb_{6f}[red] – Pc[ox] likely reflect the reversible electron transfer between
168 tip-attached Pc and surface-attached cytb_{6f} during the dwell time of the tip on the cytb_{6f} complex. In
169 addition the unbinding force distributions presented in Fig. 3B – E also show that the unbinding forces
170 present in the pre-ET (cytb_{6f}[red] – Pc[ox]) and post-ET (cytb_{6f}[ox] – Pc[red]) states were nearly
171 identical, both displaying two lower populations, with most probable unbinding forces (taken from
172 the mean of Gaussian fitting) of 219 ± 64 pN and 311 ± 72 pN for cytb_{6f}[red] – Pc[ox], and 223 ± 63 pN
173 and 313 ± 47 pN for cytb_{6f}[ox] – Pc[red]. The higher harmonics at ca. 400 and ca. 650 pN in Fig. 3B,C
174 represent simultaneous unbinding events between two separate Pc molecules on the probe binding
175 to two binding sites on the cytb_{6f} on the surface (either both sides of the dimer or two independent
176 cytb_{6f} complexes), which is much less likely than individual unbinding events given the low density of

177 *cytb_{6f}* complexes on the surface and of Pc on the probe. Differences observed in the distribution of
178 forces between pre and post ET states are simply the result of different distributions between single
179 and double events. The forces observed for the *cytb_{6f}*[red]–Pc[red] and *cytb_{6f}*[ox]–Pc[ox] states are
180 similar - 216 ± 68 pN and 314 ± 170 pN, and 212 ± 81 pN and 301 ± 155 pN, respectively, but the
181 probability of such interactions is ca. 5 fold lower than for complementary redox pairs (Fig. 3D,E). The
182 higher of the two interaction forces for ET partners in the pre-and post-ET states (311 ± 72 pN and 313
183 ± 47 pN, respectively) corresponds well with our previous study on *cytb_{6f}* in the grana membrane
184 (Johnson et al., 2014), which found a single most probable force of 312 ± 5 pN. The present study
185 identifies another probable unbinding force at a lower value (ca. 220 pN).

186

187 **Effect of ionic strength on interaction frequency.**

188 In order to investigate the origin of the two unbinding forces (ca. 220 pN and ca. 310 pN) further, we
189 measured the interaction of *cytb_{6f}*[red]–Pc[ox] as a function of the ionic strength of the medium. It
190 has been shown previously that the initial steering and formation of the encounter complex is
191 determined by several long-range electrostatic interactions (Modi et al., 1992b; Lee et al., 1995;
192 Sigfridsson, 1998; Hope, 2000; Illerhaus et al., 2000). In the context of our PF-QNM experiments
193 increasing the dielectric constant of the surrounding medium (the imaging buffer in our experiments)
194 should lead to a decrease of the interaction frequency due to a decrease of the Debye length, which
195 in this case reflects the extent of the electrostatic effect exerted by the charged residues at the
196 surfaces of *cytb_{6f}* and Pc. Such an experiment could also provide some more information on the
197 bimodal distribution of unbinding forces, for example in Fig. 2A; it is possible that the lower ca. 220
198 pN peak represents the encounter complex where electrostatic interactions dominate, while the ca.
199 315pN peak represents the more tightly bound productive ET complex where hydrophobic
200 interactions may further stabilize the complex in addition to the electrostatics. Thus, a range of salt
201 concentrations could help to discriminate between two binding states.

202 We conducted a series of PF-QNM measurements using the *cytb₆f*[red] – Pc[ox] combination, while
203 varying the ionic strength of the imaging buffer by increasing the salt concentration from the initial
204 value of 5 mM NaCl up to 250 mM NaCl, thus covering a broader range than the physiologically
205 relevant of salt concentrations (100-200 mM according to Izawa and Good 1966; Staehelin 1976). We
206 note however that there appear to be no reports that measure the ionic strength of the luminal space
207 occupied by Pc *in vivo*. Interestingly, the ratio of the ca. 310 pN and ca. 220 pN peaks in the distribution
208 of unbinding forces histogram in Fig. 4A remained broadly unchanged (following One-way ANOVA
209 examination) over the entire range of salt concentrations, but the interaction frequency decreased
210 significantly at higher salt concentrations (> 50 mM NaCl). This trend is evident in Fig. 4B, where the
211 frequency of interaction is plotted against the salt concentration and the data follow an exponential
212 decay. At the highest concentration of 250 mM NaCl the interaction frequency is decreased by a factor
213 of ca. 5. Since similar pattern was observed for both force populations, the results suggest that the
214 lower force peak is unlikely to arise specifically from the encounter complex.

215

216 **Influence of the surface immobilisation chemistry and comparison to *cytb₆f* in native grana** 217 **membranes.**

218 To find out if the immobilisation chemistry might exert an influence on the measured unbinding forces,
219 in particular the presence of two peaks in the unbinding force histograms, we performed control
220 experiments where the *cytb₆f* complexes were immobilised on a different substrate, epitaxial gold,
221 using a thiol-based chemistry (Supplementary Fig. 4C). This new substrate and attachment chemistry,
222 and the use of mixed monolayers, allowed more control over the surface density of the immobilised
223 molecules. The surface density of the *cytb₆f* complexes on the functionalised epitaxial gold surface
224 was found to be in the range 200 – 300 molecules per μm^2 . A topography image (Fig. 5A) was recorded
225 at a modulation frequency of 0.5 kHz, in imaging buffer (45 mM KCl, 10 mM HEPES pH 7.4) and
226 individual *cytb₆f* complexes can be clearly seen on the gold substrate with an average height of around

227 8 nm (Fig. 5B), consistent with the expected size of the *cytb₆f* complex (Stroebel et al., 2003) and taking
228 into account convolution effects arising from the larger radius of curvature of the functionalised AFM
229 probe. Fig. 5C shows the distribution of unbinding forces for the *cytb₆f*[red]- Pc[ox] interaction. This
230 combination again gave a bimodal distribution of unbinding forces, with the most probable forces of
231 204 ± 49 pN and 307 ± 175 pN. The forces observed for the *cytb₆f*[red] –Pc[red] combination, Fig. 5D,
232 were 212 ± 107 pN and 321 ± 62 pN, while the interaction frequency decreased by a factor of 3. All
233 these results are consistent with the results obtained on SiO_x substrates.

234 Another possible explanation for the presence of the peak at ca. 220 pN could be the orientation of
235 the *cytb₆f* complex when immobilised on the substrate. In the experiments described so far, for Figs
236 1-5, lysine side chains were used to tether the *cytb₆f* complex to the sample surface. This method lacks
237 specificity in terms of presenting the luminal face of the complex to the probe-attached Pc, because
238 there are lysine residues on both sides of the complex. It was assumed in the initial experiments that
239 *cytb₆f* complexes tethered with their stromal face distal to the surface, thus with a hidden Pc binding
240 site, would not interact with the Pc probe.

241 To investigate the influence of *cytb₆f* orientation we tested native grana membranes, where the *cytb₆f*
242 complexes have a single uniform orientation that favours the formation of an ET complex. We
243 performed measurements on grana membranes prepared for AFM imaging as previously described
244 (Johnson et al., 2014) under exactly the same conditions used for the surface-immobilised *cytb₆f* and
245 using the same AFM probes functionalised with pre-oxidised Pc, except the imaging buffer contained
246 divalent Mg²⁺ ions to enhance the immobilization of the grana membrane to the substrate.
247 Simultaneously acquired topographic (Fig. 6A) and adhesion maps (Fig. 6C), were used to quantify the
248 interaction between the *cytb₆f* in the grana membrane and pre-oxidised Pc on the AFM tip. In addition,
249 a control measurement was performed, where an excess of free pre-oxidised Pc in the imaging buffer
250 was used to block the docking site on the *cytb₆f* complex. Comparison of Fig. 6C and D shows a
251 significant decrease in the number of the recorded interaction events in the presence of excess Pc[ox],
252 and statistical analysis of the interaction frequency revealed that excess Pc[ox] caused an expected
253 drop (decrease by a factor of ca. 3, Fig. 6E, F) in the cumulative interaction frequency. Furthermore,
254 there is no indication of a ca. 220 pN peak in the force distribution histogram; the most probable
255 unbinding force was found to be 328 ± 69 pN, in agreement with our results from previous

256 experiments (Johnson et al., 2014). These findings imply that the presence of second peaks (at lower
257 values of ca. 220 pN) in the unbinding force histograms recorded with surface-immobilised *cytb₆f*
258 might be related to poor orientation of *cytb₆f* on the sample substrate.

259

260 **Discussion**

261 We have performed a serial, molecule-by-molecule study of the interaction of Pc with its native
262 electron donor *cytb₆f*. The parameters obtained in bulk, ensemble studies that would describe such
263 an interaction, such as dissociation constants, have no counterpart at the single molecule level, and
264 instead we obtain binding frequencies and unbinding forces in pN. The behaviour of single molecules,
265 rather than the collective billions studied in bulk measurements, are arguably a valid representation
266 of intermolecular processes that occur *in vivo*. Our previous AFM study of the Pc-*cytb₆f* interaction
267 used grana membranes isolated from spinach, adsorbed onto a mica surface (Johnson et al., 2014).
268 Although native membranes have the advantage of consistently presenting the luminal face of the
269 *cytb₆f* complexes to incoming Pc molecules borne on the AFM probe, many thousands of *cytb₆f*
270 complexes are required for statistical analysis. The present study, which uses purified *cytb₆f* complexes
271 immobilised on silicon or gold, makes large numbers of molecules available for our PF-QNM approach
272 and records a much larger number of unbinding events within a given scan area compared to *cytb₆f* in
273 membranes. This approach also ensures that the Pc targets only *cytb₆f* complexes rather than the
274 heterogeneous surface of a native photosynthetic membrane. However, there is a likely mixed
275 orientation of *cytb₆f* complexes, arising from lysine attachment points on both the stromal and
276 luminal faces of the complex, and tethered complexes could pivot around their flexible attachment
277 point on the silicon or gold substrate, presenting a tilted binding site that could lower the frequency
278 of interaction with tip-bound Pc. Nevertheless, our use of purified proteins did produce robust
279 statistics for the unbinding force of ca. 310 pN required to pull apart the *cytb₆f*–Pc complex. There is
280 good agreement with the previous measurements and those in Fig 6 in the present work, also on
281 membranes, which yielded ca. 310 pN (Johnson et al., 2014). Using purified *cytb₆f* complexes

282 dispersed over a planar surface also yielded a second, lower unbinding force (ca. 220 pN), which we
283 attribute to tilted *cytb₆f* complexes pivoting about their tether and poorly oriented for presenting the
284 full complement of surface charges on *cyt f* to the tip-attached Pc (see Fig. 7). Such a transient complex
285 might be expected to be easier to separate, when compared with the fully established *cytb₆f* – Pc ET
286 complex. Also, we cannot exclude the possibility that the ca. 220 pN unbinding force reflects a non-
287 native association of Pc with the stromal surface of *cytb₆f* complexes tethered in an inverted
288 orientation, e.g. via interaction with a putative ferredoxin binding site.

289 The use of Pc with its attachment site on the AFM probe distal to the Pc-*cyt f* contact region steers the
290 Pc toward its binding site on *cyt f*. Once brought into contact, the probe tethered Pc and the *cytb₆f* at
291 the surface have a limited time to form an interaction prior to the retraction of the probe. The Pc
292 docking site on the *cytb₆f* complex is located on the luminal domain of *cyt f*. The initial loose
293 electrostatic interaction is guided by a basic patch comprised of K58, K65, K66 and K187 on *cyt f* and
294 two acidic patches on Pc comprised of residues 42-45 and 59-61 (Hope 2000; Ueda et al., 2012) (Fig
295 7). The ionic strength dependence of the interaction frequency shows that this initial electrostatic
296 encounter shows significant screening at 250 mM NaCl, so the physiologically relevant range of 100-
297 200 mM, even to ca. 350 mM (Izawa and Good 1966; Staehelin 1976; Kaiser et al., 1983) is far from
298 being optimized for *cytb₆f*-Pc binding. Site-directed mutagenesis of the five lysines comprising the
299 putative Pc binding site of *cyt f* in *Chlamydomonas reinhardtii*, roughly corresponding to the Lys
300 residues highlighted in Fig. 7, had only a limited effect on *cyt f* oxidation, much less than expected
301 from *in vitro* mutagenesis studies (Soriano et al 1996; 1998), so electrostatic forces might not play a
302 large role in mediating encounters between *cytb₆f* and Pc *in vivo*.

303 A kinetic study conducted using recombinant Pc and native *cytb₆f* complexes from spinach used site
304 directed mutagenesis to alter Pc residues on the face that encounters the luminal surface of *cyt f*;
305 acidic patches D42, E43, D44, E45 and E59, E60, D61. Y83, A90 and L12 in the flat hydrophobic surface
306 region were also examined, and most of these residues are represented in Fig. 7. This study pointed

307 to an initial encounter where there is a pre-orientation of Pc mediated by transient, long-range
308 electrostatic forces involving the acidic patches of Pc (red in Fig. 7), and complementary basic residues
309 (blue in Fig 7) in *cyt f* (Illerhaus et al 2000). A more recent study used NMR to study the membrane-
310 bound *cytb₆f* and Pc from spinach and concluded that hydrophobic interactions (L12, and A90 on Pc
311 and Y1, Y4, Y160 on *cyt f*) form the more tightly bound electron transfer complex (Ueda et al 2012).

312 The AFM experiments described in the present study, with probe-attached Pc brought into transient
313 contact with surface-tethered *cytb₆f* complexes, is intended to measure the forces at the single
314 molecule level that stabilise the ET complex, and to examine the influence of the oxidation/reduction
315 state of the reactants. Thus, the Pc, descending towards a tethered *cytb₆f* complex must locate its
316 binding site on *cytb₆f* via translational and rotational movements, guided by loose, complementary
317 electrostatic forces. The ca. 500 μ s dwell-time of the AFM probe, longer than the 70-130 μ s *cytb₆f*-Pc
318 ET is sufficient to allow formation of a tight ET complex where the close, hydrophobic contact zone
319 has been established. Our method quantifies the unbinding force, ca. 310 pN, required to disrupt this
320 hydrophobic interaction.

321 We also measure the interaction frequency, a relative measurement of the formation of the Pc- *cytb₆f*
322 complex that is indicative of the association rate of the complex. It is the interaction frequency, rather
323 than the unbinding force, that shows a redox dependency; the unbinding forces required to disrupt
324 the interactions between all reduced/oxidized combinations are similar, and it is just the frequency of
325 interaction that changes. Whereas previous bulk phase measurements of the Pc-*cytb₆f* interaction
326 showed no clear redox dependency (Modi et al., 1992), our previous single-molecule *cytb₆f*-Pc study
327 using membranes suggested some selectivity (Johnson et al., 2014). The present study finds similarly
328 high interaction frequencies for the *cytb₆f* [red]-Pc[ox] and *cytb₆f* [ox]-Pc[red] pairs (Fig. 3). Thus,
329 complementary redox states for the *cytb₆f* and Pc are essential for bringing about a highly probable
330 interaction but once the association is established, likely through short-range hydrophobic contacts,
331 the same level of force is required to disrupt it, regardless of redox states. The PF-QNM experiments

332 therefore show that formation of the docking interface is under redox state control, as also found for
333 the MD simulations of the analogous *cyt bc₁* - *cyt c₂* interaction (Singharoy et al., 2016).
334 Complementary charges on the *cytb_{6f}* and Pc cofactors must contribute to the electrostatic forces that
335 initiate formation of the ET complex. Continued turnover of the *cytb_{6f}* complex will regenerate *cytb_{6f}*
336 [red] and produce a strongly disfavoured *cytb_{6f}* [red]-Pc[red] pair. However, other factors probably
337 contribute to the undocking of Pc[red], including encroaching water molecules destabilising the
338 hydrophobic interface as envisaged for *cyt bc₁* - *cyt c₂* (Singharoy et al., 2016).

339 The thermodynamic driving force for the *cyt c₂*- *cyt bc₁* and Pc-PSI ET reactions is much larger (100-150
340 mV) than that for Pc-*cytb_{6f}* ET (30 mV, Haehnel et al., 1980). It therefore seems likely that the small
341 driving force for the *cytb_{6f}* to Pc ET prolongs the lifetime of the bound state since the reaction is readily
342 reversible and this could provide an explanation for the relatively slow ET rate (ca. 70-130µs)
343 compared to that reported for Pc to PSI (ca. 10 µs) (Haehnel et al., 1980). Interestingly however, the
344 interaction frequencies for the *cytb_{6f}* [ox]-Pc[ox] and *cytb_{6f}* [red]-Pc[red] pairs were around 5 times
345 lower (Fig. 3A). Therefore, once the two proteins have dissociated the arrival of another electron on
346 *cyt f* will significantly lower the probability of *cytb_{6f}*[red] rebinding the just-reduced Pc molecule.
347 Nature likely uses this phenomenon to avoid 'product inhibition', in other words unproductive
348 encounters between Pc and *cytb_{6f}* molecules in the same redox state, ensuring the efficiency and
349 directionality of the electron transport process. Future studies can now focus on testing the roles of
350 specific residues within the binding locus on the unbinding force and interaction frequency.

351

352

353 **Materials and Methods**

354 **Purification of the cytochrome *b₆f* complex**

355 Spinach was purchased from the local market. Thylakoid membranes were extracted as in Dietrich and
356 Kuhlbrandt (1999). Following this, the membranes were diluted to 2mg/ml chlorophyll, with a final
357 concentration of 1% 6-O-(N-heptylcarbamoyl)-methyl- α -D-glucopyranoside (HECAMEG) in 40 mM
358 Tricine pH 8.0, 10 MgCl₂, 10 mM KCl (Reconstitution buffer) for 3 minutes. HECAMEG was used for its
359 ability to selectively solubilise *cytb₆f* (Pierre et al., 1995, Dietrich et al., 1999). The mixture was then
360 diluted 1.4 x and spun at 95,000 x g for 30 minutes. The supernatant was concentrated and applied to
361 a 10-35% sucrose gradient, made in reconstitution buffer, with 0.8% HECAMEG, and centrifuged at
362 125,000 x g for 16 hours. The dark band seen in figure S2A (left) was taken and loaded onto a 5 ml
363 Ceramic Hydroxyapatite (CHT) column (Bio-Rad) equilibrated in 20 mM Tricine pH 8.0, 25 mM
364 HECAMEG. A wash buffer of 100 mM ammonium phosphate pH 8.0 was then used to remove any
365 green colour. The brown band on the column was then eluted using 400 mM ammonium phosphate.
366 The CHT column product was then placed on another 10-35% sucrose gradient in 50mM HEPES pH
367 8.0, 20 mM NaCl, 0.3 mM trans-4-(trans-4'-propylcyclohexyl)cyclohexyl- α -D-maltoside (tPCC- α -M),
368 and centrifuged at 125,000 x g for 16 hours. The resulting band (S2A, right) was extracted then applied
369 to a Superdex™ 200 FPLC gel filtration column equilibrated in 50 mM HEPES pH 8.0, 20 mM NaCl, 0.2
370 mM tPCC- α -M. The single peak from the elution was pooled and concentrated down to 66 μ M using
371 a Merck™ Centriprep® 50K Centrifugal filter. Sucrose was added to a final concentration of 10% as a
372 cryoprotectant. This was then frozen at -80 °C until use.

373 *Cytb₆f* concentration was measured using an extinction coefficient of 20mM⁻¹ (Breyton et al., 1997)
374 for cytochrome b₆ in dithionite-reduced minus ascorbate-reduced difference spectra.

375 **Purification of Plastocyanin**

376 Spinach leaves were briefly blended in 50mM sodium phosphate pH 7.4, 5 mM MgCl₂, 300 mM
377 sucrose. Following this, leaves were filtered through two layers of muslin cloth, and then repeated
378 through muslin and cotton. The solution was centrifuged for 15 minutes at 4000 x g. The pellet was
379 resuspended in 10 mM Tricine pH 7.4, 5 mM MgCl₂ (Buffer 2) and left on ice for 1 minute. This was
380 then diluted 2x with Buffer 2 containing an additional 400 mM sucrose and centrifuged again at 4000x
381 g for 15 minutes. The pellet was resuspended to a chlorophyll concentration of 2 mg/ml in 10 mM
382 HEPES pH 7.6, 5 mM NaCl, 5 mM EDTA, and sonicated for 10 minutes, at 30 second intervals. The
383 solution was then centrifuged at 200,000 x g for 1 hour to pellet any large unbroken material. The
384 supernatant was then applied to 4 x 5 ml GE Healthcare Hi-TRAP Q FF anion exchange columns chained
385 together, equilibrated in HEPES pH 8, 5 mM NaCl. A gradient of 0.005-1 M NaCl was used for elution,
386 with Pc eluting around 200 mM. Pc-containing fractions were identified by the blue colour upon
387 addition of potassium ferricyanide. These fractions were pooled, concentrated in a Vivaspin 3 kDa cut
388 off spin concentrator, and loaded onto a Superdex™ 200 FPLC column, equilibrated with 20 mM HEPES
389 pH 8, 20 mM NaCl. The resulting Pc fractions were pooled, concentrated, and frozen at -80 °C until
390 use.

391

392 **Surface functionalisation**

393 Silicon Wafers and Si/Si₃N₄ AFM probes (Bruker's model SNL or Olympus BioLever Mini AC40) were
394 initially submerged in piranha solution (3:5 v/v H₂O₂:H₂SO₄) for an hour to clean and oxidise the Si
395 surface. Following this, both tips and surfaces were cleaned extensively in H₂O, dried under a stream
396 of argon and left in a desiccation chamber overnight under vacuum to remove H₂O from the surface.
397 Tips and surfaces were then placed into another chamber, which had been purged with argon for 10
398 minutes. 2 x 15 µL (3-mercaptopropyl)trimethoxysilane (MPTMS) in Eppendorf™ 1.5ml lids were
399 placed into chamber, followed by another 5 minute purge. The chamber was then evacuated and left
400 for ca. 24 hours to allow the deposition of the MPTMS self-assembled monolayer. Tips and surfaces

401 were then washed with H₂O to remove any MTPMS not covalently attached to the Si, and dried under
402 a stream of argon. Si wafers were then incubated with 5mM sulfosuccinimidyl 4-[N-
403 maleimidomethyl]cyclohexane-1-carboxylate (SMCC, ThermoFisher Scientific) in linkage buffer
404 (10mM HEPES pH 7.6) for 45 minutes, followed by incubation with 700 nM *cytb_{6f}*. Surfaces were then
405 washed thoroughly in imaging buffer (10 mM HEPES pH 7.4, 10 mM NaCl, 0.05% GDN (w/v)) and stored
406 in imaging buffer until use. Tips were incubated with 1mM (succinimidyl-[(N-
407 maleimidopropionamido)-tetracosaehtyleneglycol] ester (SM(PEG)₂₄, ThermoFisher Scientific), a ca.
408 10 nm long linker for 45 minutes in linkage buffer. Following this they were incubated in 750nM
409 plastocyanin in 20 mM HEPES pH 7.4, 20 mM NaCl. Tips were then washed in imaging buffer and stored
410 in the same buffer until use.

411 Epitaxially grown Au [111] thin layers (PHASIS, Switzerland) were functionalised, as received and
412 without further treatment, with mixed EG₃ ((11-Mercaptoundecyl)tri(ethylene glycol), Sigma-Aldrich)
413 and NH₂-thiol (Sigma-Aldrich) mixed at a ratio of 1:200 (mol/mol)/ self-assembled monolayer. Sparse
414 amines were then reacted with dimethyl suberimidate (DMS, ThermoFisher Scientific) for 45 minutes.
415 Following washing with H₂O the surfaces they were then incubated with 500nM *cytb_{6f}* for 45 minutes.
416 Surfaces were then washed with imaging buffer and stored in the same buffer until use.

417

418 **Atomic Force Microscopy**

419 Atomic force microscopy and DFS measurements were carried out on a Bruker Multimode 8, using
420 Nanoscope version 9.2. Deflection sensitivities were obtained using a clean Si surface to measure the
421 deflection, whilst the spring constants were determined using Bruker's inbuilt thermal tune function;
422 the obtained spring constants for the cantilevers used were in the range 0.08 – 0.14 N m⁻¹. Force
423 measurements were taken using the Bruker Peak Force Quantitative Nanomechanical Mapping (PF-
424 QNM) mode, with the peak contact force being kept between 50 and 250 pN. The modulation
425 frequency was kept at 0.5 kHz, with an image size of 128 pixels², over an area of 500 nm². Imaging was

426 performed in imaging buffer unless stated otherwise. Reduction of both proteins was achieved by a
427 15-minute incubation with 1mM sodium ascorbate, and oxidation by the same incubation with
428 potassium ferricyanide. For redox states; a baseline experiment was performed, placing both proteins
429 in the pre-ET state (*cytb_{cf}* [red]-Pc[ox]), and 5 images were taken using these conditions. The probe
430 and tip were then re-incubated in the conditions given by the experiment, and further images were
431 obtained. As a result, the adjusted frequencies represent a comparison of the new condition to the
432 pre-ET state, as such the pre-ET state itself has an adjusted frequency of 1 (Fig 3A). For salt
433 concentrations, a similar baseline was taken however for 5mM NaCl, and images obtained at other
434 salt concentrations using the same tip were adjusted as such. In addition, the order of these was also
435 changed, so that 5mM NaCl was not always the first experiment performed but was always included
436 to allow the adjusted frequency to be calculated. Performing the experiments in different orders was
437 found to have no effect.

438 **Data processing**

439 All the AFM data was analysed by using Gwyddion v 2.51 (open source software covered by GNU
440 general public license, www.gwyddion.net), Nanoscope Analysis v 1.80 (Bruker), MATLAB 2017a and
441 OriginPro 2016 (OriginLab Corp.) software. Analysis of force curves was performed using a homemade
442 MATLAB script, utilising Bruker's Nanoscope MATLAB tools to open the files directly, and display each
443 force curve that met predetermined parameters**. Manual data selection from these force curves
444 was then performed via MATLAB GUI. The extracted data was then analyzed in Origin. The most
445 probable values for the unbinding forces were obtained from the maximum of the Gaussian fit to the
446 force distribution combined in a statistical histogram. **Data reduction (positive identification of
447 specific rupture events) was based on the identification of rupture events greater than 90pN (for initial
448 experiments, later raised to 150pN), and occurring at tip-sample separation in the range 5 – 50 nm;
449 all force-distance curves not exhibiting these properties were not pre-selected for further review.

450 **Membrane purification**

451 Spinach grana membranes were prepared for AFM imaging according to Johnson et al., (2014).

452

453

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554

555 **Figure Legends**

556 **Figure 1. Single molecule force spectroscopy of the interaction between *cytb₆f* immobilised on a**
557 **SiOx surface and Pc attached to the AFM probe.** (A) Cartoon depicting the principle of the PF-QNM
558 experiment. The AFM probe, functionalised with Pc proteins attached via a flexible 10-nm-long SM
559 (PEG)₂₄ linker, laterally images the topography of the *cytb₆f* complexes immobilised on a SiOx substrate
560 via a tapping motion, while simultaneously recording probe-sample unbinding forces for every pixel.
561 In the first panel one of the Pc proteins attached to the AFM probe binds to a *cytb₆f* complex on the
562 surface; in the second panel the flexible linker is extended to its full 10 nm length as the AFM probe is
563 withdrawn from the surface during the upward part of the tapping motion; in the third panel the force
564 applied has exceeded the protein interaction force and has ruptured the Pc-*cytb₆f* interaction. (B) AFM
565 topograph showing the distribution of *cytb₆f* complexes on the SiOx surface. (C) Examples of height
566 profiles of individual *cytb₆f* complexes; the measured height is ca. 8.0 nm. (D) AFM adhesion image

567 showing areas where adhesion exceeds 100pN (See scale); this comprises both specific or non-specific
568 interactions. (E) 3D rendering of a subsection (250 nm²) of the AFM height and adhesion data, reveals
569 a coincidence between the regions of high unbinding force (pink) and the protrusions corresponding
570 to *cytb₆f* molecules. (F) Examples of force-distance curves representing no interaction (green, top two
571 curves), specific interaction (pink middle two curves) and non-specific interaction (pink, bottom two
572 curves). See results description for details.

573

574 **Figure 2. Distribution of unbinding forces for the Pc-*cytb₆f* interaction and effect of free Pc** (A)
575 Distribution of forces measured for probe-sample separation distances centered on 10 nm, in the
576 presence (cyan) and absence (red) of 90 μM free Pc in the imaging buffer. The cumulative frequency
577 of each histogram was adjusted to reflect the interaction frequency plotted in (B), giving the adjusted
578 frequency. (B) Mean frequency for the number of interactions observed between *cytb₆f* and Pc in the
579 presence (cyan) and absence (red) of 90 μM free Pc in the imaging buffer. Frequency adjusted to buffer
580 sample being a value of 1. Error bars show the standard error of the mean.

581

582 **Figure 3. Redox dependence of the interaction between *cytb₆f* and Pc.** (A) Mean cumulative
583 frequency for the interaction at different redox states (O – Oxidised, R – Reduced) represented as a
584 bar chart. Adjusted frequencies represent the total number of interactions occurring, adjusted to the
585 pre-ET transfer state (*cytb₆f*[red], Pc[ox]). (B-E) Force distributions (most probable forces obtained
586 from the Gaussian fits, red curves) for the interaction between; (B) *cytb₆f*[red] – Pc[ox], (C) *cytb₆f*[ox]
587 and Pc[red], (D) *cytb₆f*[red] – Pc[red], and (E) *cytb₆f*[ox]–Pc[ox]. For each, the total area of the
588 histogram reflects the adjusted frequency seen in (A). For B-E, the red line on each plot shows the
589 multiple peak fitting Gaussian for the plot, and the black lines show the underlying single peak fits
590 comprising the multipeak fit.

591

592 **Figure 4. Effect on unbinding forces and interaction frequency of varying ionic strength** (A) Multiple
593 overlaid histograms representing the distributions of unbinding forces for separating *cytb_{6f}*[red] –
594 Pc[ox] measured at different salt concentrations, with the cumulative sum adjusted relative to 5 mM
595 NaCl being 1. (B) Interaction frequency of each salt concentration adjusted relative to 5mM NaCl being
596 1. Error bars display the standard error of the mean.

597

598 **Figure 5. Single molecule force spectroscopy of the interaction between *cytb_{6f}* immobilized on a**
599 **epitaxial gold surface and Pc attached to the AFM probe.** (A) AFM height image showing distribution
600 of *cytb_{6f}* complexes on the gold substrate; (B) Height profiles of individual *cytb_{6f}* complexes from (A)
601 showing good agreement with the expected height of the complex derived from its structure. (C) Force
602 distribution for the interaction between *cytb_{6f}*[red] and Pc[ox] and (D) *cytb_{6f}*[red] – Pc[red]. The data
603 in (C) and (D) were normalized by using the surface density of the immobilised *cytb_{6f}* complexes on
604 the substrate and assuming uniform distribution of the Pc on the AFM probe. For C and D, the red line
605 on each plot shown the multiple peak fitting gaussian for the plot, and the black lines shows the
606 underlying single peak fits comprising the multipeak fit.

607

608 **Figure 6. Single molecule force spectroscopy of the interaction between *cytb_{6f}* in spinach grana**
609 **membranes and Pc attached to the AFM probe.** (A) Topographic AFM image of the luminal side of a
610 native grana membranes from spinach and the corresponding adhesion map in (C), showing the
611 interaction events occurring between Pc[ox] attached to the probe and *cytb_{6f}*[red] in the membranes.
612 (B) and (D) Corresponding topography and adhesion images after the grana membrane had been
613 incubated with an excess of Pc[ox] to block the available *cytb_{6f}* binding sites. (E) The cumulative
614 interaction frequency for the *cytb_{6f}*[red] and Pc[ox] with (blue) and without (red) excess of free

615 oxidised Pc. Error bars show the standard error of the mean. (F) Comparison between the distributions
616 of the unbinding force for *cytb₆f*[red] and Pc[ox] with (blue) and without (red) an excess of free Pc[ox].
617 The data in (E) and (F) were normalized by using the estimate number of *cytb₆f* complexes in the grana
618 and assuming uniform distribution of the Pc on the AFM probe.

619

620 **Figure 7. Binding interface between spinach plastocyanin and cytochrome b₆f.** Plastocyanin and
621 *cytb₆f* binding interfaces shown in Pymol from the plastocyanin structure (pdb:1AG6) and modelled
622 cytochrome f structure (Swiss model (Waterhouse et al., 2018), using cytochrome *f* – Spinach
623 sequence, mapped to structure of *C.reinhardtii* cytochrome *f* – pdb:1Q90). Acidic residues are
624 highlighted in blue, basic in red and hydrophobic regions are in orange.

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