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**Title:****Nanodomains of Cytochrome  $b_6f$  and Photosystem II Complexes in Spinach Grana Thylakoid Membranes****Authors:**

Matthew P. Johnson\*, Cvetelin Vasilev, John D. Olsen and C. Neil Hunter

**Affiliations:**

Department of Molecular Biology and Biotechnology, University of Sheffield, Firth Court, Western Bank, Sheffield, S10 2TN, United Kingdom.

\* Corresponding author: Matthew P. Johnson ([matt.johnson@sheffield.ac.uk](mailto:matt.johnson@sheffield.ac.uk))

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**Running Title:**

Nanodomains of  $cytb_6f$  and PSII

**Keywords:**

Electron transport / cytochrome  $b_6f$  / photosystem II / thylakoid membrane/ atomic force microscopy

**Synopsis:**

Co-localization of cytochrome  $b_6f$  and photosystem II complexes creates nanodomains that facilitate rapid exchange of the lipophilic electron carrier plastoquinone in the crowded thylakoid grana membrane.

**Abstract:**

The cytochrome  $b_6f$  (cyt $b_6f$ ) complex plays a central role in photosynthesis, coupling electron transport between photosystem II (PSII) and photosystem I (PSI) to the generation of a transmembrane proton gradient used for the biosynthesis of ATP. Photosynthesis relies on rapid shuttling of electrons by plastoquinone (PQ) molecules between PSII and cyt $b_6f$  complexes in the lipid phase of the thylakoid membrane. The relative membrane location of these complexes is thus crucial, yet remains unknown. Here, we exploit the selective binding of the electron transfer protein plastocyanin (Pc) to the luminal membrane surface of the cyt $b_6f$  complex by using a Pc-functionalized atomic force microscope (AFM) probe to identify the position of cyt $b_6f$  complexes in grana thylakoid membranes from *Spinacia oleracea* (spinach). This affinity-mapping AFM method directly correlates membrane surface topography with Pc-cyt $b_6f$  interactions allowing us to construct a map of the grana thylakoid membrane that reveals nanodomains of co-localized PSII and cyt $b_6f$  complexes. We suggest that the close proximity between PSII and cyt $b_6f$  complexes integrates solar energy conversion and electron transfer by fostering short-range diffusion of PQ in the protein-crowded thylakoid membrane, thereby optimizing photosynthetic efficiency.

## INTRODUCTION

Photosynthesis in plants, algae and cyanobacteria begins with the capture and trapping of solar energy by photosystem I and II (PSI and PSII). The cytochrome  $b_6f$  ( $cytb_6f$ ) complex acts as the electrical connection between these two photosystems by oxidizing the PSII electron acceptor plastoquinone (PQ) and reducing the PSI electron donor plastocyanin (Pc; reviewed in Cramer et al., 2011). The electron transfer reactions carried out by  $cytb_6f$  are coupled to the generation of an electrochemical proton gradient across the chloroplast thylakoid membrane, which is harnessed by ATP synthase to form ATP. The membrane location and organization of  $cytb_6f$  complexes is crucial to their function; it is known that the thylakoid membrane in higher plants is divided into two domains, the grana stacks, which are enriched in PSII, and the interconnecting stromal lamellae, which are enriched in PSI and ATP synthase complexes (reviewed in Albertsson, 2001; Dekker and Boekema, 2005; Daum and Kühlbrandt, 2011; Nevo et al., 2012). The distribution of the  $cytb_6f$  complexes between the grana and stromal lamellae is much less clear. Biochemical evidence obtained on thylakoids fractionated either mechanically or with the detergent digitonin suggested that  $cytb_6f$  was distributed fairly evenly between the grana and stromal lamellae (Boardman and Anderson, 1967; Sane et al., 1970; Cox and Andersson, 1981; Anderson, 1982; Dunahay et al., 1984). This even distribution was supported by immunogold labeling of the  $cytb_6f$  complex in intact thylakoids and by freeze-fracture electron microscopy studies comparing wild-type and a  $cytb_6f$ -less mutant of the green alga *Chlamydomonas reinhardtii* (Allred and Staehelin, 1985; Olive et al., 1986; Vallon et al., 1991; Hinshaw and Miller, 1993). By contrast, fractionation with the detergents Triton X-100 or n-dodecyl- $\alpha$ -D-maltoside ( $\alpha$ -DM) suggested that the grana were devoid of  $cytb_6f$  and that this complex was confined to the stromal lamellae or grana margins (Berthold et al., 1981; Dunahay et al., 1984; Morissey et al., 1986; van Roon et al., 2000). Exclusion of  $cytb_6f$  from the grana would have significant consequences for photosynthetic electron transport since it would require long-range diffusion of PQ over hundreds of nanometers from PSII in the grana to  $cytb_6f$  in the stromal lamellae. Yet, a range of evidence suggests that PQ diffusion is greatly restricted in the thylakoid membrane: first, PQ diffusion in thylakoid membranes is 1000-fold slower than in pure liposomes (Blackwell et al., 1994); second, spectroscopic data show that two pools of PQ with different rates of photoreduction by PSII exist in the thylakoid membrane and that equilibration between them is very slow (Joliot et al., 1992); finally, the rate of  $cytb_6f$  reduction declines

almost linearly in response to inhibition of PSII activity with the herbicide 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU)(Kirchhoff et al., 2000), suggesting that each cytb<sub>6</sub>f complex is connected to only a limited number of PQ molecules (Joliot et al., 1992; Kirchhoff et al., 2000). Percolation theory suggests that protein crowding in the densely-packed grana membrane, with ~80% occupancy, is the likely cause of impaired PQ diffusion (Kirchhoff et al., 2002). These findings led to the hypothesis that ‘microdomains’ of PSII, PQ and cytb<sub>6</sub>f facilitate rapid diffusion of PQ over short (<20 nm) distances in the grana (Joliot et al., 1992; Kirchhoff et al., 2000). Crucially however, there is no direct structural evidence for such domains since no techniques are available to visualize the local arrangement of PSII and cytb<sub>6</sub>f complexes in the same membrane. Here, we describe the application of an affinity-mapping atomic force microscopy (AFM) technique (Vasilev et al., 2014) to uniquely identify and locate cytb<sub>6</sub>f complexes and to determine their proximity to PSII in grana membranes.

## RESULTS

### **Two major types of topographic feature, distinguished by their heights, can be observed in AFM images of digitonin- and sonication-derived grana membranes**

Thylakoid grana membranes were prepared by three different methods for interrogation by AFM: by limited solubilization with the detergents  $\alpha$ -DM (van Roon et al., 2000) or digitonin (Boardman and Anderson, 1966), or alternatively mechanical fragmentation by sonication (Albertsson et al., 1994). Each of these methods is known to produce ‘inside-out’ (i.e. lumen-side exposed) appressed pairs of membranes derived from the thylakoid grana stacks (Andersson et al., 1978; Dunahay et al., 1984; van Roon et al., 2000). SDS-PAGE analysis (Supplemental Figure 1 online) and chlorophyll a/b ratios (Table 1) confirmed that the three different methods produced membranes relatively enriched in PSII and LHCII and depleted in PSI and ATP synthase compared to intact thylakoids. The chlorophyll a/b ratio and PSI contents of the  $\alpha$ -DM grana ( $\alpha$ G) were slightly lower than for grana prepared by digitonin (DG) or sonication (SG), consistent with previous reports (Boardman and Anderson 1966; Albertsson et al., 1994; van

Roon et al., 2000) (Table 1). AFM images of the three grana preparations recorded under liquid revealed appressed double membranes 400-500 nm in diameter and 18-20 nm in height. The grana membranes are covered with topographic features protruding 3-5 nm from the surrounding membrane surface that are 16-20 nm in width, consistent with the structure of the luminal protrusions of the PSII core complex (Zouni et al., 2001; Ferreira et al., 2004; Loll et al, 2005; Guskov et al., 2009; Umena et al., 2011; Sznee et al., 2011) (Figure 1A-C). The holes and frayed edges often evident in  $\alpha$ G membranes (Figure 1A) suggested a level of 'damage', as observed previously (van Roon et al., 2000; Sznee et al., 2011). Higher resolution images obtained on these membranes (Figure 1D-F) showed that the individual topographic features seen in Figure 1A-C are often resolved as two protrusions with maxima separated by  $\sim$ 8 nm. Immunoblotting with an antibody specific to the PetC subunit of  $\text{cytb}_6\text{f}$  indicated that this complex is not present in the  $\alpha$ G membranes but is found in DG and SG membranes (Figure 1G). Spectroscopic assays confirmed this result; the DG and SG membranes contained 1.4 mmol  $\text{cytb}_6\text{f}$ / mol chlorophyll while none was detected in  $\alpha$ G membranes (Table 1). The PSII/  $\text{cytb}_6\text{f}$  ratio calculated from the spectroscopic data (Table 1) for the SG and DG membranes was 2.3, suggesting that some of the topographic features in these membranes likely belong to  $\text{cytb}_6\text{f}$  which, like PSII, protrudes into the lumen space (Kurisu et al., 2003; Stroebel et al., 2003). The crystal structures of the dimeric  $\text{cytb}_6\text{f}$  and PSII complexes reveal a similar lateral size and shape, with their long axes 16 and 20 nm respectively, and with both dimers having two luminal protrusions separated by  $\sim$ 8 nm (Kurisu et al., 2003, Ferreira et al., 2004) (Figure 1H-J). These similarities hamper discrimination between these complexes in AFM images, given that probe sharpness limits the lateral resolution of the AFM to a typical value of  $\sim$ 4-5 nm. However, the much higher ( $\sim$ 0.1 nm) z-axis resolution of the AFM should resolve the  $\sim$ 1 nm height difference between the luminal protrusions of PSII and  $\text{cytb}_6\text{f}$  complexes (Figure 1K). Accordingly, we measured the height of every protruding feature in these thylakoid membranes. The histograms of the height distributions clearly show two populations of topographic features (Figure 1L-N), a major peak centered on  $4.0 \pm 0.1$  nm present in all the grana preparations and a minor peak corresponding to a  $3.0/3.1 \pm 0.1$  nm height, which was only present in the DG and SG membranes. Assuming that we measure from the luminal face of the surrounding LHCII complexes, which cover most of the grana membrane surface and protrude by  $\sim$ 0.5 nm (Liu et al., 2004; Standfuss et al., 2005; Daum et al., 2010) (Figure 1K), the major and minor peaks are consistent with the predicted

heights of the PSII and cytb<sub>6</sub>f complexes respectively. The area ratios of the 4.0 and 3.0/3.1 nm height distributions (Figure 1 M, N) were 2.5:1, in reasonable agreement with the PSII/ cytb<sub>6</sub>f ratio of 2.3 calculated by spectroscopy (Table 1). The absence of this secondary peak from analyses of AFM topographs for αG membranes correlates with the absence of cytb<sub>6</sub>f from the immunoblot data in Figure 1G and the spectroscopic data in Table 1, supporting the assignment of the 3-nm height features as cytb<sub>6</sub>f complexes.

### **Affinity-mapping AFM using Pc-functionalized AFM probes allows the position of cytb<sub>6</sub>f complexes to be mapped in grana membranes**

To positively confirm that these 3 nm protrusions correspond to the cytb<sub>6</sub>f complex we exploited the recently described affinity-mapping AFM technique (Vasilev et al., 2014), this time utilizing the native interaction between the small soluble electron transfer protein Pc, which was attached to the AFM probe, and the exposed luminal face of the cytb<sub>6</sub>f complexes in grana membranes adsorbed to a mica support. This method allows simultaneous recording and correlation of topographic and force (probe-sample adhesive interaction) data. The rationale of this experiment is shown in Figure 2A and B. Pc proteins purified from spinach thylakoids (Supplemental Figure 2A-C online) were attached by a flexible, 10 nm-long SM(PEG)<sub>24</sub> linker molecule to an AFM probe (Figure 2A). No adhesive interactions should be observed when the Pc-functionalized probe interacts with a part of the membrane surface devoid of cytb<sub>6</sub>f complexes or with a PSII complex (Figure 2A). However, when the Pc-functionalized probe encounters a cytb<sub>6</sub>f complex at the membrane surface, a specific Pc-cytb<sub>6</sub>f interaction can occur; subsequently the upward movement of the probe fully extends the linker molecule before rupturing this interaction (Figure 2B). To validate detection of specific Pc-cytb<sub>6</sub>f interactions using our affinity-mapping AFM method we nanopatterned 300 nm-wide lines of purified cytb<sub>6</sub>f complexes on a glass substrate. The correspondence between the topographic and adhesion images in Supplemental Figure 3E and F online, respectively, shows that a Pc-functionalized probe, pre-oxidized in 10 mM potassium ferricyanide, reliably detects multiple, specific Pc-cytb<sub>6</sub>f associations, as well as retaining the ability to record surface topography. Pc-functionalized probes were then used to investigate the organization of cytb<sub>6</sub>f complexes in purified grana membranes illuminated under

white light with a power density of approximately  $11 \text{ W m}^{-2}$  (measured at the sample surface) to ensure PSII-mediated formation of reduced PQ and thus reduction of cytb<sub>6</sub>f. When DG membranes were scanned with the pre-oxidized Pc-functionalized probes, the majority of the topographic features on the membrane surface were still well resolved (Figure 2C). Simultaneously with the topology image the forces required to rupture probe-sample interactions were recorded as an adhesion image (Figure 2D). We observed a number of locations ('interaction hotspots') on the membrane surface where high (>50 pN) unbinding forces were recorded between the pre-oxidized Pc probe and the sample (Figure 2D). Crucially, when the DG membranes were scanned with Pc-functionalized probes pre-reduced in 10 mM sodium dithionite, the topographic features were still observed (Figure 2E), but now there were very few interaction hotspots observed on the membrane surface in the adhesion image (Figure 2F). The force-distance curves associated with such interaction hotspots on the membrane surface (see example in Figure 3A, cyan curve) show unbinding forces in the ~200-500 pN range. The probe-sample rupture length of ~10 nm, obtained from inspection of the horizontal axis of Figure 3A (cyan curve), indicates that the linker fully extended before rupture of this interaction. Figure 3A also shows a typical force-distance curve for the majority of the membrane area (red curve), where we observed very low unbinding forces between the probe and the membrane surface that were comparable to the noise level of the measurement. Finally, Figure 3A shows an example of a force-distance curve recorded on the mica surface (orange curve) where we observed unbinding forces in the 300-500 pN range but importantly with a rupture length very close to 0 nm. Such events correspond to a non-specific and direct interaction between the apex of the AFM probe and the mica, with no involvement of the linker or the attached Pc.

In order to obtain robust statistics for the interaction hotspots on the membrane surface we analyzed 276 force-distance curves recorded on 14 different DG samples interrogated with pre-oxidized probes. The graph in Figure 3B shows a most probable unbinding force of  $312 \pm 5$  pN (Figure 3B) and an average rupture length of  $10.0 \pm 0.3$  nm, consistent with the length of the flexible SM(PEG)<sub>24</sub> linker molecule used to attach the Pc to the AFM probe. A second smaller peak in the force distribution was centered on  $607 \pm 15$  pN, which likely results from the rupture of the interaction between two Pc molecules on the functionalized probe and two cytb<sub>6</sub>f complexes on the surface.

## **The frequency of interaction between the Pc-functionalized AFM probe and cytb<sub>6</sub>f complexes on the membrane surface depends on the redox conditions**

The estimated number of cytb<sub>6</sub>f complexes determined by spectroscopy (Table 1) suggests that the DG and SG membranes contain 51.5 dimeric complexes per 500 nm diameter circular membrane disc, i.e.  $\sim 262$  cytb<sub>6</sub>f dimers/ $\mu\text{m}^2$ . The average number of interaction hotspots detected with pre-oxidized Pc functionalized probes was  $158 \pm 12$  / $\mu\text{m}^2$ , suggesting around 60% of cytb<sub>6</sub>f dimers are detected by our method (Figure 4A). When the experiment was performed with pre-reduced Pc functionalized probes the number of interaction hotspots on the membrane surface decreased by 96%, indicating the expected preference for oxidized Pc to interact with cytb<sub>6</sub>f (Figure 4A). When PSII was inhibited by addition of 10  $\mu\text{M}$  DCMU to lower the production of reduced PQ and the membrane imaged 10 minutes later, the number of interaction hotspots detected on the membrane surface decreased by 44%, indicating that the cytb<sub>6</sub>f redox state also affects the number of cytb<sub>6</sub>f complexes detected (Figure 4A-C). Addition of 50  $\mu\text{M}$  excess oxidized Pc (twice the reported  $K_D$  of 23  $\mu\text{M}$  (Meyer et al., 1993)) to the sample prior to scanning, thus blocking potential cytb<sub>6</sub>f binding sites for the oxidized Pc-functionalized probe, lowered the number of interaction hotspots by 72% (Figure 4A, D and E). Affinity-mapping AFM of  $\alpha\text{G}$  membranes was instructive since the immunoblot and spectroscopic assays had indicated that these membranes contained little or no cytb<sub>6</sub>f. Scanning  $\alpha\text{G}$  membranes with the oxidized Pc probes gave 91% fewer interaction hotspots than the DG membranes, consistent with the virtual absence of the cytb<sub>6</sub>f complex from the  $\alpha\text{G}$  preparation (Figure 4A, F and G). As expected, the SG membranes gave a very similar number of interaction hotspots as the DG membranes (Figure 4A, H and I).

## **Pc-cytb<sub>6</sub>f interactions detected by affinity-mapping AFM specifically label the 3-nm high topographic features in AFM images of grana membranes**

The interaction hotspots recorded on the surface of DG and SG membranes with pre-oxidized Pc-functionalized probes reliably identified topographic features that protrude  $\sim 3$  nm from the

membrane surface at their highest point (Figure 5A and B), supporting the assignment of these features as  $\text{cytb}_6\text{f}$  complexes. By contrast, the 4-5 nm protrusions from the membrane surface, consistent with the height of PSII, showed no high unbinding force signals, but were found in close proximity to the 3-nm topographic features (Figure 5A and B). These assignments led us to fit a representative area of our highest resolution AFM image (Figure 5C) with the crystal structures of the dimeric  $\text{cytb}_6\text{f}$  and PSII complexes (Figure 5D). The resulting membrane model in Figure 5D suggests that  $\text{cytb}_6\text{f}$  complexes (magenta) are often surrounded by PSII complexes (green) in small domains as originally suggested (Joliot et al., 1992; Kirchhoff et al., 2000).

## DISCUSSION

### **Affinity-mapping AFM is a useful tool for mapping membrane organization at the nanoscale**

Our affinity-mapping AFM approach shows how a biological membrane can be mapped on the nanoscale, by exploiting the specific binding of a membrane-extrinsic probe onto a membrane-bound target protein. Here, we made use of a transient interaction between two electron transfer proteins, the extrinsic electron acceptor Pc and the  $\text{cytb}_6\text{f}$  donor, an integral membrane protein complex. When oxidized Pc, attached to an AFM probe by a 10-nm flexible linker, scans across the thylakoid membrane surface it specifically detects reduced  $\text{cytb}_6\text{f}$  complexes. By simultaneously recording surface topology and Pc- $\text{cytb}_6\text{f}$  interactions we were able to distinguish between  $\text{cytb}_6\text{f}$  and PSII by their extents of protrusion from the membrane and by transient protein-protein interactions. The co-location and organization of the PSII and  $\text{cytb}_6\text{f}$  complexes in grana thylakoid membranes in our combined topographic and affinity maps reveals a close association between these complexes, and shows how the excitation energy and electron transfer events of photosynthesis are linked in the thylakoid membrane.

### **Affinity-mapping AFM using Pc-functionalized AFM probes allows Pc- $\text{cytb}_6\text{f}$ interactions to be quantified at the single-molecule level under near native conditions**

The large number of interaction hotspots recorded in each force map also produced robust statistics for in situ quantification of the Pc-cytb<sub>6</sub>f interaction unbinding force, at the single-molecule level. The unbinding force of  $312 \pm 5$  pN compares with a value of 480 pN measured by the same method, albeit in vitro, for the interaction between probe-attached cytochrome c<sub>2</sub> (cytc<sub>2</sub>) and surface-bound reaction center (RC) complexes from the purple photosynthetic bacterium *Rhodobacter sphaeroides* (Vasilev et al., 2014). The latter value agreed with theory-based estimates of the cytochrome c<sub>2</sub>-RC interaction, which calculated an unbinding force of 600-1000 pN (Pogorelov et al., 2007). By measuring an in situ membrane protein complex stabilized by the membrane bilayer and by interactions with neighbors, repeated measurements are possible that report functional interactions, with no extraction or unfolding of the cytb<sub>6</sub>f complex. Such stabilizing influences allow measurement of the ~300 pN unbinding forces, which are comparable with those obtained using conventional force spectroscopy for a range of other interactions including 340-454 pN for streptavidin-biotin (Lee et al., 1994; Steven et al., 2002), 507 pN for the hCG-anti-hCG pair (Steven et al., 2002), 512 pN for the MPAD-anti-MPAD interaction (Kaur et al., 2004), and 420 pN for citrate synthase-GroEL (Vinckier et al., 1998). The forces we measure likely sample the binding interaction prior to electron transfer, given the requirement for both reduced cytb<sub>6</sub>f and oxidized Pc, as well as the similarity of the dwell time (approx. 160  $\mu$ s at 1.0 kHz force curve repetition rate) of the AFM probe on the membrane surface to the ~70-130  $\mu$ s electron transfer between Pc and cytb<sub>6</sub>f (Haehnel et al., 1980; Delosme, 1991). Thus, multiple Pc-cytb<sub>6</sub>f encounters enable the mapping of most of the cytb<sub>6</sub>f complexes in the thylakoid membrane.

Earlier in vitro studies using site-directed mutants of Pc and the soluble domain of cytochrome f or the purified cytb<sub>6</sub>f complex concluded that interaction between these proteins depends largely on electrostatic interactions between the partners (Lee et al., 1995; Kannt et al., 1996; Illerhaus et al., 2000). However, mutagenesis of cytochrome f in *Chlamydomonas* found that electrostatic interactions exerted a much weaker influence on cytochrome f oxidation kinetics in vivo (Soriano et al., 1996; 1998), and other factors undoubtedly come into play in chloroplasts such as restricted diffusional space and the higher osmotic strength found in the thylakoid lumen. A nuclear magnetic resonance (NMR) study of the complex between cytochrome f and plastocyanin from the cyanobacterium *Nostoc* showed that the binding interface involves hydrophobic residues with charged residues at the edge (Díaz-Moreno et al.,

2005). A more recent NMR study of the cytb<sub>6</sub>f-Pc interaction using spinach components, including SG membranes, showed that there is an initial, loose electrostatic interaction and the subsequent formation of an electron transfer complex relies on hydrophobic interactions (Ueda et al., 2012).

The single-molecule affinity-mapping AFM approach taken in the present work approximates the *in vivo* situation due to our use of cytb<sub>6</sub>f *in situ* in native membranes, rather than solubilized complexes. Thus, we expect that electrostatic interactions are involved in the initial encounter phase of the cytb<sub>6</sub>f-Pc interaction and that hydrophobic interactions then guide the formation of an electron transfer complex. The use of Pc tethered to an AFM probe with its attachment site distal to the Pc-cytf contact region steers the tethered Pc towards its binding site, likely bypassing the multiple binding/unbinding encounters and reorientations possible when two soluble components interact in solution. Rather, our AFM measurements reflect the aftermath of cytb<sub>6</sub>f-Pc complex formation, and the ~300 pN unbinding forces we measure (Figure 3) reflect the factors that stabilize the cytb<sub>6</sub>f-Pc complex and quantify the forces required to disrupt it. Preparing the reduced cytb<sub>6</sub>f and oxidized Pc reactants prior to bringing them into contact might accelerate formation of a stable electron transfer complex, and increase the likelihood that we have to apply a ~300 pN unbinding force to pull it apart. As already noted, the duration of the 160- $\mu$ s dwell time of Pc at the membrane surface might be sufficient to allow electron transfer, which takes ~70-130  $\mu$ s (Haehnel et al., 1980; Delosme, 1991); thus our experiments could reflect the stability of the cytb<sub>6</sub>f (reduced)-Pc(oxidised) complex just prior to electron transfer or of the cytb<sub>6</sub>f (oxidised)-Pc(reduced) complex following electron transfer. In this respect, the present work differs from the forces required to disrupt the *in vitro* interaction between probe-attached cytc<sub>2</sub> and surface-bound RC complexes (Vasilev et al., 2014); in this case the cytc<sub>2</sub>-RC electron transfer is much faster, in the low microsecond range (Overfield et al. 1979; Moser and Dutton, 1988). Thus, the reduced cytc<sub>2</sub> is rapidly oxidised and the unbinding events we measured likely reflected the interaction between reduced RC and oxidised cytc<sub>2</sub>, i.e. the aftermath of the electron transfer event.

Our preparations contain PSI, so it is possible that some of the interaction hotspots we detect with our affinity-mapping AFM method could correspond to Pc-PSI interactions. However we consider this to be unlikely for several reasons: first the crystal structure of PSI (Amunts et al., 2007) shows that the luminal protrusions of this complex are <1 nm whereas the interaction

hotspots we detect consistently label features 3 nm in height, a difference easily resolved by the 0.1 nm z-resolution of the AFM; second, in the event of binding to PSI we would predict that the pre-reduced Pc-functionalized AFM probes would detect a similar number of interactions when in fact they were greatly decreased compared to the pre-oxidized probes. The fact we observed very few interactions of pre-reduced Pc-functionalized probes with the PSI complexes in DG and SG membranes, suggests we could not maintain a photo-oxidized state, likely because we did not include a PSI electron acceptor.

### **Cytb<sub>6</sub>f complexes are distributed throughout the grana membranes amongst PSII complexes**

The co-location of the cytb<sub>6</sub>f and PSII complexes has important implications for photosynthesis, and provides direct evidence for the microdomain hypothesis (Joliot et al., 1992; Kirchhoff et al., 2000). The proximity of these complexes, in regions more appropriately termed ‘nanodomains’, allows PQ diffusion over short (<20 nm) distances. Affinity-mapping AFM shows that cytb<sub>6</sub>f complexes are interspersed throughout the grana regions, in the vicinity of PSII complexes but not tightly bound with fixed stoichiometry in a supercomplex. Recent cryo-electron microscopy studies on DG membranes and intact thylakoids revealed the presence of numerous smaller densities alongside those assigned to PSII, which the authors suggested could belong to monomeric PSII or cytb<sub>6</sub>f complexes (Daum et al., 2010; Kouřil et al., 2011). Given the data reported in this paper and the fact that PSII exists almost entirely as a dimer in the thylakoid membrane (Dekker and Boekema, 2005), these unassigned densities likely belong to cytb<sub>6</sub>f. As in our data these densities were distributed throughout the PSII complexes in grana membrane (Daum et al., 2010; Kouřil et al., 2011). However, it is likely that cytb<sub>6</sub>f complexes are excluded from the 5-10% of grana membranes in wild-type plants that form ordered semi-crystalline PSII arrays (Dekker and Boekema, 2005). It is reasonable to conclude that the granal cytb<sub>6</sub>f complexes we have co-located with PSII facilitate linear electron transport while the remaining 40-50% located in the stromal lamellae facilitate cyclic electron transport as previously suggested (Albertsson et al., 2001). Indeed, there is evidence that the distribution of cytb<sub>6</sub>f complexes is dynamically responsive to the PQ redox state, with an increase in the proportion

found in the stromal lamellae under conditions where the PQ pool is reduced (Vallon et al., 1991). The absence of cytb<sub>6</sub>f complexes from Triton- and  $\alpha$ -DM-derived grana membranes may be due to their selective solubilization by Triton and  $\alpha$ -DM detergents which could explain the holes frequently observed in these membranes and their generally ragged appearance (Dunahay et al., 1984; van Roon et al., 2000; Sznee et al., 2011). Indeed, cytb<sub>6</sub>f was solubilized from the membrane at lower concentrations of Triton than used for either PSI or PSII complexes (Morrissey et al., 1986).

### **Nanodomains of cytb<sub>6</sub>f and PSII complexes ensure rapid exchange of PQ molecules and thus efficient electron transport**

There are some parallels between the co-location of cytb<sub>6</sub>f and PSII complexes shown in the present work and recent study that used EM and AFM to demonstrate proximity between the dimeric cytochrome bc<sub>1</sub> and dimeric RC complexes in the photosynthetic membrane of the bacterium *Rhodobacter sphaeroides* (Cartron et al., 2014). The close proximity of cytochrome complexes and Type II (quinone acceptor) RCs, such as PSII, may therefore be advantageous for facilitating rapid exchange of quinones. Evolution has reconciled the high density of pigment binding proteins required to ensure the efficient transfer and trapping of solar energy with the need for sufficient diffusion space for quinone traffic between RC and cytochrome complexes; the solution revealed by this work is the distribution of cytb<sub>6</sub>f complexes amongst PSII supercomplexes in the grana, minimizing the average distance traversed by PQ molecules in the crowded membrane environment and promoting rapid exchange of PQ between cytb<sub>6</sub>f and PSII complexes.

## **METHODS**

### **Membrane / protein isolation.**

Spinach thylakoid membranes (Albertsson et al., 1994),  $\alpha$ G membranes (van Roon et al., 2000), DG membranes (Boardman and Anderson, 1966), SG membranes (Albertsson et al., 1994), Pc (Morand and Krogmann, 1993) and cytb<sub>6</sub>f (Zhang et al., 2001) were prepared as previously described. SDS-PAGE and immunoblot analysis was performed as previously described (Ruban et al., 2006).

### **Spectroscopic determination of chlorophyll and cytochrome contents.**

Spectroscopic assays of P700 and cytb<sub>6</sub>f content were performed according the methods of Melis and Brown (1980) and Bendall et al. (1971) respectively. The obtained value of 1.4 mmol cytochrome f / mol chlorophyll is similar to that calculated using this method by Albertsson et al. (1994). To calculate number of cytb<sub>6</sub>f complexes per 500 nm grana disc, the following chlorophyll composition was assumed based on the antenna sizes and PSI/PSII ratios given in Danielsson et al., (2004) for grana derived by the same method from thylakoid membranes with similar chlorophyll a/b ratio: 120 C<sub>2</sub>S<sub>2</sub> type PSII (120 x 204 = 24480 Chls), 602 LHCI trimers (602 x 42 = 25284 Chls), 122 PSI-LHCI (122 x 168 = 20496 Chls), 240 CP24 (13 x 240 = 3120); total Chls = 73380, hence 0.0014 x 73380 = 102.7 cytochrome f molecules per single 500 nm grana disc (~51.5 dimers).

### **AFM / Affinity-mapping AFM.**

Grana membranes were adsorbed to the mica substrate as previously described (Sznee et al., 2011) and imaged in a buffer containing 10 mM HEPES-NaOH pH 8.0, 5 mM NaCl, 5 mM MgCl<sub>2</sub>. Imaging was performed by PeakForce quantitative nanomechanical mapping (PF-QNM) mode using a Bruker Multimode AFM. The spring constants of the AFM cantilevers used were individually determined to accurately quantify the force data and varied between 0.06 and 0.24 Nm<sup>-1</sup>. The Z-modulation amplitude was adjusted to values in the range 15 – 25 nm to allow enough probe-sample separation in order to fully stretch the SM(PEG)<sub>24</sub> linker molecule on the

AFM probe and to separate the Pc from the cytb<sub>6</sub>f molecules during each ramp cycle. Before starting the measurements, the Pc molecules on the AFM probes were either pre-oxidized by incubation in oxidizing buffer (imaging buffer supplemented with 10 mM potassium ferricyanide) or pre-reduced with reducing buffer (imaging buffer supplemented with 10 mM sodium dithionite) with a subsequent wash in imaging buffer. The contact probe-sample force was kept in the range 100 – 200 pN and the imaging rate was adjusted (depending on the scan size and pixel density of the scan) in a way that ensured a 256 x 256 pixel PeakForce Capture image could be acquired at a modulation frequency of 1 kHz. During the affinity mapping the sample was illuminated from a white light source through an optical fibre (Fiber-Lite MI-150, Dolan-Jener) and the power density of the illumination at the sample surface was measured with a Newport 842-PE meter (Newport Corp.). In addition, some partial illumination most probably occurred from the 670 nm laser used in the optical lever detection system for the AFM. For the control experiments, excess Pc or DCMU were added directly to the liquid cell by pipette at the concentrations described in the text. Subsequent image analysis was performed using Bruker Nanoscope Analysis v1.42 and OriginPro v8.5.1 (OriginLab Corp.) software. The Nanoscope Analysis was used for the extraction and analysis of the PF-QNM force spectroscopy data and OriginPro 8.5 was used for the statistical analysis of all the force spectroscopy data and for all the calculations and curve fitting. Data reduction (positive identification of specific rupture events) was based on the analysis of the rupture events with probe-sample separations in the range 3 – 30 nm. The most probable values for the unbinding force and the rupture length were obtained from the maximum of the Gaussian fit to the force and rupture length distribution combined in a statistical histogram. The unbinding forces and rupture lengths of 276 rupture events were compiled in force or length distribution histograms.

### **Preparation of functionalized probes for affinity-mapping AFM.**

Hybrid AFM probes, Si tips mounted on Si<sub>3</sub>N<sub>4</sub> rectangular or triangular cantilevers, model BL-AC40TS (Olympus Probes) or SNL (Bruker) respectively, were first cleaned by washing in acetone (HPLC grade from Fisher Scientific) and then cleaned in a home- built UV/Ozone cleaner (LSP035 Pen-Raylight source, LOT-Oriel Ltd.) for 45 min. Immediately after the cleaning step the AFM probes were placed into a glass desiccator purged with pure nitrogen for

10 min and then 20  $\mu$ l (3-mercaptopropyl)trimethoxysilane (MPTMS) was introduced into the desiccator. After another 5 min purge, the desiccator was evacuated down to a pressure of approximately 0.3 kPa using a dry mechanical pump (Welch, model 2027), and then sealed for 6 – 8 h to facilitate the deposition of the self-assembled monolayer (SAM). The next step in the functionalization of the AFM probes, immediately after the SAM formation, was to attach the Pc proteins. An amine-to-sulfhydryl heterobifunctional cross-linker with a 9.5-nm polyethylene-glycol (PEG) spacer arm, terminated at one end with N-hydroxysuccinimide (NHS) ester group and at the other with maleimide (MAL) group (SM(PEG)<sub>24</sub>, Pierce Biotechnology) was used in order to attach an exposed lysine residue on the surface of the Pc molecule to the AFM probe. Both the amine-targeted and thiol-targeted reactions were accomplished simultaneously in sodium phosphate buffer pH 7.4 at a final SM(PEG)<sub>24</sub> concentration of 1 mM and a final Pc concentration of 50 nM for 40 min. Then, the AFM probes were gently washed (4 times) in 10 mM HEPES pH 8.0 buffer and stored in the same buffer for further use.

### **Preparation of patterned surfaces of cytb<sub>6</sub>f.**

The linear patterns of cytb<sub>6</sub>f were prepared by chemical patterning of glass substrates using a combination of reverse nanoimprint lithography (RNIL) and a wet lift-off and transfer of a thin polymer film replica together with a SAM of silanes to which the cytb<sub>6</sub>f molecules were selectively attached.

### **Accession Numbers**

Structure data used in this work is available at the Protein Data Bank under the following accession numbers: Pc (1YLB), PSII (1S5L), cytb<sub>6</sub>f (1UM3), LHCII (2BHW)

### **Supplemental Data**

The following materials are available in the online version of this manuscript.

**Supplemental Figure 1.** Coomassie-stained SDS-PAGE gel of protein composition of membrane preparations.

**Supplemental Figure 2.** Preparation of Pc from spinach thylakoids.

**Supplemental Figure 3.** Preparation of cytb<sub>6</sub>f from spinach thylakoids and patterning on glass for detection by affinity-mapping AFM.

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## **AUTHOR CONTRIBUTIONS**

M.P.J. performed the purification of membranes and isolated complexes and characterized them by spectroscopy. C.V. prepared the functionalized AFM probes. M.P.J. and C.V. performed the AFM experiments and data analysis. J.D.O. provided advice and support to AFM experiments. M.P.J, C.V. and C.N.H. designed the study and wrote the manuscript. All authors discussed the results and commented upon the manuscript.

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## FIGURE LEGENDS

**Figure 1.** Atomic Force Microscopy analysis of grana thylakoid membranes isolated from spinach.

AFM topographs of **(A, D)**  $\alpha$ G membranes, **(B, E)** DG membranes, **(C, F)** SG membranes; the scale bars in **(A), (B)** and **(C)** are 100 nm and in **(D), (E), (F)** are 35 nm. **(G)** Immunoblot of the  $\text{cytb}_6\text{f}$  (PetC) content of thylakoids and grana membrane preparations (thy. = intact thylakoids). Atomic models viewed from the thylakoid lumen membrane side of **(H)** dimeric PSII (Protein data bank (PDB) code: 1S5L), **(I)** dimeric  $\text{cytb}_6\text{f}$  (PDB code: 1UM3) and **(J)** overlay comparison. **(K)** Atomic models of dimeric PSII, dimeric  $\text{cytb}_6\text{f}$  and trimeric LHCII (PDB code: 2BHW) viewed parallel to the thylakoid membrane. The dashed lines show the height difference between PSII and  $\text{cytb}_6\text{f}$  of  $\sim 1$  nm and their heights above the LHCII complex, which protrudes by  $\sim 0.5$  nm from the luminal membrane surface. **(L)** Histogram of maximum heights of protruding topological features above the surrounding area in  $\alpha$ G, **(M)** DG and **(N)** SG membranes.

**Figure 2.** Affinity-mapping AFM of digitonin-derived grana (DG) membranes isolated from spinach.

**(A)** Cartoon depicting the principle of affinity-mapping AFM. The AFM probe laterally images the topography of the thylakoid lumen membrane surface via a tapping motion (dashed black arrow), while simultaneously recording probe-sample interaction forces for every pixel; the AFM probe is functionalized with Pc proteins attached via a flexible 10 nm long SM(PEG)<sub>24</sub> linker. **(B)** Cartoon depicting the specific interaction of one of the Pc proteins attached to the AFM probe with a  $\text{cytb}_6\text{f}$  complex on the membrane surface; when the AFM probe is withdrawn from the surface during the upward part of the tapping motion the flexible linker is extended to its full 10 nm length before Pc- $\text{cytb}_6\text{f}$  unbinding occurs. The AFM probe was functionalized with pre-oxidized Pc: **(C)** topography image, **(D)** adhesion image. AFM probe functionalized with pre-reduced Pc: **(E)** topography image, **(F)** adhesion image. ‘Interaction hotspots’ where high ( $>50$  pN) unbinding forces were recorded on the membrane surface are highlighted with circles. The scale bars in **(C)** and **(E)** are 100 nm.

**Figure 3.** Analysis and quantification of force-distance data obtained on grana thylakoid membranes isolated from spinach by affinity-mapping AFM.

(A) Typical retract regions of the force-distance curves recorded using pre-oxidized Pc-functionalized probes on three different regions of the DG membrane sample attached to the mica. The orange curve was recorded on the mica surface with a peak unbinding force (uf) of ~450 pN at a probe-sample rupture length (rl) of ~0.1 nm indicating a non-specific interaction between the main body of the AFM probe and the mica. The red curve was recorded on the membrane surface and shows no distinct peak, indicating that there is little interaction with the AFM probe as it is retracted. The cyan curve was recorded at one of the ‘interaction hotspots’ on the membrane surface where a high (>50 pN) unbinding force was recorded, in this case ~300 pN, at a probe-sample rupture length of ~10 nm, which is consistent with the length of the SM(PEG)<sub>24</sub> linker with which the Pc is attached to the probe, indicating the detection of a specific Pc-cytb<sub>6</sub>f interaction. (B) Peak unbinding force versus rupture length for 276 force-distance curves recorded at membrane locations with high (>50 pN) unbinding force (red squares), and associated histograms for force (purple) and rupture length (blue) with average values shown using pre-oxidized Pc-functionalized probes on DG membranes.

**Figure 4.** Affinity-mapping AFM of grana membranes prepared under different conditions using a probe functionalized with Pc.

(A) Estimated percentage of cytb<sub>6</sub>f dimers detected by affinity-mapping AFM using various conditions as indicated on the graph (±S.E.). DG membranes were incubated with 10 μM DCMU to lower production of reduced PQ, (B) topography image, (C) adhesion image. DG membranes were incubated with 50 μM pre-oxidized Pc to block cytb<sub>6</sub>f binding sites on the membrane surface, (D) topography image, (E) adhesion image. αG membranes, (F) topography image, (G) adhesion image. SG membranes, (H) topography image, (I) adhesion image. ‘Interaction hotspots’ where high (>50 pN) unbinding forces were recorded on the membrane surface are highlighted with circles. The scale bars are 100 nm.

**Figure 5.** cytb<sub>6</sub>f/PSII nanodomain structure revealed by affinity-mapping AFM of DG membranes isolated from spinach.

(A) Topography image overlaid with the simultaneously recorded adhesion image, the interaction hotspots due to Pc-cytb<sub>6</sub>f binding are shown as green squares. The colored lines denote topographic cross-sections, also shown in (B), and the green squares denote the interaction hotspots along these cross-sections; the topographic features are assigned to either cytb<sub>6</sub>f (c) or PSII (PSII). (C) High resolution AFM image of a cytb<sub>6</sub>f/PSII nanodomain and (D) fitted with the membrane-extrinsic parts of the crystal structures of the cytb<sub>6</sub>f (purple) and PSII (green) dimers (PDB codes: 1S5L and 1UM3 respectively). The scale bar in (A) is 50 nm and in (C) is 50 nm.

**Table 1. Chlorophyll and cytochrome content of thylakoid membrane preparations. P700**

(PSI)/ chlorophyll cytochrome f/ chlorophyll and molar chlorophyll a/b ratio in grana membranes prepared by different techniques ( $\pm$ S.E.).

Membrane preparation	chlorophyll a/b ratio	P700 (PSI)/ chlorophyll ratio (mmol/mol)	cytochrome f/ chlorophyll ratio (mmol/mol)
$\alpha$ -DM Grana	2.23 $\pm$ 0.1	0.75 $\pm$ 0.1	n.d.
Digitonin Grana	2.41 $\pm$ 0.1	1.49 $\pm$ 0.1	1.41 $\pm$ 0.1
Sonication Grana	2.38 $\pm$ 0.1	1.42 $\pm$ 0.1	1.43 $\pm$ 0.1
Intact thylakoids	3.33 $\pm$ 0.1	2.63 $\pm$ 0.2	1.33 $\pm$ 0.1