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Probing the local lipid environment of the Rhodobacter sphaeroides cytochrome $bc_1$ and Synechocystis sp. PCC 6803 cytochrome $b_6f$ complexes with styrene maleic acid

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Abstract

Intracytoplasmic vesicles (chromatophores) in the photosynthetic bacterium *Rhodobacter sphaeroides* represent a minimal structural and functional unit for absorbing photons and utilising their energy for the generation of ATP. The cytochrome *bc*$_1$ complex (*cytbc$_1$*) is one of the four major components of the chromatophore alongside the reaction centre-light harvesting 1-PufX core complex (RC-LH1-PufX), the light-harvesting 2 complex (LH2), and ATP synthase. Although the membrane organisation of these complexes is known, their local lipid environments have not been investigated. Here we utilise poly(styrene-alt-maleic acid) (SMA) co-polymers as a tool to simultaneously determine the local lipid environments of the RC-LH1-PufX, LH2 and *cytbc$_1$* complexes. SMA has previously been reported to effectively solubilise complexes in lipid-rich membrane regions whilst leaving lipid-poor ordered protein arrays intact. SMA solubilises *cytbc$_1$* complexes with an efficiency of nearly 70%, whereas solubilisation of RC-LH1-PufX and LH2 was only 10% and 22% respectively. This high susceptibility of *cytbc$_1$* to SMA solubilisation is consistent with this complex residing in a locally lipid-rich region. SMA solubilised *cytbc$_1$* complexes retain their native dimeric structure and co-purify with 56 ± 6 phospholipids from the chromatophore membrane. We extended this approach to the model cyanobacterium *Synechocystis* sp. PCC 6803, and show that the cytochrome *b$_6$f* complex (*cyt*$_6$f$_6$) and PSII complexes are susceptible to SMA solubilisation, suggesting they also reside in lipid-rich environments. Thus, lipid-rich membrane regions could be a general requirement for *cytbc$_1$/cyt*$_6$f$_6$ complexes, providing a favourable local solvent to promote rapid quinol/quinone binding and release at the Q$_0$ and Q$_i$ sites.
Keywords

SMA; cytochrome $bc_1$; cytochrome $b_6f$; Rba. sphaeroides; Synechocystis; quinone pool

Highlights

- SMA preferentially solubilises cyt$bc_1$ from chromatophore membranes
- Solubilised cyt$bc_1$ SMALPs contain dimeric complexes co-purified with 56 lipids
- SMA-resistant fractions contain RC-LH1-PufX and LH2 rich membrane patches
- The Rba. sphaeroides cyt$bc_1$ complex is likely to reside in a lipid-rich environment
- Similar results for Synechocystis suggest cyt$bc_1/b_6f$ may be universally lipid-rich

Abbreviations

Rhodobacter sphaeroides (Rba. sphaeroides); cytochrome $bc_1$ complex (cyt$bc_1$); cytochrome $b_6f$ complex (cyt$b_6f$); reaction centre light harvesting complex 1 PufX core (RC-LH1-PufX); light harvesting two complex (LH2); ubiquinone 10 (UQ$_{10}$); cytochrome $c_2$ (cyt$c_2$); proton motive force (PMF); atomic force microscopy (AFM); transmission electron microscopy (TEM); Synechocystis sp. PCC 6803 (Synechocystis 6803); photosystem I (PSI); photosystem II (PSII); poly(styrene-alt-maleic acid) (SMA); styrene-maleic acid lipid particles (SMALPs); n-Dodecyl-beta-Maltoside ($n$-DDM); Mass average molecular weight ($M_w$); wild-type (WT); column volume (CV); thin layer chromatography (TLC); high performance liquid chromatography (HPLC); phosphatidylglycerol (PG); phosphatidylethanolamine (PE); cardiolipin (CL); phosphatidylcholine (PC); sulfoquinovosyl diacylglycerol (SQDG)
Intracytoplasmic vesicles (chromatophores) in the photosynthetic bacterium *Rhodobacter (Rba.)* sphaeroides (Fig. 1A) are spherical invaginations of the membrane. These structures represent a minimal structural and functional unit for absorbing photons and utilising their energy to produce ATP via an efficient energy generation mechanism [1,2]. The cytochrome $bc_1$ complex (cyt$bc_1$) is one of the four major components of the chromatophore alongside the reaction centre-light harvesting 1-PufX core complex (RC-LH1-PufX), the light-harvesting 2 complex (LH2), and ATP synthase [1]. The stoichiometry and organisation of these complexes, revealed by mass spectrometry and atomic force microscopy (AFM) [1,3], appears to be optimised for photosynthetic growth at low light intensities below 50 $\mu$mol m$^{-2}$ s$^{-1}$ [1,2,4], when *Rba. sphaeroides* cells contain up to 1500 intracytoplasmic vesicles [5], observed both as single and budded structures [6,7]. Chromatophores house short rows of dimeric RC-LH1-PufX complexes surrounded by tens of LH2 complexes [1,2], which provide a variable antenna that can adjust to changeable light levels [5]. Close packing between these complexes promotes rapid and efficient energy transfer and trapping. The overall trapping time of ~60 ps [8] is accounted for by energy migration between LH2 complexes, and from LH2 to LH1, on a picosecond timescale [9–11]. Transfer from the encircling LH1 ring to the special pair (P) BChl dimer within the RC (Fig. 1B) [12] drives picosecond charge separation within the RC [13].

Following two rounds of charge separation, and re-reduction of oxidised P (Fig. 1B, red arrow) by cytochrome $c_2$ (cyt$c_2$), a ubiquinone 10 (UQ$_{10}$) molecule bound to the Q$_b$ site becomes fully reduced, and binds two protons from the cytoplasm to generate a quinol (Fig. 1B) (see [13] for a comprehensive review of the RC). The reduced quinol leaves the RC, traverses the surrounding LH1 antenna through a pore created by the PufX polypeptide [14], and diffuses within the membrane to the cyt$bc_1$ complex [15] via the free quinone pool (Fig. 1B black arrows). The cyt$bc_1$ generates a proton motive force (PMF) via a modified Q-cycle [16] in which quinol binds to the Q$_b$ site releasing two protons into the lumen of the chromatophore (Fig. 1B orange arrows). One electron is transferred along the high potential chain via the Rieske Fe-S subunit to reduce an oxidised cyt$c_2$, which diffuses to a photooxidised RC to reduce P, completing the cyclic electron transfer chain. The second electron enters the low potential chain via two $b$-type cytochromes to reduce a second UQ$_{10}$ molecule bound at the Q$_i$ site. Arrival of another quinol and a repeat of the above reactions generates an ‘extra’ quinol at the Q$_i$ site, so this Q-cycle mechanism effectively amplifies the yield of protons for each absorbed photon [16]. The PMF generated by these processes is utilised by ATP synthase to generate ATP from ADP and inorganic phosphate. There are approximately 4 cyt$bc_1$
complex dimers present in each chromatophore, and this number is rate limiting for photosynthetic ATP formation [1,2].

Cartron et al. [1] demonstrated that dimeric cyt$\text{bc}_1$ complexes and dimeric RC-LH1-PufX complexes are found in close proximity, but not necessarily in direct contact, an arrangement that minimises the cycling time for diffusion of $Q/QH_2$ and cytochrome $c$ between them. It was also shown that the cyt$\text{bc}_1$ complex was much more susceptible to detergent solubilisation than RC-LH1-PufX and LH2 suggesting that its local environment may be lipid-rich, consistent with a proposed quinone- and lipid-rich phase surrounding the cyt$\text{bc}_1$ complexes [17]. The cytochrome $b$$_6$f (cyt$\text{b}_6$f) complex in cyanobacteria, algae and plants [18], which performs an analogous function to function cyt$\text{bc}_1$ in purple bacteria, could also sit in a lipid and quinone-rich nanoenvironment. AFM of plant thylakoids showed that cyt$\text{b}_6$f complexes are found within 20 nm of photosystem II (PSII) RCs, again consistent with predictions from kinetic studies showing confinement of quinones in the local PSII- cyt$\text{b}_6$f environment [19–21].

New approaches are required to characterise the membrane environments of the photosynthetic complexes described, and poly(styrene-alt-maleic acid) (SMA) polymers represent one such tool. In recent years there has been growing interest in SMA for membrane-protein solubilisation; once converted to the acid form 2:1 or 3:1 styrene to maleic-acid ratio polymers can effectively solubilise biological membranes and their constituent protein complexes [22–27]. Unlike detergents, SMAs do not remove the annular lipids of membrane protein complexes, instead forming nanodiscs in which the protein is embedded in a belt of lipids from the source membrane stabilised by the polymer. These structures are termed native nanodiscs or styrene-maleic acid lipid particles (SMALPs) and have been found to improve the stability of complexes as well as give more native-like biophysical properties when compared to preparations using detergents [22,23,28–33]. The resultant SMALPs are amenable to both biophysical analysis and characterisation of the co-purified lipids, providing insight into the local lipid environments of the proteins within [28,31,34]. A recent investigation of the properties of SMA has revealed that these polymers effectively preserve even weak protein-protein contacts, such as those involved in formation of RC-LH1-PufX arrays, allowing for enrichment of native large-scale protein architectures [27]. To this end we have utilised SMA as a tool to probe the local lipid environment of the RC-LH1-PufX, LH2 and cyt$\text{bc}_1$ complexes from $Rba.$ sphaeroides chromatophores. As hypothesised by previous studies we find that the cyt$\text{bc}_1$ complex resides in a lipid-rich environment whereas the RC-LH1-PufX and LH2 complexes reside in relatively lipid-poor domains. By purifying cyt$\text{bc}_1$ SMALPs we further characterised the local lipid environment of this complex. We extended this methodology to membranes of the model oxygenic photosynthetic
cyanobacterium *Synechocystis* sp. PCC 6803 and find that the PSII and cyt*bf* complexes are highly susceptible to SMA solubilisation, indicating they also reside in lipid-rich environments. Together these data suggest that lipid-rich membrane regions provide a favourable local solvent to promote rapid quinol/quinone binding and release at the Q₀ and Qᵢ sites of cyt*bc₁* and cyt*bf* complexes.

**Fig. 1.** A: Molecular model of the chromatophore comprised of LH2 (green), LH1 (Red), PufX (beige), the reaction centre (RC, blue), cyt*bc₁* (purple) and ATP synthase (yellow). This figure was produced from a model featured in refs [1,2]. B: Schematic representation of LH2, RC-LH1-PufX and cyt*bc₁* as a cross-section of part of the chromatophore vesicle in panel A. Two successive photons (red) incident on an LH2 complex, then excitation energy transfer (wavy yellow arrows) to LH1 then a RC, drives two charge separations eventually
producing a quinol (QH$_2$), which migrates via the quinone pool to a nearby cyt$_{bc1}$ complex. Electron holes at the RC are filled by reduced cyt$_c$ which receives electrons from the cyt$_{bc1}$ complex (red arrows), completing this cyclic electron transfer process. These protons are finally utilised by the ATP synthase to generate ATP from ADP and inorganic phosphate (P$_i$). Diffusion of mobile electron carriers at the membrane surface and Q/QH$_2$ within the membrane bilayer are shown with black arrows. Orange arrows denote movement of protons; for every four turnovers at the RC, six protons accumulate in the lumen; the diagram shows the average for two RC turnovers, i.e. for each quinol produced.

2. Materials and Methods

2.1. Preparation of SMA

Xiran SZ30010 polystyrene-maleic anhydride (2:1 ratio styrene to maleic anhydride, 10 kDa mass average molecular weight (M$_w$)) was a generous gift from Polyscope EU. The polymer was converted to the acid-form by refluxing in excess KOH as previously described [27] at a concentration of 5% w/v. The pH was adjusted to 8.0 with solid KOH prior to use to avoid dilution of the solution.

2.2. Growth of *Rhodobacter sphaeroides*

The construction and growth of a *Rhodobacter sphaeroides* mutant containing a His$_{10}$ tagged cyt$_{bc1}$ c-subunit (PetC) was described previously [1]. For solubilisation assays cells were grown in M22+ medium either photosynthetically in 1 L of medium with illumination from ORSAM CLASSIC 116 W halogen light bulbs under “medium light” (approximately 100 µmol m$^{-2}$ s$^{-1}$ illumination for 16 hr) or “low light” (approximately 30 µmol m$^{-2}$ s$^{-1}$ illumination for 48 hr) conditions, or semi-aerobically in 1.6 L of medium in the dark for 48 hr. Cells (8 l) for protein preparations were grown photosynthetically under approximately 30 µmol m$^{-2}$ s$^{-1}$ illumination for 72 hr. Cells were harvested by centrifugation at 4 000 RCF for 20 min and stored at −20 °C until required.

2.3. Generation of histidine-tagged cyt$_{bf}$ in *Synechocystis* sp. PCC 6803

A DNA sequence encoding the thrombin cleavable His$_{10}$ tag from pET52b (Novagen) was added in frame to the end of the petA gene (sll1317), which encodes apocytochrome $f$. This is the same tag added to the C-terminus of the *Rba. sphaeroides* cytochrome $c_1$ component of the cyt$_{bc1}$ complex described in Section 2.2 [1]. The tag was followed by a stop codon and the first 25 bp of the chloramphenicol acetyl transferase (*cat*) cassette from pACYC184. This construct was synthesised (Integrated DNA Technologies) and the *cat* cassette and 500 bp of DNA immediately downstream of the petA gene were amplified separately by PCR, from pACYC184 with primers catF/catR or from *Synechocystis* sp. PCC 6803 genomic DNA with primers petA_ds_F/petA_ds_R, respectively. These
three fragments were joined by overlap extension PCR using primers petA_F and petA_ds_R, and the
resulting product was cloned into the BamHI and HindIII sites of pUC18. The sequence verified (GATC
Biotech) fragment was excised with BamHI/HindIII and introduced to wild-type (WT) Synechocystis
as described previously [35]. Transfomants were selected using 5 µg mL⁻¹ chloramphenicol and
gene copies segregated by sequential doubling of the antibiotic concentration to 40 µg mL⁻¹.
Segregation was confirmed by PCR screening with oligos petA_screen_F and petA_screen_R,
resulting in a product of 1941 bp for transformants, compared to 1076 bp in the WT. The petA locus
amplified from transformant genomic DNA was sequenced to ensure the His-tag was in frame with
the petA gene.

2.4. Preparation of Rba. sphaeroides chromatophore membranes

Cells were suspended in 20 mM Tris pH 8 containing a few crystals of DNase1 and Roche cOmplete
EDTA-free protease inhibitors. Cells were lysed either by two passes through a French press
(AminCo, USA), or a single pass through a cell disruptor (Constant systems), both at 20 000 PSI.
Insoluble material was removed by centrifugation at 18 459 RCF (avg) for 15 min at 4 °C. Soluble
material was loaded onto 40/15% w/w sucrose step gradients and centrifuged at 107 400 RCF (avg)
for 10 hr at 4 °C. Membranes were harvested from the 40-15% sucrose solution interface and stored
in aliquots at -20 °C.

2.5. Preparation of Synechocystis sp. PCC 6803 thylakoid membranes

Synechocystis sp. PCC 6803 cells producing C-terminally His-tagged PetA were grown to an OD₇₅₀ of
approximately 1 in a volume of 16 L BG11 medium with aeration under 100 µmol m⁻¹ min⁻¹
illumination from ORSAM CLASSIC 116 W halogen light bulbs [36] and harvested by centrifugation at
17 700 RCF (avg) at 4 °C for 20 min. Pellets from 8 L of culture were washed and re-suspended in
thylakoid buffer (25 mM sodium phosphate pH 7.4, 5 mM MgCl₂ and 200 mM NaCl and cOmplete
EDTA-free Protease Inhibitors [Roche]) and mixed with an equal volume of 0.1 mm glass beads
(BioSpec).

To prepare membranes for SMA solubilisation assays the cells were broken in a Mini-Beadbeater-16
(BioSpec) for eight 20 s cycles with samples cooled on ice between each cycle. The cell lysate was
pelleted at 38 000 RCF (avg) at 4°C for 30 min before being re-suspended in thylakoid buffer.
Aliquots of 3 mL cell lysate were loaded onto multiple sucrose step gradients consisting of 2 mL 50%
w/w sucrose and 8 mL 30% w/w sucrose and centrifuged for 30 min at 111 000 RCF (avg) in a
Beckman SW41 Ti rotor for 1 hr at 4 °C. The bands containing thylakoid membranes were harvested
from the interface between the 30 and 50% sucrose steps.
To prepare membranes for the purification of His<sub>10</sub>-tagged cyt<sub>b<sub>6</sub>

beating using ten 55 s cycles. Soluble and membrane proteins were separated by centrifugation at

38 000 RCF (avg) at 4°C for 30 min and the membranes were re-suspended in 100 mL 20 mM Tris pH

8 containing 200 mM NaCl.

2.6. Solubilisation assays

The solubilisation assay was adapted from the method described by Swainsbury et al. [27]. 2 mL Rba. spphaeroides membranes with an OD<sub>850</sub> of 3 (1 cm pathlength) were prepared in solutions of 20 mM Tris pH 8 containing 200 mM NaCl and either 2.5% w/v SMA, 3% w/v n-Dodecyl-beta-Maltoside Detergent (β-DDM) or without solubilising agents. Synechocystis sp. PCC 6803 membranes were

solubilised at OD 5 (680 nm, 1 cm pathlength) in 25 mM sodium phosphate buffer pH 7.4 containing 200 mM NaCl, 5 mM MgCl<sub>2</sub> and 2.5% SMA in a total volume of 2 mL. Samples were incubated at room temperature in the dark for 1 hr.

A volume of 1 mL was centrifuged at 100 000 RCF for 2 hr at 4 °C. After this, 0.9 mL was removed with care taken not to disturb the pellet; this sample will be termed the “soluble fraction”. The remaining 1 mL of each sample, hereafter referred to as the “total fraction”, was stored at 4 °C in the dark until required.

Spectra of the “total” and “soluble” fractions were collected in the short (0.33 cm) path of a quartz semi-micro cuvette between 1000 and 400 nm. Samples were then transferred into 1 mL disposable cuvettes and spectra were collected before and after the addition of a few grains of sodium dithionite.

Spectra collected on the 0.33 cm path were processed by scatter correcting and deconvoluting the contributions of RC-LH1-PufX and LH2 using a modified version of an Excel spreadsheet by Prof O’Haver available at [https://terpconnect.umd.edu/~toh/spectrum/CurveFittingB.html](https://terpconnect.umd.edu/~toh/spectrum/CurveFittingB.html). The spreadsheet adds spectra of the two complexes and a scatter curve to achieve a best fit to the data and returns spectra for the three components according to their fitted contributions. The scatter curve used was calculated using $\Lambda^{-2.6}$, and the RC-LH1-PufX and LH2 reference spectra were produced from proteins purified as described elsewhere [37,38]. The reference spectra used are shown normalised to their maxima in Supplementary Fig. 1 panel E. The extraction efficiency of RC-LH1-PufX and LH2 was calculated from the difference in absorbance at 875 and 850 nm respectively, in the “total” and “soluble” fractions. The 1 cm pathlength spectra were processed by generating oxidised minus reduced spectra and using the difference at 560 nm between the “total” and “soluble” fractions to estimate extraction efficiency.
For *Synechocystis* sp. PCC 6803 membranes the same procedure was used, except samples had an OD$_{680}$ of 5; the short pathlength UV/Vis spectra were collected in a 0.2 cm quartz cuvette and reference spectra for PSI and PSII were used for deconvolution using the procedure described above for RC-LH1-PufX and LH2 using the reference spectra presented in Supplementary Fig. 5C [39].

The extractions of cyt$bc_1$ and cyt$bf$ were also estimated by haem blot [40]. 20 µL volumes from “soluble” and “total” fractions were separated on SDS PAGE gels, transferred to a polyvinylidene difluoride (PVDF) membrane and the cytochrome $c/f$ subunits were visualised using Westar 2.0 solution (Cyanagen). Band intensities were integrated using ImageJ [41] and extraction efficiencies were calculated by comparison of the band intensity of “total” and “soluble” fractions.

### 2.7. Native PAGE electrophoresis

*Synechocystis* sp. PCC 6803 membranes with an OD$_{680}$ of 6, 12 and 24 were solubilised in 2.5 % w/v SMA as described in section 2.6. 15 µL of each sample was incubated in 4% β-DDM for 1 hr at room temperature in the dark. Samples were then diluted 2-fold in either clear native buffer (125 mM Tris pH 6.8, 30% glycerol) or blue native buffer (125 mM Tris pH 6.8, 30% v/v glycerol, 0.01% w/v bromophenol blue). Samples were run on NuPAGE Tris-acetate 3-8% gels (Novex) at 150 v for ~2 hr. For clear native page the upper buffer was supplemented with 0.05% w/v deoxycholate and 0.04% β-DDM and blue native gels were supplemented 1 mL blue native additive.

Gels were imaged using an Amasham Imager 600 in colour. Clear native gels were also imaged by fluorescence with excitation at 460 nm monitoring emission using the cy5 filter with 12 s exposure. Bands were assigned according to [42].

### 2.8. Purification of cyt$bc_1$ and cyt$bf$ complexes

Membranes from a 4 L culture of photosynthetically grown *Rba. sphaeroides* cells, or 8 L *Synechocystis* sp. PCC 6803 cells were solubilised in 20 mM Tris pH 8 containing 200 mM NaCl and 1.5% w/v SMA at room temperature for 1 hr in the dark. Insoluble material was removed by centrifugation at 113 000 RCF (avg) for 1 hr at 4 ºC. Solubilised cyt$bc_1$ or cyt$bf$ complexes were bound to a 20 mL HisPrep FF Ni-NTA column (GE Healthcare) pre-equilibrated with 5 column volumes (CV) binding buffer (20 mM Tris pH 8 containing 200 mM NaCl and 20 mM imidazole) by recycling overnight at 5 mL min$^{-1}$. The column was washed with 20 CV binding buffer followed by 10 CV binding buffer containing 40 mM imidazole. The cyt$bc_1$ or cyt$bf$ complexes were eluted in 20 mM Tris pH 8 containing 200 mM NaCl and 250 mM imidazole. For *Rba. sphaeroides* cyt$bc_1$ further purification and buffer exchange were performed by size-exclusion chromatography on a HiPrep Sephacryl 16/60 S-300 column (GE Healthcare) in 20 mM Tris pH 8 containing 200 mM NaCl.
Fractions with an $A_{415}/A_{280}$ above 1.25 were concentrated and stored at -80 °C until required. The concentrations of cyt$bc_1$ haem $b$ and haem $c$ in the preparation were determined using dithionite reduced samples with extinction coefficients of $\varepsilon_{561-575}$ of 22 mM$^{-1}$ cm$^{-1}$ and $\varepsilon_{551-540}$ of 19 mM$^{-1}$ cm$^{-1}$, respectively, as described in [1].

2.9. Thin layer chromatography

Thin layer chromatography (TLC) was performed according to [43] with some modifications. Lipids were extracted from ~0.5 nmol cyt$bc_1$ SMALPs or ~0.4 OD$_{850}$ units of membranes in 50 µl 1:1 methanol:chloroform. Samples were loaded alongside pure lipid standards on Whatman Partsil Diamond LK6DF TLC plates. Plates were developed in either 85:15:10:3.5 or 85:25:10:3.5 chloroform:methanol:acetic acid:water (by volume) for 30 min. Lipids were visualised by incubating the plate in 50% v/v $H_2SO_4$ for 30 s followed by heating at 160 °C for 60 min. Plates were imaged and band intensities were integrated in Image J [41]. Data from both conditions were combined to give six data-sets for membranes and eight for SMALPs and all data were normalised to the intensity of the phosphatidylglycerol (PG) band.

2.10. Ubiquinone-10 quantification

Ubiquinone-10 (UQ$_{10}$) was quantified according to “Determination of Coenzyme Q10 by High Pressure Liquid Chromatography” customer application brief #101 by Dionex and the method in [44] with several modifications. Standards were prepared by dissolving pure UQ$_{10}$ in 50:50 chloroform:methanol containing 0.02% w/v ferric chloride. Lipids were extracted from 0.5 nmol cyt$bc_1$ in the same solvent. Samples were analysed by high performance liquid chromatography (HPLC) using aBeckman Coulter ODS 4.6 mm x 2.5 cm C18 column and eluted at 1 mL min$^{-1}$ in 80:20 v/v methanol:2-propanol over 45 min. Peaks at 29.5 and 32 min were integrated and used for calculations. Three samples of cyt$bc_1$ were analysed and the calculated concentrations of UQ$_{10}$ were averaged.

2.11. Lipid quantification

Phospholipids were quantified according to [1]. A total of 0.46 nmol cyt$bc_1$ was dissolved in approximately 50 µl chloroform in pre-cleaned glass test-tubes. Phosphate standards were prepared using solutions of 0, 0.25, 0.5, 1, 5, 10, 20, 50, 75, 100 and 200 nMol KH$_2$PO$_4$ dissolved in 50 µL chloroform. All samples were dried at 140 °C for 20 min. Lipids were hydrolysed by adding 0.15 mL perchloric acid and incubating at 180 °C for 2 h. Phosphate was visualised colorimetrically by adding 0.5 mL $H_2O$, 1.25% w/v ammonium hepta-molybdate and 5% w/v 0.2 mL ascorbic acid followed by
incubation at 100 °C for 5 min. Spectra were collected and the absorbance at 800 nm was used to determine phosphate concentrations.

2.12. Preparation of nanogold labelled membranes

Membranes were labelled with nanogold according to Cartron et al. [1]. Membranes (OD₈₅₀ of 10) were incubated for 1 hr in 20 mM Tris pH 8, or 20 mM Tris pH 8 containing 0.02 % β-DDM or 2.5 % SMA for 1 h in a final volume of 2.4 mL. After incubation 0.6 mL 0.5 mM 5 nm Ni-NTA-Nanogold (Nanoprobes) was added and samples were incubated for an additional 1 hr. Samples were loaded onto gradients of 50/40/30/20/15 % w/v sucrose in 20 mM Tris pH 8 and centrifuged at 178 305 RCF (avg) for 2 hr at 4 °C. Fractions were collected from the gradients in 1 mL volumes with a peristaltic pump and spectra were collected between 1000 and 400 nm. A second set of membranes were prepared as above without the addition of nanogold. The relative cytbc₁ content of gradient fractions was estimated by haem blot as described for solubilisation assays (section 2.6).

2.13. Transmission electron microscopy

Samples, either 5 µL 1 µM cytbc₁ or OD₈₅₀ 0.3 membranes, were incubated on glow-discharged carbon-coated copper grids for 30 s followed by washing with deionised water and staining with 0.75 % w/v uranyl formate for 30 s. Grids were imaged in a Phillips CM100 TEM equipped with a Gatan Ultrascan 667 camera at between 27 000 and 52 000 x magnification.

2.14. Dynamic light scattering

Solutions containing 1 µM cytbc₁ were prepared in 20 mM Tris pH 8 containing 200 mM NaCl and aggregates were removed by centrifugation at 15 000 RCF in a benchtop centrifuge for 10 min. Samples were filtered through a 0.2 µm syringe filter prior to measurement in a Zetasizer nano ZS in a 1 mL cuvette collecting three sets of ten x 10 s measurements at 25 °C.

3. Results

3.1. Solubilisation of chromatophore membranes with SMA

SMA has been shown to preferentially solubilise proteins within lipid rich membrane environments whilst leaving lipid-poor domains, such as densely packed antenna arrays, intact [27]. In order to probe the local lipid environment of the cytbc₁ complex we performed solubilisation trials with SMA and measured its ability to solubilise RC-LH1-PufX, LH2 and cytbc₁. We selected 10 kDa Mₘ SMA with a 2:1 styrene to maleic acid ratio for this study. This polymer has previously been shown to efficiently solubilise proteins residing in lipid-rich environments from the Rba. sphaeroides
membranes whereas RC-LH1-PufX arrays are left in-tact [27]. Thus, this SMA should allow the best
possible discrimination of lipid-rich complexes and lipid-poor domains. As the level of LH2 expression
can affect the morphology of chromatophore membranes, solubilisation trials were performed on
membranes prepared from cells grown either semi-aerobically in the dark, or photosynthetically
under ~30 µmol m⁻² s⁻¹ (low light) for 48 hr, or ~100 µmol m⁻² s⁻¹ (medium light) for 16 hr. By
deconvolution of the spectra the molar ratios of LH2 to RC-LH1-PufX in these membranes were
estimated to be 2.5:1, 2.4:1 and 2.0:1, respectively.

To estimate the percentage of each complex extracted, UV/Vis spectra of SMA solubilised
membranes were collected before and after ultracentrifugation. The contributions of RC-LH1-PufX
and LH2 complexes, and of light scattering, were deconvoluted by fitting spectra of pure complexes
and a calculated scatter curve to a spectrum of the SMA treated chromatophores. Example spectra
and their deconvoluted components are shown in Supplementary Fig. 1. The deconvoluted spectra
were used to estimate the solubilisation efficiencies for RC-LH1-PufX and LH2, shown in Fig. 2 with
blue and green bars respectively, calculated by the differences in absorbance before and after
removal of the insoluble material. The solubilisation of cytbc₁ was estimated by comparing the
change in absorbance at 560 nm upon dithionite treatment before and after ultracentrifugation (Fig.
2, magenta bars). A second estimate was made by detection of the covalently linked c-type
cytochrome of the 30 kDa cyt[c₁ subunit by haem staining [40] and comparing the intensity of the
bands in the total and soluble fractions (Fig. 2, shaded bars). Example haem blots and difference
spectra are shown in Supplementary Fig. 2. These blots also contained a 24 kDa band, consistent
with the apparent mass of the membrane-associated cyt[c [45], which was omitted from these
calculations.
Fig. 2. Extraction efficiencies of RC-LH1-PufX (blue), LH2 (green) and the cytbc1 complex (magenta) in membranes prepared from cells grown under semi-aerobic (SA), or low light (LL) and medium light (ML) photosynthetic conditions. The left panel shows solubilisations in 2.5% w/v SMA, the centre panel shows low-light membranes solubilised in 3% w/v β-DDM, and the right panel shows results where no solubilising agents were added. Solid bars show values obtained by spectroscopy and hatched bars show values for cytbc1 by haem staining. Error bars show standard error of the mean for three replicates.

Treatment of chromatophore membranes with 2.5% w/v SMA revealed that the major photosynthetic complexes are not solubilised with uniform efficiency. As shown in Fig. 2, the RC-LH1-PufX complexes are highly resistant to SMA solubilisation, and are extracted with 3.3 ± 1.1 % efficiency from membranes from semi-aerobically grown cells. This value is similar to that previously determined by Swainsbury et al. [27] for RC-LH1-PufX containing membranes lacking LH2, from cells grown under the same conditions. The low solubilisation efficiency arises because SMA is unable to disrupt the highly ordered and closely packed arrays formed by these complexes [27]. The solubilisation efficiency increases slightly in membranes from low and medium light grown cells at 6.4 ± 0.2 and 9.5 ± 0.3 %, respectively. The LH2 complex is slightly less resistant to SMA solubilisation with efficiencies of 7.8 ± 1.7, 17.9 ± 0.3 and 22.0 ± 0.4 % for membranes from semi aerobically grown, low light and medium light grown cells, respectively. This level of LH2 solubilisation is similar to that in trials performed on membranes from strains lacking the RC-LH1-PufX complex (data not shown). It should be noted that the increase in solubilisation efficiency correlates with the reduction...
of the LH2 to RC-LH1-PufX ratio. This is presumably a consequence of the somewhat different arrangements of the RC-LH1-PufX and LH2 within these membranes. Estimated solubilisation of cytb_{c1} was much more efficient at 64 ± 4, 62 ± 9 and 70 ± 7 % for semi aerobic, low light and medium light membranes respectively, as determined by reduced-oxidised spectra shown in Fig. 2 by the solid magenta bars (see Supplementary Fig. 2 for raw spectra). Haem blots also demonstrate that cytb_{c1} is very effectively solubilised by SMA with estimated efficiencies of 92 ± 2, 87 ± 12 and 88 ± 24 % for semi aerobic, low light and medium light membranes respectively, as shown by hatched bars in Fig. 2 (see Supplementary Fig. 2 for raw data). Additionally, there was no apparent trend in relation to the LH2 to RC-LH1-PufX ratio. To demonstrate that this effect is specific to SMA, control solubilisations were performed using 3 % w/v \text{\textit{\text{n}}}-DDM. Solubilisation efficiency was above 74% under all growth conditions for all complexes (only data from low light membranes are shown for clarity). In the absence of solubilising agents, solubilisation was less than 2 % for the RC-LH1-PufX and LH2 complexes and below the limit of detection for cyt_{bc1}.

3.2. Characterisation of insoluble material after SMA treatment

To examine RC-LH1-PufX and LH2 after treatment of chromatophores with SMA, solubilisation mixtures were fractionated on sucrose density gradients then analysed by UV/Vis spectroscopy, haem blotting and negative-stain TEM imaging. For comparative purposes we also analysed chromatophores treated with sub-solubilising concentrations (0.02 % w/v) of \text{\textit{\text{\textbeta}}}DDM to generate flattened membrane patches, which are more amenable to nanogold labelling and TEM imaging than untreated chromatophores [1]. This \text{\textit{\textbeta}}DDM concentration allows imaging of membrane patches containing all of their native complexes. Panel A in Fig. 3 shows sucrose gradients of \textit{Rba. sphaeroides} chromatophores after these treatments. The untreated membranes form an abundant band at the 20/30% w/w sucrose interface (at the boundary of fractions 4 and 5 in Fig 3A). After treatment with 0.02 % w/v \text{\textit{\text{\textbeta}}}DDM this band is still present suggesting the membranes remain intact, although the band appears more diffuse suggesting the population has become more heterogeneous. After treatment in fully solubilising concentrations of SMA, a band remains at the 20/30% sucrose interface, suggesting that much of the membrane remains intact after SMA solubilisation. These gradients were separated into 1 mL fractions, with fraction 1 being at the bottom and 11 at the top of the tube. The membrane bands (fraction 5 for all samples) have distinctive absorbance spectra for RC-LH1-PufX and LH2 demonstrating that the integrity of these complexes has been maintained. As shown in Fig. 3 panel B, haem blotting reveals that for the untreated and \text{\textit{\text{\textbeta}}}DDM-treated samples the cyt_{c1} subunit remained associated with the RC-LH1-PufX and LH2 containing membrane band (raw data are shown in Supplementary Fig. 3). However, after SMA solubilisation the cyt_{c1} subunit is found in the upper portion of the gradient and no longer co-
migrates with the remaining membrane material, showing that cytbc₁ has been solubilised and separated from the RC-LH1-PufX and LH2 complexes.

Fig. 3. A: Images of 15-50% w/w sucrose density gradients of untreated, 0.02% DDM treated and SMA-solubilised chromatophore membranes. The scale on the right-hand side shows the approximate positions of the fractions taken from the gradient. B: UV/Vis absorbance profiles of fractions from the sucrose gradients shown in panel A. Absorbance of LH1/LH2 (squares connected by lines, based on absorbance at 850 nm) and relative intensities of the cytbc₁ subunit from haem blots (bars) are shown.

In a second set of gradients Ni-NTA nanogold was used to selectively label the His₁₀-tag of the cytbc₁ complexes. As shown in Supplementary Fig. 4A, a significant band was observed at the 40/30% w/v interface, below the position of the major band in Fig. 3A at the 20/30% interface, for untreated and β-DDM treated samples. Free nanogold migrated to the bottom of the tube (data not shown). For the SMA solubilised sample the lower band (40/30% interface) was very faint with the upper band (20/30%) being prominent, suggesting a reduced degree of membrane labelling. TEM images of grids prepared from each band revealed that the upper bands of untreated and β-DDM treated membranes contain mostly unlabelled circular objects with diameters of approximately 50 nm (shown in Supplementary Fig. 4B and C, zoomed images on the left show typical particles), consistent with the expected morphology of intact closed chromatophores. The lower band of the untreated membranes also contained similar structures with a low level of nanogold labelling, as shown in Supplementary Fig. 4D. In contrast the lower band of the β-DDM treated complexes contained large patches of 100-200 nm in diameter (shown in Fig. 4A and Supplementary Fig. 4E),
with multiple nanogold beads often appearing in pairs. Measurement of the patch area and number of nanogold beads gave a labelling density of $0.41 \pm 0.19$ beads per square micron (from 5 patches with an average area of $17500 \pm 10400 \mu m^2$, errors represent standard deviations of values for the individual patches). This result is consistent with the expected dimeric structure of the cyt$bc_1$ complexes and is similar to images obtained by Cartron et al. [1] following the same sample preparation procedure. Together with the UV/Vis spectra we conclude that the membrane patches in the β-DDM-treated lower band are flattened chromatophores containing physiological quantities of cyt$bc_1$ complexes. The TEM images of the SMA-solubilised and β-DDM treated samples are clearly different. Membrane patches in the upper band were smaller than those from the β-DDM treated samples at 50-100 nm in size and did not resemble intact chromatophores (shown in Fig. 4B and Supplementary Fig. 4E). These patches contained between 0 and 2 nanogold labels on average and each label was observed in isolation with a density of $0.17 \pm 0.11$ beads per square micron (from five patches with an average area of $5900 \pm 1300 \mu m^2$). The lower SMA band also contained 50-100 nm membrane patches, which had a slightly higher degree of nanogold labelling ($0.3 \pm 0.23$ beads per square micron, from 5 patches with average area of $8100 \pm 5200 \mu m^2$) than the upper band, although this fraction accounted for only a minor proportion of the pigmented material. Together, data from the sucrose gradients in Fig. 3 and TEM imaging suggest that the majority of membrane fragments remaining after SMA solubilisation (seen in the upper SMA band, Fig 4B) are smaller than those expected for flattened intact chromatophore membranes (DDM lower band, Fig 4A), and have a greatly reduced cyt$bc_1$ content. This is likely to be the result of preferential solubilisation of lipid rich regions containing cyt$bc_1$ by SMA, so they no longer co-migrate with the protein-rich RC-LH1-PufX and LH2 arrays as shown in Fig. 3.
Fig. 4. Low resolution TEM images of typical membrane patches from nanogold-labelled chromatophores treated with 0.02% β-DDM (A, left panels) or after complete solubilisation in 2.5% SMA (B, right panels). Wide-field images are shown in Supplementary Fig. 4. Scale bars in the lower right of each image are 50 nm.

3.3. Purification and characterisation of cytbc₁ SMALPs

Multiple reports of solubilisation of proteins with SMA have demonstrated that the annular lipids of membrane complexes are co-purified and amenable to analysis [28,31,34,46]. To this end we purified cytbc₁ SMALPs by nickel-affinity chromatography. Eluted fractions had a red colour and UV/Vis spectra revealed strong absorbance at 415 nm, which is indicative of haem containing proteins. Further purification by size exclusion chromatography removed residual RC-LH1-PufX and LH2, which eluted in the void volume. As shown in Fig. 5A, SDS PAGE of purified cytbc₁ SMALPs revealed four major bands with masses corresponding to those of the cyt b (50 kDa), cyt c₁-His (30.6 kDa), the Rieske Fe-S (20 kDa) and Subunit IV (15 kDa) polypeptides [46]. A fifth faint band is observed at ~25 kDa; identification of the gel band by tryptic digestion and mass spectrometry showed that both cyt c₁ (presumably a degradation product) and the zinc transporter ZnuC (UniProtKB ID: Q3IWB5) are present. ZnuC has a molecular weight of 26.5 kDa and eight histidine
residues in its N-terminal 15 amino acids. This unusually high histidine content may give ZnuC a natural affinity for the nickel resin. Nevertheless, the lack of RC-LH1-PufX and LH2 in the final preparation demonstrates that the cyt\textsubscript{bc1} complex has been effectively separated from the other major components of the chromatophore. Spectra of the pure complexes are shown in Fig. 5B and C. Upon reduction of the complexes with dithionite signals for both \textit{b-} and \textit{c-} type haems are observed. Analysis of reduced minus oxidised spectra estimated that the haem \textit{b} to \textit{c} ratio was 1.7:1 demonstrating that the cyt\textit{bc1} complexes are intact and fully functional.

\textbf{Fig. 5.} A: SDS-PAGE gel of cyt\textit{bc1} SMALPs. * denotes a fifth contaminating band (see text). B: UV/Vis absorbance spectra of SMA purified cyt\textit{bc1} complexes either as prepared (black) or after reduction with sodium dithionite (red). Labels correspond to prominent peaks in the reduced spectrum. C: Reduced minus oxidised spectra calculated from spectra in panel B. The cyt\textit{c1} to cyt\textit{b} ratio was calculated as 1.7:1 (see Materials and Methods for details).

TEM images of the purified cyt\textit{bc1} SMALPs (Fig. 6A) shows objects with a roughly elliptical shape. The average dimensions of the selected objects shown in the right-hand panels of Fig. 6A are $14 \pm 2 \times 9 \pm 1$ nm, roughly consistent with the size of the crystal structure of the \textit{Rba. sphaeroides} cyt\textit{bc1} complex shown in Fig. 6B [47]. Analysis of the wide-field images (left) shows that larger structures are not present, demonstrating the preparation contains discrete cyt\textit{bc1} dimers rather than small membrane
fragments, such as those in Fig. 4 B. The presence of cytbc₁ dimers is further supported by dynamic light scattering of the cytbc₁ SMALPs, which detected particles with hydrodynamic diameters of 24 ± 10 nm corresponding to 97.9% of the material by mass.

Fig. 6. Panel A: Negative stain TEM images of purified cytbc₁-SMALPs. Left: wide field image with selected complexes highlighted. Scale bar is 50 nm. Right: Zoomed images of six selected objects. The average dimensions were 14 ± 2 x 9 ± 1 nm. Inset scale bars are 10 nm. Panel B. Surface views of the cytbc₁ crystal structure [47] (PDB ID: 2QJP) from the end, side, or top. Images have been approximately scaled to match the sizes of zoomed objects in panel A. The two cytbc₁ monomers are shown in blue and red. Grey ovals show the area in which the lipid disc is expected to be located.

As the cytbc₁ SMALPs are expected to contain lipids and quinones in addition to the identified polypeptides, analysis of the lipid content was performed. Thin layer chromatography of methanol/chloroform extracts of cytbc₁ SMALPs shows that five lipid species co-purified with the protein, identified as cardiolipin (CL), phosphatidylethanolamine (PE), phosphatidylglycerol (PG) and phosphatidylcholine (PC) by the matched migration of pure standards. The fifth lipid, sulfoquinovosyl diacylglycerol (SQDG), was assigned based on data from Swainsbury et al. [22,27] where the same five components were identified in Rba. sphaeroides RCs prepared from membranes lacking both LH1-PufX and LH2 complexes. It should be noted that recent publications on the rapid exchange of lipids in SMALPs suggest that conclusions cannot be drawn on particular lipid enrichment of the
target complexes unless they are tightly bound to the protein [48,49]. Nevertheless, lipid extracts of bulk chromatophores show a similar lipid profile suggesting SMALP lipids originated from the source membrane and remained associated with the complex during preparation. Quantification of the phospholipids estimates that there is an average of 56 ± 6 phospholipids per cytbc\textsubscript{1} dimer. To further characterise the co-purified membrane environment we used HPLC to show that 0.96 ± 0.18 UQ\textsubscript{10} molecules are associated with each cytbc\textsubscript{1} complex. Pre-treatment of the membranes with antimycin A prior to purification yielded complexes where UQ\textsubscript{10} was not detected (data not shown) suggesting any co-purified quinone is associated with the tight-binding Q\textsubscript{i} site [47]. It should be noted that the aforementioned lipid exchange between SMALPs, or some as yet unexplored effect of SMA, may preclude enrichment of UQ\textsubscript{10} molecules that natively reside within the lipid annulus of the cytbc\textsubscript{1} complex. This may explain why only a single, tightly bound UQ\textsubscript{10} molecule was co-purified with each cytbc\textsubscript{1}, but we also note that in the case of cytbc\textsubscript{1} there is no physical confinement of quinones by a defined protein environment surrounding the complex. In contrast, a pool of 10-15 UQ\textsubscript{10} molecules/RC is sequestered within the solubilised and purified RC-LH1-PufX complex prepared using \(\alpha\)-dodecyl maltoside [43], but in this case the quinones are effectively trapped between the RC and the encircling LH1 ring.

![Graph](image-url) **Fig. 7.** Left panel (A): Lipid profiles of chromatophore membranes (blue) and cytbc\textsubscript{1}-SMALPs (grey) determined by thin layer chromatography for phosphatidylethanolamine (PE), cardiolipin (CL), phosphatidylglycerol (PG), phosphatidylserine (PS), and sulfolipid (SQDG).
phosphatidylcholine (PC) and sulfoquinovosyl diacylglycerol (SQDG). All band intensities are normalised to PG, labelled with asterisks. Error bars show standard error of the mean from seven (membranes) or eight (cytbc1-SMALPs) samples. Right panel (B): Representative TLC lanes for membranes and cytbc1-SMALPs.

3.4. Extraction of cytbf from Synechocystis sp. PCC 6803 membranes

To further examine the ability of SMA to sample the lipid environments of complexes in photosynthetic membranes we attempted to replicate the study of Rba. sphaeroides cytbc1 with the cytbf complex of the model cyanobacterium Synechocystis sp. PCC 6803 (hereafter Synechocystis). Using a procedure similar to the one described in section 3.1, we found that SMA efficiently solubilises the cytbf complex. We estimated an extraction efficiency of 67 ± 11% by comparison of reduced minus oxidised spectra in total and soluble fractions, and 85 ± 2% by haem staining, as shown in Fig. 8A by the solid and hatched magenta bars, respectively (see Supplementary Fig. 5 for raw spectra). By deconvolution of the UV/Vis spectra of the total and soluble fractions following SMA treatment we could also estimate the extraction efficiency of both photosystems, PSI and PSII. This gave values of 2 ± 0.5% and 61 ± 11%, respectively (Fig. 8A, solid red and blue bars, and Supplementary Fig. 5). It should be noted that the values here are subject to large errors given their strongly overlapping absorption spectra. Nevertheless, as shown in Supplementary Fig. 5 panel D, the shift of the chlorophyll absorbance from 678 nm in the total fraction to 673 nm after removal of insoluble material clearly demonstrates that PSII is the dominant species solubilised by SMA whilst PSI mostly remains insoluble. This conclusion is supported by analysis of the soluble and total fractions by native PAGE. From blue native PAGE gels we observe that PSI complexes are depleted from the soluble fraction whereas PSII is not (Supplementary Fig. 5 E). Fluorescence images of clear native PAGE gels also showed a strong signal for PSII in both the total and soluble fractions (Supplementary Fig. 5 E). Integration of the fluorescence from three solubilisation trials gave an estimated PSII extraction efficiency of 58 ± 4% as shown in Fig. 8A by the blue hatched bar. This value is in good agreement with the estimate by deconvolution of the spectra.

To further analyse the solubilised cytbf complexes we enriched cytbf SMALPs by IMAC purification. As shown in Fig. 8B the UV/Vis absorbance spectrum is typical of this complex showing a 670 nm chlorophyll peak, a strong haem Soret band at 420 nm and carotenoid absorbance between 400 and 550 nm [50]. Upon dithionite reduction two distinct peaks at 565 and 559 nm for the b- and c-type haems become apparent, suggesting that the cytbf complex is intact within SMALPs with all of its cofactors present. This spectrum matches that of pure cytbf reported previously [50]. These spectra also suggest that the cytbf complex does not co-purify with other pigmented components of the
photosynthetic membranes as PSI or PSII absorbance features were not observed. We further analysed these complexes by negative stain TEM as shown in Fig. 8C. The typical wide-field image shown is free of large objects indicative of membrane patches or aggregates. The six zoomed objects, shown in the Fig. 6D, are consistent with cyt$b_6f$ viewed from multiple angles. The objects have average dimensions of $12 \pm 3 \times 8 \pm 1$ nm and are similar in shape to the roughly scaled images of the *Nostoc* sp. PCC 7120 crystal structure shown in Fig. 8 E [51]. Taken together the analysis of the pure cyt$b_6f$ SMALPs shows that they are solubilised as discrete dimeric complexes rather than as part of larger membrane fragments. They also do not co-purify with other membrane proteins.

**Fig. 8.** Cytochrome $b_6f$ extraction from *Synechocystis* thylakoids with SMA. (A): Estimated extraction efficiencies of PSI (solid red) and PSII (solid blue) from deconvoluted absorbance spectra. PSII solubilisation was also estimated from clear native page in-gel fluorescence (hatched blue bar). Solubilisation estimates for cyt$b_6f$ from reduced minus oxidised spectra and, haem staining are shown with solid and hatched magenta bars, respectively. Error bars show standard errors of the mean. Raw data are shown in Supplementary Fig. 5. (B): UV/Vis spectra of purified cyt$b_6f$ either as prepared in buffer (black line) or after reduction with dithionite.
(red line). Labels show wavelengths of prominent peaks of the reduced spectrum. (C): Wide field negative stain TEM images of purified cytbf-SMALPs. (D): Zoomed images of selected complexes highlighted in panel C. Scale bar is 50 nm. The average dimensions were 12 ± 2 x 9 ± 1 nm. (E): Surface views of the cytbf crystal structure from Nostoc sp. PCC 7120 (PDB ID: 4OGQ) from the end, side and top. Images have been approximately scaled to match the sizes of zoomed objects in panel D. The two cytbc1 monomers are shown in blue and pink. Grey ovals show the area in which the lipid-disc is expected to be located.

4. Discussion

SMA preferentially solubilises cytbc1 complexes from Rba. sphaeroides chromatophore membranes, leaving RC-LH1-PufX and LH2 complexes largely unaffected. SMA acts efficiently on membrane complexes that sit in lipid-rich environments, as shown by work on several proteins including the bacterial SecYEG translocon and potassium channel KcsA [23,28]. Conversely, arrays of well-ordered and/or closely packed protein complexes such as RC-LH1-PufX and bacteriorhodopsin are poorly solubilised [27,52]. The liberation of cytbc1 complexes from chromatophores by SMA fits with the proposed architecture of chromatophore membranes [1] in which LH2 and RC-LH1-PufX complexes form closely packed protein-rich arrays, whereas cytbc1 complexes are found in lipid-rich regions of the membrane, which is ideal for the production of SMALPs due to the abundance of lipids from which a nanodisc can be formed.

There are several compelling reasons for cytbc1 complexes residing in lipid rich domains, with no tight, stoichiometric interaction with another complex such as RC-LH1-PufX. First, strong association of the cytbc1 with RC-LH1-PufX could lower the number of potential interactions between antenna complexes. Second, cytbc1 complexes residing in lipid rich domains are able to utilise quinols from any source such as from Complex I or succinate dehydrogenase rather than those in strongly localised pools. In some bacteria cytbc1 complexes in lipid rich domains might also improve its potential to interact with the membrane-associated electron carrier cytc1, which is utilised alongside cytc2 during aerobic and semi-aerobic growth for electron transfer between the cbb3-type cytcd the and the cytbc1 complex [45]. A final advantage to residing in a lipid-rich domain may be ready access to a local pool of reduced quinols, in this case fed by turnover at nearby RCs. It is known that quinones form strongly localised pools around sub-populations of RC to improve the efficiency of quinone reduction [17], so it is therefore not unreasonable to suggest that such pools may also form around the cytbc1 complexes. Such an arrangement would allow accumulation of quinols during high-light growth where reduction of quinone at the RC exceeds the capacity of the cytbc1 complexes to oxidise them, which occurs at just 5% of full-sunlight intensity [2]. In general,
cytbc₁ complexes are the limiting factor in the overall conversion of absorbed solar energy to ATP by *Rba. sphaeroides* [1,2].

The benefits of placing the cytbc₁ complex in a lipid-rich domain may not just be limited to *Rba. sphaeroides*. This theory is supported by the finding that the cytbf complex of the model cyanobacteria *Synechocystis* sp. PCC 6803 is susceptible to SMA solubilisation with similar efficiency to that of the *Rba. sphaeroides* cytbc₁, along with PSII. These data support the hypothesis that PSI resides in highly ordered, protein-rich arrays whereas PSII and cytbf are likely to reside in less ordered regions of the thylakoid membrane [35]. The *Synechocystis* cytbf complex has been described to reside at the “crossroad of photosynthetic electron transport pathways” [18] and thus requires plasticity in terms of its molecular interactions to accept quinols and oxidised cytochrome c₆ or plastocyanin from a variety of sources [18,53,54]. Finally, it is possible that cytbf is also found in lipid rich environments in the thylakoid membranes of chloroplasts; this is supported by the finding of Bell *et al.* [32] who solubilised spinach thylakoids with SMA and found that the insoluble PSI containing fraction was devoid of cytbf and PSII complexes. This suggests that, like in *Synechocystis*, cytbf and PSII are susceptible to SMA solubilisation. Johnson *et al.* [55] have also observed that treatment of spinach thylakoid membranes with the α isomer of DDM specifically removes cytbf leaving the PSI and PSII complexes in the membrane, also consistent with a lipid-rich environment for this complex.

5. Conclusion

Using the unique ability of SMA to preferentially solubilise protein complexes from lipid-rich regions of native biological membranes, we have demonstrated that the cytbc₁ complex resides in lipid-rich regions of the otherwise protein-rich chromatophore. These lipid-rich regions have a similar lipid composition to the bulk chromatophore membrane, with around 56 lipids co-purifying with the dimeric cytbc₁ complex. The observation that the insoluble fraction contains large membrane patches of RC-LH1-PufX and LH2, whereas cytbc₁ is purified from the soluble fraction in the form of discrete dimers in small nanodisc structures, further highlights the ability of SMA to remove some complexes from a specific membrane environment while preserving protein arrays.

Together these data provide support for the model of chromatophore membranes whereby light-harvesting and RC complexes reside in densely packed arrays for optimal energy transfer efficiency, whereas the cytbc₁ complexes occupy lipid-rich areas [1]. Placing cytbc₁ complexes in lipid rich domains rather than in arrays or supercomplexes may allow them to accept quinols from a range of
electron transport chains to provide both protons and electrons to drive efficient energy generation under a wide-range of growth conditions.

We have extended our study to show that the cyt_{b6}f complex of the cyanobacterium *Synechocystis* is highly susceptible to SMA solubilisation suggesting that it also resides in a locally lipid rich environment. Our data, along with other published data regarding SMA solubilisation of energy producing membranes, suggest that cyt_{bc1}-type complexes universally reside in lipid-rich membrane regions to aid their critical functions for proton translocation during cellular energy generation under a variety of growth conditions.

### Author contributions

DJKS and CNH conceived the study. DJKS, MSP, AH and CNH designed the experiments. DJKS, MSP, AH, MLC, PQ, ECM and PJJ prepared samples, performed experiments and/or analysed data. JM and SPA aided in the preparation of SMA. DJKS, MSP, AH and CNH contributed to preparation of the manuscript.

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New insights into the photochemistry of carotenoid spheroidenone in light-harvesting


Supplementary Fig. 1. Representative spectra utilised for calculations of extraction efficiency in Fig. 2 for the low light samples for RC-LH1-PufX and LH2. Spectra for total fractions are shown on the left for SMA (A) and DDM (B), and for soluble fractions after ultracentrifugation on the right for SMA (C) and DDM (D). Raw spectra are shown in black. The fitted spectra for the scatter curve, RC-LH1-PufX and LH2 are shown with dashed lines in grey, red and purple, respectively. A combined calculated spectrum is shown in blue. Reference spectra used for the deconvolution are shown in panel E.
Supplementary Fig. 2. Data used for cytb\(_c\) extraction efficiency calculation in Fig. 2. The left panels show averaged oxidised-reduced spectra for the total (black lines) and soluble fractions after centrifugation (grey, dashed lines) for low-light membranes solubilised with SMA (A, top) or DDM (B, lower). The right panels show cytc detection by haem staining. Alternate lanes contain total and soluble fractions. The right-most lane contains purified cytb\(_c\) as a control. Data are shown for SMA (C, top) and DDM (D, lower) solubilised samples.

Supplementary Fig. 3: Haem blots of fractions from 50-15% sucrose gradients shown in Fig. 3A and B. Intensities of the upper band (~30 kDa) were quantified with Image J and the values were used to plot cytb\(_c\) migration in the gradient in Fig. 3 panel C.
Supplementary Fig. 4. Wide-field TEM images of bands from sucrose gradients in Fig. 3 with two examples of individual objects for each. (A) 15 – 50% sucrose gradients from which TEM samples were produced. The positions of upper and lower bands are marked with dashed boxes. Approximate locations of the sucrose step interfaces are also marked on the right hand side. (B) TEM image of the upper band from the untreated membrane (mem) gradient. (C) TEM image of the upper band from the β-DDM treated membrane (DDM) gradient. (D) TEM image of the lower band from the untreated membrane (mem) gradient. (E) TEM image of the lower band from the β-DDM treated membrane (DDM) gradient. (F-G) TEM images of the upper band (F) and lower band (G) from the SMA solubilised membrane (SMA) gradient.
**Supplementary Fig. 5.** Deconvolution of UV/VIS spectra of SMA solubilised *Synechocystis* membranes before (A - Total) and after (B - Sol) centrifugation to remove insoluble material. Reference spectra utilised for the deconvolution normalised to a maximum absorbance of 1 are shown in panel C. PSI and PSII spectra were taken from reference [39]. Panel D shows raw spectra of the total and soluble fractions that were deconvoluted, normalised to an absorbance of 1 at the chlorophyll Qy maximum. E: Blue-native PAGE and in-gel fluorescence from clear-native PAGE gels of total (T) and soluble (S) fractions. Assignment of PSI trimer (3), PSI monomer (1) and PSII bands are based on data in reference [42]. F: oxidised-reduced spectra of the total (black) and solubilised (grey dashed) fractions. G: Haem blots of total and soluble fractions.