Integration of Apo-α-Phycocyanin into Phycobilisomes and Its Association with FNRL in the Absence of the Phycocyanin α-Subunit Lyase (CpcF) in Synechocystis sp. PCC 6803

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Abstract
Phycocyanin is an important component of the phycobilisome, which is the principal light-harvesting complex in cyanobacteria. The covalent attachment of the phycocyanobilin chromophore to phycocyanin is catalyzed by the enzyme phycocyanin lyase. The photosynthetic properties and phycobilisome assembly state were characterized in wild type and two mutants which lack holo-α-phycocyanin. Insertional inactivation of the phycocyanin α-subunit lyase (ΔcpcF mutant) prevents the ligation of phycocyanobilin to α-phycocyanin (CpcA), while disruption of the cpcA/Cpc2/C1 operon in the CK mutant prevents synthesis of both apo-α-phycocyanin (apo-CpcA) and apo-β-phycocyanin (apo-CpcB). Both mutants exhibited similar light saturation curves under white actinic light illumination conditions, indicating the phycobilisomes in the ΔcpcF mutant are not fully functional in excitation energy transfer. Under red actinic light illumination, wild type and both phycocyanin mutant strains exhibited similar light saturation characteristics. This indicates that all three strains contain functional allophycocyanin cores associated with their phycobilisomes. Analysis of the phycobilisome content of these strains indicated that, as expected, wild type exhibited normal phycobilisome assembly and the CK mutant assembled only the allophycocyanin core. However, the ΔcpcF mutant assembled phycobilisomes which, while much larger than the allophycocyanin core observed in the CK mutant, were significantly smaller than phycobilisomes observed in wild type. Interestingly, the phycobilisomes from the ΔcpcF mutant contained holo-CpcB and apo-CpcA. Additionally, we found that the large form of FNR (FNRL) accumulated to normal levels in wild type and the ΔcpcF mutant. In the CK mutant, however, significantly less FNR was accumulated. FNRL has been reported to associate with the phycocyanin rods in phycobilisomes via its N-terminal domain, which shares sequence homology with a phycocyanin linker polypeptide. We suggest that the assembly of apo-CpcA in the phycobilisomes of ΔcpcF can stabilize FNRL and modulate its function. These phycobilisomes, however, inefficiently transfer excitation energy to Photosystem II.

Introduction
The primary photoreactions of oxygenic photosynthesis are catalyzed by two major membrane protein complexes, Photosystem I (PS I) and Photosystem II (PS II). While these photosystems both have internal chlorophyll antennae, productive photosynthesis requires additional light-harvesting components. In cyanobacteria, as well as the eukaryotic classes Rhodophyta and Glaucophyta, these are the phycobilisomes. Phycobilisomes are large, highly structured peripheral water-soluble complexes consisting of an allophycocyanin core which is attached to multiple oriented rods. These rod elements are composed of phycocyanin, phycocerythrin, and phycocerythrocyanin (the exact composition being species-dependent) and their associated linker polypeptides. Recently, it has been demonstrated that phycobilisomes can physically associate with, and transfer excitation energy to, both Photosystem II and I [1,2]. Covalent attachment of the phycobilin chromophores to specific cysteinyl residues of the apophycobiliproteins via a thioether bond is catalyzed by phycobilin lyases [3,4]. Synechocystis sp. PCC 6803, henceforth Synechocystis, has a relatively simple phycobilisome structure containing only the allophycocyanin core and phycocyanin-containing rods. The heterodimeric CpcE/CpcF lyase is specifically responsible for phycocyanobilin attachment to the α subunit of phycocyanin (CpcA) [5,6].

In both the cyanobacteria and chloroplasts, the role of ferredoxin: NADP⁺ oxidoreductase (FNR) is to catalyze the final step of photosynthetic electron transport, providing reducing equivalents in the form of NADPH for CO₂ fixation and other reductive metabolic pathways. Higher plants contain multiple...
FNR genes encoding a variety of different isoforms of this enzyme. In cyanobacteria, however, FNR is encoded by a single gene, petH. The FNR in most phycobilisome-containing cyanobacteria has an N-terminal conserved domain which shares high sequence similarity with the phycocyanin rod linker polypeptide CpcD. This domain appears to be responsible for attachment of FNR to the peripheral rods in phycobilisomes [7,8]. Recent studies indicate that in cyanobacteria two FNR isoforms (FNR\(_L\) and FNR\(_S\)) are produced from the single petH gene via alternative transcriptional start points and, consequently, different translation initiation sites [9,10]. Under normal photosynthetic conditions, FNR\(_L\) is present as the major isoform, while FNR\(_S\) is induced under a variety of stress conditions such as iron starvation [9]. An FNR\(_L\)-phycocyanin complex has been purified from Thermosynechococcus elongatus and Synechocystis and the enzymatic activities have been characterized in vitro [11,12].

In this work, we characterized two phycocyanin defective strains of Synechocystis, a phycocyanin \(\tau\)-subunit lyase mutant (\(\Delta\)pet\(F\)) and a cpcB/A/C2/C1 operon deletion mutant (CK\(_a\)) to address how the assembly of phycobilisome into the phycobilisome regulates photosynthetic performance via light energy absorption and downstream energy utilization. Interestingly, we find that apocCpcA appears to assemble into phycobilisomes containing holopCpcB, linker polypeptides and the allophycocyanin core. While these mutant phycobilisomes inefficiently transfer excitation energy to the photosystems, they can associate with FNR\(_L\) and stabilize this component.

**Material and Methods**

**Strains and cell culture conditions**

A glucose-tolerant strain of Synechocystis [13] was used as the wild-type strain for this study. The phycocyanin \(\tau\)-subunit lyase inactivation mutant \(\Delta\)pet\(F\) was constructed by insertion of a kanamycin-resistant cassette at the position 193 of sl11051. After selection, segregation of the mutant allele was verified by PCR (data not shown). The insertion mutation was tested for polar effects both down- and up-stream of the sl11051 gene; none were identified (data not shown). The WT, \(\Delta\)pet\(F\), and phycocyanin-null mutant, CK\(_a\), were generally grown in BG-11 medium buffered with 10 mM TES-KOH (pH 8.2), except for low CO\(_2\) growth conditions where Na\(_2\)CO\(_3\) was omitted from the BG-11 recipe and the medium was buffered with 20 mM HEPES-NaOH (pH 7.5). The mutant strains were maintained in the presence of 10 \(\mu\)g/ml kanamycin. The strains were grown either photoautotrophically, photomixotrophically, or photoheterotrophically. For photomixotrophic growth, 5 mM glucose was added to the basal BG-11 medium. For photoheterotrophic growth, BG-11 medium was supplemented with both 5 mM glucose and 10 \(\mu\)M 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU). The cells were grown either fractionated into 30, 400 \(\mu\)l fractions, or the blue fractions were centrifuged, washed, and resuspended in fresh growth medium at a chlorophyll concentration of 10 \(\mu\)g/ml. Chlorophyll concentration was determined as described by Williams [13]. The measurements were performed with either actinic white light or 650 nm red light. Net photosynthesis was measured in the presence of 10 mM NaHCO\(_3\) as a terminal electron acceptor. The PS II electron transfer rates from H\(_2\)O to quinone were measured in the presence of 0.5 mM 2,6 dichloro-p-quinone (DCBQ) +1 mM K\(_3\)Fe(CN)\(_6\). 

**Fluorescence measurements**

Chlorophyll fluorescence induction was monitored with a dual-modulated fluorometer FL 3000 (PSI Instruments). The cells were diluted to a chlorophyll concentration of 5 \(\mu\)g/ml. Measurements were taken after 5 min dark adaptation, followed by illumination with the highest level of actinic light (1000 \(\mu\)mol photons m\(^{-2}\) s\(^{-1}\), 625 nm). The oxidation state of the plastoquinone pool was modulated by the addition of either 15 \(\mu\)M DCMU or 20 \(\mu\)M 2,5-dibromo-3-methyl-6-isopropylbenzoquinone (DBMIB) in the fluorometer cell after dark adaptation. Data were collected with measuring light in a logarithmic series between 10 \(\mu\)s to 1 s. The fluorescence curves were plotted directly or normalized according to the equation Fluorescence = \((1-F_\text{F}/F_\text{v})/(F_\text{m}/F_\text{v})\) [14].

Data were analyzed using Origin version 8.1 and proprietary software provided by Photon Systems Instruments.

**Isolation of membrane and soluble fractions**

Synechocystis membrane and soluble fractions were isolated according to [15]. The cell cultures (150 ml OD\(_{730}\) =1) were pelleted at 4°C by 5 min centrifugation at 5,000 g. The cell pellet was washed twice with wash buffer containing 50 mM MES-NaOH pH 6.5, 50 mM NaCl, 10 mM MgCl\(_2\), 5 mM CaCl\(_2\), and resuspended in 1.5 ml wash buffer containing 25% (v/v) glycerol (break buffer). An equal volume of glass beads (Sigma, 150-212 \(\mu\)m diameter) was added to the cell suspension, which was chilled on ice. The cells were broken by manually vortexing eight times at maximal speed for 1 min at 4°C with 1 min cooling on ice between the cycles. The glass beads and cell debris were removed by centrifugation at 2,000 xg for 5 min, and the supernatant was collected. The glass beads were washed with 1 ml break buffer and centrifuged for 5 min. The supernatant was combined with the previously collected samples. The membranes and soluble fractions were separated by centrifugation at 30,000 xg for 30 min.

**Isolation of phycobilisomes**

Phycobilisomes were isolated according to [16]. The cells were harvested from 0.5–1 L culture at OD\(_{730}\) >2 and washed twice with 0.8 M potassium phosphate buffer, pH 7.0 (KP). The cell pellets (1 g in fresh weight) were resuspended in 3 ml KP in the presence of 1 mM 4-[2-aminoethyl]benzenesulfonyl fluoride hydrochloride (AEBSF). The cells were broken as described above by vortexing with glass beads. The broken cell extracts were incubated with Triton X-100 at a final concentration of 2% (v/v) for 20 min at room temperature in darkness with occasional gentle shaking. The unbroken cells and membrane debris were removed by centrifugation at 30,000 xg for 20 min at 15°C. The supernatant was loaded onto a 10–35% (v/v) linear sucrose gradient in 0.8 M KP with 1 mM AEBSF, and centrifuged at 130,000 xg for 24 h at 15°C. After centrifugation, the gradients were either fractionated into 30, 400 \(\mu\)l fractions, or the blue bands were collected directly with a syringe. The samples from the sucrose gradient were precipitated by the addition of an equal volume of 20% trichloroacetic acid and incubated on ice for

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**Assembly of Apo-Phycocyanin and FNR\(_L\) in Synechocystis sp. PCC 6803**
Electrophoresis and immunoblotting

Gradient (12.5–18%) LiDS-PAGE was used to analyze denatured protein samples according to Delepelaire and Chua [17]. Soluble protein complexes were analyzed by BN-PAGE as described in Zhang et al. [18] on a 6–13.5% gradient polyacrylamide gel. Samples were prepared at 4°C under low light illumination. No solubilization procedure was required for the soluble fraction samples. The protein samples were diluted with one third volume of a buffer containing 100 mM Bis-Tris, pH 7.0, 80% (w/v) glycerol, 0.08% n-dodecyl β-D-maltoside (DM), and 4 mM AEBSF prior to electrophoresis. After electrophoresis, the proteins or protein complexes were electroblotted onto a polyvinylidene fluoride membrane (PVDF) and labeled with protein-specific antibodies followed by chemiluminescent detection. Protein expression was semi-quantified by comparison with a dilution series of WT samples (5–75 μg protein). Signal intensities were analyzed by ImageJ [19]. These are shown in Fig S1.

Results

Growth of wild type and phycocyanin mutant stains with different carbon sources

To examine the cell viability and growth rate, wild type (WT) and the two phycocyanin-defective mutants, ΔcpcF and CK, were grown on BG-11 plates in the presence or absence of glucose and DCMU. All three strains had similar growth rates in the presence of glucose under either phototrophic or mixotrophic conditions. Under autotrophic growth conditions (≈500 ppm CO2), however, both the ΔcpcF and CK mutants grew slower than WT (Fig. 1). The slower growth rates of the mutants suggested a defect in photosynthesis. These strains were then examined during autotrophic growth under both low (<220 ppm) and elevated (>5000 ppm) CO2 concentrations. Both the WT and ΔcpcF strains exhibited enhanced growth at a high CO2 concentration. Interestingly, the CK mutant grew very poorly under both high and low levels of CO2 (Fig. 1).

Photosynthetic light response curves of WT and phycocyanin mutants

To examine the overall photosynthetic property of these strains, oxygen evolution measurements were determined at different intensities under both white and red light (650 nm) illumination conditions. Under white actinic light illumination, the maximal oxygen evolution capacity from water to CO2 of both phycocyanin mutants is lower than observed in WT (Fig. 2A), suggesting that the overall photosynthetic capacity of the ΔcpcF and CK mutants is diminished in these strains. The light compensation point (Fig. 2A, insert) and examination of the light saturation curves (Fig. 2A and C) indicated that the mutant strains saturate photosynthesis at somewhat higher light intensities than does WT. These results indicate that the white light absorption efficiency is lower in mutant cells than in WT. PS II oxygen evolution rates using DCBQ as an artificial electron acceptor were also examined (Fig. 2B and D). Both mutant strains exhibited higher maximal rates of oxygen evolution than WT. This was not unexpected, as mutants depleted in phycobilisomes accumulate increased amounts of PS II [20]. The normalized PS II light saturation curves (Fig. 2D) indicate that the ΔcpcF and CK mutants are nearly identical (Fig. 2D) suggesting that the mutants have similar white light-trapping efficiencies. The light intensity providing ½ maximal PS II activity was 500 μmole photons-m⁻²sec⁻¹ for both mutants and 300 μmole photons-m⁻²sec⁻¹ for WT. This indicates that WT is significantly more efficient at energy capture than the ΔcpcF and CK mutants when white light is used as an illumination source. These results suggest that, as expected, both phycocyanin mutants have defective light absorption properties when compared to the WT strain. These defects probably arise from the lack of functional phycocyanin-containing rod elements in both mutants.

Additionally, we measured photosynthetic activity using a red LED actinic light source (650 nm). The allophycocyanin core of the phycobilome preferentially absorbs at this wavelength. The maximal rate of oxygen evolution, using CO2 as a terminal electron acceptor under this illumination condition, was higher in WT than in either of the phycocyanin mutants (Fig. 2E), similar to what was observed in white light (Fig. 2A). The normalized light saturation curves of the mutants, however, were very similar to that of WT (Fig. 2G). Examination of PS II activity using DCBQ as an electron acceptor again demonstrated that the mutants exhibited higher rates of electron transport than WT, as was observed under white light illumination (Fig. 2B). The normalized light saturation curves indicated that WT and the phycocyanin mutants were equally efficient at harvesting 650 nm red light.

![Figure 1. Growth of WT, ΔcpcF and CK strains on BG-11 agar plates with different carbon sources.](image-url)
These results indicate that WT and both mutants appear to have functionally equivalent light absorption properties in red light, suggesting that all three strains have fully functional allophycocyanin cores associated with their phycobilisomes. We also examined the chlorophyll fluorescence kinetics of WT and the phycocyanin deficient strains over the first 1.0 s of actinic illumination (Fig. 3, Table 1, Fig. S2). WT exhibited an OJIP fluorescence transient curve typical of those observed previously in WT strains [21,22]. While the variable fluorescence (F_V) of the ΔpcF and CK mutants were similar, and smaller than that of WT, their initial fluorescence levels (F_0) were quite different. ΔpcF exhibited a higher F_0, while the CK strain had a much lower F_0 than that observed for WT (Fig. 3A, Table 1). To examine the effect of the redox state of the plastoquinone pool on the OJIP transient, DCMU (Fig. 3B) and DBMIB (Fig. 3C) were supplied to the dark-adapted cell suspensions prior to collection of the fluorescence transient data. In the presence of DBMIB, there was no significant change observed in the F_0 levels for any of the strains examined. In the presence of DCMU all of the strains exhibited higher F_0 values. Both DCMU and DBMIB slightly increased F_M levels of WT and ΔpcF cells, while the F_M of CK appeared insensitive to these treatments (Fig. 3A, B and C, Table 1). Normalization of these fluorescence transients to [F = (1-F_0/F(V))/F_V/F_M)] [14] indicated that the OJ transition occurred slower in the ΔpcF (T_J = 2.63 ms) and CK (T_J = 3.19 ms) mutants than in WT (T_J = 1.67 ms) and were of lower magnitude (Fig. 3D, Table 1). The differences in the OJ transients were not eliminated in the presence of DCMU (Fig. 3E, Table 1) or DBMIB (Fig. 3F, Table 1) suggesting that the phenomenon arose from the donor side of PS II. In all cases the T_J for the mutants were significantly slower than that observed for WT. Decreasing the actinic light intensity for WT samples nearly eliminated the observed alterations in both the magnitude of the OJ transition and its kinetics (Fig. S2). Consequently, we hypothesize that the observed differences in the OJ transitions are due to the mutants having a smaller optical cross-section brought about by alterations in the phycobilisome antennae. Treatment with DBMIB increased the J peak amplitude of WT and the CK mutant, while the J peak amplitude was essentially unchanged in the ΔpcF strain. (Fig. 3F). Interestingly, the T_P values for WT and the ΔpcF mutant were nearly identical in both the absence and presence of DBMIB. The T_P value for the CK mutant was lower under both control conditions and in the presence of DBMIB treatment (Table 1). These results indicated that the plastoquinone pool became reduced somewhat more rapidly in the CK mutant than in WT and the ΔpcF strain. This may result from defective NADP^+ reduction in the CK strain (see below).

Expression and assembly of phycobiliproteins and ferredoxin-NADP^+ reductase in WT and the phycocyanin mutants

To examine the possibility that proper incorporation of phycocyanobilins into the phycocyanin protein may affect not only its function but also the stability, assembly and accumulation of other phycobilisome proteins, we examined the expression of a number of phycobiloproteins in WT and the phycocyanin mutant strains. These included CpcA, CpcB, and ApcB. CpcA and CpcB are major pigment-binding components of the phycobilisome rods, and ApcB is a component of the allophycocyanin core of the phycobilisome. CpcA is the substrate for the heterodimeric CpcE/CpcF phycocyanin and ApcB is a component of the allophycocyanin core of the phycobilisome. CpcA is the substrate for the heterodimeric CpcE/CpcF phycocyanin α subunit lyase. We also examined the expression of FNR in these strains since cyanobacterial FNR_L has shown high affinity to the phycobilisomes, and its association with phycocyanin modulates FNR enzymatic activity [12].

The protein expression of the phycocyanin and allophycocyanin proteins as well as FNR are shown in Fig. 4. In the ΔpcF mutant, the total amount of immunodetectable CpcA is slightly larger than that found in WT (Fig. 4A and C). However, the vast majority of this protein is found in a lower apparent molecular mass band, with only a small fraction migrating at the position of CpcA in WT. This appears to arise from the accumulation of apo-CpcA since, in the absence of the phycocyanin α-subunit lyase in the ΔpcF mutant, the phycocyanobilin chromophores would be inefficiently attached to the protein. This is apparent in the unstained panel, as no major blue band is observed at this location in the ΔpcF mutant. The amount of CpcB observed in the ΔpcF

Figure 2. Light saturation curves of oxygen evolution for WT, ΔpcF and CK strains under white and red actinic light illumination. Oxygen evolution rates were measured with cell suspensions at a chlorophyll concentration of 10 µg/ml in BG-11 medium using different intensities of either white light (A–D) or red light (E–G). Net photosynthesis was measured in panels A, C, E and G (H2O to O2). PS II activity was measured in B, D, F and H (H2O to DCBQ). The upper panels A, B, E and F show the absolute oxygen evolution rates observed. In Panels C, D, G and H the data were normalized to the maximum rates observed in Panels A, B, C and D, respectively. The error bars represent ±1.0 standard deviation (SD) and in some instances were smaller than the symbols; each experiment is the average of 3–4 independent measurements. doi:10.1371/journal.pone.0105952.g002
mutant is significantly lower than that observed in WT (Fig. 4A and C) and appears to be fully pigmented. As expected, neither CpcA nor CpcB were present in the CK mutant, which lacks the genes encoding these two components. The ∆cpcF mutant accumulated slightly more ApcB than WT which is consistent with the presence of the allophycocyanin core of the phycobilisome in both of these strains; the CK mutant, however, exhibited substantially more ApcB than either WT or the ∆cpcF mutant (Fig. 4A and C). Analysis of the two forms of FNR in these strains indicated that the amount of FNR L was significantly decreased in the CK mutant. Additionally, lower mass bands were observed which may represent degradation products of this component. Similar results have been reported previously for this strain [9].

Table 1. Chlorophyll fluorescence parameters during the OJIP transition.

| Strain | F0     | FM     | FV     | FV/FM  | Tj (ms) | Tp (s) |
|--------|--------|--------|--------|--------|---------|--------|
| No addition |        |        |        |        |         |        |
| WT     | 0.75±0.01 | 1.67±0.01 | 0.92±0.01 | 0.55±0.01 | 1.67±0.01 | 0.41±0.01 |
| ∆cpcF  | 0.93±0.01 | 1.54±0.01 | 0.61±0.01 | 0.39±0.01 | 2.63±0.04 | 0.39±0.01 |
| CK     | 0.26±0.01 | 0.85±0.01 | 0.59±0.01 | 0.69±0.01 | 3.19±0.02 | 0.24±0.01 |
| + DCMU |        |        |        |        |         |        |
| WT     | 0.99±0.01 | 1.70±0.01 | 0.72±0.01 | 0.42±0.01 | 4.00±0.05 | NA     |
| ∆cpcF  | 1.06±0.01 | 1.58±0.01 | 0.52±0.01 | 0.33±0.01 | 6.33±0.30 | NA     |
| CK     | 0.40±0.01 | 0.85±0.01 | 0.44±0.01 | 0.52±0.01 | 4.66±0.18 | NA     |
| + DBMIB|        |        |        |        |         |        |
| WT     | 0.75±0.01 | 1.74±0.01 | 0.99±0.01 | 0.57±0.01 | 1.83±0.01 | 0.37±0.01 |
| ∆cpcF  | 0.93±0.03 | 1.56±0.01 | 0.63±0.01 | 0.40±0.01 | 2.73±0.04 | 0.36±0.01 |
| CK     | 0.27±0.01 | 0.87±0.01 | 0.60±0.01 | 0.69±0.01 | 3.22±0.02 | 0.26±0.01 |

The WT and mutant strains were examined either with no addition or with the addition of 10 μM DCMU or 20 μM DBMIB. The values shown represent the mean ±1.0 SD; each experiment is the average of 3 independent measurements. NA, Not Applicable.

doi:10.1371/journal.pone.0105952.t001
measurements. Symbols: *, FNRL; o, FNRS;
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this semi-quantification are shown in Fig. S1. The error bars represent
the degraded FNR observed, respectively. The standard curves used in
amounts of total FNR, the large form of FNR, the small form of FNR and
in both WT and the
condition, a putative phycobilisome subcomplex of about 80 kDa,
components of the phycobilisome partially dissociate. Under this

In order to investigate the interaction of FNRL and CpcA and
CpcB proteins, we used Blue-Native –PAGE (BN-PAGE) followed
by immunodetection with FNR and Cpc antibodies (Fig. 4B). Due
to the low phosphate concentration in the isolation buffer, the
components of the phycobilisome partially dissociate. Under this
condition, a putative phycobilisome subcomplex of about 80 kDa,
which contained the CpcA, CpcB and FNR proteins, was detected
in both WT and the \( \Delta \text{cpcF} \) mutant. Additionally, a higher
molecular mass band (\( \approx 500 \) kDa), which reacted with the FNR
antibody, was also observed in these strains. In the CK strain a
subcomplex which appeared to contain ApcB was observed at an
apparent molecular mass of about 90 kDa. No bands were found
to react with FNR antibodies in this strain. This appears to
indicate that phycocyanin was required for the formation and/or
stability of these FNR complexes (Fig 4B). Importantly, these
results imply that in the \( \Delta \text{cpcF} \) mutant, apo-CpcA can assemble
with FNR and CpcB to form a putative phycobilisome subcomplex and that this association appears to stabilize the FNR.

To examine the assembly state of the phycobilisomes in
the phycocyanin mutants, intact phycobilisomes were resolved by
succrose density gradient centrifugation in the presence of a high
phosphate concentration (Fig. 5A). Several blue bands were
observed in the sucrose gradients (Bands I, II', II, and III). Because of its position near the top of the gradient, Band I from all
samples was contaminated with multiple proteins which did not
penetrate the sucrose gradient (Fig. 5C). Since in all likelihood,
Band I contained unassembled phycobilin and allophycocyanin
proteins, we focused on the compositions of Band II and Band III.
The absorption spectrum of cells (Fig. 5B, top) and the lowest
band from each sample, Band III for WT and the \( \Delta \text{cpcF} \) mutant
and Band II for the CK strain, are shown (Fig. 5B, bottom). These
represent the largest phycobilisome assemblies present in these
strains. Compared to WT, the phycocyanin absorption peak of the
phycobilisome was significantly decreased in the \( \Delta \text{cpcF} \) mutant
strain and absent in the CK mutant. It should be noted that the
phycocyanin absorption peak which we observed in the \( \Delta \text{cpcF} \) mutant was much larger than that reported for the Synechocystis
sp. PCC 7002 \( \Delta \text{cpcE}/\text{F} \) mutants [6]. The phycobilisome which
assembles in the \( \Delta \text{cpcF} \) strain was smaller in size than that of WT
(Fig. 5A). Lithium dodecyl sulfate-PAGE (LIDS-PAGE) analysis
revealed that the phycobilisome in \( \Delta \text{cpcF} \) had relatively lower amounts of CpcA, CpcB and the linker polypeptide CpcC1, an
apparent absence of CpcC2, and more abundant CpcG1
(Fig. 5C). Interestingly, CpcC2 was also absent from phycobilis-

domes isolated from a Synechocystis phycocyanin \( \beta \)-subunit lyase
(\( \text{cycT} \)) mutant [23]. The absence of CpcC2 would suggest that no

core-distal phycocyanin hexamers were present in the assembled
phycobilisomes from the \( \Delta \text{cpcF} \) strain. Due to lack of the
phycocyanin proteins, Band III was absent in the CK strain
and Band II contained the allophycocyanin core. This band migrated
to a position which overlapped with Rubisco, which was also
enriched in the WT Band II' (Fig. 5D). It should be noted that
dissociated allophycocyanin core did not accumulate in the WT
strain; Band II in WT was larger than that observed in the CK
mutant and appeared to consist mainly of partially assembled
phycobilisome rod elements. The Band II in the \( \Delta \text{cpcF} \) mutant
was also smaller than that observed in WT. It contained phycocyanin
proteins and a significant amount of ApbB was also detected.

Consequently, Band II in \( \Delta \text{cpcF} \) appears to be a mixture of phycocyanin rods and allophycocyanin core components
(Figs. 5C–E).

In addition to the phycobilisome proteins, Bands II and III from
both WT and \( \Delta \text{cpcF} \) contained FNRL (Figs 5C and D). Band II in
the CK mutant did not exhibit any FNR protein (Fig. 5D). FNRL
seemed to be located at the top of the sucrose density gradient in
all of the examined strains and did not appear to be associated
with assembled phycobilisomes (Fig. 5D and E). These results
indicate that the phycobilisomes present in the \( \Delta \text{cpcF} \) mutant

can associate with, and stabilize, FNRL. Additionally, these phycobilisomes appear to contain apo-CpcA. Apparently, integration of
apo-CpcA can support the assembly of modified phycobilisomes in
this strain.

Figure 4. Expression and assembly of FNR and phycobiliproteins in WT, \( \Delta \text{cpcF} \) and CK strains. Soluble proteins were separated
on either (A) denaturing LIDS-PAGE or (B) BN-PAGE. Proteins were
visualized either by Coomassie stain or immunodetection. (C) The grey
scale levels of the bands obtained from the LIDS-PAGE immunoblots
were semi-quantified and plotted. FNRL, FNRS, FNRA and FNRC are
the amounts of total FNR, the large form of FNR, the small form of FNR and
the degraded FNR observed, respectively. The standard curves used in
this semi-quantification are shown in Fig. S1. The error bars represent
\( \pm 1.0 \) SD; each experiment is the average of 3 independent
measurements. Symbols: *, FNRL; o, FNRS; ■, CpcA; ▼, CpcB; ▲, ApbA;
▼, ApbB.
Proper assembly of phycocyanobilin pigments into the \( \alpha \)-subunit of phycocyanin is required for optimal growth and light-harvesting

In cyanobacteria, phycocyanin is a major component of the phycobilisome light-harvesting apparatus. Holo-phycocyanin has a characteristic blue color, which is due to the covalent attachment of phycocyanobilin chromophores. Attachment of phycocyanobilins to the CpcA and CpcB apoproteins is catalyzed by a specific phycocyanin \( \alpha \)-subunit lyase (CpcE/CpcF) and phycocyanin \( \beta \)-subunit lyases (CpcT and CpcS/CpcU), respectively [6,24,25]. It should be noted that while the attachment of phycocyanobilin can occur spontaneously, this is a very inefficient process [26]. In our study, inactivation of the *Synechocystis* 6803 cpcF gene in the \( \Delta \)cpcF mutant and the absence of CpcA and CpcB in the CK mutant resulted in major growth (Fig. 1) and light-harvesting defects (Figs. 2–3). When examined under photoheterotrophic and mixotrophic growth conditions, the WT and mutant strains grew nearly equivalently (Fig. 1). Growth under autotrophic conditions, however, was compromised in the phycocyanin mutant strains. Under ambient and low CO\(_2\) conditions, both mutants grew poorly. This had been observed previously for other phycobilisome mutants including the CK mutant and a mutant completely lacking phycobilisomes [27]. It is possible that the absence of an intact phycobilisome affects the function of the Carbon Concentrating mechanism in *Synechocystis*. Interestingly, under the high CO\(_2\) growth condition, while the \( \Delta \)cpcF mutant exhibited...
significant growth, the CK mutant exhibited no detectable growth on plates. Very slow growth had previously been observed for this strain in liquid culture [27] at high CO2 growth conditions. At this point in time we cannot provide a definitive explanation for this observation. It is possible, however, that the absence of phycobilisome-associated FNRL in this strain (see below), severely compromises linear electron transport to NADPH. Under limiting NADPH conditions and at high CO2 concentrations, Rubisco would be operating at a maximal rate but the regeneration of ribulose 1,5-bisphosphate could be problematic.

Functionally, the light saturation curves for net photosynthesis (H2O to CO2) and PS II activity (H2O to DCBQ) did not show significant differences between WT and two mutant strains when using 650 nm red actinic illumination (Fig 2G and H). This indicates that the allophycocyanin phycobilisome core is functional in providing excitation energy to PS II in WT and both phycocyanin-deficient strains. However, when using white actinic light for these measurements, both mutants required a higher saturating light intensity, particularly for PS II activity (Fig 2D). This suggested that the higher light requirement was a result of defective energy transfer from phycocyanin. Since the PS II light saturation curves of the lyase mutant ΔcpeF and phycocyanin-null mutant CK were similar (Fig. 2D), it appears that the phycocyanin which accumulates in the ΔcpeF strain is not functional in energy transfer despite the presence of holo-CpcB (Fig 4A). This assertion is supported by the fluorescence induction experiments illustrated in Fig. 3. Both of the phycocyanin mutants exhibited a slow OJ transition when compared to WT under all conditions tested (Table 1, Fig. 3D-F). This appears to be due to defective energy transfer since, when WT was examined at a lower light intensity, this difference was nearly eliminated (Fig. S2). The lack of functional phycocyanin in both mutants appears to result in insufficient light-harvesting. Consequently, both mutants had lower maximal net photosynthesis rates than WT (Figs. 2A and E), which is in agreement with a slower growth rate under autotrophic conditions (Fig. 1). Interestingly, while the amount of CpcB protein in the ΔcpeF mutant dropped to less than half of the WT, the amount of CpcA showed a slight increase, even though no bilin chromophores were attached and this protein was present as apo-CpcA (Fig. 4A and D). The apo-CpcA protein accumulated and did not appear to be subject to rapid degradation, since it appeared to be assembled into phycobilisomes in the ΔcpeF mutant (Fig. 5A, D, E).

Truncated phycocyanin rods in the ΔcpeF mutant stabilizes FNRL

The phycobilisomes of the ΔcpeF strain appeared to be similar to those found in WT, although smaller in size. They contained the allophycocyanin core subunits, CpcA and CpcB, and several linker peptides (Fig. 5A and C, Band III). While lacking CpcC2, the ΔcpeF phycobilisomes were enriched in CpcG1 (Fig. 5A and C, Band III). The lack of the CpcC2 linker may explain the smaller size of these phycobilisomes since CpcC2 appears to be required for the assembly of distal phycocyanin rod hexamers [28]. CpcC2 was also absent in a cpeT mutant which lacked the CpcB phycocyanin lyase [23]. Most interestingly, the phycobilisomes from the ΔcpeF mutant contained large amounts of CpcA (Fig. 5C and D). Since most of the CpcA present was in the unpigmented apo-CpcA form (Fig. 4A) it appears that significant quantities of apo-CpcA can assemble into phycobilisomes in the ΔcpeF strain. In an earlier study [29], the *Synechocystis* sp. PCC 6701 cpeA and cpeB genes were expressed in a *Synechocystis* sp. PCC 6803 strain, 4R, which lacks phycocyanin due to an amber mutation in the cpeB gene. While the cpeA gene is intact in this strain, no detectable CpcA accumulates [30]. A site-directed mutation converting 85C to 85A in the heterologously expressed *Synechocystis* sp. PCC 6701 CpcA protein yielded the CD3 mutant. Since the CD3 strain lacks the phycocyanobilin binding site on CpcA this mutant is functionally equivalent to the ΔcpeF strain. The CD3 mutant accumulated only a small amount of total phycocyanin in its phycobilisomes (about 10% of control) and 90% of the CpcA which was present in these phycobilisomes was derived from *Synechocystis* 6803 holo-CpcA. Consequently, the phycobilisomes isolated from the CD3 strain appear to incorporate about 1% of *Synechocystis* 6701 apo-CpcA.

The phycobilisomes which accumulate in the ΔcpeF mutant appear to function poorly in the transfer of excitation energy to the allophycocyanin core and subsequently to PS II (Fig. 2 and 3). However, they do apparently functionally associate with FNRL. The failure to detect FNRL in Band II from the CK mutant [9]; Figs. 5C and D) suggested that only the phycocyanin proteins, and not allophycocyanin, had a high affinity to FNRL. One previous study has shown that FNR was found in sucrose density gradients in association with the allophycocyanin core from a phycocyanin deletion strain [31]. These authors proposed that FNR was associated with the core-proximal allophycocyanin hexamer. We suggest that, due to the smaller size of the allophycocyanin core, it migrates to a location near the top of the sucrose density gradient which is heavily contaminated by free FNRL. This is clearly seen in Fig. 5E. While the top of the gradient contains a large amount of FNRL, only a very small amount actually comigrates with allophycocyanin. Using cryoelectron microscopy and single particle analysis, Arteni et al. [32] proposed a model suggesting that the location of FNRL is located at the interface between the phycocyanin rods and the allophycocyanin core. Our data from the ΔcpeF mutant (Figs. 4 and 5) support this model. Using BN-PAGE, we also identified a subcomplex containing FNRL-CpcA-CpcB from both the WT and ΔcpeF strains, but not from the CK mutant. This indicates that the association of FNRL and phycocyanin can be maintained even in the absence of a high concentration of phosphate, a condition in which the phycobilisome largely dissociates (Fig. 4B). Additionally this indicates that the allophycocyanin core may not be required for the interaction of FNRL with the phycocyanin. The observations of decreased FNRL accumulation and its degradation in the CK mutant ([9]; Figs. 4A, and 5D and E) indicates that the FNRL protein is not stable in the absence of assembled phycocyanin. It should be noted, however, that the functionality of the apparent degradation products of FNRL, which are significantly larger than FNRL, has not been evaluated.

FNR has been co-purified from several thylakoid membrane protein complexes, such as cytochrome b6f complex and NDH complex, and it has been suggested that it serves as an electron donor during cyclic electron flow [33,34]. In cyanobacterial PetH (FNR) the N-terminal domain shares high homology with CpcD, which apparently allows a significant amount of FNRL to be bound to purified phycobilisomes ([35]; Figs. 5C-E). Since the artificial fusion of the N-terminal domain sequence of petH with the green fluorescence protein gene led to tight association of the fusion protein with phycobilisomes [36], it appears that the N-terminal domain functions to localize FNR to the phycobilisome. However, the physiological significance of this close connection between the enzyme catalyzing the final photosynthetic linear electron transport and light-harvesting complex in cyanobacteria remains unclear. A mutant which lacked FNRL exhibited a higher NADP+/NADPH ratio [12], which supported the earlier suggestion FNRL functions as a ferredoxin-NADP+ oxidoreductase.
during linear chain electron transport, supporting autotrophic growth, while FNR₄ is a better NADPH oxidase, accumulating when linear electron transport is compromised [9]. Consequently, the association of FNR₄ with phycocyanin is beneficial for linear electron transport. It was proposed earlier that formation of the phycobilisome-PS I trimer supercomplex was mediated by interactions with FNR [37]. In this model, FNR was placed at the core-distal phycocyanin rods. Later study indicated that binding of FNR to the phycobilisome did not affect energy distribution between PSII and PSI during state transitions [36].

Others have suggested that the association of FNR to the phycocyanin rod elements would position the enzyme in close proximity to the surface of the thylakoid membrane, facilitating its interaction with reduced ferredoxin as it is produced at the stromal surface of PS I [36]. It was also reported that the membrane localization of FNR was required for FNR-dependent cyclic electron flow induced by salt stress [38]. Our study indicates that apo-CpcA assembles into phycobilisomes in the ΔcpeF mutant. Although the phycocyanin rods are inefficient in transferring excitation energy to the allophycocyanin core (Figs. 2–3), the apo-CpcA-containing phycobilisomes appear to associate with FNR, stabilizing FNR₄ (Figs. 4–5). This would yield a direct benefit to the ΔcpeF strain when grown under autotrophic growth conditions (Fig. 1).

CO₂ fixation is the major electron sink for photosynthetic electron transport, as NADPH is required to produce glyceraldehyde-3-phosphate and, consequently, to regenerate RuBP. While both the ΔcpeF and CK mutants grew poorly autotrophically under ambient CO₂ conditions, the inability of the CK strain to grow in an elevated CO₂ environment, where WT and the ΔcpeF grow robustly (Fig. 1), is puzzling and we do not fully understand this observation. We speculate that under this growth condition, where maximum rates of CO₂ fixation would be expected to occur, limitations in the amount of available NADPH in the CK strain due to instability of FNR₄ could lead to a failure in the regeneration of the high amounts of ribulose 1,5-bis-phosphate required to support the high rate of CO₂ fixation. Additionally, inadequate glyceraldehyde-3-phosphate production could have an overall detrimental effect on photoautotrophy.

Conclusions

In the absence of a functional δ-phycocyanin subunit lyase in the ΔcpeF mutant apo-CpcA accumulates and is integrated into modified phycobilisomes. While these phycobilisomes were unable to efficiently transfer excitation energy to PS II, they were able to bind and stabilize FNR₄. These studies highlight the important role played by phycocyanin both in energy transfer and in the modulation of efficient linear electron transport via FNR.

Supporting Information

Figure S1 Quantification standards for the CpcA, CpcB, ApcB and FNR Proteins. The indicated amounts of wild-type protein samples were resolved by LIDS-PAGE, electroblotted onto PVDF membranes, blocked and probed with antibodies against CpcA, CpcB, ApcB or FNR. After incubation with an anti-rabbit IgG-horseradish peroxidase conjugate, the blots were developed using chemiluminescence and detected by exposure to X-ray film. The X-ray film was scanned and the protein amounts were semi-quantified by comparison of the integrated optical density with a dilution series of wild-type samples (5–75 μg protein). Signal intensities were analyzed by ImageJ [19]. The R² values for the linear regression of each standard curve are shown. Both FNR₄ and FNR₅ were detected with the anti-FNR antibody.

Figure S2 Chlorophyll fluorescence induction of WT and ΔcpeF under low light illumination. The cells were grown autotrophically at ambient CO₂. Chlorophyll fluorescence induction of dark-adapted cells at a chlorophyll concentration of 5 μg/ml in BG-11 medium was measured using orange-red light (625 nm) at either 1000 μmol photons/m²s (100%) or 500 μmol photons/m²s (50%). The chlorophyll fluorescence was presented either as (A) the raw fluorescence traces or (B) normalized using the equation Fluorescence = (1-F₀/F(t))/(F₀/FM). The samples for the measurements contained no additions. The error bars represent ±1.0 SD and in some instances were smaller than the symbols; each experiment is the average of 3 independent measurements.

Acknowledgments

We are grateful to Professor Ghada Ajlani from Centre National de la Recherche Scientifique for supplying the CK mutant and the FNR antibody. We also thank Professor Wendy M. Schluchter from the University of New Orleans for kindly providing antisera against CpcA, CpcB and ApcB. James V. Moroney from Louisiana State University provided the antibody directed against the large subunit of Rubisco. We also thank Dr. Johnna L. Roose and Professor Wendy Schluchter for critical reading of the manuscript.

Author Contributions

Conceived and designed the experiments: PZ TMB. Performed the experiments: PZ. Analyzed the data: PZ TMB. Contributed reagents/materials/analysis tools: TMB LKF. Contributed to the writing of the manuscript: TMB PZ LKF.

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