A novel high light-inducible carotenoid-binding protein complex in the thylakoid membranes of *Synechocystis* PCC 6803

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Exposure to high-intensity light (HL) adversely affects the photosynthetic performance, cell growth, and viability of photosynthetic organisms. The damage is largely attributed to oxygen-dependent destruction of the photosynthetic apparatus and other cellular components1,2.

Oxygenic photosynthetic organisms synthesize stress-associated proteins during exposure to HL. These proteins are often important for the acclimation of cells to HL. A family of HL-inducible genes, called *hli* or *scp* genes3,4, encoding high light-inducible polypeptides (HLIP) with similarity to the light-harvesting chlorophyll a/b-binding proteins (LHCP) of plants, was shown to be critical for the survival of cyanobacteria under HL conditions5. All four HLIPs in *Synechocystis* PCC 6803, i.e., HliA, B, C, and D, are associated with photosystem I (PSI), and stabilize PSI trimers specifically under HL conditions. HliA and HliB also interact with Slr11286.

Another stress-associated protein, the iron stress-inducible chlorophyll-binding protein IsiA, stabilizes PSI trimers under HL conditions. Interestingly, IsiA, which is expressed at low levels under normal growth conditions and is induced by both HL and oxidative stress, is critical for the formation of PSI trimers in low-intensity light7. This is in contrast to the expression of Psal and Psal, both of which are essential for the formation of PSI trimers8, but are not strongly modified by stress conditions as compared to IsiA or HLIPs.

Even though the PSI complexes of green plants and cyanobacteria have similar structures and carry out similar functions, i.e., mediating the electron transfer between luminal plastocyanin (or cytochrome c6) and stromal ferredoxin (Fd; or flavodoxin), there are several differences among them. One of the differences is that higher plants contain only monomeric PSI, whereas cyanobacteria harbor both monomeric and trimeric PSI9–11. It is unclear why PSI trimers exist in cyanobacteria; however, studies in *Spirulina platensis* provided some insight into this phenomenon. In this cyanobacterium, the absorption spectra of the chlorophyll (Chl) components of the PSI trimers and monomers differ12; only the PSI trimers contain the extremely red-shifted Chl (the so-called red chlorophyll), which absorbs at 735 nm and gives rise to a 760-nm fluorescence emission peak (F760) at 77 K. The PSI monomer shows emission peaks only at 725–730 nm under the same conditions. In addition, the 760-nm emission band is only visible under reducing conditions, directly reflecting the redox state of P70012,13. Red Chl is
thought to funnel light energy to P700\textsuperscript{14–16}, thereby increasing the cross-section of light absorption, or to dissipate excess energy into heat, thereby protecting PSI against photodestruction\textsuperscript{17}.

The orange carotenoid protein (OCP), a 35-kDa two-domain soluble protein, was shown to trigger fluorescence quenching of phycoobilisome (PBS) under blue-green light illumination\textsuperscript{18}, and was proposed to be a photoactive protein that senses light intensity and triggers photoprotection\textsuperscript{19}. The protein, which was first discovered by Holt and Krogman\textsuperscript{20}, contains one non-covalently bound carotenoid molecule\textsuperscript{20–24}. While the OCP is conserved among cyanobacteria except Prochlorococcus\textsuperscript{7}, the carotenoid differs in different species, being zeaxanthin in Anacystis nidulans and Lyngbya wholei\textsuperscript{19}, 3’-hydroxyechinenone in Arthrospira maxima, and a glycoside derivative of 3’-hydroxyechinenone in Synechocystis PCC 6803\textsuperscript{21,23,24,26}. Water-soluble OCP may function as a dimer in cells\textsuperscript{22,26}.

Here, we report the discovery of a novel high light-inducible carotenoid-binding protein complex (HLCC). This complex is normally concealed by trimeric PSI on a sucrose gradient, but was readily detected in a Synechocystis strain lacking Psal (i.e., ΔPsal) and hence lacking PSI trimers. The HLCC contains Slr1128, IsiA, PsaD, and HliA/B as its intrinsic protein components, binds zeaxanthin and myxoxanthophyll, and is critical for the survival of cyanobacterial cells in HL conditions.

Results

Induction of a novel carotenoid-binding membrane protein complex by HL treatment. Thylakoid membranes were isolated from Synechocystis PCC 6803 cells grown in LL or HL, solubilized in mild detergent, and fractionated by sucrose gradient ultra-centrifugation (Fig. 1). Three pigmented fractions were detected in the samples of wild-type cells grown in LL. The uppermost orange fraction, F1, contained mainly free pigments; the middle green fraction, F2, contained PSI and PSII monomers; and the lower dark fraction, F1, contained mainly free pigments; the middle green fraction indicates the presence of carotenoids. We further purified the fraction by Blue-Native PAGE. As shown in Fig. 2A, a single orange band was detected, indicating that there is one carotenoid-containing protein complex in the sucrose gradient fraction. We named this orange protein complex high light-inducible carotenoid-binding complex (HLCC). The orange Blue-Native band was then excised, denatured, and separated on a Tricine-SDS-PAGE gel (Fig. 2B), and four protein bands were revealed in the complex by silver staining. The bands were excised and identified by mass spectrometry. The three larger bands (from largest to smallest) contained the hypothetical protein Slr1128, the iron stress-induced chlorophyll-binding protein IsiA, and PsaD, a subunit of PSI, respectively. The fourth band at the lower end of the gel was identified as HliA and HliB.

To determine the pigment composition of the complex, the pigments were extracted and separated by HPLC according to\textsuperscript{29}. As shown in Fig. 3, myxoxanthophyll and zeaxanthin accounted for most of the pigment present in the HLCC, and there was a negligible amount of Chl a.

Figure 2 | Separation of the HLCC by Blue-Native PAGE followed by second dimension Tricine-SDS-PAGE. The HLCC fraction was carefully collected from the sucrose gradient and separated by BN PAGE (A), and the resulting single band was excised and denatured in 1.5 X SDS sample buffer and further separated on a 12–20% Tricine-SDS gel with 6 M urea (B). The proteins were visualized by silver staining and identified by mass spectrometry.

Slr1128, IsiA, PsaD, and HliA/B are intrinsic components of the HLCC. To characterize the HLCC further, we investigated whether IsiA, PsaD, and HliA/B are intrinsic components of the complex using anti-PsaD immuno-coprecipitation. Briefly, the sucrose gradient fraction containing the HLCC was collected and incubated first with anti-PsaD antibody and subsequently with protein G Sepharose beads. The beads were then loaded on an empty column and the protein complex was eluted and separated by SDS-PAGE. In addition to the heavy and light chains of the antibody, the anti-PsaD antibody identified Slr1128, IsiA, PsaD, and HliA/B as the major components of the HLCC (Fig. 4a).

We then confirmed these results by crosslinking analysis using SEAD (sulfosuccinimidyl-2(7-azido-4-methylcoumarin-3-acetamido)-ethyl-1,3’-dithiopropionate), a crosslinker with a fluorescent spacer arm that fluoresces brightly when the disulfide bond between the crosslinked proteins is cleaved. Briefly, the non-covalent protein-com
plex was collected from F3 of the sucrose gradient, and the crosslinker was covalently incorporated into the complex by chemical bonding to free amino groups, mainly from lysine residues. The proteins of the complex were subsequently crosslinked by a photochemical reaction of the crosslinker’s azido group. Cleavage of the disulfide group in the crosslinker by reduction released the fluorescently labeled proteins and removed the fluorescent group from proteins that were not cross-linked. The fluorescently labeled components of the protein complex were then separated by SDS-PAGE and detected by fluorescence imaging. As shown in Fig. 4B, when crosslinked with SEAD and separated by SDS-PAGE, the HLCC complex shows four fluorescent bands under UV light, which correspond to Slr1128, IsiA, PsaD, and HliA/B, further demonstrating that these proteins are intrinsic components of the HLCC.

The HLCC is induced by high-intensity light, iron starvation, and oxidative stress. We previously showed that one of the major components of the HLCC discovered in the current study, IsiA, could be induced by iron depletion, HL stress, and rose bangal (RB) treatment7. To study the transcript induction of the other major component of the HLCC, i.e., Slr1128, we cultured both the wild-type and ΔPsaL strains in BG11, with or without iron, to the mid-logarithmic growth phase (OD$_{730}$ ~ 0.8), and then diluted the cultures to OD$_{730}$ ~ 0.1 with fresh medium and subjected them to

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**Figure 3** | HPLC separation of the pigments in the sucrose gradient fractions. Pigments were identified by comparing their retention times and absorption spectra as described previously (Takaichi et al., 2001). Peaks are 1, myxoxanphyll; 2, zeaxanthin; 3, hydroxyechinenone; 4, chlorophyll a; 5, echinenone; and 6, β-carotene.
The HLCC collected from the sucrose gradient was (A) incubated with anti-PsaL and then Protein G resin, and the eluted proteins were separated by SDS PAGE, and (B) allowed to react in the dark with SEAD, quenched with lysine, photoactivated with UV light, and separated by SDS PAGE.

Deletion of slr1128 resulted in increased sensitivity to oxidative stress due to membrane peroxidation. As shown above, all components of the HLCC were induced by oxidative stress (Fig. 6). We previously showed that PsaL, Slr1128, and IsiA are all important for *Synechocystis* survival upon exposure to HL. To evaluate the function of Slr1128 in oxidative stress conditions, we treated the deletion strain ΔSlr1128 and the wild type with RB and MV in the presence and absence of the chloroplast protein synthesis inhibitor chloramphenicol (Cm) for 12 h. We then measured the D1 protein level by immunoblot analysis as an indicator of oxidative stress. The ΔSlr1128 mutant strain was more sensitive than the wild type to both RB and MV in the dark with SEAD, quenched with lysine, photoactivated with UV light, and separated by SDS PAGE.

Deletion of slr1128 resulted in impaired state transition. State transition is a physiological adaptation in cyanobacteria that balances the distribution of light energy absorbed by phycobilisomes between PSI and PSII. Prompted by the previous observation that deletion of isiA altered the state transition capacity, we examined whether state transition was also altered in the ΔSlr1128 strain. As shown in Fig. 8, the wild type has a fully functional state transition, as indicated by the fast and full relaxation of maximum fluorescence after high actinic light illumination (Fig. 8B and D). Therefore, the photoinhibitory effect of HL (400 μmol photon m⁻² s⁻¹) on the wild type is negligible. By contrast, the ΔSlr1128 mutant exhibited a very low state transition level and resulted in substantial photoinhibitory quenching, as indicated by its poor recovery of maximum fluorescence (Fig. 8A and C).

**Discussion**

In the current study, we report the discovery of the novel high light-inducible carotenoid-binding protein complex (HLCC), which is normally concealed by trimeric PSI on a sucrose gradient, in the ΔPsaL strain of *Synechocystis* PCC 6803 subjected to HL stress. The HLCC contains Slr1128, IsiA, PsaD, and HliA/B as its intrinsic components, and binds to zeaxanthin and myxothanxophyll.

Slr1128 is a hypothetical protein highly conserved in cyanobacteria; the amino acid sequence identity between homologs of this protein in various species of cyanobacteria is generally above 80%. It also appears to be present in plants, with homologs in plants sharing 59–66% amino acid sequence identity with cyanobacterial Slr1128. Slr1128 has a transmembrane domain at its N-terminus and is likely an integral membrane protein, as it was also identified as a thylakoid membrane protein by other researchers. Furthermore, Slr1128 exhibits high levels of similarity (around 35% amino acid sequence identity) to human and animal stomatins, which are thought to regulate an associated ion channel, and to a bacterial protease (HflC), but the similarity between Slr1128 and stomatins or HflC is not as high as that between the counterparts of these proteins in other cyanobacteria or plants. We previously reported that Slr1128 was associated with HliA/B, and suggested that it functions in photoprotection. Here, we found that Slr1128 was also present in the HLCC, and appears to be closely associated with the HL-inducible proteins, HliA, and HliB. This latter finding correlates well with our previous work.

The peripheral PsaD subunit is located at the stromal side of photosystem II, and is highly conserved in all photosynthetic organisms (including bacteria with Fe-S-type reaction centers). With PsaL, PsaD plays a critical role in the assembly and stability of PSI and PSII, and mediates Fd electrostatic guidance and docking on PSI. Superoxide is formed by Fd at PSI through the Mehler reaction.

In this study, we showed that PsaD, Slr1128, IsiA, and HliA/B formed a novel protein complex (the HLCC) in the cyanobacterium *Synechocystis* PCC 6803. Our earlier results showed that HliA and HliB stabilize PSI trimers, interact with Slr1128, and protect cells under HL conditions. As our recent findings also showed that IsiA, which was identified here as one of the major components of the HLCC, forms a supercomplex with PSI and PSII, IsiA is important both for maintaining trimeric PSI integrity and for acclimation to HL and may protect cyanobacteria from HL and oxidative stresses through state transition. Interestingly, we now found that the other major component of the HLCC, Slr1128, was also important for the...
response to oxidative stresses and state transition (Fig. 7 and 8). These results suggest that Slr1128 interacts with IsiA to form the HLCC and protects cyanobacteria from oxidative stresses by mediating state transitions.

Thus, we propose that, by interacting with the peripheral PSI protein PsaD, which provides a docking site for Fd and interacts with PsaB, PsaC, PsaE, and PsaL, the HLCC complex stabilizes trimeric PSI and protects the photosystems, especially PSI, from HL and...
oxidative stresses, probably through (1) direct scavenging of the superoxide and/or other reactive oxygen species (ROS) produced at Fd; and (2) mediating state transitions that are bridged by IsaA and Slr1128. The bound carotenoids, especially zeaxanthin, may function as antioxidants and directly dissipate excessive excitation energy.

**Methods**

**Growth conditions and HL treatment.** Synechocystis cells were cultivated in BG-11 medium with 10 mM TES, pH 8.2, at 30°C. The culture was bubbled with air under low-intensity light (L1: 40 μmol of photon m⁻² s⁻¹) or high-intensity light (L2: 400 μmol of photon m⁻² s⁻¹) conditions. For the experiments conducted under HL, cells in the mid-logarithmic growth phase (OD₇₅₀ ~ 0.8) were diluted with fresh medium to OD₇₅₀ ~ 0.1 and exposed to HL at 30°C.

**Thylakoid membrane preparation and fractionation of membrane protein complexes.** Thylakoid membranes were prepared as previously described with some modifications. Briefly, cell pellets derived from cells grown to the mid-logarithmic phase were resuspended in ice-cold SMN thylakoid buffer (50 mM 3-(N-morpholino)-propanesulfonic acid (pH 7.0), 0.4 M sucrose, 10 mM NaCl, 5 mM Mgl₂, and 1 mM freshly made phenylmethylsulfonyl fluoride). An equal volume of glass beads pre-wetted with thylakoid buffer was added to the cell suspension, and the cells were broken in a Bead-Beater with an ice-jacketed sample chamber in six glass beads pre-wetted with thylakoid buffer. The homogenate was centrifuged at 1,800 g for 10 min to remove unbroken cells, cellular debris, and glass beads. The membranes in supernatant were then pelleted by centrifugation at 50,000 g at 4°C for 60 min. After washing with 2 mM dodecyl maltoside to remove any remaining phycobilisomes, the membranes were washed twice and resuspended in thylakoid buffer to a chlorophyll a concentration of 1 mg/ml. The chlorophyll a concentration was estimated from the dimethylfluoride extract by the previously reported formula:

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\text{Chlorophyll a (μg/ml)} = 12.1 \times \text{OD664} - 0.17 \times \text{OD625} \quad \text{(Eq. 1)}
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To fractionate membrane protein complexes, 150 μl of 10% dodecyl maltoside was added to the thylakoid membrane to achieve a detergent to chlorophyll ratio of 15:1. The membrane was solubilized at 4°C for 30 min before it was loaded onto a 10–30% (w/w) step sucrose gradient and centrifuged at 160,000 g for 16 h at 4°C. Pigmented fractions were collected and stored at −80°C until use.

**Blue-Native, Tricine-PAGE, and immuno-Blotting.** Blue-Native PAGE was performed as described for electrophoresis in the second dimension, the single band of the blue native gel was excised and denatured in 1.5 X SDS sample buffer (50 mM Tris-HCl (pH 6.8), 3% SDS, 150 mM DTT, and 0.01% bromophenol blue) for 30 min at room temperature. The gel slice was then laid onto a 12–20% Tricine-SDS gel with 6 M urea as described. The proteins were visualized by silver staining.

Poly peptides resolved by SDS-PAGE were electro-transferred onto a PVDF membrane. After the blocking step, the membranes were incubated with polyclonal primary antibodies (Agrisera) followed by horseradish peroxidase-conjugated secondary antibody (Sigma-Aldrich). The reactive bands were detected using enhanced chemiluminescence detection reagents (GE Healthcare). The bands were quantified using ImageJ.

**Database searching.** All MS/MS samples were analyzed using Mascot (Matrix Science, London, UK; version 2.2.03). Mascot was set up to search the CIBInr_061307 database (selected for Bacteria, unknown version, 2419804 entries) assuming the digestion enzyme trypsin. Mascot was searched with a fragment ion mass tolerance of 0.5 Da and a parent ion tolerance of 2.0 Da. The isoelectric isomer derivative of cysteine was specified in Mascot as a fixed modification. S-carbamoylmethylcysteine cyclization (N-terminus) of the N-terminus, oxidation of methionine and acetylation of the N-terminus were specified in Mascot as variable modifications.

**Criterias for protein identification.** Scaffold (version Scaffold-01_07_00, Proteome Software Inc., Portland, OR) was used to validate MS/MS-based peptide and protein identifications. Peptide identifications were accepted if they could be established at greater than 80.0% probability as specified by the Peptide Prophet algorithm. Protein identifications were accepted if they could be established at greater than 95.0% probability and contained at least three identified peptides. Protein identifications were assigned by the Protein Prophet algorithm. Proteins that contained similar peptides and could not be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony.

**SEAD crosslinking.** The complex collected from the sucrose gradient was allowed to react in the dark with a 10-fold molar excess of freshly prepared SEAD. The reaction was quenched by adding a 30-fold molar excess of lysozyme, and the derivatized complex was separated from free reagent by gel filtration through spin chromatography columns (Bio-Rad, Hercules, CA) filled with porous polyacrylamide (Bio-Gel P-2 from Bio-Rad). Photoactivation was carried out for 15 min by exposing the sample to a black-ray, long-wave, 100-W ultraviolet lamp (Ultra Violet Products Inc., San Gabriel, CA). Then 4X Tris-glycine SDS reducing sample buffer was added at a ratio of 1:3 to the reaction mix, and the mix was incubated for 30 min at 22°C and then heated for 5 min at 75°C before electrophoresis.

**Anti-PsaD Immno-Coprecipitation.** The complex collected from the sucrose gradient was allowed to react with anti-PsaD (Agrisera) at 0.5–1 mg protein sample/1 μg antibody on ice for 90 min with occasional tube inversion. The mixture was then added to the SMN-equilibrated Protein G Resin (Sigma-Aldrich), and allowed to react on ice for 1 h with occasional inversion. The Protein G Resin was loaded on a column and washed with 5X bed volume of SMN with 0.1% n-dodecyl-beta-D-maltoside (DM, Sigma-Aldrich). The protein complex was then eluted with elution buffer (Citri acid 0.1 M, pH 2.0) and immediately neutralized with 1 M Tris-HCl (pH 8.5 to pH 7.4), and dialyzed against 20 mM Tris-HCl, pH 7.4. The whole procedure was performed at 4°C and in dim light or in darkness, whenever possible.

**Analysis of Carotenoid Composition by HPLC.** Pigments extracted with methanol were subjected to HPLC analysis essentially as described. Pigments were identified by comparing their retention times and absorption spectra.

**Analysis of state transition.** Wild-type and mutant cells in the mid-logarithmic growth phase (OD₇₅₀ 0.6–0.8) were collected and resuspended to an OD₇₅₀ of 0.6 with fresh BG-11 medium. State transition was analyzed by Chl fluorescence using a Dual-PAM-100 P700 & Chl Fluorescence Measuring System (Heinz Walz, Germany) as described previously. A far-red light was first applied to dark-adapted cyanobacterial cultures to fully oxidize PSI, which was followed by a series of saturation flashes to determine the S1 level. An actinic light of 100 μmol of photon m⁻² s⁻¹ was subsequently turned on to induce photosynthesis. The S2 level was then assessed by a saturation pulse before the far-red light was applied again to drive the S2-S1 transition.
Additional information
Competing financial interests: The authors declare no competing financial interests.

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