

Structural organization of an intact phycobilisome and its association with photosystem II

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Phycobilisomes (PBSs) are light-harvesting antennae that transfer energy to photosynthetic reaction centers in cyanobacteria and red algae. PBSs are supermolecular complexes composed of phycobiliproteins (PBPs) that bear chromophores for energy absorption and linker proteins. Although the structures of some individual components have been determined using crystallography, the three-dimensional structure of an entire PBS complex, which is critical for understanding the energy transfer mechanism, remains unknown. Here, we report the structures of an intact PBS and a PBS in complex with photosystem II (PSII) from *Anabaena* sp. strain PCC 7120 using single-particle electron microscopy in combination with biochemical and molecular analyses. In the PBS structure, all PBP trimers and the conserved linker protein domains were unambiguously located, and the global distribution of all chromophores was determined. We provide evidence that ApcE and ApcF are critical for the formation of a protrusion at the bottom of PBS, which plays an important role in mediating PBS interaction with PSII. Our results provide insights into the molecular architecture of an intact PBS at different assembly levels and provide the basis for understanding how the light energy absorbed by PBS is transferred to PSII.

Keywords: phycobilisome; photosynthesis; phycobiliprotein; electron microscopy

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Introduction

Phycobilisomes (PBSs) are attached to the cytoplasmic surface of thylakoid membranes and are responsible for the majority of light capture for photosynthesis in cyanobacteria and red algae [1-3]. A hemidiscoidal PBS, which is the most common PBS type, has two parts: the core and the peripheral rods. The core of this type of PBS consists of two, three, or five cylindrical substructures, and the peripheral rods radiate from the core. PBSs from *Anabaena* sp. strain PCC 7120 (*Anabaena* 7120, also known as *Nostoc* sp. PCC 7120) have a five cylinder

core and eight peripheral rods with a molecular mass of ~6 000 kDa [4-6].

PBSs are composed of phycobiliproteins (PBPs) and linker proteins. PBPs are brilliantly colored, water-soluble proteins bearing different numbers of chromophores, which are open-chain tetrapyrroles covalently bound to cysteine residues via thioether bonds [22]. Based on the bilin energy level, PBPs are mainly categorized into three types: phycoerythrins (PEs) or phycoerythrocyanins at the core-distal ends of rods (which absorb high-energy light), phycocyanins (PCs) at the core-adjacent portions of rods (which absorb intermediate-energy light), and allophycocyanins (APCs), the major components of the core (which absorb low-energy light) [1]. Although PBPs have different absorption spectra due to different bilin energy levels, they have similar crystal structures [7, 8] and assemble into PBSs through a common hierarchical organization. Two different subunits of PBPs, α and β , initially form a heterodimer ($\alpha\beta$), conventionally called the (β) monomer, which subsequently assembles into the ($\alpha\beta$)₃ trimer. The trimers are the fundamental assembly

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unit of PBSs, and stack face to face to form a hexamer with or without linker proteins in its central cavity [2, 9]. APCs in the core have a few variants, such as α^{LCM} (the α domain in L_{CM} ; L_{CM} is coded by *apcE*), α^{APC} -like variant (denoted $\alpha^{\text{AP-B}}$, coded by *apcD*), and β^{APC} -like variant (denoted $\beta^{18.5}$, coded by *apcF*) (Supplementary information, Figure S1A) [10].

Linker proteins are mostly colorless polypeptides that are involved in the assembly of PBSs at different levels [2]. These proteins include L_{R}/CpcC (PecC; involved in rod extension), $L_{\text{RT}}/\text{CpcD}$ (for distal termination of rods), $L_{\text{RC}}/\text{CpcG}$ (mediating the association between rods and the core), L_{C} (participating in the assembly of the core), and $L_{\text{CM}}/\text{ApcE}$ (involved in the assembly of the core and interactions between the core and the thylakoid membrane) [2, 11]. There are two types of conserved domains in linkers: Pfam00427 and Pfam01383. The Pfam00427 domain makes up the N-termini of L_{R} and L_{RC} . L_{CM} contains two to four (depending on the species) repetitive (REP) domains, which are homologs of the Pfam00427 domain. In *Anabaena* 7120, four copies of REP domains are present in L_{CM} [6]. The Pfam01383 domain constitutes the majority of L_{RT} and L_{C} , and the C-terminus of L_{R} (Supplementary information, Figure S1). The crystal structures of some isolated components of PBSs from different species have been determined. These include PEs [12, 13], PCs [8, 14-17], APCs [10, 17-19], and linker proteins [10, 20] (Supplementary information, Figure S1). The structural layouts of various PBSs have also been suggested in electron microscopic images [21, 22]. However, the precise position of PBPs and the linker proteins in the PBS remains largely unclear.

How light energy is transferred from PBS to the reaction center is another important unknown question. PBS can transfer energy to both photosystem II (PSII) and photosystem I (PSI) [2], and energy distribution between PSII and PSI is regulated by a process called state transition [23, 24]. Early reports showed that state transitions occurred in a time range of subseconds to seconds [25] and that this process is controlled by the redox state of electron carriers between PSI and PSII [26]. Liu *et al.* [27] recently reported the isolation of a PBS-PSII-PSI mega-complex, and suggested that PBS does not need to change its spatial position during state transitions. In this study, we investigated the three-dimensional (3D) structures of the intact PBS and PBS-PSII complexes from *Anabaena* 7120 using single-particle electron microscopy (EM) analysis.

Results

Overall structures of the PBS and PBS-PSII complexes

PBSs from *Anabaena* 7120 were purified (Supplementary information, Figure S2A-S2E) and negative-stain EM imaging of the samples showed clear side, top, and oblique views of the isolated PBS (Supplementary information, Figure S2F). The end-on-view averages indicate a two-fold symmetry, which was verified using a rotational cross-correlation coefficient analysis (Supplementary information, Figure S3A-S3B). We reconstructed the 3D structure of the intact PBS (Figure 1A-1C) using 32 966 particles, achieving a 21 Å resolution according to the FSC_{0.5} criterion (Supplementary information, Figure S3C-S3F). The hemidiscoidal structure has a length of 49.0 nm, height of 33.7 nm, and thickness of ~20.0 nm (Figure 1A-1C). All PBP trimers of the core and the rods could be unambiguously distinguished and located in the map (Figure 1A-1C and 1I). The core consists of three complete cylinders A1, A2, and B (each formed by four layers of APC trimers) and two half cylinders C1 and C2 (each formed by two APC trimers). Two basal core cylinders A1 and A2 are arranged in an antiparallel fashion (Figure 1C) and core cylinder B is located on top of the two A cylinders (Figure 1A). The two half core cylinders C1 and C2 are unique. In contrast to half core cylinders arranged in a side-by-side manner with other core cylinders in early models [5], C1 and C2 are arranged perpendicularly to the other core cylinders (Figure 1A). In addition, these half core cylinders have connections to all other PBS components (core cylinders A1, A2, and B, and all peripheral rods), suggesting that they are important in the overall organization of this PBS.

Eight peripheral rods emanate out of the hemidiscoidal plane from the core at departure angles between 5° and 30° (Figure 1A-1C and Supplementary information, Figure S4), which could increase the cross-section areas of PBS in light harvesting. Each rod in the bottom pair (Rb and Rb'), top pair (Rt and Rt'), and one of the side pairs (Rs1 and Rs1') has two hexamers, whereas the densities of the other pair of side rods (Rs2 and Rs2') were weak and small and each rod likely contains only one hexamer (Figure 1A). The length of the core cylinder (A1, A2, or B) formed by four APC trimers is 14.4 nm, which is longer than that of the peripheral rod formed by four PC trimers (12.4 nm; Figure 1B and 1C), reflecting differences in their organization. In both the bottom and side views, the trimeric discs of the two bottom core cylinders (A1 and A2) are stacked in a staggered manner, with disk 3 protruding towards the thylakoid membrane (Figure 1C and 1D). This structural feature is important for energy transfer from PBSs to PSII complexes that are embedded in the membranes (see Discussion further below).

We noted that a portion of the raw particles and two-dimensional (2D) class averages showed additional

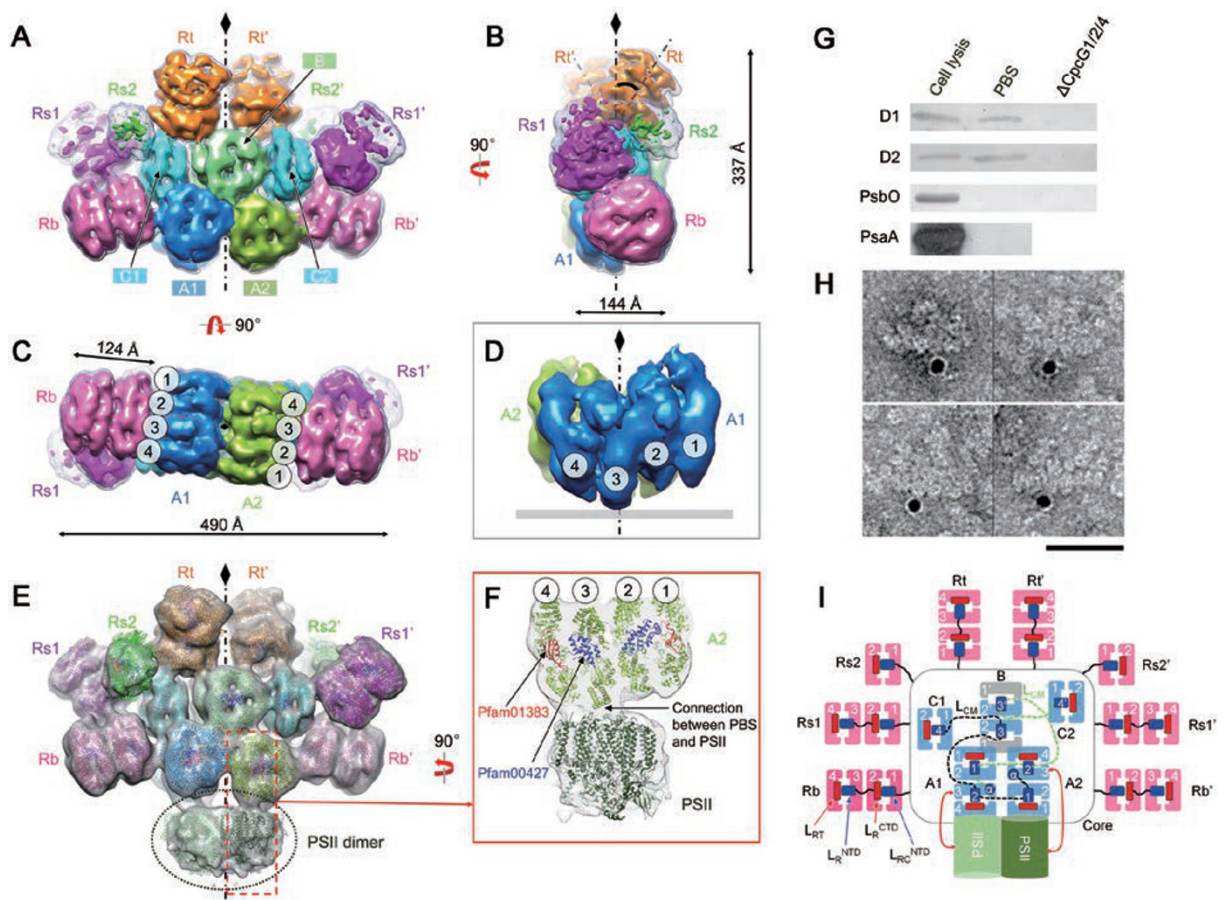


Figure 1 Overall structures of the intact PBS and the PBS-PSII complex. **(A-C)** EM density map of the PBS from front (**A**), side (**B**), and bottom (**C**) views. Cylinders of the core (A1, A2, B, C1 and C2) and rods (Rb, Rs1, Rs2 and Rt) are shown in different colors. The low-pass filtered (to 20 Å) map is shown in mesh at the lower threshold to show the side rods, Rs1 and Rs2. The tilted angle of Rt with the symmetry axis is indicated at 35° (**B**). **(D)** Extracted density map of core A1 and A2 showing the protrusion of disc 3 to the membrane. **(E)** EM density map of the PBS-PSII complex. Densities corresponding to PSII are circled. Available crystal structures of APC or PC trimers, linker domains (Pfam00427 and Pfam01383) and the PSII dimer are docked into the density map. **(F)** Zoomed view of the connection between PBS and PSII through disc 3 of the A2 cylinder. **(G)** Immunoblotting demonstrating that the reaction center subunits D1 and D2 of PSII can be detected in the PBS sample. Lane 1, cell lysis as a positive control; lane 2, PBS sample used for EM analysis in this study; lane 3, PBS sample taken from the Δ cpcG1/2/4 mutant (in which all core-rod association-related genes are deleted) as a negative control. **(H)** Immunogold labelling EM of PBS preparation using antibody against PSII subunit D1. Scale bar, 500 Å. **(I)** Schematic model of the PBS-PSII architecture. Dashed lines indicate connections between the REP domains of L_{CM}, which have not been experimentally verified. Blue box: the Pfam00427 domain; red box: the Pfam01383 domain.

density attached to the bottom of the PBS in a position where PSII should bind [28-32] (Supplementary information, Figure S5A). We screened these particles and obtained a 3D reconstruction, in which we can observe a clear density with a volume in agreement with the size of a PSII dimer (Figure 1E and Supplementary information, Figure S5B). To test whether the density is contributed by PSII, we first performed an immunoblotting experiment, which showed that the PBS sample contained the reaction center subunits (D1 and D2) of PSII [33, 34] but no PsaA (a PSI protein) or PsbO (a PSII peripheral

protein required for normal oxygen evolution) (Figure 1G). Second, we performed immunogold labelling experiments using antibody against D1 and observed gold particle labelling at the bottom of the PBS, which is consistent with the assignment of the density as PSII (Figure 1H). Assigning this density as PSII was further supported by a satisfactory docking of the crystal structure of PSII [34] into the 3D map (Figure 1E-1F). The above evidence suggests that these particles represent PBS-PSII complexes. Notably, the PSII density directly connects to the protrusions of disc 3 of the core substructures A1 and

A2 (Figure 1E and 1F). Coincidentally, the protrusion of disc 3 fits well with the cavity at the PSII surface (Supplementary information, Figure S5C and S5D).

Core and rod assembly

The crystal structure of APC in complex with the linker protein Lc (the Pfam01383 domain) from *Mastigocladus laminosus* (PDB code: 1B33) [10] and a recently reported crystal structure of the N-terminal domain of linker L_R (the Pfam00427 domain) from *Synechocystis* sp. PCC 6803 (PDB code: 3NPH) [20] were used for docking analysis of the core. The crystal structures docked well into the density maps of the core, with a cross-correlation coefficient of ~0.8 (Figure 2A, Supplementary information, Table S1 and Movie S1). Layers 1-2 of the four trimeric discs of cylinder A1 formed a

face-to-face hexamer, and layers 3-4 did the same. The two hexamers were further assembled in a back-to-back orientation to form the cylinder. All hexamers in the core are similarly packed, as suggested via crystallography (Supplementary information, Figure S6A and S6B). Within each cylinder A, all of the trimers have additional densities occupying their central holes. The central density of the trimer of disc 3 is significantly larger than that of the other three trimers (Figure 2B and 2E), suggesting different components in this trimer. Together with the fact that disc 3 directly connects to PSII, which suggests that α^{LCM} is located in disc 3, we hypothesize that the extra density in disc 3 could be partially contributed by the loop between α^{LCM} and REP1 and/or the inserted PB loop in α^{LCM} [2, 35] (Figure 2E and Supplementary information, Figure S1A). Six Pfam01383 domains from L_C

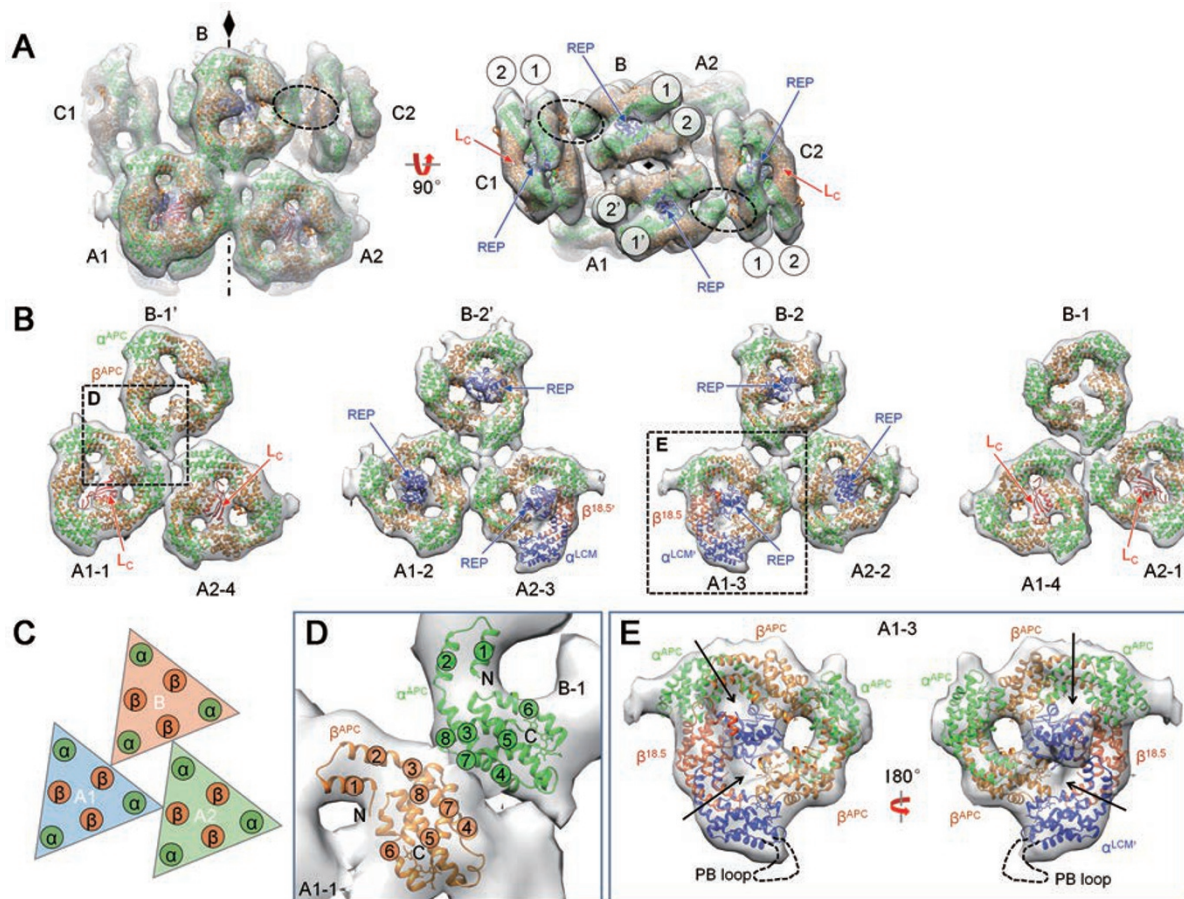


Figure 2 Assembly of the core. **(A)** Extracted density map of the core with fitted crystal structures shown in side and top views. Connections between cylinders C and B via α - β subunit interactions are circled. **(B)** The inter-cylinder interaction in cylinders A1, A2 and B is shown layer by layer. **(C)** Schematic model showing the interaction of the inter-cylinder trimers. **(D)** Zoomed view of α - β interaction mediating the interaction of inter-cylinder trimers. Helices of each subunit are numbered from the N- to C-terminus. **(E)** Details of disc 3 docking, where the terminal emitter from L_{CM} locates. The position of PB loop is indicated by a dashed line. The filled holes of the trimer are indicated with arrows.

linker proteins can be well fitted into the central densities of the outer APC trimers of cylinders A (four copies) and C (two copies). In contrast, no such central densities corresponding to the Pfam01383 domain were present in cylinder B (Figure 2A and 2B). Eight Pfam00427 domains, likely derived from two L_{CM} proteins as each L_{CM} contains four REP domains (Supplementary information, Figure S1A), can be fitted into the densities of the inner APC trimers of cylinders A (four copies), B (two copies), and C (two copies) (Figure 2A and 2B). The connections of these REP domains mediated by loops [36] cannot be clearly observed at the current resolution. The Pfam01383 domains are located inside the trimer plane, as in the crystal structure [10], whereas the Pfam00427 domains protrude from the trimer plane (Figure 2B and Supplementary information, Figure S6A). APC trimers in the core (cylinders A1, A2, and B) are associated via a specific pattern, in which a corner of the triangle-shaped trimer directly contacts the edge of another trimer (Figure 2B and 2C). This corner-edge association pattern, which

is the basic connection mode found in the core, is likely mediated by the interaction between α and β subunits (Figure 2D).

For the docking analysis of the peripheral rods, the crystal structures of PCs (PDB code: 1JBO) [14], the Pfam01383 domain (isolated from the APC-trimer-Lc complex, PDB code: 1B33) [10], and the Pfam00427 domain (PDB code: 3NPH) [20] were used (Figure 3A and 3B). The shape of the density maps of the PC trimer was similar to that of the APC trimer with a small difference due to an additional loop in the PC [17] that can be distinguished in the density map (dashed circles for Rb-1 in Figure 3B). A typical rod cylinder in our reconstruction contains two PC hexamers associated with two Pfam00427 domains and two Pfam01383 domains. The PC hexamer is formed by a face-to-face interaction of PC trimers in the same manner as in the crystal structure of the PC hexamer (Figure 3C and Supplementary information, Figure S6C and S6D).

In the core, the Pfam01383 domains were found

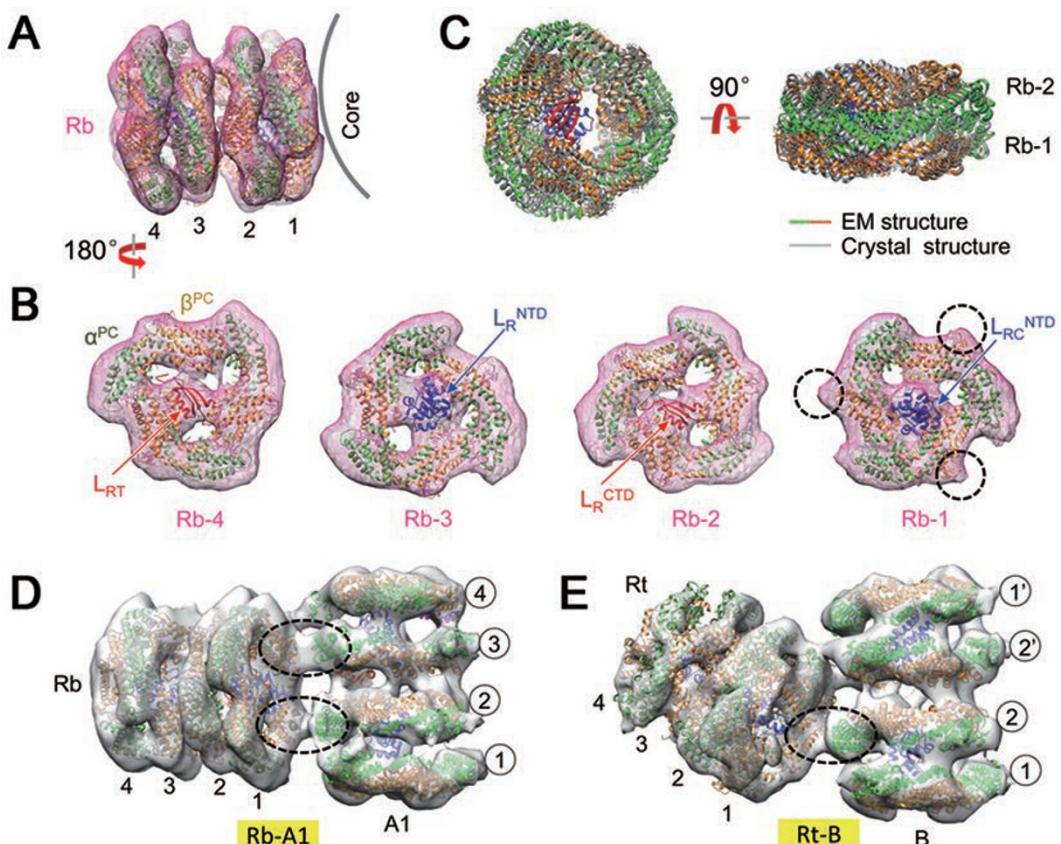


Figure 3 Assembly of the rods. **(A, B)** Docking of the bottom rod (Rb). Additional loops in the crystal structure of PC trimers (compared with the APC trimer) are distinguishable in the EM density map and are indicated by dashed circles for Rb-1. **(C)** Comparison of EM and crystal structures of the PC hexamer. **(D, E)** Connections between the bottom/top rod (Rb/Rt) and the core cylinder (A1/B). The α - β inter-trimer interactions, which mediate rod-core connections, are circled.

in the outer trimers and the Pfam00427 domains were found in the inner trimers. In contrast, the Pfam01383 and Pfam00427 domains in the rods appear alternatingly from the terminal trimer (Figure 3B). L_{RC} binds to the innermost trimer (Rb-1) through its N-terminal Pfam00427 domain (Figure 3B), whereas its C-terminus (L_{RC}^{CTD}) is likely anchored to the core. However, the density corresponding to L_{RC}^{CTD} cannot be clearly distinguished in the map, indicating that the C-terminus of L_{RC} does not fold into a globular domain as does its N-terminus. L_R connects the two PC hexamers by binding to the external hexamer (Rb-3) through its N-terminal Pfam00427 domain and to the internal hexamer (Rb-2) through its C-terminal Pfam01383 domain (Figure 3B). L_{RT} (the Pfam01383 domain) terminates the rod extension by binding to the outermost PC trimer (Rb-4) (Figure 3B).

A careful analysis of the interactions between rods Rt and Rb and the core components suggests that they also use the corner-to-edge interaction mode for rod-core assembly (Figure 3D and 3E). The corner-to-edge interaction pattern, which is likely mediated by the inter-trimer α - β subunit interaction [19, 37], may also provide a major force holding rods and core together. Although Rb and Rt are similarly geometrically oriented relative to their interacting core cylinders A1 and B, respectively (Figure 3D and 3E), this geometric orientation is not found between Rs1/Rs2 (and Rs1'/Rs2) and their associated half core cylinders C1 and C2, suggesting a variable association geometry between rods and the core [38].

Interactions between the rods and the core

We next investigated the roles of the linker protein L_{RC} in the rod-core association. L_{RC} s are encoded by *cpcG* genes (Supplementary information, Figure S1A). Four *cpcG* genes (*cpcG1-4*) exist in *Anabaena* 7120. CpcG3 (also named cpcL [39]) is equivalent to CpcG2 from *Synechocystis* sp. PCC 7002 and plays a role in the formation of a special PBS complex [40, 41]. A recent study demonstrated that CpcG3 is important for the association of a special PBS with PSI in *Anabaena* 7120 [39]. CpcG1, CpcG2, and CpcG4 of *Anabaena* 7120 are functionally similar to CpcG1 of *Synechocystis* 7002 and are responsible for the rod-core association [4, 5].

To study the specific roles of these L_{RC} proteins, we generated a series of *cpcG* mutant strains, which covered all combinations of deletions of three linker genes involved in the rod-core association, i.e., *cpcG1*, *cpcG2*, and *cpcG4* (Figure 4A and Supplementary information, Figure S7A-S7C). We isolated PBS complexes from these mutant strains and examined their assembly (Figure 4A) and energy transfer ability (Figure 4B). The role of each L_{RC} was assigned based on the EM images of PBSs

isolated from the strains mentioned above (Figure 4A and Supplementary information, Figure S7E). CpcG4 is responsible for the connections of both the top and bottom pairs (Rt and Rb) of rods to the core. CpcG2 is necessary for connection of the larger side rod Rs1 to the core, whereas CpcG1 is required for the association of the smaller side rod Rs2 to the core (Figure 4A). Thus, these L_{RC} proteins are responsible for recognizing the specific sites on the core, and each individual rod may interact with the core independently.

The EM images of PBSs from the $\Delta cpcG1/2/4$ mutant strain, which had all three *cpcG* genes deleted, showed that no rods were attached and that the densities of the half core cylinders C1 and C2 became faint and flexible (Supplementary information, Figure S7D), suggesting that rod binding may help stabilize the half core cylinders. Figure 4B shows fluorescence emission spectra of the wild-type and mutant PBSs. The spectrum of the $\Delta cpcG1/2$ mutant, in which no peripheral rods were attached to the half core cylinders C1 and C2, had an additional peak at 670 nm, indicating that some energy presumably escaped from C1 and C2. In wild-type and mutant PBSs, no peripheral rods were found at the bottom side of the PBS core. However, when we studied the PBS from a mutant lacking the key core gene *apcF* (encoding $\beta^{18.5}$), we frequently observed one or two peripheral rods attached to the bottom of the cores (Figure 4C). This result suggests that an extra site for peripheral rod attachment might be provided in the $\Delta apcF$ mutant, where ApcF ($\beta^{18.5}$) is possibly replaced by ApcB (β^{APC}).

Pigment arrangement and energy transfer

The global distribution of chromophores in the PBS was investigated using results of docking analyses (Figure 5A and Supplementary information, Movie S1). The Rb and Rt rods have structural connections with the core cylinders A and B, respectively. They also have connections with the half core cylinders C1 and C2. The side rods Rs1 and Rs2, however, are only spatially attached to the half core cylinders C1 and C2. Based on the analysis of distances between bilins (Supplementary information, Figure S8A-S8D), energy harvested by the bottom rods Rb could be transferred to the core cylinders A directly or through the half core cylinders C1 and C2. The top rods Rt could transfer their energy to the core cylinders A through either core cylinder B or the half core cylinders C1 and C2. However, side rods (Rs1 and Rs2) could only transfer their energy via half core cylinders C1 and C2 (Figure 5B). This structural feature suggests that the half core cylinders C1 and C2 play an important role in connecting the rods with the core during energy transfer.

Pigments in the basal core cylinders (A1 and A2) form

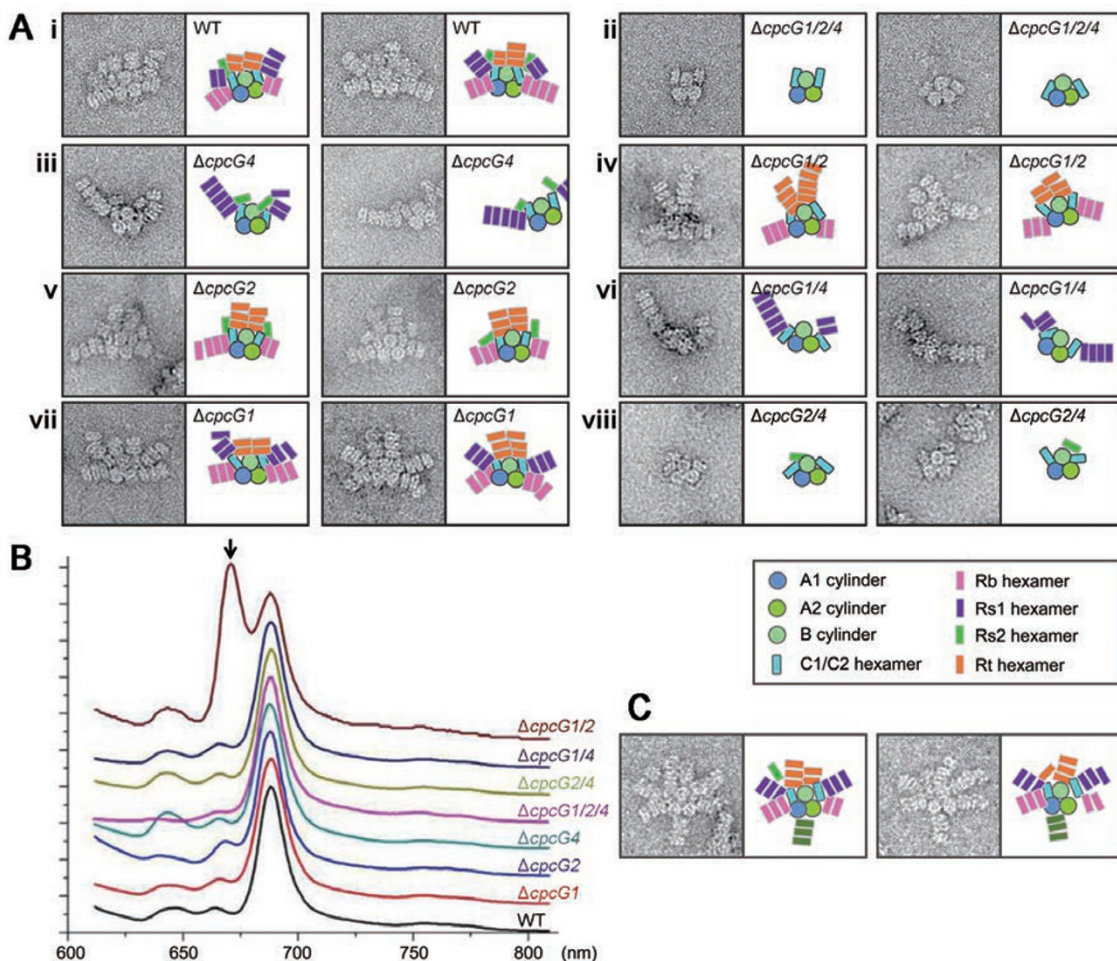


Figure 4 Interaction of rods with the core. **(A)** Typical negative-stain EM images of a wild-type and mutant PBSs. The schematic models for each sample are shown to the right of the images. **(B)** 77K fluorescence spectra of wild-type and mutant PBSs under 590 nm excitation. The unique peak at approximately 670 nm in the $\Delta cpcG1/2$ mutant (indicated with an arrow) may be caused by a leak of energy from the half core cylinders that cannot efficiently transfer the energy within the core. **(C)** Negative-stain EM images of PBSs from the $\Delta apcF$ mutant. Note that a rod is attached at the bottom of the PBS.

an arch when viewed from the face side. The two copies of chromophores bound to ApcE (terminal emitters) are located at the bottom of the arch (Figure 5A). This arrangement could efficiently transfer and converge absorbed light energy to the terminal emitters that in turn transfer energy to PSII. The other subunit containing a far-red-emitting chromophore, ApcD, which is required for energy transfer to PSI, could not be unambiguously identified in the PBS core structure. Analyses of the distance of the bilins of the PBSs closest to the chlorophylls in PSII suggest that disc 3 is spatially close to chlorophylls in the antenna protein CP43 of PSII [34] (Figure 5C). The nearest chlorophyll *a* to disc 3 (with an estimated distance of 48 Å, Supplementary information, Figure S8E) is located at the periphery of PSII and is the only chlorophyll *a* with a central magnesium atom that is

coordinated by an asparagine residue, CP43-Asn39 (others are coordinated via either water or a histidine residue [34]).

Discussion

The primary functions of PBS are the absorption of photons and the transfer of the energy to photosynthetic reaction centers. Highly ordered chromophores in PBS are very important for energy absorption and transfer. These functions are achieved via a hierarchical spatial organization of PBPs, in which linker proteins play critical roles. Extensive efforts have been made to determine the 3D structures of intact PBSs, but limited progress has been made. Adir and colleagues recently crystallized two intact PBSs; however, the overall structure of the

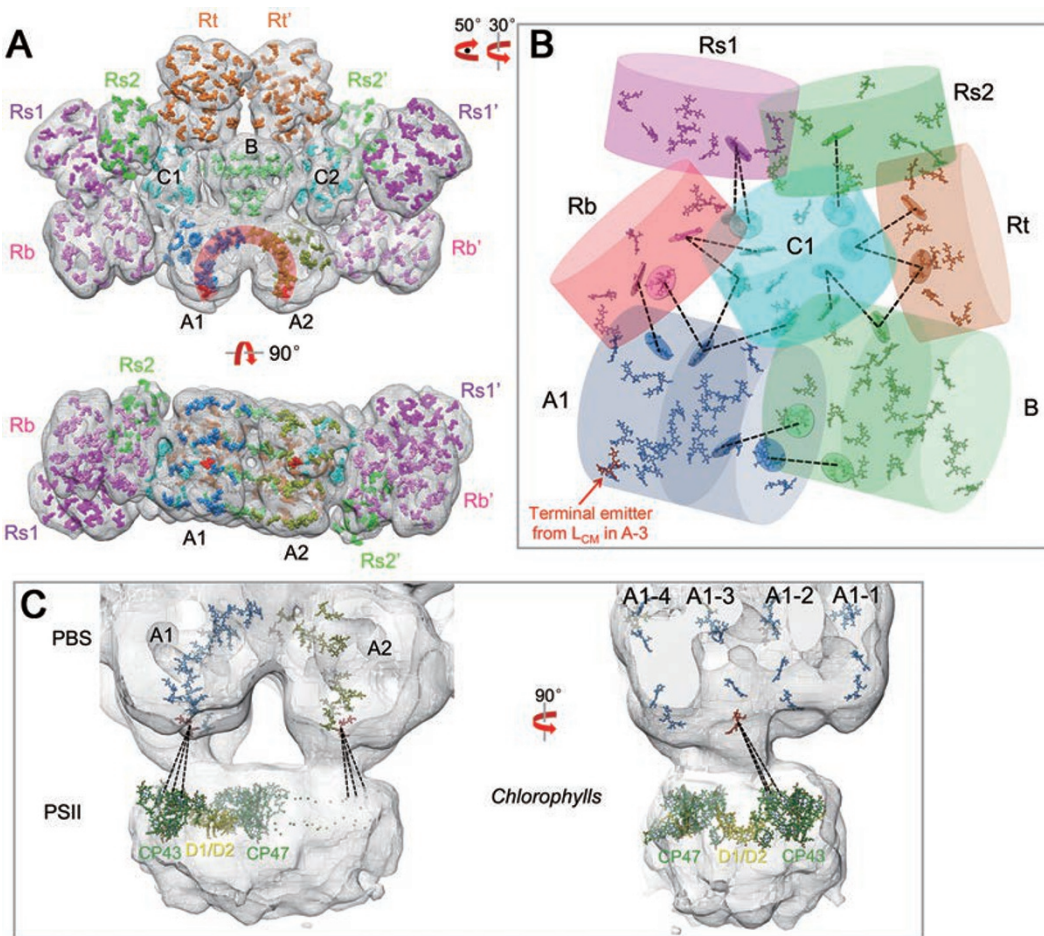


Figure 5 Pigment arrangement and energy transfer pathways. **(A)** Pigment arrangement in the intact PBS. The pigments are presented as surfaces and colored according to their cylinders shown in Figure 1A. Pigments from A1 and A2 (front view) form an inverted 'U' shape (colored dark orange), in which the terminal emitters from disc 3 are colored red. **(B)** Energy transfer from the rods to the core. All pigments in core cylinders A1, B, and C1 and the pigments from the core-connected PC trimers are shown. The distances of one or two nearest pigment (from rods to the core) are indicated by dashed lines. The disc covering a pigment is the plane defined by all the atoms of that pigment. **(C)** Energy transfer from PBS to PSII. The EM density map of the PBS-PSII complex is shown in mesh. Pigments from core cylinders A1 and A2 are presented, with the terminal emitter from disc 3 in red. Chlorophylls from one PSII monomer (35 chlorophylls) are presented as a stick model (chlorophylls from CP43/CP47 and D1/D2 are colored dark green and yellow, respectively). Close connections between A1-3 and CP43 are indicated using dashed lines.

PBS remained unsolved due to random positioning of the core components in the crystal [38]. The density maps in previous EM studies of intact PBSs were of insufficient quality to determine the accurate docking of crystal structures of PBPs and linker proteins [21, 22]. To improve PBS particle quality for EM studies, we isolated PBSs from different types of cyanobacteria. We found that PBSs from *Anabaena* 7120 yielded the best results with negative-stain EM. We also performed a cryo-EM study, but obtained poorer results due to the low contrast (possibly because of the high salt concentration in the buffer) and the preferred orientation (only the end-on

view shown in vitreous ice).

In our 3D reconstruction, the structures of the PBP trimers and the two linker protein domains (Pfam00427 and Pfam01383) can be unambiguously identified in both the core cylinders and the peripheral rods (Figure 1I), including the L_{RC} (CpcG) linker proteins that are required for the attachment of the peripheral rods to the core [42]. In hemidiscoidal PBSs, the core structures are organized by L_{CM} proteins, which have two, three, and four linker domain (REP domain) repeats in so-called two, three, and five cylinder cores, respectively. The presence of C cylinders in the core of PBS from *Anabaena* 7120 pro-

vides two additional attachment sites for peripheral rods compared with the PBSs, which have cores of three cylinders. Furthermore, two additional CpcG linkers are utilized for rod attachment. Our results suggest that CpcG4 is the primary rod-core linker that connects the majority of rods to the core, and CpcG1 and CpcG2 are likely involved in the attachment of additional rods. The fact that different linker proteins are required for different rod attachment indicates that the linker proteins play a direct role in rod attachment to the core.

Three important PBPs are located in the A cylinders of the PBS core, namely L_{CM} , $\beta^{18.5}$ and α^{AP-B} , which are encoded by *apcE*, *apcF*, and *apcD*, respectively. The density map (Figure 2) and physical position of the chromophores suggest that L_{CM} /ApcE and $\beta^{18.5}$ /ApcF in disk 3 of cylinder A are responsible for the protrusion of this trimer at the bottom of the PBS core. This protrusion fits well with the hole on the cytoplasmic side of PSII and forms a tight interaction between PBS and PSII, which is likely required for an efficient energy transfer from PBS to PSII. The tight association between PBS and PSII agrees with a recent report that demonstrated the formation of the PBS-PSII-PSI megacomplex *in vivo* via protein cross-linking and mass spectrometry analyses [27]. It should also be noted that EM imaging showed peripheral rods attached to the bottom of the cores of PBSs isolated from the $\Delta apcF$ mutant (Figure 4C). ApcF ($\beta^{18.5}$) is critical for energy transfer from PBS to PSII [41, 43] and has a C-terminal extension compared with ApcB (β^{APC}). This intriguing observation suggests that ApcB may not fully substitute for ApcF function in the PBS core.

The EM structures and the pseudo-atomic model were used to map the global distribution of chromophores and thereby provide the basis for understanding energy transfer from the rods to the core and from PBS to PSII. Many questions remain to be answered. For instance, the position of ApcD, the terminal emitter for energy transfer to PSI [43], could not be unambiguously identified in our PBS model (Supplementary information, Figure S9), and further study is needed to determine whether ApcD is located on disc 1 or 4 [44].

Materials and Methods

PBSs preparation

Anabaena sp. PCC 7120 was cultured in BG11 liquid media, bubbled with air containing 1% CO₂. Cells (3–5 g) were collected and every 1 g pellet of cells was resuspended in 8 ml buffer C (0.8 M K/Na-PO₄, pH 8.0). Samples were disrupted through a French Pressure at 20 000 psi at 4 °C for three times. Undisrupted cells and debris were removed by centrifugation at 3 300 g for 10 min. Triton X-100 (20% (w/v)) was added to the supernatant to a final concentration of 2%. The mixture was gently shaken at room tem-

perature for 30 min, and then centrifuged. The supernatant containing PBSs was loaded to the top of sucrose density gradients. The sucrose gradients were made from buffer C by adding sucrose to these concentrations: 0.25, 0.4, 0.55, 0.7, 0.85, and 1.0 M. Sucrose solution at each concentration was then manually added to a centrifuge tube layer by layer with volume of 3, 3, 3, 4, 4, and 5 ml, respectively. The samples were centrifuged at 35 000 rpm. for 6 h using a Ti-70 rotor on Beckman Optima L-80XP centrifuge and resulted in a visible bands of PBSs, with band 1 as the main layer of intact PBSs (Supplementary information, Figure S2A).

Negative-stain EM

PBSs of *Anabaena* sp. PCC 7120 (4 µl, 500 µg/ml) separated from the sucrose gradient centrifugation were applied to glow-discharged EM grids with continuous carbon film. After 30-s incubation, excess sample was blotted, and the grid was washed five to six times with buffer C to remove sucrose in the sample. After washing, the remaining liquid was removed with the filter paper. Then, 2% (w/v) uranyl acetate was added onto the grid from one side continuously, and was removed with filter paper from the other side simultaneously. This process removed the deposition formed by phosphates and uranyl acetate immediately after its formation. Deposition was thoroughly removed from the grid, and the stain was left on the grid for 30 s. Excess stains were removed and the grid was dried for EM imaging.

Samples were examined on a Tecnai F20 electron microscope (FEI) at 200 kV. Images were recorded on a Gatan 4k × 4k CCD camera at a nominal magnification of 50 000, yielding a pixel size of 2.23 Å at the sample level. Defocuses were set at about –1 200 nm.

Data analysis and 3D reconstruction

For data analysis, particles were manually selected using the EMAN [45] program boxer. In total, 32 966 particles were picked from 2 108 micrographs. The 2D classification was done using the EMAN program *refine2d.py*. Rotational cross-correlation coefficient values (Supplementary information, Figure S3B) were calculated by ‘CC C’ program in SPIDER [46]. The initial PBS model was determined using common lines method (*startAny* program in EMAN).

For reconstruction of the structure of the PBS-PSII complex, two initial models were used for multirefinement in EMAN using the *multirefine* program. The first initial model was PBS reconstructed with all the particles, while the second one was a combination of the first initial model with a PSII density map transformed from its crystal structure and added up to the bottom of PBS randomly. Both initial models were low-pass filtered to 60 Å before multirefinement. Finally, 2 363 particles were classified to the second group of PBS-PSII complex, reaching a resolution of ~34 Å by FSC = 0.5 criterion.

Docking analysis and measurements of distances, axes, and angles

For docking analyses, the crystal structure of the APC trimer (PDB code: 1B33) was fitted onto the density map using ‘fit in map’ program in UCSF Chimera [47]. Map segmentation and visualization were performed using Chimera.

Centroids of each trimer and axes of each cylinder (Supplementary information, Figure S4) were determined by ‘centroid’

and ‘define axis’ commands in Chimera, respectively, based on the docked crystal structures. Distances of pigment with pigment were measured by ‘distance’ command in Chimera. Angles (Figure 1B and Supplementary information, Figure S4C and S4D) were measured by ‘angle’ command in Chimera.

Construction of PBS mutants

In *Anabaena* sp. PCC 7120, four L_{RC} genes are located at the 3' portion of the *cpc* operon, tandemly in the sequence *cpcG1-4*. To construct Δ*cpcG1/2/4* mutant, *cpcG* flank sequences were amplified by PCRs, using the total genomic DNA as template. The 3.2-kb upper stream sequence was amplified by primer pair PN-up and PC-up (Supplementary information, Table S2), digested with *Xba*I and *Eco*RI and ligated into pBS at the same restriction sites, and resulted in pBS-Gup. The 3.3-kb downstream sequence was amplified by PN-down and PC-down, digested with *Nor*I and *Sac*I and ligated to pBS-Gup at the same restriction sites, and resulted in PBS-Gup-Gdown. A cartridge of Em' was amplified by PN-EmS and PC-Em, digested with *Eco*RI and *Nor*I, and ligated into pBS-Gup-Gdown at the same restriction sites. The result plasmid was pBS-Gup-Em-Gdown. The 7.6-kb *Xba*I-*Sac*I fragment from pBS-Gup-Em-Gdown was cloned into the pRL277 plasmid for transformation of *Anabaena* sp. PCC 7120 by conjugation (Supplementary information, Figure S7A).

In construction of other L_{RC} mutant strains, combinations of *cpcG* fragments were amplified by corresponding primers and ligated with each other and later with cartridge of Emr (Supplementary information, Table S2). The combined sequences were ligated into pBS-Gup-Gdown as former single cartridge of Emr, and later to pRL277 plasmid in the same manner as above to generate corresponding plasmids for transformation (Supplementary information, Figure S7A).

To construct *apcF* mutant strain, a 7.5-kb sequence including *apcF* gene and its flanking sequence were amplified by primer pair PN-Fup and PC-Fdown (Supplementary information, Table S2) and cloned into Promega pGEM-T easy vector to generate pTapcF-1. Primer pair PN-Fdown and PC-Fup (Supplementary information, Table S2) was used to amplify the 9.9-kb fragment by reverse PCR from pTapcF-1. A cartridge of Emr was ligated with the 9.9-kb fragment to generate pTapcF-2. The 8.4-kb *Xba*I-*Bgl*II fragment of pTapcF-2 was cloned into the pRL277 plasmid for transformation.

Segregation of mutants was confirmed by PCR analysis (Supplementary information, Figure S7B). The PBSs of mutant strains were prepared in the same way as wild-type ones, with differences in layer patterns and main layer positions for their PBSs.

Absorption and fluorescence spectra

Before spectra analysis, OD_{730 nm} of all PBS samples were adjusted to 0.8 with fresh BG11 medium. The absorption spectra between 400–800 nm were acquired by Shimadzu UV-2501PC. 77K fluorescence emission and excitation spectra were acquired with PTI fluorescence spectrophotometer. PBS samples for tests were in thermal equilibrium with liquid nitrogen in a Dewar vacuum flask.

Immunoblotting

PBSs from sucrose density gradient centrifugation were not suitable for direct use in SDS-PAGE because of high concentration

of phosphate. Desalting was performed with ultrafiltration using Milipore Amicon Ultra centrifugal filters. The desalted samples were measured by Bradford method and then diluted to 1 mg/ml. Approximately 4 µg of sample was loaded to each lane and resolved by SDS-PAGE. The samples were probed with antibodies against D1, D2, PsbO, and PsbA from *Arabidopsis thaliana*. All antibodies are rabbit polyclonal antibodies against recombinant proteins expressed from *E.coli*.

Immunogold labeling EM

Collodion/carbon coated grids (400 meshes) were glow discharged and floated on drops of PBS in buffer C for 1–2 min. PBS was fixed by floating the grids on drops of 4% paraformaldehyde for 10–15 min. After washing with 200 µl buffer C, PBS bounded to the grids was incubated with the primary antibody (antibodies against D1 subunit of PS II in *Arabidopsis thaliana*, which also recognize cyanobacteria D1 [48–50]) in buffer C for 25–30 min at the room temperature. The grids were then washed by six drops of 0.01 M PBS (pH 7.4) and incubated with 10 nm immunogold labeled second antibody (anti-rabbit IgG) for 25–30 min at the room temperature. The grids were washed with 0.01 M PBS (pH 7.4) and negative stained with 2% uranyl acetate.

Data deposition

Electron density maps have been submitted to the Electron Microscopy Data Bank (<http://www.emdatabank.org/>) under accession numbers 2821 and 2822.

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(**Supplementary information** is linked to the online version of the paper on the *Cell Research* website.)