Structure of a cyanobacterial photosystem I surrounded by octadecameric IsiA antenna proteins

Fusamichi Akita, Ryo Nagao, Koji Kato, Yoshiki Nakajima, Makio Yokono, Yoshifumi Ueno, Takehiro Suzuki, Naoshi Dohmae, Jian-Ren Shen, Seiji Akimoto, & Naoyuki Miyazaki

Iron-stress induced protein A (IsiA) is a chlorophyll-binding membrane-spanning protein in photosynthetic prokaryote cyanobacteria, and is associated with photosystem I (PSI) trimer cores, but its structural and functional significance in light harvesting remains unclear. Here we report a 2.7-Å resolution cryo-electron microscopic structure of a supercomplex between PSI core trimer and IsiA from a thermophilic cyanobacterium Thermosynechococcus vulcanus. The structure showed that 18 IsiA subunits form a closed ring surrounding a PSI trimer core. Detailed arrangement of pigments within the supercomplex, as well as molecular interactions between PSI and IsiA and among IsiAs, were resolved. Time-resolved fluorescence spectra of the PSI–IsiA supercomplex showed clear excitation-energy transfer from IsiA to PSI, strongly indicating that IsiA functions as an energy donor, but not an energy quencher, in the supercomplex. These structural and spectroscopic findings provide important insights into the excitation-energy-transfer and subunit assembly mechanisms in the PSI–IsiA supercomplex.
light-harvesting complexes (LHCs) are a family of pigment–proteins in photosynthetic organisms that play fundamental roles in harvesting solar energy and transferring it to the two photosystems, photosystem I (PSI) and II (PSII), where charge separation and electron transfer reactions are initiated. Among the photosynthetic organisms, various LHCs have been developed to capture the solar energy under different light environments. In most plants and algae, LHCs are membrane-embedded protein complexes associated with the two photosystems, and the structures of various LHC-photosystem supercomplexes have been determined from a number of different species: PSI–LHCI from pea2-3, PSI–LHCl from green algae6-8, C2S2-type PSI–LHClII from spinach9, C2S2-type PSI–LHClII from pea10, and C2S2M4N2 (C2S2M4L4) type PSI–LHClII from a green alga11.2. By contrast, cyanobacteria have evolved a completely different antenna system termed as phycobilisomes, which is a huge water-soluble pigment-protein complex attached at the stromal surface of the photosystems1. However, some cyanobacteria possess membrane-embedded light-harvesting protein complexes, prochlorophyte chlorophyll (Chl) a/b protein (Pcb) and iron-stress-inducible A protein (IsiA)5-14. IsiA has six transmembrane helices with a structural similarity to the CP43 subunit in PSI11,22, and is expressed under various stress conditions, especially under an iron starvation condition19,20. The IsiA protein is specifically associated with PSI, but not with PSII, through formations of ring structures around the PSI core. The IsiA ring can be either a single or double layer depending on the growth conditions and the species of cyanobacteria, although IsiA itself can also assemble into a single or double ring even in the absence of the PSI cores26,27.

IsiA has been reported to play a role in donating energy to the PSI core in the PSI–IsiA supercomplex28-31, whereas free IsiA is likely involved in energy quenching once IsiA is detached from PSI31–34. However, these spectroscopic results cannot exclude the possibility that energy quenching by IsiA may also occur in the PSI–IsiA supercomplex in a very early time region such as femtoseconds under physiological-temperature conditions. Recently, the overall architecture of PSI–IsiA has been determined at 3.5 Å resolution from a mesophilic cyanobacterium Synechocystis sp. PCC 6803 (hereafter designated as S. sp. PCC 6803, and PSI–IsiA from S. sp. PCC 6803 is designated as S-PSI–IsiA) by cryo-electron microscopy (cryo-EM)35. The structure showed that 18 IsiAs surround a PSI core trimer in a single ring organization, and implicated that IsiA may function in either energy harvesting or quenching35.

To examine the structure and function of IsiA in a greater detail, we solved the structure of a PSI–IsiA supercomplex from a thermophilic cyanobacterium Thermosynechococcus vulcanus (T. vulcanus) by single-particle cryo-EM analysis at a much improved resolution of 2.7 Å, which revealed the excitation energy transfer (EET) pathway and detailed protein–protein and pigment–pigment interactions between PSI and IsiAs. We also performed time-resolved fluorescence (TRF) analysis of the PSI–IsiA supercomplex to examine its EET dynamics. Together with the spectroscopic observations, the PSI–IsiA structure from T. vulcanus (T-PSI–IsiA) provides important functional insights into how IsiA contributes to the EET events in the PSI–IsiA supercomplex.

Results

Structural determination and overall structure of PSI–IsiA. We purified the PSI–IsiA supercomplex from T. vulcanus (Supplementary Fig. 1), cultured under an iron-starved condition as reported previously27. The purified PSI–IsiA retains all known cyanobacterial PSI subunits and the IsiA protein (Supplementary Fig. 1c). With this purified PSI–IsiA, we performed cryo-EM single-particle analysis (Supplementary Fig. 2). Two-dimensional classification of the cryo-EM particles revealed that octadecameric IsiA proteins surround the trimeric PSI core entirely in a single ring (Supplementary Fig. 2b). The final cryo-EM map was reconstructed from 303,983 particles taken from 20,799 micrographs at 2.74 Å resolution (Supplementary Fig. 2c and Supplementary Table 1), which was estimated by the gold-standard FSC with a 0.143 cutoff (Supplementary Fig. 3a and Supplementary Table 1). The local resolution map was shown in Supplementary Fig. 3c. The peripheral regions of PSI–IsiA have relatively low resolution, resulting in rather high numbers of clashscore and poor rotamers in the final refined structure (Supplementary Table 1). However, the Chl and Bcr molecules, which were related to energy pathways, have well-defined densities, and therefore were completely assigned. The overall structure of the PSI–IsiA supercomplex has a size of 294 Å in diameter and 110 Å in thickness, and is composed of a trimeric PSI core surrounded by 18 IsiAs with a C3 symmetry (Fig. 1), namely one asymmetric unit contains a PSI monomer and six IsiAs. The six IsiAs in an asymmetric unit are named IsiA1–6 clockwise according to their positions relative to the PSI monomer (Fig. 1a, b).

Based on the 2.74 Å resolution cryo-EM map, an atomic model was built and refined (Fig. 1, Supplementary Fig. 4). In the final structure, the PSI–IsiA supercomplex contains 12 PSI core subunits (PsaA, PsaB, PsaC, PsaD, PsaE, PsaF, PsaI, PsaJ, PsaK, PsaL, PsaM, and PsaX), 18 IsiAs, 585 Chls, 138 β-carotenes (Bcr), 6 phylloquinones (Pqn), 9 phosphatidyl-glycerol (LHG), 3 diesteraryl-monogalactosyl-diglyceride (LMG), and 12 iron-sulfur clusters (SF4) (Supplementary Table 2). IsiAs contain a total of 270 Chls which are more abundant in the stromal side than those in the luminal side, namely, 198 Chls are found in the stromal side and 72 Chls in the luminal side (Fig. 1c and Supplementary Table 2). The Chls in the stromal side are distributed evenly and form a global network over all IsiAs, whereas Chls in the luminal side are distributed radially and form distinct clusters within each IsiA monomer (Fig. 1c). This illustrates that Chls in IsiA have less interactions between adjacent IsiA subunits at the luminal side. IsiAs also contain 72 Bcrs, which surround PSI core and are distributed into two layers, namely, the inner layer and the outer layer (Fig. 1d).

The overall structure and pigment organization of T-PSI–IsiA are similar to those of S-PSI–IsiA35. However, compared with the S-PSI–IsiA structure, the T-PSI–IsiA structure has an additional PsaX subunit in the PSI core and there are some differences in the positions of several Chls associated with the PsaX subunit or located near the PsaX position between T-PSI–IsiA and S-PSI–IsiA (Supplementary Fig. 5a, b). Chl101 is coordinated by N23 of PsaX in the T-PSI–IsiA, whereas its equivalent, Chl302, is coordinated by D56 of PsaF in the S-PSI–IsiA (Supplementary Fig. 5b), and the position of Chl101 is shifted by 8.2 Å toward the stromal side relative to Chl302 (in terms of the Mg–Mg distance). The loop from residue 305 to 316 in the PsaB subunit interacts with the N-terminal loop of IsiA5 in the T-PSI–IsiA; however, the loop from residue 306 to 320 in PsaB is directed toward inside of the PSI core and does not interact with any IsiAs in the S-PSI–IsiA. Chl1240 of PsaB in the S-PSI–IsiA overlaps with PsaX in the T-PSI–IsiA, and thus has a shifted position designated as Chl834 in the T-PSI–IsiA. While Chl1240 forms a triplet Chl cluster with Chl1218 and Chl1219 in the stromal side in S-PSI–IsiA (Supplementary Fig. 5c), Chl841 in the T-PSI–IsiA forms a triplet Chl cluster with Chl834 and Chl840 in the luminal side. These differences may cause differences in the energy transfer behavior between the T-PSI–IsiA and S-PSI–IsiA.
Structures of the PSI core trimer and IsiA. Structure of the PSI core trimer in the T. PSI–IsiA was similar to its crystal structure from Thermosynechococcus elongatus (T. elongatus) solved at 2.5 Å resolution (PDB: 1JB0) with an RMSD value of 0.60 (Supplementary Fig. 5d); this is in agreement with the fact that the amino acid sequences of PSI subunits from T. vulcanus are 100% identical to those from T. elongatus. Except for PsaK, the 2.74 Å cryo-EM map has enough quality to allow assignment of the amino acid side chains (Supplementary Fig. 4). The side chains of PsaK could not be assigned unambiguously due to the limited local resolutions, and the loop regions of residues number 1–18, 39–54, and 77–83, of PsaK were disordered, but almost all chlorophylls can be unambiguously modeled in the structure. Only one Chl, Chl1601, which is coordinated by R24 of PsaM, was missing in the PSI core of T. vulcanus. A triplet Chl cluster (Chl1231, Chl1232, and Chl1233) and a dimer Chl cluster (Chl1218 and Chl1219) in the lumenal and stromal sides in the PSI core of T. elongatus, respectively.

Similarly, a triplet Chl cluster (Chl834, Chl840, and Chl841) and a dimeric Chl cluster (Chl821 and Chl822) in the PSI core of T. vulcanus were found. In contrast, the triplet Chl cluster in the photosystem II (PSII) core became a dimeric Chl cluster (Chl1218 and Chl1219), and the dimer Chl cluster in the stromal side became a triplet cluster (Chl1218, Chl1219, and Chl1240), in the PSI core of Synechocystis sp. PCC 6803 (Supplementary Fig. 5c). These results suggest that the energy transfer and dissipation system may be different between the thermophilic species Thermosynechococcus and mesophilic species Synechocystis.

The structure of the IsiA monomer (length: 358 residues) was built from residues number 20–350, which is composed of six long transmembrane helices (I–VI), five short helices (N, I', I'', II', and IV') and four β-strands (β1, β2, β3, and β4) arranged in a short anti-parallel β-sheet (Fig. 2a). Five loops connecting the transmembrane helices are designated as A to E loops from the N- to C-termini (Fig. 2a). Each IsiA monomer contains 17 Chls and 4 Bcrs (Fig. 2b and Supplementary Table 2). The central magnesium atoms of Chls are mainly coordinated by His residues, although some Chls are coordinated by Gln or Asn residues or by backbone carbonyls (Supplementary Table 3). Furthermore, two Chls, Chl404 and Chl407, have no direct
coordinating amino acid ligands, and thus they may be coordinated by water molecules and may have relatively higher excitation-energy levels than other Chls.

The structure of IsiA resembles that of PSII CP43 (PDB: 3WU2) as expected by their sequence similarities (Supplementary Fig. 6). However, PSII CP43 has more extended N- and C-terminal residues than IsiA located in the stromal side (1–11 residues in the N-terminal region and 453–473 residues in the C-terminal region) and an additional domain in the lumenal side (305–422 residues). The short additional stromal part of CP43 interacts with neighboring subunits, whereas the large lumenal part maintains the stability of the Mn₄CaO₅ cluster and protects it.

**Fig. 2 Structure of an IsiA monomer and its pigment organization.**

- **a** Ribbon diagram of an IsiA monomer. Helices and sheets are classified by Greek number.
- **b** Arrangement of the pigments (Chls and Bcrs) within the IsiA monomer. Left-side: a side-view represented by stick and light-colored cylinder model. Helices and sheets are classified by Greek number. Chlorophylls are colored by aquamarine (Chl401), gray (Chl402), yellow (Chl403), red (Chl404), coral (Chl405), forest green (Chl406), gold (Chl407), magenta (Chl408), pink (Chl409), brown (Chl410), green (Chl411), blue (Chl412), cyan (Chl413), orange (Chl414), purple (Chl415), deep pink (Chl416), and black (Chl417), respectively. Middle: Arrangement of the Bcrs. Bcrs are colored by gray. Right-side: top-view from the stromal side.
- **c** Interactions among Chls and Bcrs.
from attack by outside solute molecules. In addition, the residues 
coordinating the Chls are well conserved between IsiA and the 
PSII CP43 proteins (Supplementary Figs. 6 and 7). However, we 
found four additional Chls, Chl414, Chl415, Chl416, and Chl417 
in the IsiA protein, coordinated by H159, Q331, Q206, and 
carbonyl of I297, respectively (Supplementary Fig. 7 and 
Supplementary Table 3). These IsiA-unique Chls are conserved 
between T_PSI–IsiA and S_PSI–IsiA structures (Supplementary 
Fig. 7 and Supplementary Table 3). Chl414 and Chl416 are 
located in the peripheral part of the PSI–IsiA complex, and 
Chl415 and Chl417 are located at the interface between IsiA and 
the PSI core and hence may be involved in the EET pathways 
between the IsiAs and PSI core as described below. Some Chls in 
the PSI core and hence may be involved in the EET pathways 
Chl415 and Chl417 are located at the interface between IsiA and 
interacts with Chl405 in the stromal side at a distance of 3.5 Å, 
and may also mediate EET from IsiA to the PSI core (Fig. 2c).

Interactions and possible EET pathways among different IsiAs. 
Each IsiA interacts with their neighboring subunits in a similar 
manner (Fig. 3). The surface contact areas (buried surface areas) 
between adjacent IsiA pairs (IsiA1–2, IsiA2–3, IsiA3–4, IsiA4–5, 
IsiA5–6, and IsiA6–1) are 1395, 1421, 1353, 1412, 1327, and 
1152 Å², respectively (Supplementary Table 4), and the molecular 
interactions are formed mainly by hydrophobic interactions via 
protein–protein, protein–pigment, and pigment–pigment inter-
actions. Hydrophobic patches are observed both at the stromal 
and luminal sides. For example, a hydrophobic residue F127 of 
one IsiA interacts with hydrophobic L338 and F343 residues from 
the adjacent subunit at the stromal side, and hydrophobic L62 
and F105 residues interact with hydrophobic V293 and L295 
residues from the neighboring subunit at the luminal side (Fig. 3c).

In addition, five Chls (Chl405, Chl413, Chl415, Chl416, and 
Chl417) and two Bcrs (Bcr422 and Bcr424) are extensively 
involved in the inter-subunit interactions between the IsiAs 
(Fig. 3c, d). Chl405 coordinated by the H333 residue interacts 
with four hydrophobic residues (I113, V117, A120, and F124) 
from the neighboring IsiA and also forms a hydrophilic inter-
action between its carboxy group and R128 of the neighboring 
IsiA (Fig. 3c, right panel). Chl405 also interacts with Chl413, 
Bcr422 and Bcr424 from the adjacent subunit (Fig. 3d). Chl417 
coordinated by a backbone carbonyl of I297 interacts with 
hydrophobic residue F60 from the neighboring IsiA (Fig. 3c, 
right panel). Bcr422 interacts with Chl416 (Fig. 3d), and Bcr424 
interacts with the hydrophobic residue L330 and Chls (Chl415 
and Chl417) (Fig. 3c and d), from the adjacent subunit, 
respectively.

To find EET pathways between IsiAs, we calculated the Förster 
energy transfer rates \(^{13,12,35,38}\) between IsiA (Supplementary 
Table 4). Common EET pathways, which are defined by a half-life 
time of <20 ps, are found between all neighboring IsiA pairs 
(Fig. 3e). These include three inter-IsiA EET pathways, which 
were designated as interior, exterior, and intermediate pathways 
according to their positions on the IsiA ring relative to the PSI 
core. All of these pathways are found at the stromal side. In the 
interior pathway, which is closest to the PSI core, two possible 
inter-IsiA EET pathways are identified between Chl404 and 
Chl415, and between Chl410 and Chl417. In the exterior 
pathway, there are also two possible inter-IsiA EET pathways 
found between Chl412 and Chl405 and between Chl413 and 
Chl405. In the intermediate pathway, one possible inter-IsiA EET 
pathway is found between Chl411 and Chl405. The fastest energy 
transfer rates between the neighboring IsiAs are always observed 
between Chl411 and Chl405, and between Chl413 and Chl405, 
respectively. Thus, these pathways are considered as the main 
EET pathways between the neighboring IsiAs.

Interactions between IsiA and PSI. Interactions of each IsiA 
with the PSI core are rather diverse (Fig. 4a) which are sum-
morized in Supplementary Table 5. This can be seen in the 
surface contact areas between the PSI core and the individual IsiA 
subunits from IsiA1 to IsiA6, which are 267.4, 103.8, 317.0, 371.3, 
229.2, and 4.1 Å², respectively (Supplementary Table 5). Rela-
tively strong interactions with the PSI core are found for IsiA1, 
IsiA3, isiA4, and IsiA5 (Figs. 4b and d–f and Supplementary 
Table 5). Especially, the C-terminal region of the IsiA4 is 
extended and interacts with the PsAF subunit in the PSI core by 
hydrophobic interactions and thus exhibits the largest surface 
contact area (Fig. 4e). In contrast, IsiA2 has limited interactions 
with the PSI core (Fig. 4c), and IsiA6 has almost no contact with 
the PSI core, since IsiA6 is obviously located far from the PSI 
core with a distance over 8 Å. These results indicate that IsiA4 might 
be a key subunit for the assembly of the octadecameric IsiA ring 
around the PSI core with specific interactions at its C-terminal 
region. This is apparently different from the S_PSI–IsiA struc-
ture \(^{35}\), where the C-terminus of IsiA contains a flexible amphi-
pathic helix and adopts slightly different orientations at different 
IsiA positions. The C-terminus of IsiA1–5 in the S_PSI–IsiA is 
directed to the neighboring IsiA subunit and interacts with the N-
terminus of the adjacent monomer, whereas the C-terminus of 
IsiA6 appears to be disordered. Indeed, the amino acid sequences 
in the C-terminal region appear less conserved between 
S_PSI–IsiA and T_PSI–IsiA (Supplementary Fig. 9). Thus, the 
molecular recognition between PSI and IsiA as well as between 
different IsiAs may be different between S_PSI–IsiA and 
T_PSI–IsiA.

Possible excitation-energy transfer pathways between IsiA and 
PSI. The possible EET pathways between IsiA and the PSI core 
were investigated based on the Förster energy transfer rates 
among adjacent Chls (Fig. 4h–j and Supplementary Table 5). The 
following possible EET pathways were identified: 1As, 1Al, 2As, 
2Al, 3Al, 3Jl, 4Jl, 5Bl, 5Xl, and 6Bl, which were named according 
to that used for PSI–LHCII \(^{27,40}\). As examples of these names, 1As 
and 1Al represent the pathways identified from IsiA1 to PsA1 
at the stromal and luminal sides, respectively. The results showed 
that all IsiAs have substantial EET pathways to the PSI core, 
among which, the EET pathway from IsiA6 to PSI core is inde-
pendent of the molecular interactions between this IsiA and PSI 
core. There are five Chls in IsiAs that are involved in these EET 
pathways; they are Chl404, Chl408, Chl411, Chl415, and Chl417. 
In particular, Chl404 and Chl417 have the fastest energy 
migration rate between the IsiAs and the PSI core (Supplemen-
tary Table 5); therefore these two Chls may play the most 
important roles for the EET from IsiAs to the PSI core. Chl404 in 
IsiA5 and Chl841 in PSI had the nearest distance in all EET 
pathway (Supplementary Table 5). However, Chl841 forms
tripled Chl in PSI, and the pathway might rather work as energy quencher. Among all IsiAs, IsiA1 has three pathways with relatively faster energy migration rates between the IsiAs and the PSI core than those in other IsiAs, suggesting that IsiA1 plays an important role for EET from the IsiA ring to the PSI core.

**Fig. 3 Interactions and possible EET pathways within the IsiA ring.**

- **a** Overview of interactions between adjacent IsiAs. Squared areas by green dashed and red solid lines are enlarged in panels (c) and (e), respectively.
- **b** Superimposition of the structures of the six IsiAs. The squared region by a red dashed line indicates the C-terminal region of IsiA and is enlarged in the left bottom side. The C-terminal region of IsiA4 (cyan) are more ordered than that of other IsiAs.
- **c** Interactions between IsiA1 and IsiA2. The middle panel is an overview, and the left and right panels show the protein–protein and protein–chlorophylls interactions, respectively.
- **d** Pigment–pigment interactions between IsiA1 and IsiA2. Right panel is a top-view from the stromal side.
- **e** Possible EET pathways between IsiA1 and IsiA2.

**Time-resolved fluorescence analyses of PSI–IsiA.** To examine the function of IsiA in *T. vulcanus*, we compared TRF spectra between the PSI–IsiA supercomplex (red lines) and the PSI core trimer (black lines) (Fig. 5a). The spectra of PSI–IsiA and PSI cores exhibited a fluorescence peak at around 730 nm in the time range from 0–4.9 ps to 2.0–2.4 ns. Thus, the peak at around 730 nm was attributed to the fluorescence from Chls with lower energy in the PSI cores than the energy level of the special pair Chls P700. The existence of the low-energy Chls are confirmed by an absorption peak located at 709 nm (Supplementary Fig. 1). By contrast, a peak around 683 nm appeared only in the PSI–IsiA spectra particularly at the picosecond time range of 0–4.9 ps after the excitation. Based on its absence in the spectra of the PSI trimer and the blue-shifted Qy absorption spectrum of the PSI–IsiA (Supplementary Fig. 1), the 683-nm peak is attributed to the fluorescence from the IsiAs. The relative intensity of the 683-nm peak decreased gradually, and then almost disappeared in the spectrum in the time range of 180–210 ps. These results indicate that IsiA transfers the excitation energy to the PSI core or quenches the energy within ~180 ps.
To examine the detailed excitation-energy dynamics in the PSI–IsiA, we constructed fluorescence decay-associated (FDA) spectra (Fig. 5b). In these spectra, a pair of positive and negative peaks reflects EET from Chls with the positive peak to Chls with the negative peak. The FDS spectrum of PSI–IsiA showed a set of 685-nm positive and 730-nm negative peaks in the time range of 35 ps, indicating EET from IsiA to the PSI core. By contrast, the 50-ps FDA spectrum of the PSI core trimer showed a small positive peak at around 693 nm, indicating that the energy donor in the PSI trimer is different from that in the PSI–IsiA. The FDA spectra of the two samples in the time range of 85–470 ps exhibited only a decay component at around 730 nm, indicating either energy trapping at the RC Chls41–44 or quenching by Chl–Car interactions45–48. The forth FDA spectrum of PSI–IsiA (3.5 ns) exhibited a positive peak at 685 nm with a very tiny amplitude compared with the peak amplitudes in the first, second, and third FDA spectra (note that the scale of the Y-axis of the forth FDA is enlarged by 1000 times). This indicates that only slight amounts of IsiA appear to be dissociated from the supercomplex. Thus, the FDA analysis strongly indicated functional EET from IsiA to the PSI cores.

To verify whether IsiA is related to energy quenching in the time range of femtoseconds, we measured femtosecond fluorescence decay curves at 685 nm with an upconversion system under room-temperature49 (Fig. 5c). The fluorescence decay of the PSI–IsiA supercomplex (red line) was apparently slower than that of the PSI trimer (black line), indicating that excitation-energy quenching is not facilitated by the association of IsiA with the PSI
Thus, our TRF analyses provides strong evidence that IsiA serves as an energy donor to the PSI core.

**Discussion**

The structure and spectroscopic results presented here provide important information regarding the possible EET pathways in the PSI–IsiA supercomplex. Our structure revealed that a trimeric PSI core is encircled by 18 copies of IsiA. All IsiAs are connected with each other in similar manners and form circular EET pathways within the ring surrounding the PSI core trimer. Therefore, the inter-IsiA network may enable the PSI cores in the trimer to share the excitation energies captured by any of the antenna IsiAs efficiently (Fig. 6a). Plausible time constants for the energy migration among the 18 IsiAs should be within 35 ps, because the FDA spectrum of PSI–IsiA showed a time of 35 ps for EET from IsiA to PSI (Fig. 5b). On the other hand, the EET pathways from IsiAs to the PSI core are apparently different depending on the IsiA positions (IsiA1 to IsiA6). Although all IsiAs have possible EET pathways to PSI, the energy migration rates between the Chls in IsiA1 and the PSI core is relatively faster than those between the other IsiAs and the PSI core (Supplementary Table 5). Thus, IsiA1 has the main EET pathway to PSI between the IsiA ring and the PSI core (Fig. 6a). In addition, two IsiA-unique Chls, Chl415 and Chl417, are extensively involved in the EET pathways between IsiAs and PSI, and both of them are also involved in the inter-IsiA EET pathways. These observations imply that IsiA, probably derived from CP43, acquired these Chls for the efficient EET to PSI. By contrast, other IsiA-unique Chls, Chl414 and Chl416, are located in the peripheral part of the PSI–IsiA complex. As the PSI core is sometimes surrounded by two rings of IsiAs, these Chl414 and Chl416 may face to the possible second, exterior IsiA ring and contribute to EET between the two IsiA rings.

**Fig. 5 TRF analyses of the PSI–IsiA supercomplex.** a 77-K TRF spectra excited at 445 nm. The spectra of the PSI–IsiA and PSI trimer cores were normalized by the maximum intensity of each spectrum. The spectra of the PSI–IsiA and PSI trimer cores are depicted in red and black lines, respectively. b 77-K FDA spectra. The spectra of the PSI–IsiA and PSI trimer cores are depicted in red and black lines, respectively. c Normalized fluorescence decay curves at 296 K monitored at 685 nm. The curves of the PSI–IsiA and PSI trimer cores are depicted in red and black lines, respectively.
The PSI–IsiA structure obtained here also brings important implications on the assembly mechanism of the PSI–IsiA supercomplex. The 18 IsiAs can be divided into three units, each consisting of six IsiAs, and each of these six IsiAs binds to the PSI core in significantly different manners. This is reflected by the remarkable differences in the contact surface areas between each IsiA and the PSI core. In particular, IsiA4 has the largest contact surface area with the PSI core, and its C-terminal region is in close interaction with the PsaF subunit of the PSI core. Compared with other IsiAs, the C-terminal region of the IsiA4 subunit is well ordered and extended to the PSI core. The strongest interactions between IsiA4 and the PSI core suggest that the IsiA ring formation may be initiated by adsorption of an IsiA to the IsiA4 position on the PSI core, followed by attachment of other IsiAs. On the other hand, IsiAs have an intrinsic ability to form single and double ring assemblies in the absence of the PSI core by their lateral, inter-subunit interactions. This is consistent with the fact that all IsiAs in the PSI–IsiA supercomplex are connected with each other in similar manners. These lines of evidence suggest that the mechanism of IsiA-layer formation is like a traditional epitaxial nucleated-growth type involving an initial nucleation step, in which IsiAs recognize the IsiA4 position, followed by a series of growth steps mainly by lateral inter-IsiA interactions (Fig. 6b). In relation to this, it is interesting to note that the structures of the IsiA C-terminal regions are remarkably different between T_PSI–IsiA and S_PSI–IsiA (Supplementary Fig. 7c). Since the C-terminal region plays an important role in the assembly of the supercomplex, this may suggest a different manner of assembly between T_PSI–IsiA and S_PSI–IsiA.

In summary, this study provides a basis for the possible EET mechanisms in the PSI–IsiA supercomplex of T. vulcanus based on its 2.7-Å cryo-EM structure and spectroscopic analyses. The high-resolution structure obtained revealed the tight association of Chl molecules among neighboring IsiAs within the single ring of the PSI–IsiA supercomplex, enabling a fast energy migration within 35 ps within the IsiA ring, followed by trapping to P700 via Chls positioned at the interface between IsiA and PsaA/PsaB/PsaF. Our femtosecond TRF decay curves do not support the possibility that IsiA serves as an energy quencher, thereby emphasizing the role of IsiA in light-harvesting and energy donation to PSI under iron-stress conditions.

Some structural differences were found between the PSI–IsiA from T. vulcanus and S. sp. PCC 6803, and these structural differences may bring some functional differences of the IsiAs between the two species. This may be related with the growth environments that each species of the cyanobacterium experiences.

**Methods**

**Purification of PSI–IsiA from T. vulcanus.** Cells of T. vulcanus were cultured in two litters of an iron-free medium at 50 °C for a week to reach OD720 = 1.0. The cells were then collected by centrifugation and resuspended with five litters of the iron-free medium, and continued to grow for 2 weeks at 50 °C. FeCl3 was added to a final concentration of 2.5 mM and the culture was further incubated for a week at 30 °C. The cells were pelleted by centrifugation at 13,700 × g for 20 min at 4 °C, followed by solubilization with a buffer containing 10 mM MgCl2, 20 mM HEPES–NaOH (pH 7.0), 1.0% (w/v) n-dodecyl–D-maltoside (β-DDM) at 4 °C for an hour. Insoluble membrane was removed by centrifugation at 20,000 × g for 10 min at 4 °C. The supernatant was loaded onto a linear trehalose gradient of 30% (w/v) in a buffer containing 10 mM MgCl2, 20 mM HEPES–NaOH (pH 7.0), 0.04% (w/v) β-DDM, and centrifuged at 180,000 × g for 2 h at 4 °C. After centrifugation, the band containing PSI–IsiA was collected, and then pelleted with PEG 1500 (at a final concentration of 13% w/v) for 20 h at 4 °C. The PSI–IsiA containing band was collected and applied to cryo-EM grids for data set A. For the cryo-EM data set B, trehalose in the sample solution was removed by
pelleting the PSI–IsiA containing band with PEG 1500 (at a final concentration of 13% w/v), and resuspended to a buffer of 50 mM HEPES–NaOH (pH 7.0), 0.04% β-DDM before being applied to the cryo-EM grids.

**Cryo-EM data collection.** Two image data sets (data set A and B) were acquired. The data set A contains 4920 images and was acquired from holey carbon grids covered with amorphous carbon film, and the data set B contains 15,897 images and acquired from holey carbon grids without amorphous carbon film. For cryo-EM experiments of data set A, a 4-ml aliquot of the PSI–IsiA sample (30 μg of Chl mL⁻¹) in a buffer containing 50 mM HEPES–NaOH (pH 7.0), 10 mM MgCl₂, 0.04% β-DDM, 30% trehalose was applied to glow-discharged holey carbon grids (Quantifoil R1.2/1.3, Mo 200 mesh) covered with 5–10 nm amorphous carbon film. The grids were incubated for 30 s in the chamber of FEI Vitrobot Mark IV at 4°C and 100% humidity, and then washed once with 3 μl of a wash buffer containing 50 mM HEPES–NaOH (pH 7.0) and 0.04% β-DDM Remaining trehalose. This wash process removes trehalose dramatically and enhanced the image contrast of particles as described previously. The washed grids were immediately blotted with filter papers for 3 s and plunged into liquid ethane cooled by liquid nitrogen and then transferred into a cryo-electron microscope (Titan Krios, Thermo Fischer Scientific) equipped with a field emission gun, a Cs corrector (CEOS GmbH), and a direct electron detection camera (Falcon 3EC, Thermo Fischer Scientific) and operated at 300 kV. Image movies were recorded using the Falcon 3EC in a linear mode with a nominal magnification of x59,000, which results in a final pixel size of 1.113 Å. Each exposure of 2.5 s was dose-fractonated into 32 movie frames, leading to a total of 80 electron doses per Å². The diameter of the defocus range was set to −2.0 to −4.0 μm. For cryo-EM experiments of data set B, a 2.5-μl aliquot of the PSI–IsiA sample (105 μg of Chl mL⁻¹, a higher concentration than that used for the sample for the data set A) in a buffer containing 20 mM HEPES–NaOH (pH 7.0), 10 mM MgCl₂, and 0.04% β-DDM was applied to glow-discharged holey carbon grids (Quantifoil R1.2/1.3, Mo 300 mesh). The grids were incubated for 30 s in the chamber of FEI Vitrobot Mark IV at 4°C and 100% humidity. The grids were blotted with a filter paper manually and another 2.5-μl aliquot of the PSI–IsiA (103 μg of Chl mL⁻¹) was applied to the grids in order to increase the number of the particles in the holes. Then, the grids were immediately plunged into liquid ethane cooled by liquid nitrogen. Movies from the data set B were combined and used for the structure analysis in the same conditions as that for the data set A. All of the image data sets (total 20,799 micrographs) were finally combined and used for the structure analysis.

**Cryo-EM image processing.** Movie frames were aligned and summed using the MotionCor2 software to obtain a final dose weighted image. Estimation of the contrast transfer function (CTF) was performed using the CTFFIND4 program. All of the following processes were performed using RELION3.0. For structural analyses of PSI–IsiA, 1,391,531 and 3,109,982 particles were automatically picked from 4,902 and 15,897 micrographs in data sets A and B, respectively, and then transferred into a cryo-electron microscope (Titan Krios, Thermo Fischer Scientific) equipped with the field emission gun, a Cs corrector (CEOS GmbH), and a direct electron detection camera (Falcon 3EC, Thermo Fischer Scientific) and operated at 300 kV. Image movies were recorded using the Falcon 3EC in a linear mode with a nominal magnification of x59,000, which results in a final pixel size of 1.113 Å. Each exposure of 2.5 s was dose-fractonated into 32 movie frames, leading to a total of 80 electron doses per Å². The diameter of the defocus range was set to −2.0 to −4.0 μm. For cryo-EM experiments of data set A, a 2.5-μl aliquot of the PSI–IsiA sample (105 μg of Chl mL⁻¹, a higher concentration than that used for the sample for the data set A) in a buffer containing 20 mM HEPES–NaOH (pH 7.0), 10 mM MgCl₂, and 0.04% β-DDM was applied to glow-discharged holey carbon grids (Quantifoil R1.2/1.3, Mo 200 mesh). The grids were incubated for 30 s in the chamber of FEI Vitrobot Mark IV at 4°C and 100% humidity. The grids were blotted with a filter paper manually and another 2.5-μl aliquot of the PSI–IsiA (103 μg of Chl mL⁻¹) was applied to the grids in order to increase the number of the particles in the holes. Then, the grids were immediately plunged into liquid ethane cooled by liquid nitrogen. Movies from the data set B were combined and used for the structure analysis in the same conditions as that for the data set A. All of the image data sets (total 20,799 micrographs) were finally combined and used for the structure analysis.

**Statistics and reproducibility.** No statistical method was used to predetermined the sample size, and the experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment. The purification of PSI–IsiA was repeated over three times, which showed same results. The spectroscopic anaysis was performed at least two times, and similar results were obtained. The cryo-EM data was collected from four grids. Individual images with bad ice were excluded from the data set by visual inspection. Data collection, processing and refinement statistics were summarized in Supplementary Table 1.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability.** Atomic coordinates and Cryo-EM maps for the reported structure of PSI–IsiA have been deposited in the Protein Data Bank under an accession code 6K33 and in the Electron Microscopy Data Bank under an accession code EMD-9908. Source data of Fig. 5 is provided as Supplementary Data. Further data supporting this study are available from the corresponding author upon reasonable request.

Received: 27 September 2019; Accepted: 17 April 2020; Published online: 11 May 2020

**References.**

1. Nelson, N. & Yocum, C. F. Structure and function of photosystems I and II. *Annu. Rev. Plant Biol.* 57, 521–565 (2006).
2. Qin, X., Suga, M., Kuang, T. & Shen, J.-R. Photosynthesis. Structural basis for energy transfer pathways in the plant PSI-LHCI supercomplex. *Science* 348, 989–993 (2015).
3. Qin, X., Borovkova, A., Caspy, I. & Nelson, N. Structure of the plant photosystem I supercomplex at 2.6 Å resolution. *Nat. Plants* 3, 17014 (2017).
4. Pi, X. et al. Unique organization of photosystem I-light-harvesting supercomplex revealed by cryo-EM from a red alga. *Proc. Natl Acad. Sci. USA* 115, 4423–4428 (2018).
5. Antoschwil, M., Caspy, I., Hippler, M. & Nelson, N. Structure and function of photosystem I in *Chlorella variabilis* merosome. *Photosynth. Res.* 139, 499–508 (2019).
6. Su, X. et al. Antenna arrangement and energy transfer pathways of a green algal photosystem-LHI supercomplex. *Nat. Plants* 5, 273–281 (2019).
7. Suga, M. et al. Structure of the green algal photosystem I supercomplex with a decameric light-harvesting complex I. *Nat. Plants* 5, 626–636 (2019).
8. Qin, X. et al. Structure of a green algal PSI in complex with a large number of light-harvesting complex I subunits. *Nat. Plants* 5, 263–272 (2019).
9. Wei, X. et al. Structure of spinach photosystem II–LHCII supercomplex at 3.2 Å resolution. *Nature* 534, 69–74 (2016).
10. Su, X. et al. Structure and assembly mechanism of plant C₃S₅M₂-type PSI-LHCCI supercomplex. *Science* 357, 815–820 (2017).
11. Shen, L. et al. Structure of a C₃S₅M₂-type PSI-LHCCI supercomplex from the green alga *Chlamydomonas reinhardtii*. *Proc. Natl Acad. Sci. USA* 116, 21246–21255 (2019).
12. Sheng, X. et al. Structural insight into light harvesting for photosystem II in green algae. *Nat. Plants* 5, 1320–1330 (2019).
13. Zhang, J. et al. Structure of phycoerythrin from the red alga *Griphellopsis pacifica*. *Nature* 551, 57–63 (2017).
14. La Roche, J. et al. Independent evolution of the prochlorophyte and green plant chlorophyll a/b light-harvesting proteins. *Proc. Natl Acad. Sci. USA* 93, 15244–15248 (1996).
15. Bibby, T. S., Mary, I., Nield, J., Partensky, F. & Barber, J. Light-leaf-adapted Prochlorococcus species possess specific antennae for each photosystem. *Nature* **424**, 1053–1054 (2003).

16. Bibby, T. S., Nield, J., Chen, M., Larkum, A. W. & Barber, J. Structure of a photosystem II supercomplex isolated from *Prochloron didemni* retaining its chlorophyll a/b light-harvesting system. *Proc. Natl Acad. Sci. USA* **100**, 9050–9054 (2003).

17. Berka, E. J. A giant chlorophyll-protein complex induced by iron deficiency in cyanobacteria. *Nature* **412**, 745–748 (2001).

18. Pakrasi, H. B., Goldenberg, A. & Sherman, L. A. Membrane development in cyanobacterium membranes of cyanobacteria during iron starvation. *Plant Physiol.* **79**, 290–295 (1985).

19. Burnay, R. L., Trojan, T. & Sherman, L. A. The highly abundant chlorophyll-protein complex of iron-deficient Synechocystis sp. PC6804 (CP43) is encoded by the isiA gene. *Plant Physiol.* **103**, 893–902 (1993).

20. Falk, S., Samson, G., Bruce, D., Huner, N. P. & Laudenbach, D. E. Functional analysis of the iron-stress induced CP 43′ polypeptide of PS II in the cyanobacterium *Synechocystis* sp. PCC 7942. *Photosynth. Res.* **45**, 51–60 (1995).

21. Leonhardt, K. & Straus, N. A. An iron stress operon involved in photosynthetic electron transport in the marine cyanobacterium *Synechococcus* sp. PCC 7002. *J. Gen. Microbiol.* **138**, 1613–1612 (1992).

22. Laudenbach, D. E. & Straus, N. A. Characterization of a cyanobacterial iron stress-induced gene similar to pbc1. *J. Bacteriol.* **170**, 5018–5026 (1988).

23. Vinniemeier, J., Kunert, A. & Hagemann, M. Transcriptional analysis of the isiA operon in iron-stressed cells of the cyanobacterium *Synechocystis* sp. PCC 6803. *FEBS Microbiol. Lett.* **169**, 323–330 (1998).

24. Havaux, M. et al. The chlorophyll-binding protein IsiA is inducible by high light and protects the cyanobacterium *Synechococcus* PCC6803 from photooxidative stress. *FEBS Lett.* **579**, 2289–2293 (2005).

25. Guikema, J. A. & Sherman, L. A. Organization and function of chlorophyll in membranes of cyanobacteria during iron starvation. *Plant Physiol.* **73**, 250–256 (1983).

26. Yeremenko, N. et al. Supramolecular organization and dual function of the IsiA chlorophyll-binding protein in cyanobacteria. *Biochemistry* **43**, 10308–10313 (2004).

27. Chauhan, D. et al. A novel photosynthetic strategy for adaptation to low iron aquatic environments. *Biochemistry* **50**, 686–692 (2011).

28. Andrizhiyevskaya, E. G. et al. Spectroscopic properties of PSI–IsiA supercomplexes from the cyanobacterium *Synechocystis* PCC 7942. *Biochim. Biophys. Acta, Bioenerg.* **1556**, 265–272 (2002).

29. Andrizhiyevskaya, E. G., Frolov, D., van Grondelle, R. & Dekker, J. P. Energy transfer and trapping in the Photosystem I complex of *Synechocystis* PCC 7942 and in its supercomplex with IsiA. *Biochim. Biophys. Acta, Bioenerg.* **1656**, 104–113 (2004).

30. Melkozernov, A. N., Bibby, T. S., Lin, S., Barber, J. & Blankenship, R. E. Time-resolved absorption and emission show that the CP43 antenna ring of iron-stressed *Synechocystis* sp. PCC6803 is efficiently coupled to the Photosystem I reaction center. *Biochemistry* **42**, 3893–3903 (2003).

31. Chen, H. S., Liberton, M., Pakrasi, H. B. & Niedzwiedzki, D. M. Reevaluating the mechanism of excitation energy regulation in iron-starved cyanobacteria. *Biochim. Biophys. Acta, Bioenerg.* **1858**, 249–258 (2017).

32. Ihalaenen, J. A. et al. Aggregates of the chlorophyll-binding protein IsiA (CP43) dissipate energy in cyanobacteria. *Biochemistry* **44**, 10864–10873 (2005).

33. Berera, R., van Stokkum, I. H. M., Kennis, J. T. M., van Grondelle, R. & Dekker, J. P. The light-harvesting function of carotenoids in the cyanobacterium iron-stress-inducible IsiA complex. *Chem. Phys.* **373**, 65–70 (2010).

34. Berera, R. et al. A mechanism of energy dissipation in cyanobacteria. *Biophys. J.* **96**, 2261–2267 (2009).

35. Toporkit, H., Li, J., Williams, D., Chiu, P. L. & Mazor, Y. The structure of the stress-induced photosystem I-IsiA antenna supercomplex. *Nat. Struct. Mol. Biol.* **26**, 443–449 (2019).

36. Jordan, P. et al. Three-dimensional structure of cyanobacterial photosynthetic I-IHCL monomers: implications for the structural model of the major plant antenna. *Biophys. J.* **75**, 3064–3077 (1998).

37. Suga, M., Qin, X., Kuang, T. & Shen, J.-R. Structure and energy transfer pathways of the plant photosystem I-LHCl supercomplex. *Curr. Opin. Struct. Biol.* **39**, 46–53 (2016).

38. Wlodarcyzk, L. M., Denc, E., Croce, R. & Dekker, J. P. Excitation energy transfer in Chlamydomonas reinhardtii deficient in the PSI core or the PSI core under conditions mimicking state transitions. *Biochim. Biophys. Acta, Bioenerg.* **1857**, 625–633 (2016).

39. Gradinaru, C. C. et al. The flow of excitation energy in LHCl monomers: implications for the structural model of the major plant antenna. *Biophys. J.* **75**, 3064–3077 (1998).

40. Gobets, B. & van Grondelle, R. Energy flow and trapping in photosystem I. *Biochim. Biophys. Acta, Bioenerg.* **1507**, 80–99 (2001).

41. Zabban, A. V., Johnson, M. P. & Duffy, C. D. The photosynthetic molecular switch in the photosystem II antenna. *Biochim. Biophys. Acta, Bioenerg.* **1817**, 167–181 (2012).

42. Ahn, T. K. et al. Architecture of a charge-transfer state regulating light harvesting in a plant antenna protein. *Science* **320**, 794–797 (2008).

43. Ruban, A. V. et al. Identification of a mechanism of photoprotective energy dissipation in higher plants. *Nature* **450**, 575–578 (2007).

44. Barr, J. et al. The protein fusion CooT: a visualization system for exploratory research and analysis. *J. Comput. Chem.* **25**, 1605–1612 (2004).

45. Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot. *Acta Crystallogr. Sect. D., Biol. Crystallogr.* **66**, 486–501 (2010).

46. Kelley, L. A., Mezulis, S., Yates, C. M., Wass, M. N. & Sternberg, M. J. E. The Phyre2 web portal for protein modeling, prediction and analysis. *Nat. Protoc.* **10**, 845–858 (2015).

47. Adams, P. D. et al. PHENIX: a comprehensive Python-based system for macromolecular structure solution. *Acta Crystallogr. Sect. D., Biol. Crystallogr.* **66**, 213–221 (2010).

48. Chen, V. B. et al. MolProbity: all-atom structure validation for macromolecular crystallography. *Acta Crystallogr. Sect. D., Biol. Crystallogr.* **66**, 12–21 (2010).

49. Barad, B. A. et al. EMRinger: side-chain directed model and map validation for 3D cryo-electron microscopy. *Nat. Methods* **12**, 943–946 (2015).

50. Collaborative Computational Project, N. The CCP4 suite: programs for protein crystallography. *Acta Crystallogr. Sect. D., Biol. Crystallogr.* **50**, 2152–2153 (1994).

51. van Grondelle, R., Dekker, J. P., Gillbro, T. & Sundström, V. Energy transfer and trapping in photosynthesis. *Biochim. Biophys. Acta, Bioenerg.* **1187**, 1–65 (1994).

52. van Grondelle, R. Excitation energy transfer, trapping and annihilation in photosynthetic systems. *Biochim. Biophys. Acta, Bioenerg.* **1181**, 147–195 (1985).
Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s42003-020-0949-6.

Correspondence and requests for materials should be addressed to F.A., J.-R.S., S.A. or N.M.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020