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Plants lacking the main light-harvesting complex retain photosystem II macro-organization

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Photosystem II (PSII) is a key component of photosynthesis, the process of converting sunlight into the chemical energy of life. In plant cells, it forms a unique oligomeric macrostructure in membranes of the chloroplasts. Several light-harvesting antenna complexes are organized precisely in the PSII macrostructure—the major trimeric complexes (LHCII) that bind 70% of PSII chlorophyll and three minor monomeric complexes—which together form PSII supercomplexes. The antenna complexes are essential for collecting sunlight and regulating photosynthesis, but the relationship between these functions and their molecular architecture is unresolved. Here we report that antisense Arabidopsis plants lacking the proteins that form LHCII trimers have PSII supercomplexes with almost identical abundance and structure to those found in wild-type plants. The place of LHCII is taken by a normally minor and monomeric complex, CP26, which is synthesized in large amounts and organized into trimers. Trimerization is clearly not a specific attribute of LHCII. Our results highlight the importance of the PSII macrostructure: in the absence of one of its main components, another protein is recruited to allow it to assemble and function.

The central unit of PSII is the core complex, in which light is used to oxidize water to molecular oxygen, to reduce plastoquinone and to generate a transmembrane proton gradient. In plants, each core dimer binds two copies of the minor, monomeric light-harvesting proteins CP29, CP26 and CP24 (the Lhcb1, Lhcb5 and Lhcb6 gene products), respectively, and two–four LHCII trimers (made up of the Lhcb1, Lhcb2 and Lhcb3 gene products) to form a PSII supercomplex. Additional LHCII trimers have been located in membrane domains deficient in PSII supercomplexes, although they still can be involved in efficient light-harvesting. These trimers consist of the Lhcb1 and Lhcb2 gene products, and can partition, depending on the phosphorylation state of LHCII, between the grana and stromal thylakoids to transfer energy to PSI and PSII, respectively.

This model for PSII macrostructure suggests that each light-harvesting protein has a unique position and role, but little is known about the structure–function relationships involved. In addition, the possibility cannot be excluded that there is some redundancy.
among these similar light-harvesting proteins\textsuperscript{17}. We have taken a genetic approach to investigating these complexes. Expression of a full-length \textit{Lhcb2} antisense construct in \textit{Arabidopsis thaliana} abolishes the expression of both \textit{Lhcb2} and \textit{Lhcb1}, owing to stretches of identical nucleotide sequences in these two genes\textsuperscript{10}. The plants (called as\textit{Lhcb2}) are almost completely devoid of the two main proteins of trimeric \textit{LHCII}, \textit{Lhcb1} and \textit{Lhcb2} (ref. 10). Western blot analysis of PSII membranes prepared from wild-type and as\textit{Lhcb2} plants showed that \textit{Lhcb1} is completely absent and the amount of \textit{Lhcb2} is less than 5\% of that found in wild type (Fig. 1a). When normalized to the amount of the PSII core protein \textit{PsbA}, the quantities of \textit{Lhcb3}, \textit{Lhcb4} and \textit{Lhcb6} were constant or less than twofold higher, whereas the amount of \textit{Lhcb5}, the apoprotein of \textit{CP26}, increased by more than sixfold (Fig. 1b).

The absorption spectra showed a strong reduction in the chlorophyll \textit{b} band at 650 nm in the antisense membranes (Fig. 2a, curves 1 and 2), which is consistent with the measured chlorophyll \textit{a/b} ratio of 4.3, as compared with 3.3 in the wild type, and is explained by the depletion of \textit{LHCII} and the increase in \textit{CP26} (\textit{CP26} has a considerably higher chlorophyll \textit{a/b} ratio than has \textit{LHCII}\textsuperscript{18}). The similarity between the fluorescence excitation spectrum and the absorption spectrum in both membranes (Fig. 2a, dotted lines) shows that the extra \textit{CP26} functions as an efficient light-harvesting antenna for PSII. Together with the identical photosynthetic activity of the wild-type and antisense plants\textsuperscript{10}, this indicates that PSII activity is the same in the wild-type and antisense plants.

Chromatographic analysis of detergent-solubilized PSII membranes\textsuperscript{19} gave unexpected results. Fraction II, containing PSII supercomplexes, was still present in the antisense plants (Fig. 2b) but was highly enriched in \textit{CP26}, containing over three times more \textit{Lhcb5} than present in the wild-type fraction (Fig. 1c, d). The quantity of \textit{Lhcb3} was also higher, whereas that of \textit{Lhcb4} and \textit{Lhcb6} was unchanged. These data indicate that the missing \textit{Lhcb1} and \textit{Lhcb2} in the supercomplex are replaced by \textit{CP26}, which prompts an

![Figure 2](image-url)

Figure 2. PSII membrane spectra. a, Absorption spectra of PSII membranes (curves 1 and 2) and gel-filtration fraction \textit{V} (curves 3 and 4) prepared from wild-type (curves 1 and 3) and as\textit{Lhcb2} (curves 2 and 4) plants. Dotted spectra show a fragment of the excitation spectrum of PSII fluorescence emission (685 nm) for the PSII membrane samples. b, Gel-filtration elution profiles of solubilized PSII membranes from wild-type (1) and as\textit{Lhcb2} (2) samples, detected at 670 nm. The main fractions contain PSII supercomplexes (\textit{II}), PSII (\textit{III}), PSII cores (\textit{IV}) and monomeric light-harvesting complexes (\textit{VI}). Curves 3 and 4 show the elution profiles of purified \textit{LHCII} trimers and a monomeric LHC fraction. c, Western blots of fraction \textit{V} prepared from wild-type (WT) and as\textit{Lhcb2} (As) plants. Details are as described in Fig. 1.

![Figure 3](image-url)

Figure 3. Average projections of the PSII complexes from \textit{Arabidopsis Lhcb2} plants. a, An \textit{Arabidopsis C}_{2}S_{2}M_{2} supercomplex\textsuperscript{19} obtained previously, depicting the \textit{S}- and \textit{M}-\textit{LHCII} trimers (blue), the monomeric complexes (green) and the detergent shell (yellow). b-d, Antisense \textit{Lhcb2} plants; e-g, wild-type plants. b, Average of 510 \textit{C}_{2}S_{2} projections; c, average of 290 \textit{C}_{2}S_{2}M_{2} projections; d, average of 411 \textit{C}_{2}S_{2}M_{2} projections; e, average of 300 \textit{C}_{2}S_{2} projections; f, average of 948 \textit{C}_{2}S_{2}M_{2} projections; g, average of 218 \textit{C}_{2}S_{2}M_{2} projections.
intriguing question: can CP26 assemble itself into a trimeric form to mimic the LHCII trimer? In fact, the gel-filtration separation procedure showed that a trimeric fraction (Fig. 2b, fraction V) was still present in the antisense plants, although its concentration was strongly reduced. Western blot analysis of this trimer showed that it contained only CP26 and Lhcb3 (Fig. 2c), whereas the wild-type fraction V contained almost exclusively Lhcb2 and Lhcb1, with some Lhcb3 (but at a much lower amount than in the antisense plant), as expected (refs 3, 7, and Fig. 2c).

The absorption spectra of the wild-type and antisense trimers were also clearly different (Fig. 2a (curves 3 and 4)), with reduced absorption at 650 nm from chlorophyll b in the antisense trimer. We calculated the chlorophyll a/b ratio of the wild-type trimer to be 1.4 as expected, but this ratio increased to 1.90–1.95 in the antisense plant. This higher ratio is explained by the chlorophyll a/b ratios of CP26 (2.0)18, which we confirmed to be the same in the CP26 from both wild-type and antisense plants, and Lhcb3 (1.75)20. The ability to assemble into trimers in vivo was previously thought to be specific for Lhcb1 and Lhcb2. Clearly, this is not so.

Of all the other Lhc genes, Lhcb5 bears the strongest sequence similarity to Lhcb1 and Lhcb2 (ref. 17), greater than 40%, as compared with roughly 30% or less for the other Lhc genes. Lhcb5 is the only gene that clusters with the Lhcb1–Lhcb3 genes17. In addition, a WYXXR motif near the amino terminus of Lhcb1, which is necessary for LHCII trimerization21, is conserved in Lhcb5, the apoprotein of CP26, but not in the apoproteins of CP29 and CP24. But another LHCII trimerization domain near the carboxy terminus (Trp 222)22 is not found in Lhcb5, which could explain the reduced efficiency of trimerization for CP26.

We also analysed the isolated supercomplex population by electron microscopy and image analysis. The same three main classes of supercomplex were found in both asLhcb2 (Fig. 3b–d) and wild-type (Fig. 3e–g) samples. Clearly, there are trimers lacking Lhcb1 and Lhcb2 that are assembled into the supercomplex in the antisense plants and, within the 2-nm resolution of the negative staining, are identical to those found in the wild type. Notably, the abundance of supercomplexes was about the same in both samples. Thus, they cannot arise from the residual Lhcb2 present in the antisense plants and cannot be due to Lhcb3, because the amount of this protein is not increased.

To find out how the supercomplexes are organized in the thylakoid membrane, electron microscopy was carried out on grana membrane fragments3,17. Ordered arrays of PSII supercomplexes were found in both wild-type and antisense plants (Fig. 4a, b). The frequency of occurrence of these arrays was judged to be about the same and, notably, the structural features were almost identical. Both images show arrays of PSII supercomplexes (Fig. 4b, red outline)16. In each case, ordered arrays of PSII cores (bright areas) are separated by areas of trimers, despite the absence in the antisense plants of the main proteins that are known to form LHCII trimers in the wild-type plants. The spacing between the rows is slightly reduced in the antisense plants, suggesting a slightly different packing of the complexes. Notably, also in the membranes from the antisense plants, the contours of the trimers are clearly apparent (Fig. 4a, arrows). Again, within the resolution of the technique, these trimers appear to be identical to those seen in the wild type.

These data show that the PSII supercomplexes have a precise molecular design in which trimers enable a normal supercomplex to be assembled and allow its correct organization in the thylakoid membrane. The observations that ordered arrays of PSII are formed in the antisense plants and that other proteins have been recruited to allow this formation show the importance of this feature of macromolecular organization of PSII. The plants attempt to compensate for the absence of LHCII by synthesizing more CP26 and assembling it into trimers, which then function as a PSII antenna and enable a normal macro-organization to be attained, including their grana structure5. This organization is essential for the high efficiency of energy usage in photosynthesis, because it allows energy exchange between PSII cores and energy migration over a large domain of chlorophyll molecules23.

Because the wild-type supercomplex contains two CP26 copies for each dimeric PSII core complex, we estimate from the western blot analysis that the supercomplex from the antisense plants contains at least four additional copies of CP26 and up to two additional copies of Lhcb3. Thus, there is sufficient CP26 and Lhcb3 to form at least two trimers, thereby explaining how the structure of the complex can be virtually identical to the wild-type even though Lhcb1 and Lhcb2 are absent. In the PSII membranes, in which there is a sixfold increase in CP26, we estimate the presence of two further CP26-containing trimers, explaining the presence of C2S2M2 supercomplexes.

The compensatory replacement of LHCII by CP26 does not give rise to a PSII unit of identical structure and function. There are small differences in the packing of the supercomplexes in the grana membrane but, more notably, the extra LHCII trimers per PSII not present in the supercomplex in wild-type plants3,17 are apparently not replaced by CP26 in the antisense plants. Thus, there is a reduction in the total content of light-harvesting pigments. In addition, CP26 does not have the capacity to be phosphorylated,
and therefore the antisense plants cannot carry out state transitions, a regulatory process that balances the excitation of the two photosystems and requires LHCII phosphorylation. These two differences almost certainly account for the phenotype of these plants, the reduced fitness in the field and the failure to grow normally under conditions of very low irradiance (J. Andersson and S. Jansson, unpublished data). Although there is some apparent redundancy between the similar Lhcb proteins, each one has specific features that confer its unique role in PSI structure and function. Lhcb1 and Lhcb2 are designed to form the essential trimeric building blocks of the supercomplex, where they form the major part of the PSI antenna, but they also provide the vital flexibility needed to regulate excitation distribution between the photosystems. When Lhcb1 and Lhcb2 are missing, CP26 can substitute but not completely. This explains why normally the precision of the composition of the supercomplexes is maintained absolutely in spite of this potential redundancy.

Do other membrane protein complexes show similar features? In cyanobacterial photosynthetic membranes, the CP43 protein, which forms the PSI core, is phosphorylated under certain stress conditions and forms an oligomeric ring-like structure around PSI. Thus, there are hitherto unrecognized features of the structure of macromolecular membrane protein complexes: the same or similar proteins may adopt different structural and functional roles in different or even the same complexes; and complexes that have the same overall structure may contain different proteins resulting in different functions.

Methods

Plant growth

Arabidopsis thaliana cv. Columbia plants were grown in growth chambers with an 8-h photoperiod and 200 μmol quanta m⁻² s⁻¹ for 8 weeks as described. The aslhcb2 line used shows almost total absence of the Lhcb1 and Lhcb2 gene products.

Sample preparation and analysis

PSII membranes were prepared as previously described, but with some modifications. In brief, leaves were homogenized three times for 5 s in a medium containing 20 mM Tricine-NaOH, pH 8.4, 0.45 M sorbitol, 10 mM EDTA and 0.1% bovine serum albumin. After pelleting and washing in 0.3 M sorbitol, 20 mM Tricine NaOH, pH 7.6, 5 mM MgCl₂, chloroplasts were osmotically shocked for 30 s with 5 mM MgCl₂, pH 7.6. Pelleted thylakoids were incubated in stacking medium (5 mM MgCl₂, 15 mM NaCl and 2 mM MES, pH 6.3) for 1 h, after which they treated with 2.5% Triton X-100 at a chlorophyll concentration of 3 mg ml⁻¹. The PSII membranes were sedimented at 30,000g for 30 min, resuspended in buffer containing 20 mM Bis-Tris (pH 6.5) and 5 mM MgCl₂, and cryo-stabilized with 2% dodecyl-D-maltoside at a chlorophyll concentration of 1 mg ml⁻¹. Large fragments were removed by centrifugation for 3 min at 9,000 r.p.m. and the supernatant was filtered promptly through a 0.45-μm filter. Absorption spectra and corrected fluorescence excitation spectra were recorded using a SPEX FluoroLog FL3-22 spectrofluorimeter.

Spectroscopic analysis

Low-temperature spectroscopy was done using an Optistat-DN LN-2 cooled bath cryostat (Lauda). Samples were diluted in a medium containing 70% glycerol (w/v), 20 mM HEPS buffer, pH 7.8, 5 mM MgCl₂, and 0.33 M sorbitol. The chlorophyll concentration for absorption was 4 μg ml⁻¹ and for the fluorescence excitation measurements it was 1 μg ml⁻¹. A poly(methyl metacrylate) PMMA 1-cm cuvette was used for all experiments. Absorption measurements were carried out using a Cary 500 UV-Vis-NIR spectrophotometer and corrected fluorescence excitation spectra were recorded using a SPEX FluoroLog FL3-22 spectrofluorimeter.

Electrophoresis and western blot analysis

Protein samples were solubilized and separated by 15% denaturing SDS-PAGE essentially as described. Roughly equal amounts of chlorophyll were loaded, about 2 μg per lane for PSII membranes and 0.5 μg for isolated supercomplexes and trimers. Proteins were transferred to Hybond-P PVDF membrane (Amersham Pharmacia) in a Mini-Trans-Blot transfer cell (Bio-Rad) at 30 mA for 12 h. Membranes were treated with specific antibodies against the proteins Lhcb1–Lhcb6 (ref. 30) and PsbA (a gift from P. Nixon). The primary antibody was detected by a horseradish peroxidase (HRP)-labelled secondary antibody using an ECL Plus kit (Amersham Pharmacia). Chemiluminescence was detected on Hyperfilm ECL (Amersham Pharmacia) photographic film. We developed the films for 20 min, dried and digitized them in 256-bit greyscale at a resolution of 600 d.p.i. using an Umax Powerlook III high-resolution scanner set in transmission mode. (Films were developed for 0.5–60 min and densities were found to be linear over the first 40 min for each antibody.) A standard Kodak photographic 21 step tablet (OD 0.05–3.05) was used to calibrate the scanner each time an image was scanned. We proceeded to images using the 1D software package of the Image Master Suite (Amersham Pharmacia). Individual lanes and bands were detected automatically by the software, and background for each lane was subtracted using the rolling disc option to give the final band density. For the antibodies against Lhcb1, Lhcb2 and Lhcb5, linear densitometric responses were found in the range of 0.5–10 μg of chlorophyll per lane for PSII membranes.

Electron microscopy

After chromatography, fractions were immediately prepared for electron microscopy. Samples were negatively stained using the droplet method with 2% uranyl acetate and were prepared on glow discharge carbon-coated copper grids as described. The membrane fragments and supercomplexes were analysed by a similar method to that used for preparations from spinach and the wild-type Arabidopsis plants. The samples were imaged in a Philips CM10 electron microscope at 50,000 magnification. Electron micrographs were digitized with a Kodak Eikonix Model 1412 CCD (charge-coupled device) camera. Single-particle projections were extracted from negatives and analysed with IMAGIC software (Image Science Software GmbH) and Groningen Image Processing (‘GRIP’) software (Groningen University Software).
Structure and catalytic mechanism of the human histone methyltransferase SET7/9


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Acetylation1,2, phosphorylation3 and methylation4 of the amino-terminal tails of histones are thought to be involved in the regulation of chromatin structure and function5–7. With just one exception8,9, the enzymes identified in the methylation of specific lysine residues on histones (histone methyltransferases) belong to the SET family10. The high-resolution crystal structure of a ternary complex of human SET7/9 with a histone peptide and cofactor reveals that the peptide substrate and cofactor bind on opposite surfaces of the enzyme. The target lysine accesses the active site of the enzyme and the S-adenosyl-L-methionine (AdoMet) cofactor by inserting its side chain into a narrow channel that runs through the enzyme, connecting the two surfaces. Here we show from the structure and from solution studies that SET7/9, unlike most other SET proteins, is exclusively a mono-methylase. The structure indicates the molecular basis of the specificity of the enzyme for the histone target, and allows us to propose a model for the methylation reaction that accounts for the role of many of the residues that are invariant across the SET family.

Many SET proteins have now been characterized biochemically and several have been the subject of X-ray structure analysis: SET7/9 from human11 and its complex with the product S-adenosyl-L-homocysteine (AdoHcy)12; Dim-5 from Neurospora crassa13; Rubisco large subunit methyltransferase (LSMT) from pea with AdoHcy14; and Clr4 from Schizosaccharomyces pombe15. SET proteins can be classified according to the lysine residues that they target on histones H3, H4 and H2A16, and it is apparent that methylation at these different sites gives rise to distinct biological effects. An additional level of complexity is that lysine residues may be mono-, di- or tri-methylated and that these distinct species lead to different signalling events. For example, in Saccharomyces cerevisiae, although di-methylation of Lys 4 on histone H3 is present at both active and inactive euchromatic genes, tri-methylation is linked exclusively to active genes17.

NMR studies (see Supplementary Information) indicated that a histone peptide containing mono-methylated Lys 4 was better ordered in complex with SET7/9 than unmodified peptide. We therefore used the products of the normal histone methyltransferase (HMT) reaction for crystallization experiments (methylated lysine peptide and AdoHcy). In our previous studies11 we obtained useful SET7/9 crystals only from constructs lacking the small carboxy-terminal segment, which is, nevertheless, essential for the catalytic activity of the enzyme. Here, we have used a catalytically active construct that contains the complete C-terminal segment. We obtained well-ordered crystals of the ternary complex of SET7/9 that diffracted to at least 1.7 Å spacing, and the structure was readily solved by molecular replacement. The C-terminal segment, the AdoHcy cofactor and most of the substrate peptide are well defined in the electron density maps, as are all the important residues around the active site. The overall structure of the ternary complex...