STRUCTURE AND MEMBRANE ORGANIZATION OF PHOTOSYSTEM II IN GREEN PLANTS

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ABSTRACT
Photosystem II (PSII) is the pigment protein complex embedded in the thylakoid membrane of higher plants, algae, and cyanobacteria that uses solar energy to drive the photosynthetic water-splitting reaction. This chapter reviews the primary, secondary, tertiary, and quaternary structures of PSII as well as the function of its constituent subunits. The understanding of in vivo organization of PSII is based in part on freeze-etched and freeze-fracture images of thylakoid membranes. These images show a resolution of about 40–50 Å and so provide information mainly on the localization, heterogeneity, dimensions, and shapes of membrane-embedded PSII complexes. Higher resolution of about 15–40 Å has been obtained from single particle images of isolated PSII complexes of defined and differing subunit composition and from electron crystallography of 2-D crystals. Observations are discussed in terms of the oligomeric state and subunit organization of PSII and its antenna components.

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INTRODUCTION

Photosystem II (PSII) is a multisubunit complex embedded in the thylakoid membranes of higher plants, algae, and cyanobacteria. It uses light energy to catalyze a series of electron transfer reactions resulting in the splitting of water into molecular oxygen, protons, and electrons. These reactions occur on an enormous scale and are responsible for the production of atmospheric oxygen and indirectly for almost all the biomass on the planet. Despite its importance, the catalytic properties of PSII have never been reproduced artificially. Understanding PSII's unique chemistry is important and could have implications for agriculture as PSII is a main site of damage during environmental stress.

The chapter reviews our current knowledge of the three-dimensional structure of PSII in higher plants, an area of research that has developed rapidly over recent years. We first outline the photochemical reactions in this photosystem and summarize in a table the main structural features of individual subunits, focusing on their likely transmembrane helical content and cofactor binding properties. Electron microscopy of PSII is then reviewed to relate the subunit and cofactor composition of PSII to its three-dimensional structure. Low-resolution structural data on PSIi, obtained from freeze-etch and freeze-fracture studies of thylakoid membranes, is reviewed initially. Such studies have provided information on the location, heterogeneity, and overall size and shape of PSII and its antenna system in the thylakoid membrane at resolutions of 40–50 Å. To obtain higher-resolution information (~15–40 Å), two other approaches have been used: single particle image averaging of detergent-solubilized PSII complexes and analysis of two-dimensional crystals. The former

1 Abbreviations: β-Car, β-carotene; BChl, bacteriochlorophyll; Chl, chlorophyll; CD, circular dichroism; CP, chlorophyll protein; cyt b559, cytochrome b559; LHCII, light-harvesting complex of PSII; LH2, light-harvesting complex of purple photosynthetic bacteria; PSI, photosystem one; PSII, photosystem two; P680, primary electron donor of PSII; Pheo, pheophytin; PQ, plastoquinone; QA and QB, primary and secondary electron acceptors of PSII; RC, reaction center; STEM, scanning transmission electron microscopy
PHOTOSYSTEM II STRUCTURE AND ORGANIZATION has yielded considerable information on the oligomeric state and subunit organization of PSII and its antenna system, whereas the latter offers the potential of a structure at atomic resolution. Results obtained from both approaches are discussed in terms of the question of whether PSII exists as a monomer or dimer in vivo. Finally, the conclusions emerging from these studies are compared with biochemical and cross-linking data.

THE PHOTOSYSTEM II SUBUNITS: STRUCTURE AND FUNCTION

Well over 20 subunits (PsbA–PsbX, Lhcb1–6) are associated with PSII of higher plants and green algae (Figure 1) and have been named after the genes.

*Figure 1* Photosystem II: an overview of subunit composition and electron transport. The PSII complex and its antenna system consists of more than 20 subunits that are either embedded in the thylakoid membrane or associated with its luminal surface. Light energy is trapped predominantly in the outer antenna, consisting of the proteins Lhcb1–6. The excitation energy is transferred to the photochemically active reaction center (D1 and D2 proteins) via CP47 and CP43, where it is used to drive the water-splitting reaction. The electrons extracted from water are passed from the luminally located four-atom Mn cluster to D1-Tyr 161 (Yz), P680+, Phe, QA and on to QB, via a nonheme iron group. This electron transport pathway is marked with arrows. The protons and molecular oxygen produced during the water-splitting reaction are released into the lumen. The plastoquinone (PQ) bound at the QB site accepts two electrons derived from water via the electron transport chain and two protons from the stroma before being released into the thylakoid membrane in the form of PQH2. The letter notation used for the subunits of the core complex reflects gene origin (e.g. A = product of psbA gene).
Table 1  The PSII subunits: primary and secondary structure, cofactor content, and function

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit</th>
<th>Mass (kDa)</th>
<th>No. of transmembrane α-helices</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>psbA</td>
<td>D1</td>
<td>38.021(S)</td>
<td>5</td>
<td>YZ &amp; binds P680, Pheo, QA</td>
</tr>
<tr>
<td>psbD</td>
<td>D2</td>
<td>39.48(S)</td>
<td>5</td>
<td>YD &amp; binds P680, QA</td>
</tr>
<tr>
<td>psbE</td>
<td>α-cyt b559</td>
<td>9.255(S)</td>
<td>1</td>
<td>Binds heme, photoprotection</td>
</tr>
<tr>
<td>psbF</td>
<td>β-cyt b559</td>
<td>4.409(S)</td>
<td>1</td>
<td>Binds heme, photoprotection</td>
</tr>
<tr>
<td>psbl</td>
<td>b protein</td>
<td>4.195(S)</td>
<td>1</td>
<td>?</td>
</tr>
<tr>
<td>psbB</td>
<td>CP47</td>
<td>56.278(S)</td>
<td>6</td>
<td>Excitation energy transfer, binds 33 kDa</td>
</tr>
<tr>
<td>psbC</td>
<td>CP43</td>
<td>50.066(S)</td>
<td>6</td>
<td>Excitation energy transfer, binds 33 kDa</td>
</tr>
<tr>
<td>psbH</td>
<td>H protein</td>
<td>7.697(S)</td>
<td>1</td>
<td>Photoprotection</td>
</tr>
<tr>
<td>psbK</td>
<td>K protein</td>
<td>4.283(S)</td>
<td>1</td>
<td>PSII assembly, PSII stability</td>
</tr>
<tr>
<td>psbL</td>
<td>L protein</td>
<td>4.366(S)</td>
<td>1</td>
<td>Involved in QA function</td>
</tr>
<tr>
<td>psbM</td>
<td>M protein</td>
<td>3.755(P)</td>
<td>1</td>
<td>?</td>
</tr>
<tr>
<td>psbN</td>
<td>N protein</td>
<td>4.722(T)</td>
<td>1</td>
<td>?</td>
</tr>
<tr>
<td>psbO</td>
<td>33-kDa ext. protein</td>
<td>26.539(S)</td>
<td>0</td>
<td>Stabilizes Mn cluster, Ca\textsuperscript{2+} &amp; Cl\textsuperscript{-} binding</td>
</tr>
<tr>
<td>psbP</td>
<td>23-kDa ext. protein</td>
<td>20.210(S)</td>
<td>0</td>
<td>Ca\textsuperscript{2+} and Cl\textsuperscript{-} binding</td>
</tr>
<tr>
<td>psbQ</td>
<td>16-kDa ext. protein</td>
<td>16.523(S)</td>
<td>0</td>
<td>Ca\textsuperscript{2+} and Cl\textsuperscript{-} binding</td>
</tr>
<tr>
<td>psbR</td>
<td>R protein</td>
<td>10.236(S)</td>
<td>0</td>
<td>Donor and acceptor side functions</td>
</tr>
<tr>
<td>psbS</td>
<td>S protein</td>
<td>21.705(S)</td>
<td>4</td>
<td>Chl chaperonin/antenna component</td>
</tr>
<tr>
<td>psbT</td>
<td>T protein</td>
<td>3.283(S)</td>
<td>0</td>
<td>?</td>
</tr>
<tr>
<td>psbV</td>
<td>V protein*</td>
<td>15.121(Sy)</td>
<td>0</td>
<td>Donor side stability</td>
</tr>
<tr>
<td>psbW</td>
<td>W protein</td>
<td>5.928(S)</td>
<td>1</td>
<td>?</td>
</tr>
<tr>
<td>psbX</td>
<td>X protein</td>
<td>4.225(S)</td>
<td>1</td>
<td>QA function</td>
</tr>
<tr>
<td>lhcb4</td>
<td>Lhcb4 (CP29)</td>
<td>29</td>
<td>3</td>
<td>Excitation energy transfer &amp; dissipation</td>
</tr>
<tr>
<td>lhcb5</td>
<td>Lhcb5 (CP26)</td>
<td>26</td>
<td>3</td>
<td>Excitation energy transfer &amp; dissipation</td>
</tr>
<tr>
<td>lhcb6</td>
<td>Lhcb6 (CP24)</td>
<td>24</td>
<td>3</td>
<td>Excitation energy transfer &amp; dissipation</td>
</tr>
<tr>
<td>psbJ</td>
<td>J protein</td>
<td>4.116(P)</td>
<td>1</td>
<td>PSII assembly</td>
</tr>
<tr>
<td>psbU</td>
<td>U protein*</td>
<td>10(Cy)</td>
<td>?</td>
<td>?</td>
</tr>
<tr>
<td>lhcb1</td>
<td>Lhcb1</td>
<td>25</td>
<td>3</td>
<td>Light harvesting</td>
</tr>
<tr>
<td>lhcb2</td>
<td>Lhcb2</td>
<td>25</td>
<td>3</td>
<td>Light harvesting</td>
</tr>
<tr>
<td>lhcb3</td>
<td>Lhcb3</td>
<td>25</td>
<td>3</td>
<td>Light harvesting</td>
</tr>
</tbody>
</table>

\*Twenty-three putative PSII proteins are encoded by the psbA–psbX genes, whereas at least six outer antenna components are encoded by Lhcb1–L–6. The subunits encoded by these genes are listed according to the complexes with which they copurify (e.g. reaction centers), and those marked (*) have only been detected in cyanobacteria. The molecular masses of the mature PsbA–PsbX proteins, except PsbU, are calculated from the protein sequences reported in the SWISSPROT data base using the MacBioSpec program (Sciex Corp., Thornhill, Ontario, Canada). The abbreviations given in brackets after the molecular masses denote the organism for which the subunit mass is given, as
## PHOTOSYSTEM II STRUCTURE AND ORGANIZATION

<table>
<thead>
<tr>
<th>Chla</th>
<th>Chlb</th>
<th>β-Car</th>
<th>Pheo</th>
<th>Lut</th>
<th>Neo</th>
<th>Viol</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

**PSII reaction center**

| 10–25 | 0 | 3 | 0 | ? | 0 | 0 |
| 9–25  | 0 | 5 | 0 | ? | 0 | 0 |

**Full PSII core**

| 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 |

**PSII core & minor CAB proteins**

| 9–10 | 3–4 | — | 0 | 1–2 | 1 | 1–2 |
| 7–9  | 4–5 | — | 0 | 2 | 0.5–1 | 0.5–1 |
| 6    | 7 | — | 2 | 1 | 1 |

**Full PSII & antenna complex**

| 8 | 6 | 0 | 0 | 2 | 0.5–1 | 0.5 |
| 8 | 6 | 0 | 0 | 2 | 0.5–1 | 0.5 |

follows: S, spinach; Sy, *Synechococcus* sp.; P, pea; T, tobacco. For the Lhcb proteins and PsbU, only the apparent molecular masses are provided. The next column lists the number of predicted transmembrane helices of each subunit. The putative functions of each subunit are given in the next column. The cofactors associated with the individual subunits are also listed as follows: Chla, chlorophyll a; Chlb, chlorophyll b; β-Car, β-carotene; Pheo, Pheophytin; Lut, lutein; Neo, neoxanthin; Viol, violoxanthin. The references reporting the information summarized in this table are given in the text.
encoding them (38, 46–49, 58, 65). The structure (22, 28, 33, 53, 65, 69, 79, 89, 97, 105, 111, 123, 129), cofactor organization (16, 22, 25, 38, 46–49, 53, 65, 69, 79, 89, 97, 108, 111, 123, 129), and function (32, 38, 46–49, 53, 62, 65, 69, 89, 97, 111) of these subunits have been studied in detail and are summarized in Table 1.

Figure 1 shows the subunit composition of PSII and its antenna system and indicates the photochemical processes that this photosystem catalyzes. The overall reaction driven by PSII is given in Equation 1, and all the cofactors involved are either bound within the reaction center proteins D1 and D2 or are closely associated with them.

\[
\text{LIGHT} \quad \text{PSII} \quad 2\text{H}_2\text{O} + 2\text{PQ} \rightarrow \text{O}_2 + 2\text{PQH}_2
\]

The light energy used to drive the water-plastoquinone (PQ) oxidoreductase reactions is captured predominantly by many molecules of chlorophyll (Chl) a and b [averaging about 250 per reaction center (70)] and carotenoid [β-carotene, lutein, neoxanthin, and violoxanthin (20, 69)] associated with light-harvesting antenna proteins, Lhcb1–6 (18, 21, 26, 46–49, 52, 65, 69, 97). The derived excitation energy is passed from the Lhcb proteins along an excitonically linked network of Chl molecules associated with CP47 (PsbB) and CP43 (PsbC) (18, 21, 22, 26, 65, 99, 100) to the PSII reaction center (RC). The RC consists of the D1 (PsbA) and D2 (PsbD) proteins (17, 91, 126), which are highly conserved between higher plant species and have a significant degree of local homology with the primary sequence of the L and M subunits of purple bacteria (16, 78, 110, 129). Ultimately the excitation energy derived from light is used to convert the primary oxidant, P680, to P680* (17, 91, 110, 111, 126). P680 is thought to consist of two Chl molecules ligated to the D1 and D2 proteins, though the excitonic coupling is much weaker than in the “special pair” of the purple bacterial reaction center (36, 111, 132). The excited state, P680*, donates a single high-energy unpaired electron to a molecule of pheophytin (Pheo), thereby forming the radical pair, P680*Pheo− (17, 91, 126, 110, 111) (Figure 1). Each time P680+ is formed, it accepts an electron from a specific amino acid residue (D1-Tyr161) and therefore is reduced to P680 (13, 25, 60, 98, 127). Illumination of PSII allows the P680, P680*, P680+ cycle to be repeated and enables the sequential extraction of electrons from D1-Tyr161 (Figure 1). As D1-Tyr161 donates an electron to P680+, it accepts another from water via a four-atom manganese (Mn) cluster, associated with the lumenal surface of PSII. Joliot et al (66) and Kok et al (68) used short light flashes to induce single turnovers of P680 and showed that the Mn cluster passed through a series of oxidation states referred to as the S0–S4 cycle. The
electrons, which are accepted by P680+, are passed along the electron transport chain (Figure 1). In this way, Pheo accepts electrons from P680 and passes them on to a plastoquinone molecule (QA), tightly bound to the D2 protein (34, 71, 78, 107, 129). QA− passes its electron on to a second plastoquinone molecule, associated with the QB site on the D1 protein (35, 59, 71, 78, 79, 96, 103, 105, 113). This electron transfer is aided by the presence of a nonheme iron located between QA and QB (16, 34, 79, 110). Each plastoquinone associated with the QB site can accept two electrons derived from water and two protons from the stroma before being released into the lipid matrix in the form of reduced plastoquinone (PQH2).

The luminal surfaces of membrane embedded CP47, CP43, and reaction center proteins of higher plants and green algae are in close contact with a number of extrinsic proteins. These include the products of the psbO (33-kDa subunit), psbP (23-kDa subunit), and psbQ (16-kDa subunits) genes, which together form the oxygen-evolving complex (OEC). The OEC also includes the Mn cluster (9, 25, 44, 89) and the three extrinsic proteins are involved in the optimization of its function in water splitting. The 33-kDa subunit stabilizes the Mn cluster (89) while the 23-kDa subunit allows PSII to evolve oxygen under both Ca2+- (43, 89) and Cl−-limiting conditions (86, 87). This has led to the suggestion that the 23-kDa subunit acts as a concentrator of these ions (89). The 16-kDa polypeptide aids PSII to evolve oxygen efficiently under severely Cl−-limiting (<3 mM) conditions (2).

Except for subunit PsbS, all other subunits associated with PSII (Figure 1, Table 1) have a mass under 10 kDa (38). The exact number of the small subunits associated with PSII in vivo is not known, primarily because some of these proteins stain poorly and/or are difficult to resolve even in high-resolution gels (73). The reader is referred to the detailed review by Erickson & Rochaix (38) and the SWISSPROT data base for further information on the low-molecular-weight subunits of PSII, because their properties and those of the other PSII subunits are only summarized here (Table 1). Table 1 lists the subunits, in a way that reflects the PSII particle type that they are associated with, the smallest being the isolated reaction center and the largest being PSII with its complete antenna. The cofactors bound by each subunit are given, together with subunit function, common name, and the number of α-helices that the subunit is predicted to contain.

We now relate the information on the subunit structure summarized in Table 1 to structural information available on PSII and its antenna system.
ARRANGEMENT OF PSII IN THE THYLAKOID MEMBRANE

Much of our understanding of the in vivo organization of PSII is based on freeze-etched and freeze-fracture images of thylakoid membranes. Such electron microscopy studies provide structural detail at a resolution of about 40–50 Å and have provided information on the localization, heterogeneity, dimensions, and shapes of PSII and its antenna system.

By thin sectioning of fixed chloroplasts, electron microscopy has revealed the overall architecture of the thylakoid membrane of higher plants. The thylakoid membrane consists of two main compartments, the grana and the stroma lamellae. The stroma membranes form unstacked (nonappressed) regions, whereas the grana membranes are mostly present as stacked (appressed) membranes. There is a marked heterogeneity in lateral distribution of the major complexes of this membrane. It is generally accepted that PSII, photosystem one (PSI), and ATP synthase are mainly laterally segregated, with PSI and ATP synthase excluded from the appressed grana membranes and PSII abundantly present in the stacked parts of the thylakoid membrane (10, 15). The total picture, however, is more complex, because there is also heterogeneity among both PSII and PSI in subunit composition and lateral distribution (3).

In freeze-etching studies, thylakoids are flash frozen before the evaporation of surface water under vacuum (typically at −100°C). This process exposes the membrane surface and so allows the visualization of extrinsic components in their near-to-native state. The organization of the membrane embedded parts of PSII and the antenna proteins have been studied using the freeze-fracture technique, which involves the cleavage of the lipid bilayer along its internal hydrophobic plane, before image analysis. The terminology (ESs, PSs, ESu, PSu) used by Staehelin (120, 121) has generally been adopted to describe the endoplasmic (E) and protoplasmic (P) surfaces (S) of stacked (s) and unstacked (u) freeze-etched thylakoid membranes. The corresponding freeze-fracture (F) planes are referred to as EFs, PFs, EFu, and PFu.

The ultrastructure of thylakoid membranes of barley (55, 80), spinach (85, 119, 120, 122), maize (84), lettuce (56), soybean (67), Alocasia (7), and pea (11, 101) have all been analyzed by freeze-etch and freeze-fracture electron microscopy and show marked similarities (see 80, 121). The protein complexes detected in these analyses were named according to the surfaces with which they were associated (e.g. ESs or PFu particles). They differed in size and shape, and the constituent components of many were identified by the analysis of mutant membranes. For example, the analysis of PSI-deficient mutant thylakoid membranes showed that this photosystem formed part of the
large PFu particles (117). Localization of complexes in the thylakoid membranes was further facilitated by antibody labeling (95).

**Localization of PSII and LHCII**

Comparative freeze-etch and freeze-fracture studies of wild-type and PSII-deficient mutants of tobacco showed that the thylakoid membranes of the latter were depleted of ESs and EFs particles (81). From these results, it was concluded that the ESs and EFs particles corresponded to the extrinsic and internal parts of PSII, respectively. Support for this conclusion came from parallel studies of PSII-deficient barley mutants (*xantha*-b12, *viridis*-c12, *viridis* e-64, and *viridis* zd69), which showed that their granal membranes were also greatly depleted of EFs particles (e.g. 115, 121). The antenna proteins were located using Lhcb protein-deficient mutants (e.g. barley mutants *xantha*-135 and *viridis*-k23 and *chlorina* f2) and by comparing thylakoid membranes from light- and dark-grown plants, the latter being depleted in these antenna proteins (12, 83, 116, 118). These studies showed that membranes depleted of Lhcb proteins lacked the PFs particles found in wild-type membranes. Freeze-etch images of membranes depleted of Lhcb protein also showed the ESs particles to be smaller. Together, these results suggested that the PFs particles contained the antenna proteins and that they were closely associated with PSII (EFs and ESs) particles.

The close association of PSII and the Lhcb proteins was characterized in more detail by the analysis of 2-D crystalline arrays of ESs complexes. Images of such ordered arrays showing a section of their freeze-etch surface (ESs surface) and a part of the protoplasmic fracture face (PFs surface) were presented by Miller (80) and Simpson (116). They showed that the Lhcb proteins (PFs particles in the PFs fracture face) fitted in register into the grooves between the PSII complexes (ESs particles in the freeze-etched ESs surface).

**Heterogeneity of PSII In Vivo**

The ESs and EFs particles that contain PSII are not evenly distributed within the thylakoid membrane. Freeze-etch and freeze-fracture studies have shown that ESs and EFs particles, respectively, can be organized into 2-D arrays or be more randomly dispersed within the grana. PSII-like particles are also observed in the stroma lamellae. It has been calculated that 85% of PSII particles are located in the granal stacks. They are called PSIIα. The remaining 15% of PSII complexes are located in the stroma and are called PSIIβ (3, 10). PSIIα complexes have approximately twice the antenna size of the PSIIβ complexes (8) and are more efficient at reducing ferricyanide and duroquinone (57).
Because PSIIα complexes are more abundant and active than PSIIβ and because all the available structural data on PSII have been collected from such types of complexes, we focus on them in the sections below.

The PSIIα population in the grana can itself be divided into subpopulations of differing antenna size (4, 5). Interestingly, freeze-etch studies have shown that some ESs particles within the grana form 2-D arrays, whereas others are more randomly dispersed and so form at least two subpopulations (112, 116).

Several publications provide the dimensions of arrayed ESs particles. Wild-type ESs arrays in barley have unit cell dimensions of 175 × 247, 180 × 225, and 180 × 240 Å (82, 115). ESs arrays in spinach had an almost rectangular lattice with spacings of 175 × 204 Å (112), while those in the grana of two fatty acid desaturase mutants of Arabidopsis thaliana had unit cell dimensions of 190 × 230 Å and 180 × 230 Å (131). Closer analysis of the arrayed ESs particles showed them to have a tetramer-like appearance on their lumenal (ESs) surface and a height of 82 Å perpendicular to the membrane plane (112, 116). Sequential removal of the 16, 23, and 33 kDa proteins of the OEC resulted in a height reduction to 78, 74, and 61 Å, respectively (112). The removal of all three polypeptides exposed the membrane-embedded portion of the PSII-LHCII supercomplex and revealed it to have a dimer-like structure. This led to the conclusion that the ESs particle is a dimeric PSII-LHCII supercomplex and that the tetramer-like appearance of its lumenal surface reflects the organization of two copies of the OEC. Dimeric PSII-LHCII supercomplexes (Figure 2D) isolated by solubilization in detergent have been biochemically and structurally characterized (21, 52, 93). The structure of the isolated complex can be computationally aligned to form an array with a spacing of 190 × 210 Å and a similar lumenal surface topology to ESs particles (21, 52, 112). It is therefore possible that ESs-type arrays consist of dimeric PSII-LHCII supercomplexes of the type reported by Boekema et al (21).

One argument that could be raised against the proposal that the ESs particles consist of dimeric PSII-LHCII supercomplexes of this type is that, on average, PSII reaction centers in the grana are associated with about 250 Chl molecules (70). Thus a dimeric complex of PSII would be expected to bind about 500 Chl as opposed to the 200 Chl per RC found for the isolated PSII-LHCII supercomplex (21, 93). There is insufficient space within the unit cell of the constructed ESs lattice to accommodate the extra Lhcb subunits that would be required to bind an additional 300 Chl molecules. However, it should be stressed that the value of 250 Chl:RC obtained for granal membranes is an average value and that the randomly organized ESs particles within the grana are spaced sufficiently widely to allow them to be associated with much larger antenna systems (>250 Chl:RC). This hypothesis fits both the finding that the
Figure 2  Top view projection maps of spinach PSII complexes obtained from nonperiodic (single particle) averaging of electron microscopy images obtained after negative staining. (A) A monomeric PSII core complex (240 kDa) consisting of CP47, CP43, the D1 and D2 proteins, cyt b559, the 33-kDa extrinsic protein, and associated low-molecular-weight polypeptides; (B) a dimeric PSII core complex (450 kDa) of the same subunit composition as the PSII core monomer; (C) a PSII-LHCII supercomplex consisting of a PSII core dimer and one set of Lhcb proteins (Lhcb1, 2, 4, and 5); (D) a PSII-LHCII supercomplex consisting of a PSII core dimer and two sets of Lhcb proteins (Lhcb1, 2, 4, and 5). The outline of the PSII core dimer shown in (B) is superimposed upon the PSII-LHCII supercomplex (D) to show the central location of the PSII core dimer. All the images are similar to those published by Boekema et al (21), but they differ in that they are the averages of larger data sets and so have an improved resolution (approximately 20 Å). Note: a twofold rotational symmetry has been imposed on images B and D. The scale bar measures 5 nm.
grana contain 250 Chl:RC on average and that the PSIIα population consists of subpopulations differing in antenna size (4, 5).

STRUCTURE DETERMINATION BY ELECTRON MICROSCOPY

X-ray crystallography is a powerful technique for elucidating the structure of proteins and has been particularly successful with photosynthetic membrane proteins. Atomic structures exist for the reaction center (6, 27–29) and the light-harvesting complex (LH2) (77) of photosynthetic purple bacteria. A map of near-atomic resolution also exists for PSI (42). Recently, the structure of the mitochondrial cytochrome b-c1 complex has been solved at atomic resolution, thus providing structural information relevant to the related photosynthetic cytochrome b6f complex (137). Despite these striking successes, the X-ray approach has not revealed the structure of PSII. Adir et al (1) and Fotinou et al (40) have grown 3-D crystals of PSII cores, but they were too small and insufficiently ordered for high-resolution analyses.

In the absence of highly ordered 3-D crystals suitable for X-ray diffraction analyses, electron microscopy offers an alternative approach. Two techniques are available: single particle image averaging and electron crystallography using 2-D crystals. Both have been applied to elucidate the structure of PSII and in principle could yield maps at atomic resolution (54). To date, both techniques have yielded information on the oligomeric state (monomers vs dimers) of PSII in vivo and on its subunit organization.

Single Particle Image Averaging

Single particle analysis has been used to study the structure of a wide range of biological molecules, either imaged in negative stain or in vitreous ice (cryo-electron microscopy). Data obtained from biological molecules in vitreous ice correspond most clearly with the native protein structure and typically have a resolution up to 15 Å. Conventionally negatively stained samples (air dried and imaged at room temperature) usually give lower resolution (approximately 20 Å). The images obtained are classified according to the orientation of the particle (e.g. top and side views), and members of each class are then rotationally and translationally (i.e. shifted in the X-Y plane) aligned. The procedure is more useful for large particles (>250 kDa), which can be more accurately aligned.

The single particle image averaging approach has been used to analyze a number of detergent-solubilized PSII complexes of known subunit composi-
tion and oligomeric state (i.e. monomers and dimers) under negative stain conditions. The smallest higher plant PSII complex analyzed using this approach is a monomeric PSII core (approximately 240 kDa) consisting of CP47, CP43, D1, D2, cyt b559, the 33-kDa subunit, and associated low-molecular-weight polypeptides, together with about 36 Chla molecules (21, 51, 52). This complex is depicted in its top view orientation in Figure 2A. A dimeric complex of identical subunit composition and having an apparent molecular mass of 450 kDa is shown in its top view orientation in Figure 2B. By comparing Figures 2A and 2B, it can be seen that the projection map of the monomeric complex corresponds well with each half of the PSII core dimer. The dimensions of the latter were calculated to be $206 \times 131$ Å in the presence of the detergent (dodecyl maltoside) shell. When a detergent layer of 17 Å is deducted (30), the corrected dimensions of the isolated dimer are calculated to be $172 \times 97$ Å (21). Single particle analysis has also identified a PSII core dimer of similar size isolated from the cyanobacterium *Synechococcus elongatus* (21).

Although the dimeric organization of PSII has been readily accepted for cyanobacteria, there is currently a debate about whether the PSII particles in the grana are monomeric or dimeric complexes. It has been argued that dimeric PSII complexes, such as the one shown in Figure 2B, that were isolated from granal membranes, are the product of aggregation of PSII monomers rather than an in vivo form of PSII (39, 61, 92). A considerable volume of evidence speaks against this hypothesis. Freeze-etch images of thylakoid membranes showed that PSII complexes (ESs particles) located in the grana were most probably dimers (112). Holzenburg et al (61) argued that at the relatively low resolution of such freeze-etch images and of projection maps of crystallized dimer-like complexes (19), a pseudosymmetry relating to the similarity in structure of the D1 vs D2 proteins and CP47 vs CP43 proteins could be mistaken for a real twofold symmetry indicative of a dimeric complex. Consequently, it was suggested that PSII was actually a monomeric pseudosymmetric complex. To address this point, PSII complexes derived from gently solubilized granal and stromal membranes were analyzed under nondenaturing electrophoresis conditions. The granal PSII fraction consisted predominantly of dimeric PSII complexes associated with Lhcb proteins, while their stromal counterparts were monomeric (109). This result also showed that the non-denaturating electrophoresis conditions used did not induce aggregation of monomeric PSII (109). Other non-denaturing gel electrophoresis studies of this type came to the similar conclusion that PSII in the grana is dimeric (99). Furthermore, our own data (51, 52) has shown that isolated PSII core dimers (Figure 2B) were more active in terms of oxygen evolution, are associated with higher
levels of Chla (39 Chla:2 Pheo), and have much lower levels of D1 and D2 breakdown fragments than their monomeric counterparts (Figure 2A). Based on this information it is difficult to see how two damaged monomers could aggregate to form an intact dimer of the type shown in Figure 2B (51, 52). Taken together, these and other studies (see section on “Heterogeneity of PSII In Vivo”) strongly suggest that PSII is a dimer in the grana.

Recently, grana membranes were solubilized with a single gentle solubilization step, to obtain a more native PSII complex. The solubilized mixture was then resolved by sucrose density gradient centrifugation. The largest PSII complex (approximately 700 kDa) consisted of the PSII core dimer subunits (Figure 2B) plus Lhcb1, 2, 4, and 5 proteins and contained about 200 Chl molecules (156 Chla, 4 Chlb) (21, 93). Like the PSII monomer (Figure 2A) and dimer (Figure 2B) cores, this PSII-LHCII supercomplex was active in oxygen evolution. Single particle analysis showed an unsymmetrized form of this PSII-LHCII supercomplex (Figure 2D) to have a clear twofold rotational symmetry axis around the center of the complex, consistent with a dimeric structure (21). Superimposed upon the projection map of the PSII-LHCII supercomplex (Figure 2D) is the outline of the PSII core dimer depicted in Figure 2B. This alignment of Figures 2B and 2D shows that the PSII core dimer forms the central part of the PSII-LHCII supercomplex (Figure 2D) and that the two regions flanking it must contain the Lhcb proteins. Figure 2C shows a PSII-LHCII complex from which one of the Lhcb protein sets has dissociated. This form of PSII-LHCII complex (Figure 2C) lacks the twofold symmetry of the PSII-LHCII supercomplex (Figure 2D) due to the loss of this single Lhcb set. Nevertheless the PSII core region is clearly twofold symmetric and closely resembles the averaged PSII core dimer top view (Figure 2B). Furthermore the structural features of the remaining Lhcb set (Figure 2C) correspond closely with that of the symmetric PSII-LHCII supercomplex. These findings confirm the true twofold symmetry and hence the dimeric nature of the PSII core.

After correction for the detergent layer, the PSII-LHCII supercomplex shown in Figure 2D is calculated to have dimensions of 270 × 125 Å. In its side view orientation, the particle has a thickness of about 60 Å in the Lhcb region, consistent with the height of LHCII (69, 133). Its maximal thickness, close to the center of the complex, is about 90 Å. This dimension is similar to that of the arrayed ESs particles observed in freeze-etch and freeze-fracture images of thylakoid membranes of a number of higher plant species (see section on “Heterogeneity of PSII In Vivo”). This suggests further that the ESs (EFs) particles are probably dimeric PSII-LHCII supercomplexes, as discussed above.
Electron Crystallography

The successful application of the other type of high-resolution electron microscopy (electron crystallography) in the determination of the atomic structure of trimeric LHCII demonstrates the power of this approach (69). Figure 3 shows that within each Lhcb protein monomer of LHCII, the two longest transmembrane helices are surrounded by Chla and Chlb molecules, and two molecules of lutein. The chlorophylls are arranged in two layers, toward the lumenal and stromal membrane surfaces, and the main ligands are Glu, Asn, Glu, His, and Gly. The close association of the luteins with the excitonically linked chlorophylls prevents singlet oxygen formation and the damage induced by it (69). The luteins also play a structural role in that they form a cross-brace in the center of the complex, providing a direct and strong link between the peptide

Figure 3  The structure of LHCII. This figure is taken from Kühlbrandt (69) and shows a folding model of a monomeric LHCII subunit. The LHCII monomer has three transmembrane α-helices (A–C) and a shorter helix (D) toward the lumenal surface of the membrane. The chlorophylls are arranged in two layers close to the stromal and lumenal surfaces of the membrane. Two Lut residues (L) form a cross-brace between helices A and B and are excitonically linked to the chlorophyll (Chl) network. Sequence homology between the subunits Lhcb1–6 suggests that all their structures will be similar.
loops at both surfaces (69). Based on the high degree of sequence homology, it is likely that the atomic structures of the other Lhcb proteins will also be similar to that of the LHCII monomer.

Isolated PSII complexes have also been crystallized when reconstituted with detergent solubilized lipid (31, 51, 90, 130). Another 2-D crystallization approach involves a detergent-induced delipidation step of granal membranes enriched in PSII and Lhcb proteins. This forces the complexes in the granal membranes into close contact and has yielded small 2-D crystals of PSII (19, 39, 61, 74–76, 109).

To compare the projection maps obtained using single particle image averaging (Figure 2) and electron crystallography (Figure 4), the isolated PSII core dimer shown in Figure 2B (51) was reconstituted with thylakoid lipid and crystallized, though the 33-kDa extrinsic protein dissociated during the process. The top view projection map, obtained from this D1-D2-cyt b559-CP47-CP43 complex, under negative stain conditions, is shown in Figure 4A. The first important point to note is that the unit cell of the crystallized PSII core dimer (a = 117 Å x b = 173 Å, γ = 110°) compares well with the size of the single particle averaged image of this complex (97 x 172 Å). The second is that the structural features of the two images are almost identical. The most prominent features are two densities (marked with * in Figure 4A), which are positioned on either side of a central hole. Although the projection map shown in Figure 4A is unsymmetrized, it clearly has a twofold rotational symmetry around its center, confirming its dimeric structure. Image sections taken through the 3-D image of this complex show it to have twofold symmetry throughout (E Morris, B Hankamer, D Zheleva & J Barber, unpublished data).

The top view projection map of another PSII crystal, imaged under cryo conditions, is shown in Figure 4B. The crystal was produced by delipidating granal membranes (75). A comparison of Figures 4A and 4B shows that the two projection maps have very similar features, including a central hole and two predominant regions of density, on either side of it (marked with *). The two crystal forms also have very similar unit cell dimensions to that of the complex shown in Figure 4B (a = 114 Å, b = 173 Å, γ = 106.6°). Despite the similarity of the two projection maps and their unit cell dimensions, the crystals of Marr et al (75) were reported to contain Lhcb4, 5, and 6 and PsbS, in addition to the other PSII subunits of the complex shown in Figure 4A. Marr et al (75) came to this conclusion based on direct immunolabeling of their crystals. This apparent discrepancy remains to be resolved because the complex shown in Figure 4A does not contain these subunits.

Holzenburg et al (61) and Ford et al (39) also reported projection maps of crystallized PSII complexes with and without the extrinsic subunits of the
Figure 4  Projection structures of spinach PSII complexes obtained by crystallographic averaging of electron microscopy images of 2-D crystals. (A) A nonsymmetrized projection map of a crystallized PSII core dimer of the type shown in Figure 3A, consisting of CP47, CP43, the D1 and D2 proteins, cyt b559, and associated low-molecular-weight polypeptides but lacking the 33-kDa protein (E. Morris, B. Hankamer, D. Zheleva & J. Barber, unpublished results). (B) A filtered 2-D crystal projection map with twofold rotational symmetry imposed (redrawn from 75). (C) Nonsymmetrized section from the 3-D model of Ford et al (39), redrawn. (D) A nonsymmetrized filtered projection map of a 2-D crystal from Tsioitis et al (130), redrawn. In each of these four images, two prominent densities marked (*) are located in equivalent positions on either side of a central hole. The similarity in size and shape of these contoured maps suggests that all these complexes are dimeric. The scale bar measures 5 nm.
OEC. As in the case of Marr et al (75), these crystals were formed by partially delipidating granal membranes. To aid comparison with Figures 4A and 4B, the images of the crystal form lacking the extrinsic proteins are shown in Figure 4C (39). It is reported to have a unit cell ($a = 177 \text{ Å, } b = 201\text{Å, } \gamma = 91^\circ$) that is slightly larger than those of the dimeric PSII complexes shown in Figure 4A and 4B. On the basis of SDS-PAGE analysis, the authors concluded that their crystals contained both PSII core and Lhcb subunits. One of the difficulties in determining the subunit composition of a complex crystallized by the partial delipidation of granal membranes is to confirm the presence of given subunits in the crystalline fraction, when both crystalline and noncrystalline material is contained in the sample. Without direct immunolabeling of the crystals, their composition must remain unconfirmed. However, on the basis of the assumption that Lhcb proteins are associated with the crystallized complex, it was concluded that the PSII core complex must be monomeric because there was insufficient volume to accommodate a PSII core dimer and a large complement of Lhcb proteins (61). However, a comparison of the unsymmetrized images of Figures 4A, 4B, and 4C show them to be very similar. Once again, the same two prominent regions of density (*), also seen in Figures 4A and 4B, are apparent on either side of a central hole in Figure 4C. This suggests that this core complex may actually be a core dimer rather than a monomeric PSII complex and that its symmetry was mistaken as pseudosymmetry because of the relatively low resolution of the images.

Tsiotis et al (130) recently reported the projection maps of a PSII crystal (CP47, CP43, D1, D2, cyt b559, the 33-kDa extrinsic subunit, and associated low-molecular-weight polypeptides) obtained using detergent solubilized PSII core complexes (Figure 4D). This crystal was reported to have a unit cell of ($a = 162 \text{ Å, } b = 137\text{Å, } \gamma = 142^\circ$). However, if a choice of unit cell parameters is made to include all the densities shown in Figure 4D, the dimensions are actually very close to the ones reported by Hankamer et al (51) (Figure 4A) and Marr et al (75) (see Figure 4B). Furthermore, a central hole (or cavity) is visible in these complexes and two densities (*) are once again seen on either side of it, separated by about the same distance. All these points would suggest that this complex could also be a dimeric PSII core complex, although the authors concluded that the complex was a monomer, partly because of scanning transmission electron microscopy (STEM) measurements that suggested the complex had a mass of 318 kDa (130).

Recently, Nakazato et al (90) reported the crystallization of a D1-D2-cyt b559-CP47 complex, which yielded a projection map of 20-Å resolution. The projection map of this complex can be superimposed upon a PSII core monomer (Figure 3A) and comfortably fits into it (see Figure 5 and associated...
discussion). This finding confirms that the PSII complexes depicted in Figures 4A and 4B, and probably in 4C and 4D, are dimeric.

**SUBUNIT ORGANIZATION OF PSII**

This section reviews the subunit organization of PSII and its antenna system using information that has been obtained by electron crystallography, single particle image averaging, and crosslinking and other biochemical techniques. The combined data is summarized in the form of two currently favored subunit organization models (Figures 5a and 5b) because the information available is still insufficient to confirm which of these is correct.

Top (Figures 5a and 5b) and side view (Figure 5C) projection maps of the largest PSII-LHCII supercomplex structurally characterized to date are used as the framework for the two possible models of subunit organization. These contoured projection maps are very similar to those presented by Boekema et al (21) but improved in that they are the sums of larger data sets (1925 vs 500 top views, 2213 vs 80 side views) and have a higher resolution (approximately 20 Å). They also differ in that they contain the densities of two 23-kDa subunits in addition to those of the other core (CP47, CP43, D1, D2, the 33-kDa subunit, cyt b559) and antenna (Lhcb1, 2, 4, and 5) (21, 93) proteins. Both models are identical in terms of their depiction of the extrinsic and Lhcb protein components. They differ only in the attributed locations of the PSII core components, CP47, D1, D2, and cyt b559.

**Localization of D1-D2-Cyt b559-CP47 Complex and CP43**

Regions within each of the projection maps are shaded in dark, mid, and light gray. The dark-gray regions represent aligned monomeric D1-D2-cyt b559-CP47 complexes (and associated low-molecular-weight subunits) of the type reported by Dekker et al (31) and Nakazato et al (90). Together, the two dark- and two mid-gray regions (Figures 5a and 5b) depict the shape of the PSII core dimer (Figure 2b) consisting of the integral membrane protein components D1, D2, cyt b559, CP47, and CP43 and associated low-molecular-weight subunits. By elimination, it follows that the two mid-gray regions each contain CP43 (106).

**Localization of the 33-kDa Extrinsic Subunits**

Top and side view projection maps of PSII complexes (±33-kDa extrinsic subunit) were used to produce subunit difference maps (21, 30). In Figure 5 each monomeric portion of the dimer is associated with a region of density attributed to the 33-kDa subunit, and these densities overlap to form a single
Figure 5  Subunit organization models of photosystem II. This figure shows top (Figures 5a and 5b) and side view (Figure 5c) projection maps of the largest PSII-LHCII supercomplex structurally characterized to date. It provides the framework for two possible models of subunit organization (Model 1 and Model 2), based predominantly on crosslinking studies and averaged images of PSII complexes of known and differing subunit composition. Both models are identical in terms of their depiction of the extrinsic and Lhcb protein components. They differ only in the attributed locations of the PSII core components, CP47, D1, D2, and cyt b559. In Model 1, CP47 is placed between CP43 and the reaction center components. In Model 2, the RC components are placed between CP47 and CP43. The three contoured projection maps (a, b, and c) are very similar to those presented in Boekema et al (21) but are improved in that they are the sums of larger data sets (1925 vs 500 top views, 2213 vs 80 side views) and have a higher resolution (approximately 20 Å). They also differ in that they contain the densities of two 23-kDa subunits in addition to those of the other core (CP47, CP43, D1, D2, the 33-kDa subunit, cyt b559) and the antenna (Lhcb1, Lhcb2, Lhcb4, and Lhcb5) (21, 93) proteins. Superimposed upon the central regions of each of these images are areas shaded in dark, mid, and light gray. Each of the two dark-gray regions corresponds to a monomeric CP47-RC complex similar to those reported by Dekker et al (31) and Nakazato et al (90). Together the two dark- and two mid-gray regions depict the centrally located CP47-Cp43-RC core dimer (21). The four densities shaded in light gray in the top view projections (a and b) correspond to two 33-kDa and two 23-kDa subunits (21; see text). In the side view orientation (c) the two lumenally exposed 33-kDa subunits partially overlap. Positioned on either side of them are two densities attributed to the 23-kDa extrinsic proteins. The electron density map of an LHCII trimer (133) has been superimposed upon the two tips of the PSII-LHCII complex (a and b). It is suggested that these trimers consist of Lhcb1 and Lhcb2 and that they are linked to the centrally located PSII core via two monomeric Lhcb proteins (Lhcb4 and Lhcb5). The Lhcb4 and Lhcb5 proteins are depicted using the projection maps of monomeric LHCII components (133).
central protrusion in the side view (Figure 5c). The model implies that the 33-kDa subunit:PSII core monomer stoichiometry is 1:1. Some reports indicate the presence of two copies of the 33-kDa subunit per reaction center (72, 135, 136). However, the dimensions of the 33-kDa subunit protrusions shown in Figure 5 are consistent with single copies of the protein monomer (64).

**Localization of the 23-kDa Extrinsic Subunits**

When isolated in the presence of glycine betaine, PSII-LHCII supercomplexes additionally bind the 23-kDa subunit (EJ Boekema, J Nield, B Hankamer & J Barber, unpublished data). Preliminary difference mapping experiments suggested that the two lumenal protrusions in the side view projection map, located on either side of the 33-kDa components (Figure 5c), each contain a 23-kDa subunit. Similarly, positions in the top views, also identified by difference mapping, are shown in Figures 5a and 5b. These proposed positions are consistent with analysis of crystalline PSII arrays containing the 23-kDa subunit (76). Freeze-etching studies of the lumenal surface of the grana regions of thylakoid membranes (ESs surfaces) also show four lumenal protrusions (112) that could correspond to the four densities shown in Models 1 and 2 (Figures 5a and 5b; see section “Heterogeneity of PSII In Vivo”). If the PSII complexes studied, before and after the removal of extrinsic proteins by Ford et al (39), are interpreted as dimers, the positions attributed to the extrinsic subunits would be consistent with models shown in Figure 5.

**Organization of the Membrane-Embedded PSII Core Components**

Topological information is available on the organization of the subunits within the PSII reaction center complex. In its isolated form, the PSII reaction center consists of the α and β subunits of cyt b559 and PsbI (17, 91, 134). Crosslinking studies have shown that the α and β subunits of cyt b559 are closely associated with the D1 and D2 proteins (14, 88), whereas the stromally exposed N-terminus of PsbI is suggested to be in close contact with the D2 protein (128).

Sequence homology between the D1 and D2 subunits and the L and M subunits of purple bacteria suggest that the D1/D2 and L/M heterodimers are likely to have similar dimensions (approximately 70 × 30 Å) in the membrane plane (29). Cryo-electron crystallography studies have shown that the monomeric PSII RC-CP47 complex (dark-gray region) has dimensions of 81 × 75 Å (90), indicating that the estimated PSII RC size is reasonable.

Isolated RC-CP47 complexes (D1-D2-cyt b559-CP47) have also been subjected to crosslinking experiments, and the results indicate that CP47 is more
closely associated with D2 than with the other reaction center components (88). It is for this reason that CP47 is placed close to the D2 protein in both subunit organization models (Figures 5a and 5b).

There are currently no crosslinking data that help to position CP43 with respect to the D1-D2-cyt b559-CP47 complex. Consequently, at least two models can be proposed. In the first, CP47 is positioned between the reaction center components and CP43 (Model 1, Figure 5a). In the second, the reaction center components are located between CP43 and CP47 (Model 2, Figure 5b).

Recently, PSII core dimer crystals were labeled with Fab antibody fragments, specific to the C-termini of the D1 protein and cyt b559 (75). These studies suggested that the D1 and cyt b559 subunits were located farthest from CP43 but within the dark-gray region as shown in Model 1 (Figure 5a). However, crosslinking results of Seidler (114) detected interactions between the 23-kDa subunit and cyt b559, favoring Model 2 depicted in Figure 5b. The positions of cyt b559 as shown in Models 1 and 2 might allow the formation of an α-cyt b559 homodimer in crosslinking experiments carried out using D1-D2-cyt b559-CP47 complexes, assuming these preparations contained dimeric complexes (88).

Crosslinking and other biochemical studies have shown that CP43 (63) and cyt b559 (124, 125) are also closely associated with the 33-kDa subunit. Both cyt b559 and PsbI are estimated to be within 11 Å of the 33-kDa extrinsic polypeptide (37). Model 2 (Figure 5b) is more consistent with this combined data because it is difficult to see how CP43 and cyt b559 could interact simultaneously with the 33-kDa subunit in Model 1 (Figure 5a). Perhaps the most compelling evidence for the relative positioning of CP43 and CP47 on either side of the D1/D2 subunits (Model 2) comes from the comparison with the reaction center structure of PSI (42). This complex is composed of a heterodimer of PsaA and PsaB proteins each having 11 transmembrane helices. In each case, 5 of these helices are similarly arranged to the 5 helices of the L and M subunits of the purple bacterial reaction center (and presumably with the D1 and D2 proteins). The remaining 6 helices in each PSI subunit are positioned farther from the pseudo twofold axis that relates the two subunits in the heterodimer. Furthermore, the two sets of peripheral helices show sequence homology with those of CP47 and CP43. By analogy, therefore, it would seem reasonable to place CP43 and CP47 on either side of the D1 and D2 proteins.

Recently 3-D maps obtained from dimeric RC-CP47-CP43 complexes showed that the binding region for the 33-kDa subunit protrudes into the lumen (E Morris, B Hankamer, D Zheleva & J Barber, unpublished data). The D1 and D2 proteins, the α and β subunits of cyt b559 and PsbI, are all very
hydrophobic proteins with small lumenally exposed loops. In contrast, the
lumenal loops E of CP47 and CP43 are large (22, 41, 45), suggesting that they
could form part of this extrinsic region. This conclusion agrees well with the
finding that loop E of CP47 is in close contact with the 33-kDa extrinsic
polypeptide (23, 24, 41, 50, 94, 102). However, this information does not
provide a sufficiently clear distinction between Models 1 and 2. This is be-
cause in Model 1, the 33-kDa subunit is positioned directly above CP47,
allowing direct contact with loop E of the latter. However, in Model 2, CP47 is
adjacent rather than directly under the 33-kDa subunit, and it is quite conceiv-
able that loop E could fold over the lumenal surface of the centrally located
D1/D2 heterodimer and so come into contact with the 33-kDa subunit.

Localization of the Lhcb Proteins

Western blot analysis of isolated PSII-LHCII supercomplexes showed them to
be enriched in Lhcb1, Lhcb2, Lhcb4 (CP29), and Lhcb5 (CP26), but depleted
of Lhcb3 and Lhcb6 (CP24) (93). Crosslinking studies have shown that CP47
and CP43 are both in close contact with CP29 (Lhcb4) and that CP43 is also in
close contact with CP26 (Lhcb5) (26, 104). By superimposing a high-resolu-
tion electron density map of the Lhcb1/Lhcb2 heterotrimer (133) upon the
top-view projection maps shown in Figure 5a and 5b, the LHCII complex can
be seen to fit snugly into the two tips of the PSII-LHCII supercomplex.
Furthermore, because Lhcb4 and 5 are similar in mass and have strong se-
quence homology with the LHCII proteins, two monomeric electron density
maps based on the data of Wang & Kühlbrandt (133) were positioned in the
PSII-LHCII supercomplex (in both Models 1 and 2) between the PSII core and
the LHCII heterotrimer. There appears to be insufficient space to fit any
additional monomeric Lhcb proteins. This conclusion is in agreement with
pigment analyses of the isolated PSII-LHCII supercomplex, which showed it
to be associated with 156 Chla and 46 Chlb molecules (see Figure 2d and 51,
52, 93). It was suggested that Lhcb6 (CP24) might also be present in these
isolated PSII-LHCII supercomplexes (21), but this was a precautionary con-
clusion based on the detection of low levels of the protein present in early and
less pure preparations. This conclusion has been amended in the light of the
Western blot data presented by Hankamer et al (52) and Nield et al (93). The
positions of Lhcb4 and 5 between the LHCII trimer and the PSII core is in
agreement not only with the available crosslinking data (26, 104) but with the
hypothesis that Lhcb4, 5, and 6 are involved in the regulation of the rate of
excitation energy transfer from the outer Lhcb (Lhcb1, 2, and 3) proteins to the
reaction center via CP47 and CP43 (see 65). However, the depletion of Lhcb3
and 6 in these complexes with respect to granal membranes must be explained.
One explanation is that the isolated PSII-LHCII supercomplex characterized originates from the population of ESs particles equivalent to those observed as arrays in freeze-etching studies of grana regions, and that Lhcb3 and Lhcb6 are associated predominantly with the nonarrayed ESs particles in the grana (see section on “Heterogeneity of PSII In Vivo”). This hypothesis is consistent with the isolation of a Lhcb6-Lhcb4-LHCII supramolecular complex (18). It is also consistent with the finding that PSIIα complexes in the grana consist of more than one subpopulation differing in antenna size (4, 5). Furthermore, the large distances between the nonarrayed ESs particles means that they may have larger antenna systems than their arrayed counterparts and could additionally bind Lhcb3 and Lhcb6 (see section on “Heterogeneity of PSII In Vivo”).

The low-resolution subunit organizations presented in Figure 5, though incomplete, give two models that it is hoped will aid more detailed determinations of subunit organization and PSII structure in the future.

CONCLUDING REMARKS

The PSII-LHCII supercomplex is in projection possibly one of the largest integral membrane protein complexes in nature. Despite its large size, this discrete unit has escaped attention until recently. One of the reasons may be its thickness, which is only 60 Å at the outer ends. In fact, only the central parts, which are 90 Å in height, substantially protrude from the membrane and contribute to the appearance of the approximately 200 × 200 Å “particles,” already seen long ago by freeze-fracture/freeze-etching techniques. No doubt, the PSII-LHCII supercomplex will provide a further basis for structural work, although its stability after isolation will possibly prevent the growing of highly ordered crystals. It is, however, highly suited for further single particle analysis using cryoelectron microscopy to produce a more detailed 3-D map suitable for the insertion of high-resolution data. Such data will almost certainly be obtained by crystallography (2-D and 3-D) of PSII complexes with smaller numbers of subunits or of individual subunits. In this way, the complete structure of PSII at atomic resolution will be built. Whether the high-resolution data will be obtained from X-ray or electron crystallography is uncertain, but with the successful outcome of the LHCII structure, the electron microscopy approach has yielded the most information to date for PSII. The instability of this photosystem, particularly when illuminated, presents a major hurdle that is unique to this complex. The authors are confident, however, that this hurdle will be surmounted in the not-too-distant future and that PSII, like the other major complexes of the photosynthetic electron transport chain, will have its structure determined at atomic resolution.
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