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Published in:
Biochimica et Biophysica Acta

DOI:
10.1016/j.bbabio.2013.07.012

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Document Version
Publisher's PDF, also known as Version of record

Publication date:
2014

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):

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Download date: 18-03-2019
Light-harvesting complex II (LHCII) and its supramolecular organization in *Chlamydomonas reinhardtii*

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**Abstract**

LHCII is the most abundant membrane protein on earth. It participates in the first steps of photosynthesis by harvesting sunlight and transferring excitation energy to the core complex. Here we have analyzed the LHCII complex of the green alga *Chlamydomonas reinhardtii* and its association with the core of Photosystem II (PSII) to form multiprotein complexes. Several PSII supercomplexes with different antenna sizes have been purified, the largest of which contains three LHCII trimers (named S, M and N) per monomeric core. A projection map at a 13 Å resolution was obtained allowing the reconstruction of the 3D structure of the supercomplex. The position and orientation of the S trimer are the same as in plants; trimer M is rotated by 45° and the additional trimer (named here as LHCII-N), which is taking the position occupied in plants by CP24, is directly associated with the core. The analysis of supercomplexes with different antenna sizes suggests that LhcbM1, LhcbM2/7 and LhcbM3 are the major components of the trimers in the PSII supercomplex, while LhcbM5 is part of the “extra” LHCII pool not directly associated with the supercomplex. It is also shown that *Chlamydomonas* LHCII has a slightly lower Chlorophyll a/b ratio than the complex from plants and a blue shifted absorption spectrum. Finally the data indicate that there are at least six LHCII trimers per dimeric core in the thylakoid membranes, meaning that the antenna size of PSII of *C. reinhardtii* is larger than that of plants.

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**1. Introduction**

Photosystem II (PSII) is a multisubunit pigment–protein complex located in the thylakoid membrane of cyanobacteria, algae and higher plants. It captures and converts light into chemical energy, which is used to oxidize water and reduce plastoquinone in the light reactions. The structure and function of PSII has been extensively studied in cyanobacteria and higher plants. It consists of a core, which performs the light harvesting capacity of the core [5]. The PSII core complex is highly conserved in all organisms performing oxygenic photosynthesis [6–8]. The core contains the reaction center (RC, composed of D1 and D2 and cytochrome b559) that generates the redox potential required to drive water splitting [9] and the Chlorophyll (Chl) a-binding antenna complexes CP43 and CP47 [5,10]. On the luminal side of the core, several extrinsic proteins (PsbO, PsbQ, PsbP, PsbR in plants and green algae) form the oxygen-evolving complex (OEC), which supports water oxidation [9,11]. The genes encoding for the antennas of PSII of plants and green algae are members of the light-harvesting complex (Lhcs) multigenic family, which also includes the antenna complexes of PSI [12]. These proteins show structural homology [13,14]: each Lhc polypeptide has three transmembrane α-helices and coordinates Chls a, Chls b, and different carotenoid molecules. In higher plant LHCII, the most abundant light-harvesting complex is composed of 3 gene products (Lhcb1–3) organized as heterotrimers. Each LHCII apoprotien binds eight Chls a, six Chls b and four carotenoids [13,15]. The other Chlorophyll a/b-binding proteins, Lhcb4, Lhcb5, and Lhcb6, also known as CP29, CP26, and CP24, exist as monomers and electron transport chain, and the outer antenna, which increases the light-harvesting capacity of the core [5].

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http://dx.doi.org/10.1016/j.bbabio.2013.07.012
have different pigment compositions [16–18]. The major function of the outer antenna system is to capture light energy and transfer excitation energy to the RC [2]. However, in high light, when the absorbed energy exceeds the photosynthetic capacity and can be damaging for the system, it ensures photoprotection by the dissipation of excess energy as heat (via the process called non-photochemical quenching) to avoid formation of harmful radicals, e.g. [19,20].

In Chlamydomonas reinhardtii (C.r.) nine genes (LhcbM1–M9) encode LHCII proteins. Two minor antennas, CP26 and CP29, are present while CP24 is not found in the genome of this alga [21–23]. The LhcbM gene products have been divided into four groups based on their sequence similarity: Type I (LhcbM3, LhcbM4, LhcbM6, LhcbM8, and LhcbM9), Type II (LhcbM5), Type III (LhcbM2 and LhcbM7), and Type IV (LhcbM1) [22,24]. Within their types, these proteins share a very high homology, i.e. sequence analysis of the Type I or Type III proteins shows up to 99% identity. Instead, the average homology between the types is lower: sequence identity between LhcbM6 (Type I) and LhcbM5 (Type II), LhcbM2 (Type III), LhcbM1 (Type IV) is 80%, 77%, and 74%, respectively [22]. Functional studies of these complexes are available only for LhcbM2/7, LhcbM5 and LhcbM1 [25–27] and suggest that the first two are involved in state transitions, while LhcbM1 is important for NPQ.

The structures of several PSII components, including the cyanobacteria core [6,28], the plant LHCII trimer [13,15] and CP29 from plants [16] have been resolved at intermediate/high resolution. Information about the organization of the PSII–LHCII supercomplexes has been obtained by electron microscopy and single particle analysis [3,7,29–33] and at present the best map has a maximal resolution of 12 Å [3]. The LHCII trimers associated with the core can be distinguished in three types, based on their position in the PSII supercomplex and their strong (S), moderate (M) or loose (L) association with the core (C) [29]. The C2-S2-M2 supercomplex is the largest PSII–LHCII observed in A. thaliana and it is composed of a dimeric core (C2), two LHCII-S trimers, interacting directly with the core and with CP29, CP26, and two M-trimers, which are associated with the core via CP29 and CP24 [3,5,29,30,32]. The position of the loosely bound trimer LHCII-L is still unclear and complexes containing one L trimer have been observed only in spinach [29].

In C. reinhardtii a complex with C2-S2 organization was reported [7,34], as expected due to the absence of CP24, which in higher plants is essential for connecting trimer M to the core [35,36]. However, recently, larger PSII supercomplexes with up to 6 trimers per dimeric core were observed [33], showing that the absence of CP24 does not influence the association of trimer M with the supercomplex, and that an additional trimer is associated with the supercomplex on the side that in plants is occupied by CP24.

In order to obtain further information about the antenna complexes of PSII and their supramolecular organization in C. reinhardtii we have isolated different PSII sub- and supercomplexes. These complexes were characterized by combining biochemistry, spectroscopy and single particle electron microscopy. The analysis reveals the structural organization of PSII–LHCII and addresses the position and the composition of individual LHCI trimers and their role in the assembly and functioning of the supercomplex.

2. Material and methods

2.1. Strain and growth conditions

Cells of C. reinhardtii strain (JVD-1B[pGG1]) in which a hexahistidine tag has been added at the N-terminus of PorA core subunit of PSI [37] were grown in liquid Tris–acetate–phosphate (TAP) medium [38] at room temperature (25 °C) on an incubator shaker (Minitrон, INFORS HT) at 170 rpm under a continuous illumination flux of 20 μmol photons PAR m⁻² s⁻¹. In those conditions the cells were in state 1.

2.2. Thylakoid preparations

Cells were harvested by centrifugation (3500 rpm, 5 min, 4 °C) at mid-logarithmic phase (OD₇₅₀ nm ≈ 0.7). Cr. thylakoid membranes were prepared under dim light in a cold room as described by Fischer et al. [39] with modification from Drop et al. [40]. PSI enriched membranes (BBY) from A. thaliana were prepared according to Berthold et al. [41] with the modifications reported by Caffarri et al. [3].

2.3. Isolation of PSII light-harvesting antenna complexes

PSII supercomplexes isolation was modified from Drop et al. [40]. Thylakoids were pelleted, unstacked with 5 mM EDTA and washed with 10 mM Hepes (pH 7.5). Membranes were then resuspended in 20 mM Hepes (pH 7.5), 0.15 M NaCl and solubilized at a final chlorophyll concentration of 0.5 mg/ml by adding an equal volume of 0.6% α-dodecyl maltoside (α-DM). Unsolubilized material was eliminated by centrifugation (12,000 rpm for 10 min at 4 °C).

To remove His-tagged PSI complexes from the preparation, the supernatant was loaded onto a HisTrap HP Column (GE Healthcare) equilibrated with 20 mM Hepes (pH 7.5), 0.15 M NaCl and 0.03% α-DM. PSI-depleted, fraction (flow through) was loaded on a sucrose density gradient (prepared by freezing and thawing 0.5 M sucrose, 20 mM Hepes (pH 7.5), and 0.03% α-DM buffer layered over 1 ml of 2 M sucrose). PSII complexes were separated by ultracentrifugation (41,000 rpm, 14 h, 4 °C). The green bands visible on the sucrose gradient were harvested with a syringe.

The purification of PSII supercomplexes from A.t. was performed in the same conditions used for the purification of the Cr. PSII supercomplexes, with the difference that upon solubilization the membranes (BBY) were directly loaded on the sucrose gradient.

2.4. Gel electrophoresis

Proteins were analyzed by a SDS-6 M urea PAGE with Tris–sulphate buffer system prepared as discussed in Bassi [42] at a 14% acrylamide concentration. The amounts of sample loaded into each well were: 3 μg (in Chls) for thylakoids; 2.5 μg for PSI supercomplexes; and 1.5 μg for Lhcb fractions. The Coomassie stained gel was imaged with ImageQuant LAS-4000 (GE Healthcare).

2.5. Mass spectrometry analysis

2.5.1. In-gel tryptic digestion

For mass spectrometry–based protein identification, the SDS-PAGE bands were excised from the gel and treated with 10 mM DTT followed by 55 mM iodoacetamide in 50 mM NH₄HCO₃, to reduce and alkylate cysteine residues, and subsequently dehydrated by incubation for 5 min in 100% acetonitrile. The gel slices were rehydrated in 10 μl trypsin solution (Trypsin Gold, mass spectrometry grade, Promega10 ng/μl in 25 mM NH₄HCO₃), and incubated for 2 h at 37 °C. Subsequently 10 μl of 25 mM NH₄HCO₃ was added to prevent drying and the incubation was prolonged overnight at 37 °C. The tryptic peptides were recovered by three subsequent extractions with 50 μl of 35%, 50% and 70% acetonitrile in 0.1% TFA. The extracted peptides were pooled and concentrated under vacuum.

To determine protein composition of PSII supercomplexes, sucrose gradient fractions were first loaded on 10% SDS-PAGE and run about 1 cm through the resolving gel. This procedure was applied to clean up samples from detergent. The whole gel bands were then cut in three slices and tryptic digestion was performed as described above.

2.5.2. Liquid chromatography–mass spectrometry (LC–MS)

Fractions of the peptide mixtures from in-gel trypsin digestions were diluted in 5% formic acid, passed through a pre-column (EASY-Column C18, 100 μm × 20 mm, 5 μm particle size, Thermo Scientific, Bremen,
Germany) and separated on a capillary column (C18 PepMap 300, 75 μm × 100 mm, 3-μm particle size, Thermo, Thermo Scientific, Bremen, Germany) mounted on a Proxeon Easy-nLCII system (Thermo Scientific, Bremen, Germany). Solutions of 0.1% formic acid in water and a 0.1% formic acid in 100% acetonirole were used as the mobile phases. A gradient from 4 to 35% acetonirole was performed for 120 min at a flow rate of 300 nl/min. Eluted peptides were analyzed using a Linear ion Trap–Orbitrap hybrid mass spectrometer (LTQ–Orbitrap XL, Thermo Scientific). MS scans were acquired in the Orbitrap, in the range from 350 to 1800 m/z, with a resolution of 60,000 (FWHM). The 7 most intense ions per scan were submitted to MS/MS fragmentation (35% Normalized Collision EnergyTM) and detected in the linear ion trap.

2.5.3. Protein identification
The MS raw data were analyzed with Mascot (version 2.1, Matrix Science, London, UK) using the Proteome Discoverer 1.3 analysis platform (Thermo Scientific) and searched against the C. protoeke. Peptide toler-ance was set to 10 ppm and 2.0 Da for intact peptides and fragment ions respectively, using semi-trypsin as protease specificity and allowing for up to 2 missed cleavages. Oxidation of methionine residues, deamidation of asparagine and glutamine, and carbamidomethylation of cysteines were specified as variable modifications. The MS/MS based peptide and protein identifications were further validated with the program Scaffold (version Scaffold 4.0, Proteome Software Inc., Portland, OR). Protein identifications based on at least 2 unique peptides identified by MS/MS, each with a confidence of identification probability higher than 95%, were accepted.

2.6. Pigment composition
The chlorophyll concentrations of thylakoid preparations were calculated in 80% (v/v) acetone, according to Porra et al. [43]. The pigment composition of the complexes was analyzed by fitting the spectrum of the 80% acetone extracted pigments with the spectra of the individual pigments in acetone and by HPLC, as described previously [17]. As shown by Angeler and Schagerl [44], lorioxanthin was eluted just after neoxanthin, but in our experimental setup the separa-tion of these two was not possible: both carotenoids resulted in single peak. The data are the results of at least four different preparations in two replicates.

2.7. Spectroscopic analysis
Room temperature absorption spectra were recorded with a Cary 4000 spectrophotometer (Varian). The fluorescence emission spectra were recorded at 5 °C at low temper-ature (77 K) using a Fluorolog 3.22 spectrophluorometer (Jobin Yvon-Spx). For 77 K measurements a home built liquid nitrogen cooled de-vise was used. The excitation wavelengths were 440 nm, 475 nm and 500 nm and emission was detected in the 600–800 nm range. Excitation and emission slits bandwidth were set to 3 nm. All fluorescence spectra were measured at OD 0.05 at the maximum of the Qy absorption. Room temperature measurements were performed in 0.5 M sucrose, 20 mM Hepes (pH 7.5), and 0.03% α-DM buffer. For low-temperature measure-ments, samples were in 66.7% glycerol (w/v), 20 mM Hepes (pH 7.5), and 0.03% α-DM buffer.

2.8. Electrochromic shift (ECS)
PSI/PSII ratio of Chlamydomonas cells was measured with a Joliot-type spectrophotometer [45] (Bio-Logic SAS JTS-10) as described previ-ously [46,47]. A PSI/PSII ratio of 0.97 ± 0.2 was obtained.

2.9. Electron microscopy and single particle analysis
Samples were negatively stained using the droplet method with 2% uranyl acetate on glow discharged carbon-coated copper grids. Electron microscopy was performed on a Philips CM120 electron microscope equipped with a LaB6 filament operating at 120 kV. Images were recorded with a Gatan 4000 SP 4K slow-scan CCD camera at 130,000× magnification at a pixel size (after binning the images) of 2.25 Å at the specimen level with GRACE software for semi-automated specimen selection and data acquisition [48]. Single particle analysis was performed using GRIP software including multi-reference and non-reference alignments, multivariate statistical analysis, and classification, as in Boekema et al. [29].

To determine the angle of rotation of the trimers: 1) in GRIP, all three trimers were selected at their midpoints, boxed out of the com-plex projections and aligned to their 3-fold rotationally averaged projections before aligning with S-trimer as reference, and 2), in comparison, PowerPoint (Microsoft) was used to draw a triangle as a reference around the reference trimer. Copies of this triangle were then rotated to align with the other trimers and the angle by which the triangle was rotated was determined in the program.

To model the supramolecular organization of the supercomplex, the available crystal structures of the cyanobacterial PSII core [6] (3ARC), LHClI trimer [13] (1RWT) and CP29 [16] (3PL9) were used. PyMOL was used to construct an aligned complex model.

3. Results

3.1. Isolation of PSI supercomplexes
To purify the PSI–LHClI complexes of C. reinhardtii the procedure de-scribed before for higher plants [3] was modified. Since it is not possible to prepare grana membranes from C. reinhardtii, as the presence of PSI interferes with the purification of PSII supercomplexes, we have used a strain carrying a hexa-histidine tag at the N-terminus of the core subunit PsaA [37,40]. The thylakoid membranes purified from this strain were mildly solubilized, to keep the large PSI supercomplexes intact, and loaded on a Ni-Sepharose column to eliminate PSI. To obtain homoge-neous preparations of PSI sub- and super-complexes with different an-tenna sizes, the flowthrough fraction was subjected to sucrose density gradient ultracentrifugation. The gradient separation resulted in one yellow band (B1, containing free pigments), eight green bands (B2–B9), containing protein–chlorophyll complexes, and a fraction of partially unsolubilized material laying on top of the 2 M sucrose solution (B10) (Fig. 1A). The band pattern was very similar to that obtained from the solubilization of the grana membrane of A. thaliana, characterized before [3], suggesting the presence of PSI sub- and supercomplexes. As expected, the band corresponding to the At. CP24/CP29/LHClI-M complex (“band 4”) was absent in C. reinhardtii, due to the lack of CP24.

The polypeptide composition of fractions B2–B9 was analyzed by SDS-PAGE (Fig. 1B), showing the presence of PSIi components in all frac-tions. The exact protein composition of the most prominent gel bands was confirmed by mass spectrometry. The results of the analysis are summarized in Fig. 1.

Fraction B2, which mobility in the gradient corresponds to monomer-Lhcb, contains CP26 and CP29, and LHClI (mainly LhcbM1, LhcbM2/M7 and LhcbM3), while fraction B3 contains the trimeric complexes. It was suggested that the trimeric form of C. reinhardtii is less stable than that of plants and may partly dissociate into monomers upon treatment with detergents [49]. This can explain the high content of LHClI in fraction B2 and the higher abundance of monomeric complexes (B2) compared to trimeric (B3) (Fig. 1A, B). However, it should be mentioned that when solubilized thylakoids were directly loaded on the sucrose grade-intit, the trimeric fraction was far more abundant than that of monomers (data not shown) indicating that, although less stable than in plants,
almost all of the LhcbMs are present as trimers in the membrane. The LhcbM protein composition of B2 and B3 was very similar with the exception of LhcbM5, which was only present in B3, suggesting that trimers containing LhcbM5 are relatively stable. Interestingly, CP26, which in plants exists as a monomer, could be observed mainly in fraction B3. There are two possible explanations: CP26 forms homo or hetero-trimers in C. reinhardtii, or it is strongly associated with a LHCII trimer.

The polypeptide composition of fraction B4 revealed the presence of PSII core subunits but also of LHCII and CP29. It is clear that the antenna present in this band cannot be associated with the core because the molecular weight of this band corresponds to that of PSII core monomer. This suggests that the monomeric core co-migrates with an oligomer of antenna complexes. In A. thaliana a complex composed of CP24/CP29/LHCII-M could be purified. In C. reinhardtii CP24 is absent but its position in the supercomplex is occupied by a LHCII trimer (C2S) [33] (see below). It might thus be possible that a complex composed of CP29 and 2 LHCII trimers is stable enough to survive solubilization. The molecular weight of this complex would indeed be comparable to that of the monomeric core. In contrast to B4, in fraction B5 the amount of CP26 increased significantly. PSII subcomplex in B5 might consist of PSII monomeric core/LHCII-S/CP26 (CS/CP26), as was reported before for higher plants [3]. The SDS-PAGE of fractions B4 and B6 showed the presence of LHCII antenna indicating that these fractions are contaminated with PSI sub- and super-complexes, which are indeed expected to migrate at these positions [40]. The presence of PSI contamination is confirmed by the spectra which are red-shifted compared to the spectra of the other fractions (data not shown). In the upper part of the gel the bands of the α and β subunits of the ATPase are also visible. Due to their heterogeneous content bands B4–B6 were thus not analyzed further.

The protein compositions of B7, B8 and B9, which correspond to PSII supercomplexes, were very similar, while the ratio between PSII core subunits (CP47 and/or CP43) and Lhcb subunits decreased when going from B7 to B9, in agreement with an increased antenna size.

### 3.2. Electron microscopy and single-particle analysis

To determine the structural organization of PSII–LHCII, electron microscopy and single particle analysis were performed. The analysis of about 50,000 projections from fractions B7, B8 and B9 yielded six types of supercomplexes (Fig. 2A, B), in which the dimeric PSII core complex (C2) was associated with a variable number of LHCII trimers. Two LHCII trimers occupied positions equivalent to those of the S- and M-trimers in the C2S2M2 supercomplex of A. thaliana [3], while the third trimer was located in the position that in A. thaliana is occupied by CP24, as observed before [33]. Because this trimer was directly associated with the PSII core, without involvement of any monomeric antenna, it is named trimer-N (naked). We prefer this notation to “trimer L” for two reasons: (i) trimer L has been observed only in spinach where it occupies a different position; and (ii) “L” is short for “loosely bound” [29], which is not completely appropriate for this trimer because its association to the supercomplex survives purification.

Fractions B7 and B8 contain mainly PSII dimeric core complexes with one S trimer (C2S) and two S trimers (C2S2) respectively. Fraction B9 contains three types of PSII–LHCII particles, which beside LHCII-S and LHCII-M also include LHCII-N trimers: C2SMN, C2S2MN and the largest complex composed of 6 LHCII trimers surrounding the dimeric, C2S2M2N2. If we consider all the particles from the data set, and not only those added in the final figures because of their good image quality, the ratio of the three largest particles in B9 was about 4.5 (C2SMN):4 (C2S2MN):1 (C2S2M2N2). This means that in the lower band the average number of trimers per dimeric core complex is about 3.7.

The projection of the C2S2MN supercomplex was obtained at a 13 Å resolution. This resolution allows to identify the major structural features of the individual complexes and to use them to fit the X-ray high-resolution structures of LHCII [13], CP29 [16] and PSII core [6] into the projection map of the particle. The obtained model of the supramolecular organization of the PSII supercomplex is shown in Figs. 3 and 4.

The projection maps of the C2S2 supercomplexes of C. reinhardtii and A. thaliana are very similar, suggesting that CP26 and CP29 occupy the same positions, with CP26 located close to CP43 and CP29 next to CP47 [3,29,32]. Trimer S is also located in the same position and it

![Fig. 1. Purification of PSII sub- and supercomplexes. Panel A: fractionation of PSII sub/super-complexes by sucrose density gradient. Panel B: SDS-PAGE analysis of fractions B2–B9 from sucrose gradient. Proteins identified by mass spectrometry are indicated.](Image)
has the same orientation with respect to the core in the two supercomplexes. This is not the case for trimer M, which in *C. reinhardtii* is rotated to 45° compared to its orientation in *A. thaliana*. N-trimer is associated with the core, CP29 and trimer M and it is rotated to 25° with reference to trimer S (Fig. 4B). These values are within an accuracy of 5–10° because the trimer features are not precisely outlined by the uranyl acetate negative stain. This contrasting agent does not penetrate much inside the membrane, where the bulk of the protein is located.

### 3.3. Protein composition of the supercomplexes

To get information about the protein composition of the PSII supercomplexes, fractions B7–B9 were analyzed by mass spectrometry. A shotgun proteomic approach revealed the presence of all major core subunits (D1, D2, CP43, CP47) as well as of the oxygen evolving complex subunits (PsbP, PsbO, PsbR) and of some of the small core subunits (PsbE, PsbF, PsbH, PsbW) (Table S1). The other core subunits are likely to be present in PSII supercomplexes, but unidentifiable by tryptic digestion, because the resulting peptides would be too short and would not be detectable in our experimental setup. The analysis also showed the presence of CP26 (*Lhcb5*), CP29 (*Lhcb4*) and of several *LhcbM* gene products. The sequences of *LhcbM* proteins are highly homologous, and the analysis allows to identify with 100% probability (at least 2 exclusive peptides with >95% probability) *LhcbM1, LhcbM2, LhcbM7* are practically identical and it is not possible to discriminate between the two), *LhcbM3* and *LhcbM5*, while only one peptide was identified for *LhcbM8* and *LhcbM9*. However, several peptides that are common to *LhcbM4* and *M6* were also detected, indicating that at least one of these two proteins is present in the preparation.

To determine the relative abundance of each subunit in the different fractions, spectral count normalization was applied, meaning that the spectra of each subunit (Table S1) were counted and were normalized to the spectra of D1 and D2 in the same fraction to compensate for differences in sample loading [50]. The PSII core subunit content of B7–B9 is presented in Fig. 5 normalized to the abundance of the same subunit in B9 (Table S1). The data indicate that the amount of the major core subunits – D1, D2, CP43 and CP47 – remained almost unchanged, supporting the reliability of the method.

*Lhcb4* and *Lhcb5* were present in the same amount in B8 and B9 while their content was reduced in B7, in agreement with the presence...
of C2S complexes in this fraction. Because of the high sequence identity between the LhcbM proteins, it was not possible to use spectral counts to determine their relative amount in the three samples. However, the number of unique spectra matched to the different LhcbM types is very different and implies that LhcbM2, LhcbM1 and LhcbM3 are the most abundant LhcbM proteins in this alga.

3.4. Pigment composition

The pigment composition of fractions B2, B3 and B7–B9, was analyzed together with the composition of C. reinhardtii cells. The results are reported in Table 1.

A Chl a/b ratio of 1.29 and 1.28 was obtained for the fractions of monomeric (B2) and trimeric (B3) antenna complexes, similar to what was previously observed [51] slightly lower than the value of plant LHCII (1.33). The trimer coordinates four carotenoids per 14 Chls as it is the case in plants. The carotenoid composition of C. reinhardtii is considered to be similar to that of higher plants [52] with the addition of loroxanthin, which derives from lutein by hydroxylation of the methyl group at C9 of the polyene chain and is selectively bound to LHCII [53,54]. Indeed in the trimeric fraction we observed the presence of loroxanthin and a lower content of lutein as compared to higher plant LHCII [52]. Although it was not possible to fully separate neoxanthin from loroxanthin, the fact that all LhcbMs contain the tyrosine responsible for the selectivity of the N1 binding site for neoxanthin [55], suggests the binding of 1 neoxanthin per complex and thus of around 0.7 loroxanthin per monomer. Loroxanthin is probably substituting lutein in one of the two internal binding sites (L1 and L2).

3.5. Spectroscopic characterization

The absorption spectra of fractions B2–B9 were measured at room temperature and are presented in Fig. 6. The spectra of fractions B2 and B3 (Fig. 6A) show maxima at 672 nm and at 670 nm respectively. Interestingly the absorption maximum of the Cr. LHCII trimer is 4 nm blue shifted compared to that of the complex of higher plants (Fig. 6B) and it shows relatively more intense absorption around 650–653 nm, in agreement with the lower Chl a/b ratio.

The absorption spectra of fractions B7–B9 (Fig. 6C) showed maxima at 675 nm. A relative increase in intensity of the absorption in the Chl b
region (630–660 nm) was observed going from B7 to B9 (Fig. 6C), in agreement with the increased antenna size of the supercomplexes.

To check the integrity of PSII supercomplexes, fluorescence spectra from fractions B7–B9 were measured at 5 °C (Fig. 7A). The maximum emission for all fractions was at 678 nm. The perfect overlapping of the emission spectra after excitation at 440 nm, 475 nm and 500 nm, emission for all fractions was at 678 nm. The perfect overlapping of the spectrum of the supercomplexes is clearly red-shifted as compared to that of LHCII trimer, confirming that there is energy transfer between LHCII and the core.

4. Discussion

4.1. In the C. reinhardtii PSII supercomplex LHCII-N trimer substitutes CP24 and stabilizes the binding of trimer M (and the other way around)

Early work comparing a PSII supercomplex isolated from C. reinhardtii with the C2S2 supercomplex of spinach showed high similarity in shape and size [7]. This, together with the fact that CP24, which is essential for the binding of trimer M in plants [35,36], is absent in C. reinhardtii led to the conclusion that the PSII supercomplex of this alga is organized as C2S2. Recently, larger complexes were observed containing a novel trimer (here called N) in addition to trimers S and M [33]. In our study, the EM analysis of isolated C. reinhardtii supercomplexes (B7–B9) showed that they were present in C2S2M2N2, C2S2MN, C2SMN and C2S2 configurations, while differently from higher plants, no C2S2M, C2SM, C2S2M2 or C2M [3] could be found. Apparently, the lack of the neighboring N-trimer strongly influences the binding of LHCII-M, as does CP24 in plants. Indeed in higher plants a complex composed of CP24/CP29/LHCII could be isolated [56], indicating that the association between these proteins is stronger than between them and the core. The CP24/CP29/LHCII complex is obviously not present in C. reinhardtii, but the analysis of the sucrose gradient bands indicates the presence of a CP29/LHCII-M/LHCII-N complex. This suggests that the connection of trimers N and M is stronger than their association with the core and might also explain why, after detergent solubilization, only C2S2 could be detected in early works [7,34].

These results point out that the hypothesis that CP24 has evolved in land plants to increase the antenna size of PSII (or to avoid the formation of C2S2) should be reconsidered as clearly PSII–LHCII of C. reinhardtii is larger than the largest supercomplex of plants [35,36]. It has been shown that under high light stress CP24 and LHCII-M dissociate from the supercomplex [57]. It is thus likely that CP24 (and Lhcb3) has evolved not to make the antenna size of PSII larger, but more flexible, allowing for a fast regulation of the PSII antenna size.

4.2. LhcbM1, LhcbM2/7 and LhcbM3 are the main components of the PSII supercomplexes while LhcbM5 is enriched in the “extra” LHCII population

LHCII antenna in C. reinhardtii is codified by 9 different genes, LhcbM1–M9, but very little is known about their organization and occurrence. The possibility to compare the composition of supercomplexes containing different trimers (S for B7 and B8 and S, M and N for B9) with that of the LHCII trimers that were detached from the supercomplexes (B3) allows to analyze the protein composition of each trimer. Four main LhcbM bands could be resolved in SDS-PAGE, which were identified as the four types of LhcbM complexes. It is interesting to observe that, while the ratio LhcbM/core is increasing when

Table 1

| Pigment composition of: LHCII monomers (B2), trimers (B3) and Cr. PSII supercomplexes (B7–B9). The values of individual carotenoids are normalized to 14 Chls (a + b) in fractions B3 and to 100 Chls (a + b) in fractions with PSII supercomplexes (B7–B9) and thylakoids. The data are the results of at least four different preparations in two replicas. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Pigment                      | B2              | B3              | B7              | B8              | B9              | Cr. cells       |
| Chl a/Chl b                  | 1.29 ± 0.06     | 1.28 ± 0.02     | 2.94 ± 0.18     | 2.41 ± 0.17     | 2.22 ± 0.13     | 2.20 ± 0.061    |
| Chls/Cars                     | 3.79 ± 0.54     | 3.61 ± 0.11     | 5.60 ± 0.39     | 4.80 ± 0.75     | 4.67 ± 0.38     | 4.51 ± 0.421    |
| Neoxanthin/loroxanthin        | 2.41 ± 0.20     | 1.67 ± 0.22     | 5.75 ± 0.48     | 6.96 ± 0.33     | 7.09 ± 0.42     | –               |
| Violoxanthin                  | 0.55 ± 0.07     | 0.68 ± 0.05     | 1.01 ± 0.09     | 1.03 ± 0.09     | 1.12 ± 0.11     | –               |
| Lutein                        | 0.91 ± 0.04     | 1.44 ± 0.18     | 5.42 ± 0.47     | 6.43 ± 0.91     | 8.57 ± 1.44     | –               |
| β-Carotene                    | –               | –               | 5.65 ± 0.50     | 6.40 ± 1.04     | 7.11 ± 1.92     | –               |
| Neoxanthin/loroxanthin        | –               | –               | 5.42 ± 0.47     | 6.43 ± 0.91     | 8.57 ± 1.44     | –               |
| Violoxanthin                  | –               | –               | 5.65 ± 0.50     | 6.40 ± 1.04     | 7.11 ± 1.92     | –               |
| β-Carotene                    | –               | –               | –               | –               | –               | –               |

Fig. 6. Absorption spectra at room temperature of the fractions from sucrose gradient. The spectra were normalized to the maximum absorption of the Qy region. Panel A: absorption spectra of fractions B2 and B3. Panel B: comparison of the absorption spectra of Cr. LHCII and A.t. LHCII trimers. C: absorption spectra of fractions B7–B9.
4.3. The properties of C. reinhardtii LHCII differ from those of plant LHCII

Sequence analysis has shown that the LhcbM proteins of C. reinhardtii do not exactly correlate with the Lhcb1–3 proteins of higher plants [21,24] although they show a high degree of identity. All amino acids which were shown to be responsible for the Chl binding in plants [13] are conserved in the nine LhcbM proteins suggesting that they bind the same number of Chls as the complexes from plants. The Chl a/b ratio of the trimer differs only very slightly from that of plants (1.28 in C. reinhardtii vs. 1.33 in plants) indicating at most the change of affinity of one binding site. However, the absorption spectrum of C. reinhardtii LHCII trimers is different from that of A. thaliana LHCII (Fig. 6B) showing a blue shifted maximum. This difference is not due to changes in the lowest energy pigments, but rather in the environment of Chls absorbing around 674 nm, which in plants are located in the 602 and 603 binding sites [60].

4.4. Energy transfer in PSII–LHCII of C. reinhardtii

The role of LHCII is to absorb light and to transfer excitation energy to the reaction center where charge separation occurs. The connection of LHCII trimers with the core is thus essential in assuring efficient energy transfer [2]. In plants it has been shown that the overall trapping time in the PSII supercomplex is 143 ps [61] and that this very fast transfer is probably due to the presence of preferential energy transfer pathways between complexes involving mainly Chls a. The reconstituted structure of the Cr. PSII supercomplex gives the possibility to point out preferential energy transfer pathways, based on the organization of the Chls. The model of the supercomplex shows that the relative orientation of the S trimer is identical to that in higher plants, which means that chlorophylls 610–611–612 (nomenclature from [13]) of one LHCII monomer, are close to Chl 633 of the core, (nomenclature from [61]), allowing for rapid energy transfer as in plants. The orientation of trimer M is different than in A. thaliana. In this configuration Chls a 610 and 604 of LHII face directly CP29, while Chls 611 and 612 of a different monomer face trimer N. In the case of trimer N the shorter Chl (LHII)–Chl (core) distance (around 20 Å) is between Chls 611 and 612 of one LHII trimer and Chl 612 of the core. In conclusion, also in the case of C. reinhardtii Chls 610–612 seem to be involved in the transfer between complexes, although the connection between trimer M and CP29 involves Chl 610 instead of Chls 611 and 612 as in A. thaliana. These Chls represent the lowest energy sites in LHII of plants [60,62] and they are likely to be conserved also in C. reinhardtii. The minimal distance between Chls in trimer N and in the core is rather large, suggesting a slower transfer, although at this stage we cannot exclude that an additional Chl could be located in between these two complexes in C. reinhardtii.

4.5. The thylakoid membrane of C. reinhardtii harbors at least 6 LHCII trimers per monomeric PSII core complex

The fraction containing the largest PSII supercomplex has a Chl a/b ratio of 2.22, in perfect agreement with the presence of 3.7 trimers on average per dimeric core complex in this fraction. This number is very close to the Chl a/b ratio of C. reinhardtii cells which is 2.28. However, the cells also contain PSI in a close to 1:1 stoichiometry with PSII, in agreement with previous studies [46,63–65]. Considering that PSI coordinates around 240 Chls, 196 Chl a and 44 Chl b [40], the low Chl a/b ratio of the cells can only be explained with the presence of extra LHII trimers in the photosynthetic membrane of C. reinhardtii. The number can be roughly calculated on the basis of the pigment stoichiometry of core (35 Chls a) and PSI antennas [66]. We have used 14 Chls a value of 1.28 for the Chl a/b (than 7.85 Chl a and 6.15 Chl b) of all antenna complexes, although it is probably on the low side for CP29 and CP26, which means that we underestimate the number of LHII trimers. The number X of antenna (monomers) is thus calculated as \( 2.28 = (196 + 35 + 7.85X) / (44 + 6.15X) \). The result suggests the presence of 3–4 additional LHII trimers per monomeric core (meaning 6/7 in total), making the light harvesting capacity of C. reinhardtii far larger than that of plants, in which a maximum of 4.5 LHII trimers was observed [67,68]. Where are the additional trimers located? Looking at the structure of the PSII supercomplex it is clear that there is space at most for one extra trimer associated with the core. The tighter possible organization of the PSII supercomplexes of C. reinhardtii is shown in Fig. 8 and indicates that differently from the case of plants the dimeric cores cannot be directly connected but are separated by a minimum of one row of LHII. In this case we can speculate that trimer N should be able to transfer energy to two different core complexes.
It remains to be elucidated if the "extra" trimers are located in between the PSI supercomplexes and if a subpopulation of them is associated with PSI in all conditions, as it is the case in plants [58] or if they form LHClI-only domains.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.bbacbio.2013.07.012.

Acknowledgements
We thank Kevin Redding (Arizona State University, Tempe, USA) for kindly providing the C. reinhardtii strain [JVD-1B (pPG1)] used in this work and Pierre Cardol (Université de Liège) for his help with the ECP measurements. This work was supported by the ERC Starting Grant 281341 (ASAP) to RC and FOM Grant 10TBSC12-2 to EJB.