Engineering photosynthetic light capture: impacts on improved solar energy to biomass conversion

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Summary

The main function of the photosynthetic process is to capture solar energy and to store it in the form of chemical ‘fuels’. Increasingly, the photosynthetic machinery is being used for the production of biofuels such as bio-ethanol, biodiesel and bio-H2. Fuel production efficiency is directly dependent on the solar photon capture and conversion efficiency of the system. Green algae (e.g. Chlamydomonas reinhardtii) have evolved genetic strategies to assemble large light-harvesting antenna complexes (LHC) to maximize light capture under low-light conditions, with the downside that under high solar irradiance, most of the absorbed photons are wasted as fluorescence and heat to protect against photodamage. This limits the production process efficiency of mass culture. We applied RNAi technology to down-regulate the entire LHC gene family simultaneously to reduce energy losses by fluorescence and heat. The mutant Stm3LR3 had significantly reduced levels of LHCI and LHCII mRNAs and proteins while chlorophyll and pigment synthesis was functional. The grana were markedly less tightly stacked, consistent with the role of LHCII. Stm3LR3 also exhibited reduced levels of fluorescence, a higher photosynthetic quantum yield and a reduced sensitivity to photoinhibition, resulting in an increased efficiency of cell cultivation under elevated light conditions. Collectively, these properties offer three advantages in terms of algal bioreactor efficiency under natural high-light levels: (i) reduced fluorescence and LHC-dependent heat losses and thus increased photosynthetic efficiencies under high-light conditions; (ii) improved light penetration properties; and (iii) potentially reduced risk of oxidative photodamage of PSII.

Keywords: light harvesting, solar energy conversion, biomass, photosynthesis, RNAi, photoinhibition.

Introduction

Photosynthesis is of fundamental biological importance, as almost all life on Earth depends on it, either directly or indirectly via its products. Light-harvesting proteins fulfil a special role in this process as they form the intersect between the physical world (solar electromagnetic radiation) and living cells. Specifically, they capture solar energy and regulate the flow of the derived excitation energy to the photosynthetic reaction centres, in all photosynthetic organisms. The central role of light-harvesting proteins in solar energy capture also makes them key biotechnological targets in any strategy to genetically optimize the efficiency of the synthesis of bio-products and biofuels such as bio-ethanol, biodiesel, bio-H2 and biomass to liquid (BTL) (Kruse et al., 2005; Rupprecht et al., 2006; Zwart et al., 2006).

Natural antenna diversity

It is perhaps not surprising that the huge diversity of photosynthetic organisms (e.g. purple bacteria, cyanobacteria, green-sulphur bacteria, green algae, lower and higher plants)
have evolved a wide variety of light-harvesting proteins, specialized for their specific habitats (Grossman et al., 1995). In plants and a variety of microalgae (e.g. green algae), these are called light-harvesting complex (LHC) proteins. They are embedded in the thylakoid membranes of the chloroplast and act as structural scaffolds that hold chlorophylls and carotenoids in defined orientations and optimized molecular environments for light capture. Light-harvesting complexes are divided into distinct LHCI and LHCII protein families, based on their predominant association with photosystem (PS) I or II, respectively.

The LHCI protein family comprises several nuclear encoded isoforms [e.g. 6 LHCA1-6 isoforms in Arabidopsis thaliana (Jansson, 1999) and 9 LHCA1-9 genes in Chlamydomonas reinhardtii (Takahashi et al., 2004)] which exhibit high sequence homology. The LHCI proteins provide PSI with the excitation energy necessary to reduce the electron acceptor ferredoxin.

The LHCII proteins are subdivided into the major and minor proteins. The major proteins are far more abundant and largely form trimers making up the outer light-harvesting antenna system. This funnels excitation energy via the inner antenna to the PSII reaction centre. The inner antenna consists of the minor LHCII proteins that are monomeric (Boekema et al., 1995; Hankamer et al., 1997; Dekker and Boekema, 2005).

The LHCII proteins are encoded by a family of nuclear genes and their number and nomenclature vary from species to species [e.g. 9 LHCB1-3 isoforms in Arabidopsis thaliana (Jansson, 1999; Legen et al., 2001); 9 LHCBM1-11 isoforms in C. reinhardtii (Elrad & Grossman, 2004)]. The LHCII genes and proteins share a high degree of sequence similarity, not only to one another but also with the LHCI proteins. The reasons for the evolution of such a large number of highly related proteins and their individual functions are not known but are thought to reflect the great adaptive flexibility required of the photosynthetic machinery. The expression of each isoform is highly regulated from the level of mRNA transcription to protein degradation (Escoubas et al., 1995; Flachmann and Kühlbrandt, 1995; Lindahl et al., 1995; Durnford et al., 2003; Tokutsu et al., 2004; Mussgnug et al., 2005). Furthermore, the translation of individual LHCII isoforms can be regulated by the translational repressor NAB1 through mRNA sequestration (Mussgnug et al., 2005). Additionally, light energy distribution between PSI and PSII can be fine tuned via LHC state transitions by reversible phosphorylation of certain LHCII proteins (Bonaventura and Myers, 1969; Murata, 1969; Kruse, 2001; Turkina et al., 2006a,b).

Light capture and photoprotection

When solar energy levels exceed energy demand by the cell, LHC transcription is down-regulated to prevent photodamage (Adir et al., 2003). In contrast, low-light levels lead to increased transcription to maximize photon capture efficiency. Thus, the total amount of LHC proteins and their isoform composition is constantly adjusted to the actual demand for light capture and to changing irradiation conditions.

Besides their role as light energy capture proteins, LHC proteins also fulfill a second role when solar irradiation exceeds photosynthetic capacity. Under these conditions, LHCII proteins facilitate the dissipation of light energy as heat or fluorescence via a mechanism called ‘energy-dependent non-photochemical quenching’ (NPQ) of chlorophyll fluorescence, which is reported to protect the cell from oxidative damage (Müller et al., 2001). This strategy, however, has the downside that a large proportion of the absorbed photons (~80–95%) are wasted as fluorescence and heat under high-light levels (Polle et al., 2002; Polle et al., 2003), limiting the overall photosynthetic conversion efficiency (Kruse et al., 2005). Despite this, the very existence of these processes indicates that they confer a competitive advantage on the host cell in its natural environment. This is because most plants and photosynthetic microorganisms are limited in mobility and therefore had to develop strategies to adjust to seasonal, diurnal and even daily changes in environmental light levels (e.g. due to varying cloud cover). They have achieved this through the orchestrated interplay of a diverse array of short and long-term adaptation mechanisms (Dietz, 2003).

Light adaptation vs. photon capture efficiency

The cost of having these adaptation mechanisms is reflected in the low light to biomass conversion efficiency of most ecosystems (Prince and Kheshgi, 2005). However for commercial algal bioreactor systems (e.g. for phytochemical and biofuel production) that require high efficiency and in which environmental conditions can be controlled precisely, this ability to adapt to light levels can, at least in theory, largely be dispensed with. With this aim in mind the light-harvesting antenna size can be reduced to minimize efficiency losses and improve light penetration into the bioreactor (Melis et al., 1999; Kruse et al., 2005; Prince and Kheshgi, 2005).

Comparative studies of low (100 µmol/m² s) and high (2000 µmol/m² s) light adapted Dunaliella salina algal cultures (triggering large and smaller light-harvesting complexes, respectively) suggest that cells with reduced
light-harvesting antennae could exhibit two to three times higher photosynthetic capacities in mass culture than normally pigmented cell lines (Neidhardt et al., 1998; Melis et al., 1999). In these experiments, the PSII-LHCII and PSI-LHCI complexes were reported to bind 60 and 100 chlorophylls per reaction centre, respectively. In the absence of the entire antenna system, PSI and PSII are reported to bind ~40 and ~100 chlorophylls per reaction centre (Hankamer et al., 1997; Dekker and Boekema, 2005), respectively. This suggests that the high-light conditions used resulted in the almost complete down-regulation of LHCI and a marked down-regulation of LHCII. However, the limitation of this approach is that it only allows temporary down-regulation.

This paper describes the construction of a C. reinhardtii mutant in which the entire LHCI and LHCII antenna complex system has been permanently and almost completely down-regulated and reports its genetic, biochemical, physiological and ultrastructural characterization. These results therefore represent an important step forward in engineering enhanced photon-capture efficiencies in green algal systems.

Results

Screening for LHC mutants

LHCBM1-11 encode the light-harvesting proteins predominantly associated with PSII. These genes are not only highly conserved with respect to one another, but also show homologies to those encoding the LHCI proteins (LHC1-1-9) as well as the minor antenna proteins LHCB4 and LHCB5. The most highly conserved region of this gene super family is a region coding for helix 3 of the LHC proteins and shows a region of ~30 bp which is virtually identical in all antenna genes (Supplementary Figure S1). This conserved region was selected as the target for the simultaneous down-regulation of all LHCI and -II proteins using RNAi technology (Fire et al., 1998; Schroda, 2006).

In wild type C. reinhardtii some, but not all, LHCII proteins are under the translational control of the cytosolic RNA-binding protein NAB1 (Mussgnug et al., 2005). NAB1 binds and sequesters certain LHCII mRNAs, stabilizing their RNA transcripts and preventing their translation. To avoid any ambiguity caused by the interference of this regulatory pathway with RNAi-induced mRNA degradation, the NAB1 knockout mutant Stm3 was chosen as the parental strain.

To generate a LHC-specific RNAi vector, a region of 187 bp with high homology in all LHC genes (see supplementary material Figure S1) was inserted into the vector MAA7/X-IR (Rohr et al., 2004) and subsequently transformed into Stm3 cells. One hundred and twenty positive transformants were identified through selection on the antibiotic paromomycin (data not shown). These were then screened for effective RNAi through the use of increased concentrations of 5-fluorouracil (5-FL) (Rohr et al., 2004). The Maa7/X-IR vector is designed to allow direct and gradual screening for RNAi efficiency by co-silencing the endogenous tryptophan synthase gene together with the target. Tryptophan synthase converts 5-FL in the medium into the toxic tryptophan analogue 5-fluorotryptophan (Rohr et al., 2004). Thus 5-FL-tolerant mutants can be expected to show efficient RNAi for tryptophan synthase and the co-silenced target gene. More than 50% of the 120 transformants survived 30 μM 5-FL which was lethal to the parental strain Stm3, suggesting that the construct was expressed to some degree in these transformants. One highly 5-FL-resistant mutant which tolerated up to 90 μM 5-FL in the medium was selected. This strain showed a considerably lighter green phenotype compared to the other colonies (consistent with a mutant in which LHC expression is significantly down-regulated) and was assigned the name Stm3LR3 (Stm3 LHC reduction mutant 3). The fact that Stm3LR3 survived 5-FL concentrations lethal for parental strain Stm3 suggests that the light green phenotype was due to the expression of the RNAi construct and not due to a locus-specific gene inactivation caused by random insertion of the RNAi plasmid. However, it has been reported previously that mutations of genes involved in chlorophyll or carotenoid synthesis can result in severe reduction of LHC antenna size (Polle et al., 2001; Polle et al., 2002), which can result in a pale green phenotype (Tanaka et al., 1998). To eliminate the possibility that the light green phenotype of Stm3LR3 was due to a locus-specific inactivation of a gene (or genes) involved in chlorophyll or carotenoid synthesis, its pigment composition was analysed by thin layer chromatography (Figure 1). The separation of the extracted pigments and their identification by spectroscopy showed that Stm3LR3 was capable of chlorophyll-a, chlorophyll-b and indeed carotenoid synthesis which are the only pigments known to be bound by the LHC proteins. Thus, taken together with the high resistance to 5-FL, these data indicated that one or more LHC proteins were down-regulated in Stm3LR3 by RNAi.

Identification of LHCI- and LHCII-specific mRNAs and proteins

To determine whether the LHCI (LHC1-1 to LHC1-9), LHCII (LHCBM1 to LHCBM11) and LHCB4/LHCB5 mRNA levels were indeed down-regulated, each individual LHC mRNA level was determined by quantitative real-time polymerase chain
Down-regulation of LHCs in C. reinhardtii

reaction (qRT-PCR) using cell cultures in their late logarithmic growth phase (Figure 2). In all cases the LHC mRNA levels in Stm3LR3 were significantly reduced (subunit specific to 0.1–26%; all P-values < 0.016) with respect to the levels observed in the parental strain Stm3. The LHCBM9 mRNA level was already very low and at the limit of detection in Stm3 (~0.1% of the average LHCBM mRNA level), making it impossible to detect further reduction under the experimental conditions. The nuclear encoded genes for ribulose bisphosphate carboxylase/oxygenase (Rubisco) small subunit (RBCS1, RBCS2) and the chloroplast encoded gene for Rubisco large subunit (rbcL) did not show a significant difference in mRNA abundance. These results clearly indicate that the transformed RNA construct resulted in efficient repression of the expression of all LHC mRNA tested.

To verify LHC reduction at the protein level, denaturing SDS-PAGE and LHC-specific Western analyses were performed with thylakoid membranes of Stm3 and Stm3LR3, with isolated LHC proteins serving as a positive control. The Stm3 and Stm3LR3 thylakoid membranes analysed were loaded on an equal chlorophyll basis. Consequently it should be noted that, despite the Stm3LR3 sample having a higher total protein loading, the LHC bands were still markedly reduced compared to those observed in the Stm3 lane (Figure 3, Coomassie). This unambiguously shows that LHC proteins levels are greatly reduced in Stm3LR3, and this conclusion is supported by the accompanying Western blot (Figure 3, α-LHC) and is in agreement with the qRT-PCR data (Figure 2).

Thylakoid ultrastructure

LHC proteins are reported to play a central role in thylakoid stacking (Barber and Chow, 1979; Allen and Forsberg, 2001; Barber and Nield, 2002; Chow et al., 2005) (Figure 4c). Electron microscopic images (Figure 4a,b) were taken to characterize any morphological changes in thylakoid organization in the mutant Stm3LR3.
As reported earlier, Stm3 shows an increase in thylakoid membrane stacking compared to WT (Mussgnug et al., 2005). In contrast, the images of mutant Stm3LR3 indicate that membrane stacking is decreased when LHC antenna proteins are down-regulated. Here, the typical multi-membrane pseudo-granal organization (Figure 4a inset) was replaced by long stretches of parallel double bilayers (Figure 4b inset). To our knowledge, this thylakoid organization has not been described for green algae before, but is very similar to the membrane organization observed in chlorophyll-b-less wheat mutant CD3 and chlorophyll-b-less barley mutant chlorina f2 when grown under special light conditions which trigger the down-regulation of all LHC proteins (Allen et al., 1988; Krol et al., 1995).

Our results underline the importance of LHC proteins for membrane organization in Chlamydomonas. In contrast to the chlorophyll-b-deficient mutants CD3 and chlorina f2, where special light conditions had to be applied to trigger complete LHC depletion and concomitant unstacking of the thylakoid membranes, Stm3LR3 did not show stacking under constant, standard illumination of 100 µmol/m² s white light, indicating that very low levels of LHC proteins were present. Despite this, the rate of photosynthesis was clearly sufficient to support photosynthetic growth on minimal medium (not shown).

To provide a more detailed phenotypic description, the isolated thylakoid membranes of Stm3 and Stm3LR3 were

Figure 3 LHC proteins in Stm3 and Stm3LR3. Anti-LHC Western blots with Stm3 and Stm3LR3 thylakoid membranes. Left side: Coomassie blue protein stain; right side: anti-LHC Western blot. Samples were loaded on the basis of equal chlorophyll amount (4 µg for the LHC sample, 10 µg for the thylakoid membranes of Stm3 and Stm3LR3 (half the amounts for the Western blot). The black box indicates the molecular range of LHC proteins, the asterisks mark higher molecular weight LHC aggregates; the apparent molecular weight is given in kDa.

Figure 4 Electron microscopic images, models of membrane interaction and sucrose gradients of Stm3 and Stm3LR3 (3LR3). (a) Stm3 section showing extensive granal stacks, consisting of a large number of tightly packed bilayers. (b) Stm3LR3 section showing the correlation between light-harvesting complex (LHC) reduction and the disaggregation of the granal stacks, into bilayer pairs. The red inset boxes show ×2 magnified views of A or B, respectively. The black arrow in B marks a single bilayer. (c) Model of normal thylakoid stacking in the presence of LHC. The black dots indicate the involvement of LHCs in the stacking process. C: cytochrome b6/f complex, OEC: oxygen-evolving complex. (d) Illustrates membrane organization Model 1 in which bilayer pairs are formed via lumenal interactions. (e) Illustrates membrane organization Model 2 in which bilayer pairs are formed via stromal interactions with markedly increased lumenal volume. (f) Sucrose gradient separation of photosynthetic complexes form thylakoid membranes of Stm3 and Stm3LR3 (equal chlorophyll loading). Chlorophyll containing bands are indicated. LHC PIGM = LHC pigments and low levels of proteins; LHC = LHC proteins; PSIIm = PSI monomers; PSIm = PSI monomers; PSId = PSI dimers; PSIISc1 = PSI-LHCI supercomplex 1; PSIISc2 = PSI-LHCCI supercomplex 2.
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Gently solubilized with dodecyl maltoside and resolved into their component photosynthetic complexes by sucrose density gradient centrifugation (loaded on an equal chlorophyll basis). The results clearly show that the bottom two bands (PSIISc1 and PSIIsc2) are absent in Stm3LR3 (Figure 4f). The PSIIsc2 band is enriched in the classical PSII-LHCII super-complex reported by Boekema et al. (1995) and consists of a central PSII core dimer flanked by two LHC antenna sets (the PSIIsc1 lacks one of these antenna sets). The fact that Stm3LR3 lacks the PSIIsc1 and PSIIsc2 super-complexes is consistent with the down-regulation of the LHC proteins. As the Stm3 and Stm3LR3 samples were loaded on an equal chlorophyll (not equal cell) basis, it can be seen that the abundance of LHC:PSIIm (monomer) ratio is much lower in Stm3LR3 than in Stm3, consistent with an overall reduction in LHC proteins.

Chlorophyll parameters, fluorescence properties and light transmission

Optical transmission microscopy (Figure 5a) showed that at the level of the individual cell, the Stm3LR3 strain had a markedly lower chlorophyll content than the parental Stm3 control (consistent with the down-regulation of LHC gene expression). This has the effect that when cultures were adjusted to the same cell number (6 × 10^6 cells/mL), Stm3LR3 cultures had a total chlorophyll content of 32% (± 0.95 SE) of that of Stm3 (Figure 5b). Furthermore, the chlorophyll-a/b ratio in Stm3LR3 was significantly increased from 1.95 (± 0.12 SE) in Stm3 to 4.05 (± 0.1 SE), as expected for LHC-deficient strains (Hankamer et al., 1997; Melis et al., 1999; Andersson et al., 2003). The reduced chlorophyll content in Stm3LR3 led to higher light levels at the centre of 650 mL bioreactors (~290% compared with Stm3 at optical density (OD)_{750nm} = 1.0) when illuminated with 750 µmol/m^2 s white light (not shown).

To measure the effects of the down-regulation of the antenna on fluorescence, Stm3 and Stm3LR3 were next analysed using confocal laser scanning microscopy (Figure 5c). Mixtures as opposed to pure samples of the two strains were analysed to ensure their direct comparison under identical imaging conditions. To identify each cell type within the mixed population, Stm3 or Stm3LR3 were individually labelled with a green fluorescent marker (MitoTracker® FM) in two separate experiments. Panels I–IV in Figure 5(c) show the results of experiment one, in which Stm3 was labelled with the green marker. Panels V–VIII in Figure 5(c) show the results of experiment two, in which Stm3LR3 cells were labelled with the green marker. In both experiments, the mixtures were imaged under four conditions. Panels I and V show the cells imaged in the normal optical mode. Six Chlamydomonas cells are seen in panel I; five cells in panel V.

**Figure 5** Phenotypic analysis of Stm3LR3. (a) Microscopic images of Stm3 and Stm3LR3 cells. (b) Cultures of Stm3 and Stm3LR3 (3LR3) after heterotrophic growth and adjustment to equal cell numbers (6 × 10^6 cells/mL). Relative chlorophyll concentrations are indicated (Stm3 set to 100%, ± 0.95 SE). (c) Chlorophyll fluorescence of Stm3 and Stm3LR3 cells estimated by confocal laser scanning microscopy in mixed cultures. Panels I–IV: experiment one in which Stm3 cells were pre-labelled (green). Panels V–VIII: experiment two in which Stm3LR3 cells were pre-labelled (green). Panels I and V: microscopic image of all cells. Panels II and VI: identification of Stm3 (panel I) and Stm3LR3 (panel VI) cells by separate pre-labelling using MitoTracker® FM green fluorescence. Panels III and VII: chlorophyll fluorescence (red). Panels IV and VIII: merged images of green and red fluorescence. (d) Photosynthetic quantum yield (φPSII) of Stm3 and Stm3LR3. Fluorescence parameters Fm and Fm′ were recorded during actinic illumination of liquid cultures and φPSII was calculated according to Maxwell and Johnson (2000), (φPSII = (Fm′-Fm)/Fm′), a.u. = arbitrary units. Error bars = standard error.
Next the cells were imaged in a fluorescence mode which visualized the green fluorescent marker to identify which cells were Stm3 and which were Stm3LR3. In experiment one, this mode (panel II) shows that two of the six cells were Stm3 (green). Similarly, in experiment two (panel VI) three of the five cells were identified as Stm3LR3 (green). To measure chlorophyll fluorescence, the mixed samples were then excited with an actinic light (0.21 mW, 543 nm) before measuring chlorophyll fluorescence (653–718 nm pass filter). In experiments one and two (panels III and VII), only the parental Stm3 cells fluoresced strongly (red), consistent with their large antenna system. In contrast Stm3LR3 appeared to have a near ‘null fluorescence’ phenotype under these conditions.

For control purposes, panels IV and VIII display an overlaid image of the marker-derived fluorescence (green), and the chlorophyll fluorescence (red). These results unambiguously demonstrate that the reduction of the LHC antenna size in Stm3LR3 has resulted in a strong down-regulation of fluorescence.

To determine whether the observed reduction of fluorescence translates to improved photon capture efficiencies, maximum quantum yield of PSII (Fv/Fm = (Fm–F0)/Fm) and photosynthetic quantum yield (φPSII = F′m–Ft/F′m) (Maxwell and Johnson, 2000) were determined (Figure 5d). Fv/Fm was similar for Stm3 (0.711, ± 0.016 SE) and Stm3LR3 (0.767, ± 0.005 SE), indicating efficient PSII primary activity (from water to QA) in both strains. To determine the levels of fluorescence losses during illumination, photosynthetic quantum yield was measured. Fm corresponds to the maximal fluorescence while Ft corresponds to the minimal fluorescence during illumination with actinic light (815 μmol/m² s white light). In theory, if 100% of the photons captured by the LHCII antenna are used to drive the PSII photochemistry, a φPSII value of 1 is expected. In contrast if all the energy is dissipated through the process of fluorescence, φPSII would equal 0. Therefore, the parameter φPSII can give a good indication of the photon capture efficiency. Under illumination with photosynthetically active radiation at which the parental strain Stm3 had φPSII values of 0.256 (± 0.05 SE), the φPSII value of Stm3LR3 was ~81% higher at 0.464 (± 0.008 SE) (Figure 5d). These results support the confocal microscopy data and suggest that the fluorescence losses of the PSII antenna are significantly lower in the reduced LHC strain.

Reduced photoinhibition and enhanced cell growth at high-light levels

Light induced LHC antenna reduction has previously been reported not only to improve the photosynthetic quantum yield, but to decrease photoinhibition under high-light conditions (Baroli and Melis, 1998). To determine the impact of antenna down-regulation in Stm3LR3, photoinhibition trials were conducted using 1400 μmol/m² s high light. The level of photoinhibition was determined by measuring the maximal oxygen evolution capacity in 20 min intervals over a period of 100 min. These experiments revealed that under photoinhibitory light conditions, PSII water-splitting activity in Stm3LR3 maintained a comparable high level of ~40% after 100 min of high-light treatment, compared to only 13% oxygen evolution activity remaining in Stm3 (Figure 6a). Thus antenna reduction in Stm3LR3 appears to confer a phenotypic decrease in photoinhibition.

Given the enhanced tolerance of Stm3LR3 to high-light levels, cultivation experiments were conducted at elevated light conditions of 1000 μmol/m² s. These revealed that Stm3LR3 has not only reduced sensitivity to photoinhibition (Figure 6a) but that cell growth and replication was also significantly faster than that of the parental strain Stm3 (Figure 6b). Both strains reached similar maximal cell densities

Figure 6 Photoinhibition and growth rates under high-light conditions. (a) O₂ production capacity of Stm3 and Stm3LR3 (3LR3) during photoinhibitory light treatment (1400 μmol/m² s for 100 min), determined with a Clark-type oxygen electrode. (b) Mixotrophic growth rates of Stm3 and Stm3LR3 (3LR3) cultures under continuous high-light conditions (1000 μmol/m² s) in TAP medium. Error bars = standard error.
(100% in Stm3; 104% in Stm3LR3) due to their common ability for mixotrophic growth in tris acetate phosphate (TAP) medium. However, peak density of Stm3LR3 cultures was detected already after 26.5 h, at which Stm3 cultures had only been able to grow to 54% of their maximal cell densities (Figure 6b, dotted line).

Discussion

Photosystem and antenna conservation

While photosynthetic core complexes (PSI and PSII) have retained a remarkable degree of conservation between higher plants, green algae and cyanobacteria (Dekker and Boekema, 2005), they have evolved quite different complexes for harvesting solar energy. This reflects their optimal adaptation to specific natural habitats (Grossman et al., 1995). This diversity is presumably linked to the fact that light-harvesting proteins are not only involved in energy capture and transfer, but also fulfil the important role of balancing excitation energy between PSI and PSII (Allen and Forsberg, 2001) and to protect the cells under conditions of excess irradiation (Müller et al., 2001). Furthermore, LHC proteins are thought to be important for maintaining the structural organization of granal thylakoid membrane stacks (Barber and Chow, 1979; Allen and Forsberg, 2001; Chow et al., 2005).

Efficient antenna down-regulation in mutant Stm3LR3

Using RNAi technology, the expression of all 20 genes encoding for LHCl, LHClI, CP26 and CP29 were simultaneously and strongly down-regulated in Stm3LR3 (0.1–26% of Stm3; Figures 2–4). This level of permanent LHC reduction has, to our knowledge, not been reported in any algal photosynthetic organism to date.

Pigment synthesis is functional in Stm3LR3

The degree of LHC down-regulation appears to be independent of the main biochemical pathways involved in photosynthetic pigment production as chlorophylls a and b, carotene and specifically lutein, violaxanthin and neoxanthin are all produced in Stm3LR3 (Figure 1). This suggests that the altered thylakoid ultrastructure observed in Stm3LR3 (Figure 4b) is the direct result of LHC down-regulation and supports previous reports that LHC proteins are involved in membrane stacking (Barber and Chow, 1979; Allen and Forsberg, 2001; Chow et al., 2005).

Thylakoid ultrastructure is altered

Despite major changes in ultrastructure (see bilayer pairs in Figure 4b), Stm3LR3 is not only capable of photoautotrophic growth but of supporting virtually identical maximal rates of O₂ evolution compared to Stm3 under saturated light conditions (not shown). This observation indicates that Stm3LR3 has a fully functional lumen, as in its absence, functional photosynthetic reactions such as the photophosphorylation which are crucial for the cells survival could not be maintained. Currently, two models that fulfil the requirement for an intact lumen, appressed bilayer pairs, and the involvement of LHC proteins in bilayer stacking can be proposed (Figure 4d,e). More experimental data will be required to unambiguously determine which of these two models most accurately describes the Stm3LR3 phenotype. However, our conclusions are consistent with the observation that Stm3 (Figure 4a) is enriched in LHCs and exhibits strongly appressed membranes. In C. reinhardtii WT strains, which contain less LHC proteins than Stm3, less tightly stacked pseudo-grana are observed (Mussgnug et al., 2005).

Chlorophyll fluorescence losses and photoinhibition are decreased in Stm3LR3 while cell growth is enhanced under high-light conditions

The confocal (Figure 5c), φPSII (Figure 5d) and oxygen evolution experiments (Figure 6a) highlight four important properties of Stm3LR3. First, LHC down-regulation in this mutant resulted in a decrease in the dissipation of captured light energy through the process of fluorescence (and possibly heat dissipation, although this was not measured). Second, this would be expected to lead to an increase in photosynthetic quantum yield and this was indeed observed (Figure 5d). Third, reducing antenna size reduced the sensitivity of the system to photoinhibition (Figure 6a). Fourth, despite such major antenna reductions the remaining functional photosynthetically active pigments (mainly associated with the PSI and PSII core complexes) were still clearly sufficient to drive photosynthesis efficiently and promoted increased cell growth and replication compared with the parental strain cell cultures (Figure 6b) at elevated light conditions of 1000 µmol/m² s.

Biotechnology applications

The Stm3LR3 phenotype has important implications for biotechnological applications. Photosynthetic microorganisms are increasingly used in large-scale algal culture for the
production of pigments (e.g. β-carotene and astaxanthin), lipids (e.g. polyunsaturated fatty acids) polysaccharides and biomass for biofuel generation (e.g. biodiesel, BTL and bio-H₂) (Pulz and Gross, 2004; Rupprecht et al., 2006). Compared with higher plants, photosynthetic microorganisms have a number of advantages. Biomass is built up rapidly and production facilities (e.g. open and closed bioreactors) can be sited on non-arable land (eliminating competition with crops required for food production). Closed bioreactors offer the additional advantage that cultures can be maintained under highly controlled conditions (e.g. light, pH, nutrients, temperature) to improve productivity, and in the case of the synthesis of medical products, ensure high quality standards.

The ability to control light conditions in bioreactors is particularly relevant for the work presented here and indeed complements it, as it offers a potential way to increase photosynthetic efficiency levels from ~1–2% (in many ecosystems) towards 10% (Kruse et al., 2005). Specifically, regulating the light levels in both open and closed bioreactors by engineering means, reduces the dependence of the organism to regulate excitation energy transfer to the photosynthetic reaction centres and the need for photoprotection strategies via wasteful fluorescence and heat dissipation, which can account for up to ~80–95% of energy loss from the system (Polle et al., 2002, 2003), most of which is lost at the illuminated surface of the bioreactor. In this context it should also be noted that for typical wild-type green algae, outdoor conditions very rapidly exceed light utilization capacity during the course of a day (e.g. at mid-latitudes during a cloudless spring day in the Northern Hemisphere, light intensity exceeds light utilization capacity between 7.00 and 17.00 hours (Melis et al., 1999)). The use of small LHC antenna mutants in light-regulated bioreactors is therefore expected to reduce energy losses and has further benefits in terms of increasing the overall photosynthetic efficiency of the algal culture. In particular, reducing fluorescence and heat losses of this magnitude should result in a considerable improvement in light penetration into the bioreactor. Consequently, a much larger proportion of the cells in the culture can be optimally illuminated by the incident solar irradiation, thereby increasing the overall productivity of the bioreactor. This in turn facilitates the reduction of bioreactor sizes and costs that are directly dependent on the efficiency of solar energy conversion (Prince and Kheshgi, 2005). Alternatively, higher cell densities can be used.

In conclusion, RNAi technology has enabled us to down-regulate almost the entire LHC antenna system (0.1–26%, subunit specific). Analysis of the thylakoid ultrastructure indicated that this LHC down-regulation reduces interactions between adjacent thylakoid membrane pairs within the granal stacks consistent with the loss of PSII supercomplexes (Figure 4f) which have been postulated to mediate bilayer stacking (Barber and Chow, 1979; Allen and Forsberg, 2001; Barber and Nield, 2002; Chow et al., 2005). Stm3LR3 also showed a number of high light-related phenotypes (reduced fluorescence losses, increased photosynthetic quantum yield, increased resistance to photoinhibition and faster growth rate at elevated light levels) that could have important implications for improved algal cultivation at natural light conditions. In terms of biotechnological applications, the optimization of photosynthetic light capture is central to the enhancement of photosynthetic efficiency for the improved production of photosynthetic biofuels (e.g. bio-ethanol, biodiesel, bio-H₂, BTL) and a wide range of other products of large-scale algal culture.

Experimental procedures

Strains and culture conditions

The mutant Stm3 was generated as described before (Mussgnug et al., 2005). Liquid cultures of C. reinhardtii were grown in continuous white light (100 µmolm⁻² s⁻¹). Media for mixotrophic growth (TAP) and photoautotrophic growth (high salt medium) were prepared as described (Harris, 1989). Ten milligram per litre paromomycin (Sigma) and 1.5 mM L-tryptophan (Sigma) were added for screening for pAlkk1- and mAA7/X-IR-positive transformants. Plates with 30–90 µM 5-fluorooridole (Sigma) were prepared to re-screen transformants for efficient RNAi.

RNAi vector construction and transformation

Manipulations of nucleic acids were performed following standard methods (Sambrook et al., 1989). A 187-bp region (separated by a spacer of 145 bp) was selected for RNAi to suppress gene expression of all LHC gene isoforms. The selected sequence (corresponding to nucleotides 527–713 3' from the cDNA start codon) was amplified by PCR using LHCBM1 cDNA as template (sense primers: cggaattcgcctgacacgaattccgtacc-gcgtctgaaggtgtgcagacaggtc, antisense primers: cggaattcgcctgacacgaattccgtacc-gcgtctgaacagagacaccagagacagac, XbaI and EcoRI restriction sites are in italics) cloned in sense-antisense orientation via the XbaI site and inserted into RNAi vector MAA7/X-IR (Rohr et al., 2004) via the EcoRI site. This LHC-RNAi vector (pAlkk1) was then transformed into C. reinhardtii strain Stm3 by glass bead transformation (Kindle, 1990).

RNA isolation and cDNA synthesis

Four biological replicates of total RNA were isolated from 10 mL of C. reinhardtii culture (OD₇₅₀nm = 1.0) using Promega™ SV RNA Isolation Kit. First strand cDNA was synthesized using 2 µg total RNA and the Superscript III™ RT (Invitrogen) protocol with the
following amendments: enzyme 0.5 µL, oligo-dT (0.2 µL of 100 µM) and random hexamer (0.05 µL of 3 µg/µL) primers. Following synthesis, cDNA was diluted to 5 ng/µL.

**Quantitative real-time PCR (qRT-PCR)**

qRT-analysis was carried out in ABI optical 384-well plates using an ABI PRISM® 7900 HT Sequence Detection System (Applied Biosystems) and the standard thermal profile and dissociation stage. Each reaction contained 3 µL SYBR® Green 2X Master Mix, 5 ng of cDNA, and 200 nM of each gene-specific primer pair to a final volume of 6 µL. PCR reagents were aliquoted into 384-well plates using an Eppendorf Epmotion 5075™ Liquid Handler. The PCR primer efficiency (E-value) was calculated as previously described (McGrath et al., 2005). Gene expression levels relative to 18S rRNA were calculated for each cDNA sample using the equation: relative ratio (gene) = (E(18S rRNA–Ct gene))/(E18S rRNA–Ct 18S rRNA). The average ratios of the four Stm3 and four Stm3LR3 samples were used to determine the fold-change in transcript level between Stm3 and Stm3LR3. For the primers used see supplementary material Table S1.

**Chlorophyll fluorescence measurements**

For fluorescence microscopy, 500 µL of Stm3 and Stm3LR3 cultures was harvested by centrifugation and re-suspended in 300 µL TAP. A total of 0.1 µL MitoTracker® Green FM (Molecular Probes, 35 nM final concentration) was added, and cell suspensions were incubated for 45 min in the dark. Cells were collected by centrifugation, washed twice and re-suspended in 100 µL TAP. Mixtures (1 : 1) of labelled Stm3 with unlabelled Stm3LR3 cells, or the converse, unlabelled Stm3 with labelled Stm3LR3 cells were prepared and fluorescence signals were recorded (Zeiss LSM 510 META). All chlorophyll fluorescence parameters were determined at room temperature (Mini-PAM, Walz). Fv/Fm was determined after dark incubation of cell suspension (15 µg Chl/mL) for 5 min by recording of F0 and Fm (the latter after application of a saturating light pulse) incubation of cell suspension (15 µmol/m² s). Quantum yield was determined at the end of the measurements by dark incubation and subtracted from the apparent O2 production levels. Light transmission was measured with a spherical light-meter (QSL-2101, Biospherical Instruments Inc.) in the centre of a 650 mL glass reactor after adjusting cell cultures to OD750nm = 1.0.

**Photo inhibition measurements**

Cell cultures were grown for 2 days, adjusted to OD750nm = 0.2 in TAP medium and then illuminated with 1400 µmol/m² s white light for 100 min. Samples were taken in 20 min intervals, 10 mM NaHCO3 added and apparent O2 production measured in a Clark-type oxygen electrode (1400 µmol/m² s actinic white light). Respiration was determined at the end of the measurements by dark incubation and subtracted from the apparent O2 production levels.

**SDS gel electrophoresis, thin layer chromatography (TLC) and thylakoid membrane preparation**

Proteins were separated by SDS gel electrophoresis (Laemmli, 1970) and subsequently electroblotted onto nitrocellulose membranes (Amersham). LHC-specific antibodies, raised against higher plant LHC proteins, were kindly provided by S. Jansson and used in conjunction with Anti-Rabbit IgG-AP conjugate (Sigma). Pigments were separated by TLC. One hundred and fifty-millilitre liquid cultures of Stm3 and Stm3LR3 were grown and harvested as described. Ten-milligram CaCO3 and 5 mL acetone were added to each pellet. The acetone extract was then filtered and applied on silica TLC plates (10 × 20 cm, Sigma). A petroleumisopropanol/water (2000 : 220 : 1) mixture was used as the running solvent for pigment separation. Absorption spectra were taken to confirm pigment identity by resuspending separated pigments in ethanol, centrifuging (5 min 18 000 g) and scanning the absorption spectra of the supernatant (range 350 nm to 750 nm, Varian Cary 50 UV/VIS Bio-Spectrophotometer).

For thylakoid membrane purification, 300 mL of Stm3 and Stm3LR3 cells was grown to late log phase, adjusted to equal OD750nm values, centrifuged (10 min, 2200 g, 4 °C) and washed by re-suspending in 30 mL of buffer A (25 mM HEPES pH 7.5, 1 mM MgCl2, 0.3 mM succrose). The washed cells were then pelleted by centrifugation (10 min, 2200 g, 4 °C), before being re-suspended in 8 mL of buffer A and sheared open by two passes through a French Press (2000 psi, 4 °C). The sample volume was then increased to 30 mL (buffer A) and the thylakoid membranes were precipitated through centrifugation (45 min, 20 000 g, 4 °C). To wash the thylakoid membranes they were re-suspended in 30 mL buffer B (5 mM HEPES, pH 7.5, 10 mM EDTA, 0.3 mM succrose) and pelleted (45 min, 4 °C, 48 000 g). The thylakoid membranes were then re-suspended in 2.65 mL of buffer B and 1 mL buffer C (5 mM HEPES, pH 7.5, 10 mM EDTA, 2.2 mM succrose), to adjust the final sucrose concentration (not including volume of the pellet) to 1.82 mM. This suspension was then dispersed into a Beckman SW32Ti centrifugation tube, before being overlaid with 1.75 mM sucrose solution [half the volume of tube (15.3 mL)], and a further layer (5 mL) of buffer D (5 mM HEPES, pH 7.5, 0.5 mM succrose) before centrifuging (60 min, 100 000 g, 4 °C). The centrifuged thylakoid membrane sample formed a dense
green band at the position of the sucrose cushion step. Upon harvesting, the sample was diluted with 5 volumes of buffer E (20 mM MES, pH 6.3, 5 mM MgCl2, 15 mM NaCl, 10% (v/v) glycerol) to facilitate pelleting of the purified thylakoid membranes on centrifugation (20 min, 40 000 g, 4 °C). The thylakoid membranes were then re-suspended in a minimal volume (~1 mL) of MMNB buffer (25 mM MES, pH 6.0, 5 mM MgCl2, 10 mM NaCl, 2 M Betaine) and flash frozen in liquid N2 prior to storage at –80 °C. LHC proteins were isolated from the thylakoid membranes by solubilizing the samples in 33 mM β-dodecyl maltoside (500 µL final volume) and resolving them on a sucrose density gradient (16 h, ~250 000 g, 4 °C). Sucrose gradients were formed by freeze-thawing centrifuging tubes with 25 mM Mes pH 5.5, 500 mM sucrose, 10 mM NaCl, 5 mM CaCl2, 10 mM NaHCO3, and 0.03% β-dodecyl maltoside according to Hankamer et al. (1997).

Electron microscopy of cell sections

For transmission electron microscopy, cells were grown in TAP medium to an OD750nm of 1.0 in continuous white light (100 µmol/m2s). Samples were pelleted gently at 1000 r.p.m. in 15 mL of culture medium. An equal volume of 20% bovine serum albumin and 20% dextran in phosphate-buffered saline (PBS) was then mixed with the cell pellet. Aliquots of cells, concentrated by centrifugation in heat-sealed micro-pipette tips, were then applied to 200 micron Leica wells and high-pressure frozen using a Leica Empact 2 system. Frozen cells were freeze substituted in 2% OsO4 for a week using the Leica AF5 system. Samples were rinsed with dry fresh acetone at the end of freeze substitution and embedded in epon resin. Resin blocks were sectioned at 80 nm thickness (Leica Ultramicrotome UC6), and sections were stained with uranyl acetate/lead citrate and imaged at 300 keV (FEI F30 electron microscope, equipped with a GATAN 4 × 4K CCD camera).

Acknowledgements

We thank M. Lohr (University of Mainz) for violaxanthin spectra analysis, S. Jansson (Umeå Plant Science Centre) for the LHC-specific antibody and E. Knauth and A. Malnoë for isolating thylakoid membranes, running sucrose gradients and optimizing gel conditions. We also gratefully acknowledge the support of the DFG grant KR1586/2-4 (OK), MU2305/1, 2 (JM) and by the Australian Research Council DP0343130 and DP0452362 (BH).

References


Supplementary Material

The following supplementary material is available for this article:

**Figure S1**  Alignment (CLUSTAL W) of all LHC genes and of the region selected for RNAi. Black: 100% sequences identity, grey shades: sequence homology.

**Table S1**  List of qRT primers (5’–3’orientation).

This material is available as part of the online article from: http://www.blackwell-synergy.com/doi/abs/10.1111/j.1467-7652.2007.00285.x
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