Combined Increases in Mitochondrial Cooperation and Oxygen Photoreduction Compensate for Deficiency in Cyclic Electron Flow in *Chlamydomonas reinhardtii*

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During oxygenic photosynthesis, metabolic reactions of CO₂ fixation require more ATP than is supplied by the linear electron flow operating from photosystem II to photosystem I (PSI). Different mechanisms, such as cyclic electron flow (CEF) around PSI, have been proposed to participate in reequilibrating the ATP/NADPH balance. To determine the contribution of CEF to microalgal biomass productivity, here, we studied photosynthesis and growth performances of a knockout *Chlamydomonas reinhardtii* mutant (*pgrl1*) deficient in PROTON GRADIENT REGULATION LIKE1 (*PGRL1*)–mediated CEF. Steady state biomass productivity of the *pgrl1* mutant, measured in photobioreactors operated as turbidostats, was similar to its wild-type progenitor under a wide range of illumination and CO₂ concentrations. Several changes were observed in *pgrl1*, including higher sensitivity of photosynthesis to mitochondrial inhibitors, increased light-dependent O₂ uptake, and increased amounts of flavodiiron (FLV) proteins. We conclude that a combination of mitochondrial cooperation and oxygen photoreduction downstream of PSI (Mehler reactions) supplies extra ATP for photosynthesis in the *pgrl1* mutant, resulting in normal biomass productivity under steady state conditions. The lower biomass productivity observed in the *pgrl1* mutant in fluctuating light is attributed to an inability of compensation mechanisms to respond to a rapid increase in ATP demand.

INTRODUCTION

Oxygenic photosynthesis is a highly integrated bioenergetic and metabolic process that converts solar energy into chemical energy in land plants, microalgae, and cyanobacteria. During oxygenic photosynthesis, electron transfer reactions operating within thylakoid membranes generate both reducing (NADPH) and phosphorylating (ATP) powers, subsequently used to fuel metabolic reactions of CO₂ fixation in the chloroplast stroma. In natural conditions, photosynthetic organisms face constant environmental changes (illumination, temperature, availability of nutrients, and water, etc.), which may differentially affect the efficiency of electron transfer and metabolic reactions, possibly resulting in imbalances between the production and use of chemical energy. Any disequilibrium between energy supply and demand can damage photosynthetic cells since it can lead to overreduction of photosynthetic electron acceptors and to the generation of reactive oxygen species and finally to photodamaging stress. To avoid negative effects of environmental fluctuations, photosynthetic organisms have developed a set of cellular mechanisms allowing them to fine-tune the supply of energy to the demand (Asada, 1999; Niyogi, 2000; Peers et al., 2009; Peltier et al., 2010).

A key parameter for optimal functioning of photosynthesis is the balance between reducing power (NADPH) and phosphorylating power (ATP). It is generally considered that the reactions of linear electron transfer, which involve both photosystem II (PSII) and photosystem I (PSI) operating in series, generate less ATP than is required for the metabolic reactions of photosynthesis (Osmond, 1981; Kramer and Evans, 2011; Foyer et al., 2012). Moreover, the ATP demand varies depending on the environmental conditions and on the metabolic status. In land plants, the ATP demand increases under high light and CO₂ limitation due to the activity of photorespiration (Osmond, 1981; Munekage et al., 2008). In addition, when exposed to low CO₂ concentration, microalgae and cyanobacteria induce a CO₂ concentrating mechanism (CCM), which requires extra ATP (Karlsson et al., 1994; Fridlyand, 1997; Duanmu et al., 2009; Lucker and Kramer, 2013).

Different mechanisms have been proposed to be responsible for reequilibrating the NADPH/ATP balance and avoiding
overreduction of the NADPH stromal pool and of photosynthetic electron acceptors (Kramer and Evans, 2011). The most studied is cyclic electron flow (CEF), which generates extra proton gradient by recycling electrons around PSI, thus resulting in the supply of extra ATP. This CEF pathway, described as antimycin A sensitive, involves ferredoxin (Fd), PROTON GRADIENT REGULATIONS (Munekage et al., 2002), and PROTON GRADIENT REGULATION LIKE1 (PGRL1) proteins (DalCorso et al., 2008; Iwai et al., 2010), the latter was recently proposed to act as a Fd-quinone reductase enzyme (Hertle et al., 2013). Another CEF pathway, described as antimycin A insensitive, operates in thylakoid membranes of land plants (Joët et al., 2001; Munekage et al., 2002) and microalgae (Ravenel et al., 1994). The latter involves the multiple-subunit NADH dehydrogenase complex (NDH-1) of land plant chloroplasts (Joët et al., 2001; Munekage et al., 2004; Rumeau et al., 2005). The plastidial NDH-1 complex is absent from microalgal species, where a plastidial type II NAD (P)H dehydrogenase (NDA2) catalyzing plastoquinone (PQ) reduction has been suggested to participate in CEF (Jans et al., 2008; Desplats et al., 2009; Peltier et al., 2010).

Mitochondrial respiration may also participate in the reequilibration between the reducing and phosphorylating power within the cell. This has been experimentally supported by the effect of respiratory inhibitors on photosynthesis (Krömer and Heldt, 1991; Krömer, 1995) or by the study of mutants affected in plastidial ATPase (Cardol et al., 2009). Metabolic shuttles such as the malate-oxaloacetate shuttle would export reducing power from the chloroplast toward the cytosol and subsequently from the cytosol to the mitochondria (Scheibe, 2004; Shen et al., 2006). Oxygen photoreduction at PSI, also called Mehler reactions, has long been proposed to supply extra ATP for photosynthesis through pseudocyclic photophosphorylations (Allen, 1975).

Although Mehler reactions were widely studied in the 1970s, mostly using thylakoid preparations, their contribution in vivo is less clear (Kramer and Evans, 2011). The recent discovery of flavodiiron (FLV) proteins in cyanobacteria provided evidence at the molecular level for the existence of special types of oxygen photoreduction processes downstream of PSI (Helman et al., 2003; Allahverdiyeva et al., 2013). However, Mehler reactions and Mehler-like reactions driven by FLVs are different processes, the former generating reactive oxygen species (ROS) (Asada, 1999), but the latter not (Vicente et al., 2002; Helman et al., 2003). While FLV genes showing high homology with cyanobacterial genes are present in microalgal genomes (Peltier et al., 2010), their physiological significance and their possible involvement in Mehler-like reactions remain to be established.

From the screening of an insertional mutant library based on the analysis of chlorophyll fluorescence transients, we recently isolated a Chlamydomonas reinhardtii knockout mutant of the PGRL1 gene. The pgrl1 mutant showed decreased activity of CEF and increased capacity to produce hydrogen under anaerobic conditions (Tolleter et al., 2011). In this work, we investigated physiological adaptations occurring under aerobic conditions in response to the impairment of PGRL1-mediated cyclic electron flow. Based on biomass productivity measurements and on the analysis of different photosynthetic parameters in cells grown under different light regimes and CO2 concentrations, we conclude that under a wide range of environmental conditions, deficiency in CEF is compensated for by the concerted action of different mechanisms, including mitochondrial respiration and oxygen photoreduction, resulting in normal steady state growth. Growth retardation in the mutant was observed only in conditions of high ATP demand (low CO2) and fluctuating illumination. While dispensable in steady state growth conditions, PGRL1-mediated CEF would confer a selective advantage in response to rapid changes in the environment.

RESULTS

Photosynthetic Properties of pgrl1 Grown

Photoautotrophically under Various CO2 and Light Regimes

Photosynthetic activities of the pgrl1 mutant and its wild-type progenitor were measured by means of chlorophyll fluorescence in cells grown photoautotrophically in different conditions by changing CO2 supply (either 0.04% CO2 for “air” or 2% CO2 in air for “CO2”) and light intensity (50 µmol photons m−2 s−1 for low light [LL] and 200 µmol photons m−2 s−1 for high light [HL]). The use of a pulse-modulated amplitude fluorometer enabled the determination of both photosynthetic electron transport rate (ETR; Figures 1A to 1D) and nonphotochemical quenching (NPQ; Figures 1E to 1H). Under most growth conditions (“CO2 LL,” “air LL,” and “CO2 HL”), the ETR of pgrl1 was slightly higher than that of the wild-type progenitor line (Figures 1A to 1C).

Under “air HL” conditions, ETR was diminished in the wild-type line compared with “air LL” conditions, the decrease being much more pronounced in pgrl1 (Figure 1D). The ETR decrease observed in “air HL”-grown pgrl1 cells resulted from a decrease in the PSII yield and an increase in the electron pressure on the PSII acceptor QA monitored by the 1-qP parameter (Supplemental Figures 1C and 1D). Under most growth conditions (“CO2 LL,” “air LL,” and “CO2 HL”), NPQ was slightly lower in pgrl1 but not in the wild type (Supplemental Figures 1C and 1D). Under most growth conditions (“CO2 LL,” “air LL,” and “CO2 HL”), NPQ was slightly lower in pgrl1 than in the wild-type line (Figures 1E to 1G). This effect was previously attributed to a decrease in qE resulting from a lower pH gradient in the mutant in the absence of PGRL1-mediated CEF (Tolleter et al., 2011). Surprisingly, NPQ was strongly increased in pgrl1 under “air HL” compared with the wild-type progenitor (Figure 1H).

Transition to State 2 Is Triggered in pgrl1 under High Light and Low CO2

To determine whether the NPQ increase might result from another component of NPQ, such as state transition (qt), we performed low temperature (77K) chlorophyll fluorescence emission measurements in light-adapted algal samples (Figures 2A and 2B). Under “CO2 HL” conditions, fluorescence spectra of pgrl1 and wild-type lines were similar (Figures 2A and 2C). When cells were grown in “air HL” conditions, a large increase of the 710-nm fluorescence peak was observed in pgrl1 (Figures 2B and 2C). Such an increase in the 710-nm fluorescence peak
indicates that light harvesting complex II (LHCII) is more connected to PSI than to PSII as it is the case in State 2 in response to the phosphorylation of LHCII (Wollman and Delepeilare, 1984). To test this interpretation, we performed immunodetection of phosphorylated LHCII proteins using an antiphosphothreonine antibody (Figures 2D and 2E). When C. reinhardtii cells are in State 2, LHCII proteins CP29, CP26, and LHCP11 are phosphorylated (Fleischmann et al., 1999). A higher level of phosphorylation of LHCII proteins was observed in pgrl1 compared with its wild-type progenitor when cells were grown in “air HL” (Figures 2D and 2E). However, since the 710-nm emission fluorescence peak observed in the mutant is much stronger than previously reported in response to physiological situations (Takahashi et al., 2013), it is likely that the large increase in the 710-nm peak does not solely result from state transition, but also from a decrease in PSI (Delepeilare and Wollman, 1985). This question is addressed below.

**Biomass Productivity of pgrl1 Is Not Affected under Constant Light, but Reduced under Fluctuating Light**

To determine growth properties of the pgrl1 mutant and its wild-type progenitor in different conditions of CO2 supply and illumination, cells were cultivated in 1-liter photobioreactors operated as turbidostats. In this experimental setup, the cell density of the culture was measured by an OD probe and maintained constant by addition of fresh medium. We first determined, in this new setup, light conditions leading to similar effects as previously observed in the flask cultures (Figure 1). LL and HL incident illuminations were increased from 50 to 120 μmol photons m⁻² s⁻¹ (LL) and from 200 to 500 μmol photons m⁻² s⁻¹ (HL), respectively, to take into account differences in light paths between these two setups. Two parameters were determined, ETR capacity (measured at 400 μmol photons m⁻² s⁻¹) and the 77K chlorophyll fluorescence emission peak ratio E₆₈₅/E₇₁₀ (Supplemental Figure 2). In the presence of high CO₂, no difference was noticed between ETR capacities of pgrl1 and wild-type lines, ETR progressively increasing for both strains as the growth irradiance rose (Supplemental Figure 2A). In these conditions, no major change in the fluorescence E₆₈₅/E₇₁₀ emission peak ratio was observed, except transiently when switching from 360 to 500 μmol photons m⁻² s⁻¹ (Supplemental Figure 2C). In the presence of air CO₂ levels, both ETR and E₆₈₅/E₇₁₀ ratio remained mostly constant in the wild-type line, but progressively decreased in pgrl1 as the light intensity increased (Supplemental Figures 2B and 2D). The decrease in ETR was maximal at 500 μmol photons m⁻² s⁻¹ with full ETR activity reestablished after 1 h at 120 μmol photons m⁻² s⁻¹ (LL); a return to the initial the E₆₈₅/E₇₁₀ ratio was observed after 24 h. We conclude from this experiment that the changes in photosynthetic parameters observed in the pgrl1 mutant when grown in “air HL” conditions are reversible.

Specific growth rates were determined from the measurement of fresh culture medium added to the turbidostat to maintain a constant biomass concentration (Figures 3A and 3B). The growth rate measured under “CO₂ HL” (−3 d⁻¹) was higher than under “CO₂ LL” (−1 d⁻¹), but no significant growth difference was observed between pgrl1 and wild-type lines (Figure 3A).
Photosynthesis Is More Dependent on Mitochondrial Respiration

The similar growth performances observed at steady state for both pgr11 and wild-type lines under a wide range of environmental conditions, strongly suggest the involvement of mechanisms that compensate for ATP deficiency due to the impairment in PGRL1-mediated CEF. In the wild-type, PGRL1 amounts were higher at HL than at LL (at high CO2) and higher at low CO2 than at high CO2 (at HL), indicating an increased contribution of CEF in these conditions (Supplemental Figure 3). We observed no increase in NDA2 amounts either in the wild type or in pgr11 (Supplemental Figure 3), indicating that no compensation occurred via upregulation of NDA2 in these conditions. We then

Figure 2. State Transition Analyzed by Low Temperature (77K) Emission Spectra of Chlorophyll Fluorescence and LHCII Phosphorylation in Wild-Type and pgr11 Mutant C. reinhardtii Lines.

Cells were grown photoautotrophically in batch cultures under 200 μmol photons m⁻² s⁻¹ (HL) or 50 μmol photons m⁻² s⁻¹ (LL) in 2% CO2-enriched air. Samples were loaded at equal protein amounts based on Coomassie blue staining. LHCII proteins CP29, CP26, and LHCP11, which are phosphorylated in State 2 conditions but not in State 1, are shown (Fleischmann et al., 1999). (A) and (B) 77K chlorophyll fluorescence emission spectra; wild-type progenitor line (black line); pgr11 cells (gray line). (C) E₄₈₅/E₇₃₀ chlorophyll fluorescence emission ratios measured in light-adapted pgr11 (white bars) and wild-type (dark bars) lines grown under different light intensities and CO₂ concentrations. (D) and (E) Immunodetection of phosphorylated LHCII using an antiphosphothreonine antibody in light-adapted pgr11 and wild-type cells grown in HL in the presence of 2% CO₂-enriched air (D) or in the presence of air (E).

Under air conditions, no difference in growth rates was observed between LL- and HL-grown cells (Figure 3B), showing that growth is limited by CO₂ availability. Again, no growth difference was observed between wild-type and pgr11 lines, despite a decrease in the maximal photosynthetic capacity of the mutant (measured by chlorophyll fluorescence parameters at saturating CO₂; see Figure 1). Growth performances were also assayed on a solid medium by plating serial dilutions of a liquid culture onto Petri dishes (Figures 3C and 3D). While no difference was observed between wild-type and mutant strains under low light (50 μmol photons m⁻² s⁻¹), growth was strongly reduced in pgr11 at high light (200 μmol photons m⁻² s⁻¹), but only in the presence of low CO₂ (air) levels (Figures 3C and 3D). Such a difference in growth performances observed between liquid and solid cultures was quite surprising. It may result from differences in physiological situations experienced by cells in the two modes of cultivation. In liquid cultures, cells were grown under steady state conditions. For the solid tests, cells were plated onto a solid medium from liquid cultures, therefore experiencing a strong change in light intensity. In order to test this hypothesis, low-density liquid cultures (4 × 10⁵ cells mL⁻¹) were adapted to 50 μmol photons m⁻² s⁻¹ for 48 h and then submitted to a sudden light increase to 800 μmol photons m⁻² s⁻¹. Dilution with fresh medium was stopped in order to measure as accurately as possible the biomass increase during the transient (Figures 3E to 3H). At high CO₂ levels, the initial OD increase observed in response to HL was faster in pgr11 than in its wild-type progenitor line, but then resumed at a similar rate (Figures 3E and 3G). Under air, severe growth retardation was observed in the mutant compared with the wild-type line (Figures 3F and 3H). This suggests that growth differences observed on a solid medium are the result of a higher sensitivity of the pgr11 mutant to a LL to HL transient.

To further determine the effect of HL pulses on growth, specific growth rates were measured at 50 μmol photon m⁻² s⁻¹ and in the presence of an additional 1-min HL pulse (50/800 μmol photon m⁻² s⁻¹ 5/1 min) as recently studied in the Arabidopsis thaliana pgr5 mutant (Suorsa et al., 2012). In the presence of high CO₂ levels, both mutant and wild-type strains gained from the additional light supplied by the high light pulse by increasing their specific growth rates (Figure 3I). In the presence of low CO₂ levels (air), only the wild-type line could take advantage of the additional light pulse, the specific growth rate of the mutant remaining unchanged (Figure 3J). We conclude from these experiments that PGRL1-dependent CEF is not essential for steady state growth under a wide range of CO₂ levels and light intensities but is important when HL transients are experienced at low CO₂ concentration.
evaluated to what extent increased cooperation with mitochondrial respiration might contribute. For this purpose, the effect of respiratory inhibitors was assessed on photosynthetic activities (Figure 4). In *C. reinhardtii*, simultaneous addition of inhibitors of cytochrome *bc* and of the alternative respiration pathway is required to fully inhibit the respiratory chain (Cournac et al., 2002). When salicyl hydroxamic acid (SHAM), an inhibitor of the alternative oxidase was added, no significant change in chlorophyll fluorescence was observed. When myxothiazol, an inhibitor of the mitochondrial cytochrome *bc1* complex, was subsequently added, an increase in the stationary fluorescence level (*F*<sub>s</sub>) was observed in *pgrl1* but not in its wild-type progenitor line (Figure 4A), resulting in a drop in the PSII yield (Figures 4D and 4E). Quenching analysis revealed a strong increase of the 1-*qP* parameter in the mutant after treatment with respiratory inhibitors, indicating a higher reduction of PSII electron acceptors (Figures 4B and 4C). The effect was even more pronounced in *pgrl1* mutant cells grown in air (Figures 4C and 4E). We conclude from this experiment that photosynthetic activity of the *pgrl1* mutant is more dependent on cooperation with mitochondrial respiration than is that of its wild-type progenitor line, the cooperation being increased in conditions (such as low CO<sub>2</sub>) where the ATP demand is increased.

**Figure 3.** Growth Performances of *pgrl1* and Wild-Type *C. reinhardtii* Lines Cultivated Photoautotrophically in Liquid or Solid Media. (A) and (B) Growth performances were analyzed in liquid cultures using 1-liter photobioreactors operated as turbidostats. Cell density was measured using an absorption probe and maintained at a constant level (≥1.5 × 10<sup>6</sup> cells mL<sup>-1</sup>) by injection of fresh medium. Growth rates were measured in LL (120 μmol photons m<sup>-2</sup> s<sup>-1</sup>) or HL (500 μmol photons m<sup>-2</sup> s<sup>-1</sup>) in cells grown in the presence of 2% CO<sub>2</sub>-enriched air (A) or air (B). Wild-type progenitor line (dark bars); *pgrl1* mutant (white bars). Shown are means ± se (n = 7 in [A] or n = 3 in [B]). (C) and (D) Growth on solid medium was assessed by plating serial (1/10) dilutions of an algal culture (initial culture concentration 10<sup>6</sup> cells mL<sup>-1</sup>) on a minimal medium under constant LL (50 μmol photons m<sup>-2</sup> s<sup>-1</sup>) or HL (200 μmol photons m<sup>-2</sup> s<sup>-1</sup>) in the presence of air enriched with 2% CO<sub>2</sub> (C) or air (D). Shown are representative cultures out of three biological repeats showing similar effects. (E) to (H) Growth performances of *pgrl1* and the wild type in response to LL to HL switch. The culture was maintained at a low cell density (≥4 × 10<sup>5</sup> cells mL<sup>-1</sup>) by injection of fresh medium during a 48-h period at 50 μmol photons m<sup>-2</sup> s<sup>-1</sup>. When indicated by arrows, light intensity was increased to 800 μmol photons m<sup>-2</sup> s<sup>-1</sup>. The injection of fresh medium was stopped and the OD<sub>800</sub> increase recorded. Wild-type line (E) and (F); *pgrl1* (G) and (H). Transients shown are representative of three independent experiments. (I) and (J) Growth performances of *pgrl1* and wild-type lines under fluctuating light. Low cell density (≥4 × 10<sup>5</sup> cells mL<sup>-1</sup>) cultures were performed at 50 μmol photons m<sup>-2</sup> s<sup>-1</sup> (LL) or under a 50/800 μmol photons m<sup>-2</sup> s<sup>-1</sup> (5 min/1 min) fluctuating light regime (LL+1‘LP). Wild-type progenitor line (dark bars) and *pgrl1* (white bars). Shown are means ± se (n = 2).
Oxygen Photoreduction in a constant biomass concentration (autotrophically in 1-liter photobioreactors operated as turbidostats at 150 μmol photons m⁻² s⁻¹). Net O₂ production was mainly unaffected in all conditions tested, except in HL air conditions, where pgrl1 showed lower activity (Figure 5A). Light-dependent O₂ uptake was higher in the pgrl1 mutant than in the wild-type line when cells were grown in “air LL” (Figure 5B). To determine the contribution of mitochondrial respiration to the light-dependent O₂ uptake process, we used mitochondrial respiratory inhibitors myxothiazol and SHAM (Figures 5C and 5D).

In “HL” grown cells, net O₂ production measured in pgrl1 was more sensitive to respiratory inhibitors than in its wild-type progenitor (Figure 5C). By contrast, the light-dependent O₂ uptake process independent of mitochondrial respiration and most likely resulting from O₂ photoreduction (also called the Mehler reactions) was triggered in the mutant.

O₂ photoreduction or Mehler reactions can result from direct interaction of reduced PSI electron acceptors or reduced Fd with molecular O₂, thus producing superoxide and in turn H₂O₂ (Asada, 1999; Rutherford et al., 2012). In cyanobacteria, and possibly in microalgae, a Mehler-like reaction may also result from the action of FLV proteins that use NADPH as an electron donor and produce water (Vicente et al., 2002; Helman et al., 2003). To gain insight into the nature of the O₂ photoreduction mechanism triggered in the pgrl1 mutant, we first measured extracellular production of H₂O₂ (Figures 6A and 6B). Indeed, wild-type as well as photosynthetic mutant C. reinhardtii lines produce H₂O₂ in the extracellular medium during HL exposure (Allorent et al., 2013). This phenomenon occurs when the electron flow capacity is saturated at the level of PSI acceptors, thus triggering O₂ photoreduction. H₂O₂ production was relatively low in most conditions and increased in “air LL” (when compared with “air LL”) conditions both in the wild-type and in the mutant lines, the increase being much more pronounced in the mutant (Figure 6B). Immunoblots performed with an antibody recognizing both C. reinhardtii FLVA and FLVB proteins showed that both protein amounts are higher in the pgrl1 mutant than in wild-type progenitor in conditions of high CO₂ (Figure 6C) and in air growth conditions, higher FLVB amounts were observed in pgrl1 in comparison to the wild type, FLVA levels remaining unchanged (Figure 6D). It is concluded from these experiments that the decreased sensitivity of O₂ photoreduction to respiratory inhibitors observed in pgrl1 (Figure 5D) may result from two effects: (1) an increase in FLVs (Figures 6C and 6D) and likely in FLV-mediated O₂ photoreduction in most conditions; and (2) an additional increase in direct O₂ photoreduction (Mehler reactions) leading to the production of H₂O₂, the latter occurring essentially in air (Figure 6B).

**Figure 4.** Effect of Respiratory Inhibitors on Photosynthetic Activity of C. reinhardtii Wild-Type and pgrl1 Lines Measured by Chlorophyll Fluorescence. (A) Wild-type progenitor (black) and pgrl1 (red) lines were grown photoautotrophically in 1-liter photobioreactors operated as turbidostats at a constant biomass concentration (≈1.5 x 10⁶ cells mL⁻¹) at a light intensity of 120 μmol photons m⁻² s⁻¹ in the presence air enriched with 2% CO₂. Chlorophyll fluorescence measurements were performed under a light intensity of 150 μmol photons m⁻² s⁻¹ in the presence of 5 mM NaHCO₃. When indicated by arrows, respiratory inhibitors SHAM and myxothiazol were sequentially added at respective concentrations of 0.4 mM and 2 μM. (B) and (C) Chlorophyll fluorescence parameter (1-qP) related to the reduction of Q₀. Control extracellular production of H₂O₂ (Figures 6A and 6B). Indeed, wild-type as well as photosynthetic mutant C. reinhardtii lines produce H₂O₂ in the extracellular medium during HL exposure (Allorent et al., 2013). This phenomenon occurs when the electron flow capacity is saturated at the level of PSI acceptors, thus triggering O₂ photoreduction. H₂O₂ production was relatively low in most conditions and increased in “air LL” (when compared with “air LL”) conditions both in the wild-type and in the mutant lines, the increase being much more pronounced in the mutant (Figure 6B). Immunoblots performed with an antibody recognizing both C. reinhardtii FLVA and FLVB proteins showed that both protein amounts are higher in the pgrl1 mutant than in wild-type progenitor in conditions of high CO₂ (Figure 6C). In air growth conditions, higher FLVB amounts were observed in pgrl1 in comparison to the wild type, FLVA levels remaining unchanged (Figure 6D). It is concluded from these experiments that the decreased sensitivity of O₂ photoreduction to respiratory inhibitors observed in pgrl1 (Figure 5D) may result from two effects: (1) an increase in FLVs (Figures 6C and 6D) and likely in FLV-mediated O₂ photoreduction in most conditions; and (2) an additional increase in direct O₂ photoreduction (Mehler reactions) leading to the production of H₂O₂, the latter occurring essentially in air (Figure 6B).

**Oxygen Photoreduction in pgrl1 Is Less Sensitive to Respiratory Inhibitors**

Light-dependent O₂ exchange was then measured using a membrane inlet mass spectrometer (MIMS) and [¹⁸O]-labeled O₂ (Figure 5). This technique allows measuring O₂ uptake fluxes in the light (Dimon et al., 1988; Beckmann et al., 2009), which may result from different processes, including mitochondrial respiration, photosrespiration, or oxygen photoreduction at PSI (Badger, 1985; Peltier and Thibault, 1985). Net O₂ production was mainly unaffected in all conditions tested, except in HL air conditions, where pgrl1 showed lower activity (Figure 5A). Light-dependent O₂ uptake was higher in the pgrl1 mutant than in the wild-type line when cells were grown in “air LL” (Figure 5B). To determine the contribution of mitochondrial respiration to the light-dependent O₂ uptake process, we used mitochondrial respiratory inhibitors myxothiazol and SHAM (Figures 5C and 5D). In “HL” grown cells, net O₂ production measured in pgrl1 was more sensitive to respiratory inhibitors than in its wild-type progenitor (Figure 5C). By contrast, the light-dependent O₂ uptake process independent of mitochondrial respiration and most likely resulting from O₂ photoreduction (also called the Mehler reactions) was triggered in the mutant.

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**Adaptation Mechanisms Involved in pgrl1 during a Switch from High to Low CO₂**

Adaptation of the pgrl1 mutant to low CO₂ was then investigated during a switch at HL from high CO₂ to low (air) CO₂ concentration (Figure 7). In conditions of low CO₂, C. reinhardtii cells
induce CCM, therefore increasing the ATP demand of photosynthesis (Fridlyand, 1997; Duanmu et al., 2009; Lucker and Kramer, 2013). We first checked CCM induction by monitoring the increase in carbonic anhydrase activity of intact cells (Supplemental Figure 4A) and accumulation of the low-carbon-inducible protein LCIB (Yamano et al., 2010) (Figure 7A). Abundances of different photosynthetic and respiratory components were then determined by immunoblot analysis (Figure 7A). A transient increase in PGR1 amounts was observed in the wild-type control (between 2 and 4 h after the switch), indicating an involvement of the PGR1-mediated CEF to the supply of extra ATP for the CCM. While PSI and PSII amounts, probed by PSAC and PSBD subunits, respectively, increased in the wild type between 4 and 24 h after the switch, almost no change was observed in pgr1 (Figure 7C) (Supplemental Figure 6A). Upon transfer to low CO2, the ΔpH contribution to proton motive force increased significantly in the wild type and remained at a low level in pgr1 (Supplemental Figure 6B). An increase of ECS was observed in pgr1 (Supplemental Figure 6C), which mirrored the decrease in PSI amounts (Figure 7A), showing that the PSI centers that remain active are capable of turning over faster. Finally, the time-resolved growth rate patterns were analyzed during the transient, showing that both strains reached a similar value ~12 h after the switch (Figure 7E). However, the growth rate took longer to stabilize in pgr1 due to the existence of a strong oscillatory regime. We conclude from these data that adaptation of *C. reinhardtii* cells to low CO2 involves a complex set of mechanisms with different time responses. In the wild type, a fast and transient increase in PGR1 amounts is followed by an increase in PSI (Figure 7A). In the pgr1 mutant, the absence of PGR1 increase is compensated for by an upregulation of FLVs (Figures 7B), which is maintained at a higher level than in the wild type, resulting in a higher light-dependent O2 uptake rate. On a longer time scale, FLV amounts decrease and a different set of mechanisms is triggered, including upregulation of LHCSR3, AOX, and a decrease of the PSI/PSII ratio.

**DISCUSSION**

We have shown in this study that steady state growth and biomass productivity of the *pgr1* *C. reinhardtii* mutant are comparable to that of the wild-type progenitor line under a wide range of CO2 concentrations and light intensities. Growth retardation is observed at ambient CO2 concentrations only when the *pgr1* mutant is subjected to fluctuating light conditions. This strongly suggests that alternative mechanisms are triggered in the *pgr1* mutant, efficiently compensating for the ATP deficit resulting from the absence of PGR1-mediated CEF (Tolleter et al., 2011).
Increased Cooperation with Mitochondrial Respiration

Based on the effect of respiratory inhibitors myxothiazol and SHAM on photosynthesis, we conclude that photosynthetic activity in the pgrl1 mutant is more dependent on the activity of mitochondrial respiration than in the wild-type line. Such cooperation between photosynthesis and respiration has been widely documented in the past both in higher plants (Krömer and Heldt, 1991; Krömer, 1995) and microalgae (Lemaire et al., 1988). It has been proposed to result from the activity of metabolic shuttles, such as the malate/oxaloacetate valve, which enable the export of reducing power from the chloroplast to the cytosol, and in turn, from the cytosol to mitochondria. In C. reinhardtii, cooperation with mitochondrial respiration has been proposed to restore photoautotrophic growth in a suppressor strain of a mutant deficient in the plastidial ATPase, by converting reducing power produced in excess into ATP within mitochondria (Lemaire et al., 1988). ATP exchange between cellular compartments is also possible through ATP/ADP translocators or other metabolic shuttles (Hoefnagel et al., 1998). While the mitochondrial ATP/ADP translocator efficiently exports ATP from mitochondria to the cytosol, the DHAP/3-PGA shuttle may be involved in ATP import into the chloroplast (Hoefnagel et al., 1998). However, in contrast with previous works reporting higher COX levels in TAP-grown pgrl1 cells (Petroutsos et al., 2009; Tolleter et al., 2011), no difference in COX levels was observed between pgrl1 and wild-type cells during photoautotrophic growth (Figure 7A). While no difference in dark respiration was measured at high CO2, pgrl1 showed a 30% higher respiration rate than the wild type after 24 h at low CO2 (Supplemental Figure 4B). We therefore conclude that the increased dependence of photosynthesis on mitochondrial respiration observed at high CO2 proceeds via efficient metabolic coupling in the light, but does not result in a sufficient increase in the metabolic pools of respiratory substrates to induce a notable change in dark respiration rates. In spite of a higher dependence of pgrl1 photosynthesis upon cooperation with mitochondria, we observed under HL a decreased sensitivity of O2 photoreduction to mitochondrial respiratory inhibitors (Figure 5D). This phenomenon likely results from the activation of O2 photoreduction mechanisms in the mutant (see discussion below), therefore allowing O2 uptake to be maintained in the light upon inhibition of respiration by chemicals. We also conclude that the increase in AOX levels observed in the mutant at low CO2 (Figure 7A), which takes place much later than the stimulation of O2 photoreduction (Figures 7D), likely reflects a general stress response, as reported in land plants in response to several stress conditions (Millar et al., 2011).

Stimulation of O2 Photoreduction

Based on measurements of light-dependent O2 uptake rates and on the effect of mitochondrial respiratory inhibitors, we operated as turbidostats at a constant biomass concentration (≈1.5 × 10^6 cells mL^−1). The cultures were grown in 120 μmol photons m^−2 s^−1 (LL) or 500 μmol photons m^−2 s^−1 (HL) light intensity and in the presence of 2% CO2 in air (C) or in air (D). The FLVB protein has a size of ≈ 60 kD and FLVA shows the faint band at ≈ 70 kD. Fifteen micrograms of total protein as 100% was loaded per lane, and from the pgrl1 mutant 50 and 25% were loaded as well.

Figure 6. Production of Extracellular Hydrogen Peroxide and Accumulation of FLVs by Wild-Type and pgrl1 Lines Grown under Different Light and CO2 Conditions.

(A) and (B) Measurements were performed on cells grown photoautotrophically in batch cultures supplied with 2% CO2 in air or with air in the presence of LL (50 μmol photons m^−2 s^−1) or HL (200 μmol photons m^−2 s^−1). Hydrogen peroxide concentration was assessed in culture medium using Amplex red by measuring fluorescence emission at 580 nm. Wild-type progenitor (dark bars) and pgrl1 (white bars) lines. Shown are means ± so (n = 3).

(C) and (D) Accumulation of the FLV proteins was determined by immunoblot analysis of wild-type and pgrl1 lines grown in photobioreactors.
concluded that photosynthetic O$_2$ photoreduction mechanisms are triggered in the pgrl1 mutant. Such reactions (also called Mehler reactions) may proceed via direct reduction of O$_2$ by PSI acceptors such as Fd, resulting in the production of reduced O$_2$ forms (Figure 8) that are detoxified by enzymes of the so-called water-water cycle (Asada, 1999). Since a proton gradient is also formed by electron transfer reactions to O$_2$, this mechanism participates in the reequilibration of the ATP/NADPH balance through the functioning of pseudocyclic photophosphorylations (Allen, 1975). FLV-mediated O$_2$ photoreduction may achieve a similar function (Helman et al., 2003; Allahverdiyeva et al., 2011, 2013). Four genes encoding FLV proteins (Flv1, Flv2, Flv3, and Flv4) have been identified in Synechocystis PCC6803. While Flv1 and Flv3 are found in green algae and mosses, but not in higher plants (Zhang et al., 2009; Peltier et al., 2010). C. reinhardtii FLVA and FLVB belong to two clusters containing cyanobacterial Flv1 and Flv2 (for the FlvA cluster) and cyanobacterial Flv3 and Flv4 (for the FlvB cluster) (Zhang et al., 2009; Peltier et al., 2010). The enhanced FLVA and FLVB protein accumulation observed in the C. reinhardtii pgrl1 mutant compared with its wild-type progenitor, together with the decreased sensitivity of light-dependent O$_2$ uptake to mitochondrial inhibitors, strongly suggest that FLV proteins are involved in O$_2$ photoreduction processes downstream of PSI. The absence of FLVA upregulation in pgrl1 in “air HL” (Figure 7B) might result, as for the decrease in PSI protein amounts observed in the mutant (Figure 7A), from the instability of these proteins under photooxidative stress conditions. As shown in Synechocystis cells, FLV-mediated O$_2$ photoreduction does not result in the production of ROS (Vicente et al., 2002; Helman et al., 2003). The increased H$_2$O$_2$ production observed in pgrl1 in “air HL” may

Figure 7. Adaptation of the Photosynthetic Apparatus of pgrl1 and Wild-Type C. reinhardtii Cells to a Switch from High to Low CO$_2$.

Cells were cultivated autotrophically in photobioreactors operated as turbidostats at a constant biomass concentration ($\approx 1.5 \times 10^6$ cells mL$^{-1}$) in the presence of 2% CO$_2$-enriched air under a light intensity of 500 $\mu$mol photons m$^{-2}$ s$^{-1}$. Upon 48 h stabilization, cultures were shifted to air CO$_2$ levels (0 h). Samples were taken at 0, 2, 4, 6, and 24 h after the shift in order to perform immunodetection [A] and [B] and functional analysis [C] to [E].

(A) and (B) Different antibodies raised against PsaC (PSI), PsbD (PSII), PGRL1, NDA2, AOX1 (AOX), COXIIb (COXII), FeSOD, and FLVs were used to decorate immunoblots. Samples were loaded at equal total proteins amounts based on Coomassie blue staining (Control).

(C) PSI/PSII ratio determined from ECS measurements.

(D) Light-dependent O$_2$ uptake rates were measured using a MIMS in the presence of [$^{18}$O]-enriched O$_2$ as in Figure 5; corresponding O$_2$ production rates are shown as Supplemental Figure 5.

(E) Growth performances measured as dilution rates used to maintain the culture at a constant biomass concentration; pgrl1 (gray line) and wild-type control (black line).
therefore indicate that FLV-mediated \( O_2 \) photoreduction is over-challenged in these conditions, thus resulting in true Mehler reactions producing ROS and \( H_2O_2 \) (Figure 8).

**State Transition in \( pgr1 \)**

LHCII phosphorylation and a transition from State 1 to State 2 were observed in the \( pgr1 \) mutant when grown under “air HL” conditions (Figure 2). This phenomenon results from the reduced state of the PQ pool, causing activation of the STT7 kinase that catalyzes phosphorylation of mobile LHCII, which then migrate from PSII (State 1) to PSI (State 2) (Eberhard et al., 2008). Impairments of mitochondrial activity by respiratory inhibitors (Bulté et al., 1990) or by genetic mutations (Cardol et al., 2009) have been shown to induce a highly reduced state of the stromal pools and a transition from State 1 to State 2. A similar phenomenon likely occurs in “air HL”-grown \( pgr1 \) cells and may be explained by the fact that compensating mechanisms (malate valve/mitochondrial cooperation and \( O_2 \) photoreduction) are over-challenged and unable to dissipate the excess of reducing power generated within the chloroplast, therefore resulting in an increase in the reduced state of the PQ pool, phosphorylation of LHCII, and migration of phosphorylated LHCII from PSII to PSI. It is possible that state transition observed in these conditions has a protective role as recently proposed from the study of a \( C. \ reinhardtii \) double mutant affected both in \( \Delta q_E \) and state transition (Allorent et al., 2013). The recent finding that CEF and state transition are independent mechanisms (Terashima et al., 2012; Takahashi et al., 2013) might explain how state transition may have a role in the absence of PGRL1-mediated CEF activity.

**Control of Linear Electron Flow by the Proton Gradient Generated by CEF**

It has been shown that CEF generates a proton gradient that controls linear electron flow at the level of the cytochrome \( b/f \) complex (Avenson et al., 2005; Joliot and Johnson, 2011), thus protecting PSI from photo-inhibition (Suorsa et al., 2012). Under anaerobic conditions, hydrogen production is enhanced in the \( pgr1 \) mutant due to decreased inhibition of cytochrome \( b/f \) activity resulting from a restricted proton gradient (Tolleter et al., 2011). In these conditions, protons are used as electron acceptors thanks to the activity of the plastidial [FeFe] hydrogenase. A similar situation likely occurs in aerobic conditions, but in that case \( O_2 \) is used as an electron sink, electrons generated in excess at the PSI acceptor side being used to reduce \( O_2 \) either in chloroplasts (by direct or FLV-mediated \( O_2 \) reduction), or in mitochondria thanks to the export of reducing power by metabolic shuttles.

**ATP Requirement of Photosynthesis in Land Plants and Microalgae**

Growth and photosynthesis of \( pgr5 \) and \( pgr1 \) \( Arabidopsis \) mutants were reported to be lower than in the wild-type in plants grown in air (Munekage et al., 2002; DalCorso et al., 2008), the decrease being almost totally suppressed when \( pgr5 \) was grown in air enriched with 2% \( CO_2 \) (Munekage et al., 2008). This differs from the phenotype of the \( C. \ reinhardtii \) \( pgr1 \) mutant described here. Indeed, in liquid cultures, no growth difference was observed between the \( C. \ reinhardtii \) mutant and its wild-type progenitor line under steady state growth conditions either at high or at low \( \ CO_2 \) concentration. An explanation to such a difference may be related to the ATP requirement of photosynthesis. In microalgae, the activity of photorespiration is low due to the existence of a CCM. The CCM needs additional energy to function, most likely supplied by ATP (Karlsson et al., 1994; Duanmu et al., 2009). During \( \Delta q_E \) photosynthesis, and in the absence of photorespiration (high \( \ CO_2 \) concentration), each \( \ CO_2 \) molecule requires three ATP and two NADPH to be assimilated, corresponding to an ATP/NADPH ratio of 1.5. The ATP demand of \( \ CO_2 \) fixation is increased in conditions of photorespiration (low \( \ CO_2 \)), the ATP/NADPH ratio increasing up to 1.62 (Foyer et al., 2012). In microalgae, the CCM would require at least one extra ATP per assimilated \( \ CO_2 \) molecule (Fridlyand, 1997). Therefore, the ATP/NADPH requirement ratio would increase up to a value of two (or even more). \( PGRL1 \) amounts increased in the wild type during adaptation to low \( \ CO_2 \), indicating as recently proposed (Lucker and Kramer, 2013), a higher contribution to the increased ATP demand. Increased \( \ CO_2 \) photoreduction was also observed in the wild type during adaptation to low \( \ CO_2 \) (Figure 7D), indicating, as previously proposed (Sultemeyer et al., 1993), a contribution of pseudocyclic photophosphorylation to the energy requirement of the CCM. Note that this was not accompanied by upregulation of FLV proteins at low \( \ CO_2 \) (Figure 7B). Therefore, different mechanisms are triggered in \( pgr1 \) to compensate for the ATP deficiency, including increased cooperation with mitochondrial respiration (Figure 5) and increased \( \ CO_2 \) photoreduction associated with the persistence of high FLV protein levels (Figures 7B and 7D). In this context, the absence of FLV-mediated \( \ CO_2 \) photoreduction in land plants and/or the existence of less efficient cooperation with mitochondria would explain the defect in growth observed in \( Arabidopsis \) \( pgr5 \) (Munekage et al., 2002, 2008) or \( pgr1 \) mutants (DalCorso et al., 2008). Differences in growth phenotypes observed under fluctuating light between land plants and algal mutants may result from similar causes. Indeed, while the \( Arabidopsis \) \( pgr5 \) mutant does not grow under fluctuating light (Tikkkanen et al., 2010), growth of the \( C. \ reinhardtii \) \( pgr1 \) mutant is only slightly affected compared with that of the wild-type progenitor line (Figure 3). The growth decrease observed in the \( C. \ reinhardtii \) \( pgr1 \) mutant does not seem to result from a direct inhibitory effect, but rather from the incapacity of the mutant to take advantage of the HL pulse (Figures 3I and 3J). Additional fluctuating light experiments on the \( Arabidopsis \) \( pgr1 \) mutant and/or on the recently isolated \( C. \ reinhardtii \) \( pgr5 \) mutant (Johnson et al., 2014) will be needed to determine whether such phenotypic differences are related to the target gene (\( pgr1 \) \( v \) \( pgr5 \)) or to the recipient species (\( C. \ reinhardtii \) \( v \) \( Arabidopsis \)).

**Proposed Cascade of Events Occurring in \( pgr1 \) in Response to Increased ATP Demand**

Based on our data, we propose a scenario in which different mechanisms would be sequentially triggered in the \( pgr1 \) mutant in response to increased ATP demand to reequilibrate the
ATP/NADPH imbalance resulting from the absence of PGRL1-mediated CEF (Figure 8). When the ATP demand is relatively low, increased coupling with mitochondria and increased FLV-mediated O₂ photoreduction would allow the dissipation of excess reducing power and supply sufficient ATP, resulting in reequilibration of the ATP/NADPH ratio. However, these mechanisms would not be able to dissipate excess electrons and compensate for the ATP deficit as the ATP demand increases (“air HL”). As a consequence, the redox state of the stromal NADPH pool would increase, thus resulting in increased leakage of electrons to O₂ at PSI and in a more reduced PQ pool, which would in turn activate the STT7 kinase responsible for LHCII phosphorylation and trigger transition to State 2. Increased direct O₂ photoreduction would again reequilibrate the ATP/NADPH ratio, but this would be accompanied by an enhanced production of H₂O₂ therefore resulting in PSI photodamage and a decrease of the PSI/PSII ratio.

We conclude from this study that the CO₂ assimilation machinery is capable of great flexibility and adaptation. The deficiency of PGRL1-mediated CEF can be compensated for at steady state by a set of mechanisms, resulting in similar growth performance and biomass productivity. However, these compensation mechanisms are not perfect. First, they operate on a longer time scale, causing the mutant to exhibit a growth delay during a high light transient or oscillatory growth upon adaptation to low CO₂. Second, they have limited activity and, when overcome, may be accompanied by the production of ROS, which are detoxified to a certain extent, but may induce a decrease in PSI and an impairment of photosynthetic capacity. PGRL1-mediated CEF may therefore have been selected for during evolution as a mechanism allowing rapid adaptation of the photosynthetic electron transport chain in response to sudden changes in the environment.
**METHODS**

**Strains and Growth Conditions**

The *Chlamydomonas reinhardtii* wild-type strain CC124 (mt- nit1 nit2), progenitor of the pgr1 mutant, and the *pgr1*-ko mutant (Tolleter et al., 2011) were grown at 25°C under continuous illumination on minimal culture medium (Harris, 1989). Batch cultures were grown on a rotary shaker in Erlenmeyer flasks (100 mL) placed in a thermostated (25°C) incubator (Multifron; Inforos) under continuous illumination (50 or 200 μmol photons m⁻² s⁻¹) in the presence of air or 2% CO₂-enriched air. For continuous photoautotrophic growth experiment, cells were cultured in four autoclavable 1-liter photobioreactors (BIOSTAT Aplus; Sartorius Stedim Biotech) equipped with a biomass probe (Excell probe, Exner, for measuring OD₅₅₀ by a 2-cm light path) and operated as turbidostats. A regulation system allowed the maintenance of cultures at a constant OD₅₅₀ by injection of fresh medium (Stepdos FEM03TT18RC; KNF). The pH (Easyferm K160; Hamilton) was maintained at a constant value (pH 7.0).

**Methods**

**Pulse Amplitude Modulated Fluorometer (DUAL-PAM-100; Walz).** For Flv immunoblot analysis, total proteins were separated in SDS-PAGE (14% polyacrylamide, without urea), electrophoretically transferred to a polyvinylidene difluoride membrane (Millipore), and blocked with 5% blotting grade blocker (Bio-Rad). FlvB was detected using a purified rabbit antibody prepared against a peptide antigen mix (CKVIAEYSGRDEP and CARKKAAMSGEVAKA) conjugated with keyhole limpet hemocyanin. Due to the high homology, this antibody recognizes also FlvA protein. As a secondary antibody, anti-rabbit horseradish peroxidase was used 1:10,000 and visualized with ECL.

**77K Chlorophyll Fluorescence Spectra**

Low-temperature fluorescence spectra were measured on whole cells at 77K using a SAFAS Xenius optical fiber fluorescence spectrophotometer. Light-adapted cell suspension (1.5 mL at ~1.5 x 10⁶ cells mL⁻¹), cultured in different conditions of illumination and CO₂ supply, was frozen in a liquid nitrogen bath cryostat (Optisat DN; Oxford Instruments). The excitation wavelength was 440 nm, and excitation and emission slits were 10 and 5 nm, respectively.

**Measurement of O₂ Exchange Using a MIMS**

Oₐ exchange was measured in the presence of [¹⁸O]-enriched O₂ using a water-jacketed, thermostated (25°C) reaction vessel coupled to a mass spectrometer (model Prima i8; Thermo Electron) through a membrane inlet system (Tolleter et al., 2011). The cell suspension (1.5 mL) was placed in the reaction vessel and bicarbonate (5 mM final concentration) was added to reach a saturating CO₂ concentration. One hundred micromolars of [¹⁸O]-enriched O₂ (99% ¹⁸O₂ isotope content; Euriso-Top) was bubbled at the top of the suspension just before vessel closure and gas exchange measurements. O₂ exchanges were measured during a 5-min period in the dark, then the suspension was illuminated at 600 μmol photons m⁻² s⁻¹ for 10 min. Isotopic O₂ species [¹⁸O₂] (m/e = 36), [¹⁸O¹⁶O] (m/e = 34), and [¹⁶O¹⁶O] (m/e = 32) were monitored, and O₂ exchange rates were determined as described previously (Coumac et al., 2002). When indicated, mitochondrial respiratory inhibitors myxothiazol and SHAM were added 15 min before starting measurements at final concentrations of 2 μM and 0.4 mM, respectively.

**Production of Extracellular H₂O₂**

Extracellular H₂O₂ was detected as described (Allorent et al., 2013) using the Amplex Red reagent (Invitrogen). Cells were centrifuged once and supernatant was incubated for 30 min in the dark in the presence of the Amplex Red reagent (2.5 μM final concentration) and horseradish peroxidase (0.025 units mL⁻¹; Sigma-Aldrich) forming the fluorescent re sofuran product. Fluorescence emission at 580 nm (excitation 540 nm) was measured on a SAFAS Xenius fluorescence spectrophotometer.

**Accession Numbers**

Sequence data from this article can be found in the GenBank/EMBL databases under the following accession numbers: AOX1 (EMBL EDP02600.1), AOX2 (EMBL EDP06011.1), FLVA (EMBL EDP03485.1), FLVB (EMBL EDP09775.1), LCIB (EMBL ABG38184.1), LHCsR3 (EMBL EDP01087.1), NDA2 (EMBL EDO06450.1), PGR1L (GenBank XP_001700905), PsaC (EMBL AAB17714.1), and PsbD (EMBL P06007.1).

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Figure 1. Photosynthetic Activity of C. reinhardtii Wild-Type and pgr1 Mutant Lines Measured by Chlorophyll Fluorescence.**
**Supplemental Figure 2.** ETR Capacities and 77K Chlorophyll Fluorescence Measured in Wild-Type and pgrl1 during Transients from Moderate to High Light Performed under Two Different CO₂ Concentrations.

**Supplemental Figure 3.** Accumulation of NDA2 and PGR1 Proteins in Wild-Type and pgrl1 C. reinhardtii Lines Grown under Different Environmental Conditions.

**Supplemental Figure 4.** Effect of a High-to-Low (air) CO₂ Concentration Switch on Carbonic Anhydrase Activity and Mitochondrial Respiration Measured on Intact Cells.

**Supplemental Figure 5.** Light-Dependent O₂ Exchange Measured in Wild-Type and pgr1 Cells Shifted from High CO₂ to Low CO₂ (Air).

**Supplemental Figure 6.** Electrochromic Shift (ECS) and Proton Concentration (ΔpH) in pgr1 and Wild-Type C. reinhardtii Cells to a Switch from High to Low CO₂.

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**AUTHOR CONTRIBUTIONS**


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