Nitric Oxide–Triggered Remodeling of Chloroplast Bioenergetics and Thylakoid Proteins upon Nitrogen Starvation in Chlamydomonas reinhardtii

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Starving microalgae for nitrogen sources is commonly used as a biotechnological tool to boost storage of reduced carbon into starch granules or lipid droplets, but the accompanying changes in bioenergetics have been little studied so far. Here, we report that the selective depletion of Rubisco and cytochrome b₆f complex that occurs when Chlamydomonas reinhardtii is starved for nitrogen in the presence of acetate and under normoxic conditions is accompanied by a marked increase in chlororespiratory enzymes, which converts the photosynthetic thylakoid membrane into an intracellular matrix for oxidative catabolism of reductants. Cytochrome b₆f subunits and most proteins specifically involved in their biogenesis are selectively degraded, mainly by the FtsH and Clp chloroplast proteases. This regulated degradation pathway does not require light, active photosynthesis, or state transitions but is prevented when respiration is impaired or under phototrophic conditions. We provide genetic and pharmacological evidence that NO production from intracellular nitrite governs this degradation pathway: Addition of a NO scavenger and of two distinct NO producers decrease and increase, respectively, the rate of cytochrome b₆f degradation; NO-sensitive fluorescence probes, visualized by confocal microscopy, demonstrate that nitrogen-starved cells produce NO only when the cytochrome b₆f degradation pathway is activated.

INTRODUCTION

In most ecosystems, growth of plants and algae is restricted by nutrient availability: Nitrogen and phosphorus are often suboptimal in terrestrial and freshwater ecosystems, while primary production in oceans is limited by phosphorus near the coast and by iron in open oceans. Organisms cope with nutrient limitation by developing acclimation processes that combine general stress responses with changes in gene expression and metabolism that are specific to the limiting nutrient (Davies and Grossman, 1998; Merchant and Helmann, 2012), among which the induction of transporters to optimize the uptake of the limiting nutrient, of scavenging enzymes, and of recycling processes aimed at accessing alternative sources of this nutrient. As a paradigmatic example, cyanobacteria degrade their phycobilisomes to use them as nitrogen and carbon sources when facing the corresponding nutrient limitation (Yamanaka and Glazer, 1980; Collier and Grossman, 1994).

In Chlamydomonas reinhardtii, a unicellular green alga with great metabolic flexibility and for which there are powerful genetic tools (Grossman et al., 2007), acclimation to nutrient limitation has been extensively studied over the years (recently reviewed in Merchant and Helmann, 2012). C. reinhardtii expresses arylsulfatases (Lien and Schreiner, 1975; Schreiner et al., 1975), phosphatases (Quisel et al., 1996), and l-amino acid oxidase (LAO1; Vallon et al., 1993) when starved for sulfur, phosphorus, and nitrogen, respectively. It recycles ~85% of its chloroplast sulfolipids upon sulfur limitation (Sugimoto et al., 2007, 2010) or part of its chloroplast DNA when starved for sulfur or nitrogen (Sears et al., 1980; Yehudai-Resheff et al., 2007). As do other microorganisms facing nutrient shortage, C. reinhardtii undergoes cell cycle arrest and downregulation of photosynthesis when nutrient starved. Upon phosphorus and sulfur deprivation, decreased photosynthesis in C. reinhardtii has mainly been attributed to compromised photosystem II (PSII) activity (Wykoff...
et al., 1998), although this simple causal relationship has recently been revisited (Malnoé et al., 2014).

Nitrogen deprivation is of particular physiological significance for *C. reinhardtii* since, besides triggering acclimation to optimize nitrogen metabolism, it also induces ribosome remodeling and differentiation into sexually competent gametes (Siersma and Chiang, 1971; Martin et al., 1976; Bulté and Bennoun, 1990). Transcript profiling and proteomic studies have illustrated the extensive changes in gene expression undergone by nitrogen-starved *C. reinhardtii* (Miller et al., 2010; Boyle et al., 2012; Longworth et al., 2012), which reflect both the gamete differentiation program and the metabolic changes aimed at nitrogen recycling and carbon storage. This latter process has attracted increasing attention with the recent recognition of microalgae as a most promising source for renewable biofuel production (Wiffels and Barbosa, 2010). Macronutrient limitation is commonly used to divert *C. reinhardtii* metabolism toward biotechnologically valuable end products: Sulfur starvation triggers H$_2$ photoproduction (Ghirardi et al., 2000), while nitrogen depletion boosts the storage of reduced carbon into lipid bodies (Hu et al., 2008). The latter consist mostly of neutral lipids, namely, triacylglycerol, the accumulation of which may further increase in the absence of starch biosynthesis (Wang et al., 2009; Work et al., 2010), even if the competition between these two pathways has been recently challenged (Siaut et al., 2011). Clearly, a better understanding of the changes in the bioenergetics of nitrogen-starved cells is required to get a refined picture of carbon allocation circuits.

The photosynthetic properties of nitrogen-starved *C. reinhardtii* were studied more than two decades ago (Plumley and Schmidt, 1989; Pettier and Schmidt, 1991; Bulté and Wollman, 1992), at a time when knowledge of the thylakoid membrane protein complexity was still limited. A most noticeable feature of *C. reinhardtii* depleted in nitrogen sources in heterotrophic conditions under low light is photosynthetic downregulation due to a specific loss in cytochrome b$_{6f}$ complexes (Bulté and Wollman, 1992), rather than to PSII inactivation as reported for heterotrophic organisms (Wykoff et al., 1998; Philipps Wollman, 1992), at a time when knowledge of the thylakoid membrane protein complexes (LHCII, PSI, PSII, and ATP synthase) remained unaltered (Figure 1A). The selective loss of the cytochrome b$_{6f}$ complex and of Rubisco was not triggered by light since it was observed as well when nitrogen starvation was performed in complete darkness (Figure 2A, left panel). The kinetics of the loss remained similar whether starvation was performed in darkness or in low or high light (120 µE·m$^{-2}$·s$^{-1}$; Figure 2A). However, increasing the light intensity to 120 µE·m$^{-2}$·s$^{-1}$ generated additional photo-inhibitory processes, targeting both PSI and PSII. This is shown by the loss of PsA and D1 proteins (Figure 2A, right panel) and by the changes in the fluorescence patterns of wild-type cells deprived of nitrogen for 34 h under high light, which became similar with and without 3-(3,4-dichorophenyl)-1,1-dimethyleurea (DCMU), no longer showing any variable fluorescence upon illumination and thus demonstrating the loss of PSII activity (Figure 2B).

**RESULTS**

**Upon Nitrogen Starvation, *C. reinhardtii* Becomes Photosynthetically Inactive by Losing Both the Cytochrome b$_{6f}$ Complex and Rubisco**

As previously reported (Bulté and Wollman, 1992), thylakoids specifically lose the cytochrome b$_{6f}$ complex upon nitrogen starvation of a wild-type strain of *C. reinhardtii*, derived from strain 137c (Harris, 2009), when grown heterotrophically (e.g., in the presence of acetate) in aerobic conditions under dim light (5 to 10 µE·m$^{-2}$·s$^{-1}$). In the typical experiment shown in Figure 1A for two strains of *C. reinhardtii* that have a wild-type phenotype for photosynthesis, WT-S24, and the wild-type-like strain *tmtFHB* (for details, see Table 1), all cytochrome b$_{6f}$ subunits, as well as its associated protein PETO (Harnel et al., 2000), dramatically decreased over time, as quantified in Figure 1B for cytochrome f. The loss of the cytochrome b$_{6f}$ complex, the major intersystem electron carrier that transfers electrons from PSII to photosystem I (PSI), leads to a major change in the quantum yield of PSII ($F_{0}$,$F_{s}$)/$F_{m}$, where $F_{0}$, $F_{s}$, and $F_{m}$ represent the initial, stationary, and maximal fluorescence level, respectively; Maxwell and Johnson, 2000; Figures 1C and 1D), as previously observed (Bulté and Wollman, 1992). In parallel, LAO1, a nitrogen scavenging enzyme that we previously identified as a marker of the cell response to nitrogen deprivation (Bulté and Wollman, 1992; Vallon et al., 1993), is induced (Figure 1A). Thus, nitrogen starvation triggers a cytochrome b$_{6f}$-mediated downregulation of both linear and cyclic photosynthetic electron flows in *C. reinhardtii*. As previously observed (Majeran, 2002), Figure 1A also shows that the content in Rubisco decreases upon nitrogen starvation, with the same kinetics as that of the cytochrome b$_{6f}$ complex, further preventing photosynthetic reduction of carbon. By contrast, the accumulation of the other major photosynthetic protein complexes (LHCII, PSI, PSII, and ATP synthase) remained unaltered (Figure 1A). The selective loss of the cytochrome b$_{6f}$ complex and of Rubisco was not triggered by light since it was observed as well when nitrogen starvation was performed in complete darkness (Figure 2A, left panel). The kinetics of the loss remained similar whether starvation was performed in darkness or in low or high light (120 µE·m$^{-2}$·s$^{-1}$; Figure 2A). However, increasing the light intensity to 120 µE·m$^{-2}$·s$^{-1}$ generated additional photo-inhibitory processes, targeting both PSI and PSII. This is shown by the loss of PsA and D1 proteins (Figure 2A, right panel) and by the changes in the fluorescence patterns of wild-type cells deprived of nitrogen for 34 h under high light, which became similar with and without 3-(3,4-dichorophenyl)-1,1-dimethyleurea (DCMU), no longer showing any variable fluorescence upon illumination and thus demonstrating the loss of PSII activity (Figure 2B).

**Nitrogen Starvation Induces Overexpression of the Chlororespiratory Enzymes**

The loss of cytochrome b$_{6f}$ complexes during nitrogen starvation hampers the light-induced oxidation of the plastoquinone (PQ) pool. However, in darkness or dim light (5 to 10 µE·m$^{-2}$·s$^{-1}$), the redox status of the PQ pool is primarily determined by chlororespiration (Bennoun, 1982). This prompted us
to monitor two major chlororespiratory enzymes: the chloroplast-localized type-II NAD(P)H dehydrogenase NDA2 (Desplats et al., 2009) and PTOX2 (for Plastid Terminal Oxidase, isoform 2), the major oxidase involved in chlororespiration (Houille-Vernes et al., 2011). Both behaved opposite to the subunits of the cytochrome $b_{6}f$ complex: They increased about 4-fold over 34 h of nitrogen starvation under dim light (Figure 3A, left panel). This increase was independent of the light regime, still occurring when nitrogen starvation was performed in darkness or at 120 $\mu$E $m^{-2} s^{-1}$ (Supplemental Figure 1). The respective increases in PTOX2 and NDA2 proved largely independent from one another: NDA2 still increased about 3-fold in a $pto\!x\!2^{knockout}$ mutant starved for nitrogen (Houille-Vernes et al., 2011), while PTOX2 increased 2-fold in a knockdown $nda\!2^{RNAi}$ strain (Jans et al., 2008) (Figure 3B). These two strains lost the cytochrome $b_{6}f$ complexes with similar kinetics as in the wild type (Figure 3B). Conversely, a mutant lacking the cytochrome $b_{6}f$ complex still showed a 4-fold increase in NDA2 and PTOX2 ($\Delta pet\!B$), Figure 3A, right panel). Thus, the loss of cytochrome $b_{6}f$ complexes and the upregulation of these two enzymes are independently triggered during nitrogen starvation.

Figure 1. Cytochrome $b_{6}f$ Complexes and Rubisco Are Selectively Lost during Nitrogen Starvation under Low Light (5 to 10 $\mu$E $m^{-2} s^{-1}$).

(A) Whole-protein extracts of cells harvested at indicated time points (0, 4, 10, 24, and 34 h) after the onset of nitrogen starvation, probed with the antibodies indicated between the two panels. To detect all proteins investigated here but also in Figures 3A and 4, the samples were loaded several times on independent gels. WT-S24 (left panel) and strain tmFH8, which expresses tagged versions of MCA1 and TCA1 but is otherwise wild-type (right panel), are shown. Antibodies recognize the major cytochrome $b_{6}f$ subunits (top) or one major subunit of each other photosynthetic protein complex (bottom), whose abundance reflects that of the whole complex.

(B) Quantification of the loss of cytochrome $b_{6}f$ complex in strain WT-S24. Abundance of cytochrome $f$, expressed as a fraction of its initial amount, was quantified from phosphor imager scans of immunobLOTS revealed with $^{125}$I-protein A and normalized to the amount of the invariant CF1 subunit $\beta$ to correct for variations in loading; mean of three independent experiments ± sd.

(C) Kinetics of fluorescence induction of dark-adapted WT-S24 cells at the onset (left panel) and at the end (34 h; right panel) of the nitrogen starvation, recorded in the presence (gray curves) or in the absence (black curves) of DCMU (5 $\mu$M final). Relevant parameters ($F_{\text{m}}, F_{\text{s}},$ and $F_{\text{o}}$) are shown. Curves were normalized to the maximum fluorescence recorded in the presence of DCMU. a.u., arbitrary units.

(D) Change in photosynthetic efficiency of WT-S24 cells during nitrogen starvation assessed by the quantum yield of PSII [$\phi_{\text{PSII}} = (F_{\text{m}} - F_{\text{o}})/F_{\text{m}}$]; mean of four independent experiments ± sd.
above the tent of plastoquinol reoxidation was then measured as the area (Figure 3A). Cells, preilluminated for 2 s to fully reduce the PQ.

Our reference strain Lacks expression of NiR, NaR, and HANT; cannot grow on nitrate nor nitrite

Mutant Strains Used in This Work

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Phenotype</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT 4γ</td>
<td>NIT1 NIT2 mt+</td>
<td>Wild-type for photosynthesis and nitrogen assimilation; grows on nitrate and nitrite</td>
<td>[1]</td>
</tr>
<tr>
<td>nit1-137</td>
<td>nit1-137 mt+</td>
<td>Lacks NaR; cannot grow on nitrate, but grows on nitrite</td>
<td>[1]</td>
</tr>
<tr>
<td>nit2-124</td>
<td>nit2-124 mt+</td>
<td>Lacks expression of NaR, NiR, and HANT; cannot grow on nitrate nor nitrite</td>
<td>[1]</td>
</tr>
<tr>
<td>WT S24</td>
<td>nit1-137 nit2-124 mt−; results of multiple back-crosses of 137c mt+ and mt− strains</td>
<td>Lacks expression of NiR, NaR, and HANT; cannot grow on nitrate nor nitrite</td>
<td>[1]</td>
</tr>
<tr>
<td>21gr</td>
<td>NIT1 NIT2 mt+</td>
<td>Wild-type for photosynthesis and nitrogen assimilation</td>
<td>CC-1690</td>
</tr>
<tr>
<td>M3</td>
<td>∆NIT1 NRT2;2 NRT2;1 NAR2 NIT1::NIT1 constructed from WT-S24 and M4. The resulting strain expresses epitope-tagged versions of MCA1 and TCA1 (Raynaud et al., 2002).</td>
<td>Lacks NiR; cannot grow on nitrate nor nitrite</td>
<td>[2]</td>
</tr>
<tr>
<td>M4</td>
<td>∆NIT1 NRT2;2 NRT2;1 NAR2 NIT1::NIT1 generated by cross WT-S24 and M4. The resulting strain expresses epitope-tagged versions of MCA1 and TCA1 (Raynaud et al., 2002).</td>
<td>Lacks NiR and HANT; cannot grow on nitrate nor nitrite</td>
<td>[2]</td>
</tr>
<tr>
<td>nit4-104</td>
<td>nit4-104 NIT1 NIT2 mt+</td>
<td>Lacks the MoCo cofactor of NaR and XOR; cannot grow on hypoxanthine, nitrate, or nitrite</td>
<td>[3]</td>
</tr>
<tr>
<td>Thr5</td>
<td>nit1-305::NIT1 mt+</td>
<td>Complemented NaR mutant</td>
<td>[4]</td>
</tr>
<tr>
<td>dmm22</td>
<td>nit1-137 nit2-124 mt− [Δcob, Δnd4]</td>
<td>Lacks mitochondrial cytochrome bc1, and NADH complexes.</td>
<td>[5]</td>
</tr>
<tr>
<td>nda2-RNAi</td>
<td>nda2RNAi nit1-137 nit2-124</td>
<td>Knockdown of the NDA2 gene</td>
<td>[6]</td>
</tr>
<tr>
<td>ptxo2</td>
<td>ptxo2::aphVIII nit1-137 nit2-124</td>
<td>Knockout the PTOX2 gene</td>
<td>[7]</td>
</tr>
<tr>
<td>∆petB</td>
<td>nit1-137 nit2-124 mt− ∆petB</td>
<td>Cytochrome b,f mutant deleted for the petB gene</td>
<td>[8]</td>
</tr>
<tr>
<td>ptxo2 [ΔpetB]</td>
<td>ptxo2::aphVIII nit1-137 nit2-124 [ΔpetB]</td>
<td>The ptxo2 mutant carrying a deletion of the petB gene</td>
<td>[7]</td>
</tr>
<tr>
<td>tmF18</td>
<td>mca1-6::MCA1-HA tca1-8::TCA1-Fl nitr1-137 nit2-124-mts-</td>
<td>Expresses tagged versions of MCA1 and TCA1; otherwise wild-type for photosynthesis</td>
<td>[9]</td>
</tr>
<tr>
<td>mH fhsh1-1.2+</td>
<td>mca1-6::MCA1-HA fhsh1-1 nit1-137 nit2-124</td>
<td>Generated by cross mH mt+ × fhsh1-1 mt+</td>
<td>[1]</td>
</tr>
<tr>
<td>mH [clpP-AUU]</td>
<td>mca1-6::MCA1-HA nit1-137 nit2-124 [clpP-AUU]</td>
<td>Generated by cross mH mt+ × [clpP-AUU] mt+</td>
<td>[1]</td>
</tr>
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</table>


*Generated from the cross WT-S24 mt− × 21gr mt+ (CC-1690)

To investigate the functional consequences of the increased content in chlororespiratory enzymes upon nitrogen starvation, we compared, as described by Bennoun (2001), the rate of re-oxidation of the PQ pool in nitrogen-replete conditions or after 34 h of nitrogen starvation. This measurement is best performed in a cytochrome b,f mutant that still shows increased accumulation of chlororespiratory enzymes upon nitrogen starvation (Figure 3A). Cells, preilluminated for 2 s to fully reduce the PQ pool, were returned to darkness for a variable time and the extent of plastoquinol reoxidation was then measured as the area above the fluorescence induction curve in an ensuing illumination. The PQ reoxidation rate increased 2.4-fold after 34 h of nitrogen starvation (Figure 3C). In a ptxo2 [ΔpetB] double mutant, lacking both PTOX2 and the cytochrome b,f complex, the rate of reoxidation of the PQ pool decreased 5-fold after 34 h of nitrogen starvation (Figure 3C), in agreement with the increase in NDA2 during nitrogen starvation that was still observed in the ptxo2 mutant strain (Figure 3B).

Upon Nitrogen Starvation, Most Proteins Specifically Required for the Biogenesis of Cytochrome b,f Complexes Are Lost

Many proteins, hereafter referred to as cytochrome b,f biogenesis factors, are specifically required for the biogenesis of the cytochrome b,f complex. These include proteins involved in the apo- to holoconversion of c-type cytochromes f and b6, respectively, the CCS (for C-type cytochrome synthesis; Xie et al., 1998) and CCB (for cofactor binding on cytochrome b,f subunit PetB; Kuras et al., 2007) factors, as well as trans-acting factors governing the expression of one of its chloroplast-encoded subunits, such as MCA1 and TCA1 that govern the expression of the chloroplast petA gene (maturation/stability and translation, respectively, of the cytochrome b,f subunit encoded by the petA gene; Wostrikoff et al., 2001; Loisel et al., 2008). To detect the latter two, we resorted to the tmF18 strain that expresses epitope-tagged versions of MCA1 and TCA1 (Raynaud et al., 2007; Boulouis et al., 2011). The tmF18 strain, when starved of nitrogen under dim light, lost cytochrome b,f complexes with the same kinetics as WT-S24 (Figure 1A). Notably, the various cytochrome b,f biogenesis factors decreased significantly upon nitrogen starvation in both strains, as shown in Figure 4. MCA1 was even lost earlier than the subunits of the cytochrome b,f, which is consistent with its short half-life in N-replete conditions (Raynaud et al., 2007), but we note that its steady state accumulation increased slightly again toward the end of the experiment. CCS1, CCS5, and CCB1 disappeared with the same kinetics as cytochrome b,f subunits, while TCA1, CCB2, and CCB4 showed a more limited loss compared with other cytochrome b,f biogenesis factors. Thus, when deprived of nitrogen
sources, <i>C. reinhardtii</i> specifically eliminates most of the proteins required for cytochrome <i>b</i><sub>6</sub><i>f</i> activity, whether structural subunits of the protein complex or biogenesis factors.

The Loss of the Cytochrome <i>b</i><sub>6</sub><i>f</i> Biogenesis Factors Does Not Result from a Transcriptional Shutdown but from Their Increased Proteolytic Degradation, Largely Mediated by the FtsH Protease

Most cytochrome <i>b</i><sub>6</sub><i>f</i> biogenesis factors being encoded by nuclear genes, as is the Rieske subunit (PETC), we investigated whether their loss could originate from a decreased transcription, as a part of the extensive transcriptional changes that develop during nitrogen starvation (Miller et al., 2010; Toepel et al., 2011). The expression of LAO1 was, as expected (Miller et al., 2010), rapidly induced in nitrogen-starved cells, while the chloroplast <i>petA</i> mRNA disappeared (Figure 5A) due to the loss of its stabilizing factor MCA1 (Figure 4). By contrast, the level of PETC, CCB3, and MCA1 transcripts remained invariant during the first 24 h of nitrogen starvation (Figure 5B), although the level of their protein products already strongly decreased (Figures 1A and 3). Our results on the accumulation of the PETC mRNA during nitrogen starvation differ from those of Miller et al. (2010), which may be accounted for by differences in time points (24 versus 48 h), in the light regime and in culture conditions. Thus, the decrease in these cytochrome <i>b</i><sub>6</sub><i>f</i> complex biogenesis factors cannot be attributed to an inhibition of gene transcription in the nucleus but rather to some posttranscriptional mechanisms.

We have previously shown that the loss of cytochrome <i>b</i><sub>6</sub><i>f</i> subunits in nitrogen-starved <i>C. reinhardtii</i> cells was a posttranslational event, partially mediated by the Clp protease, as it was delayed in the <i>clpP1-AUU</i> mutant that displays a 4-fold reduced abundance.

Figure 2. Loss of the Cytochrome <i>b</i><sub>6</sub><i>f</i> Complex Does Not Depend on the Incident Light.

(A) Whole-cell protein extracts from WT-S24 deprived of nitrogen under total darkness (left panel) or high light (120 µE·m<sup>-2</sup>·s<sup>-1</sup>, right panel), analyzed as in Figure 1A.

(B) Kinetics of fluorescence induction of dark-adapted (30 min) WT-S24 cells, recorded at the 0- and 34 h-time points of nitrogen starvation in high light. The <i>F</i><sub>m</sub>, <i>F</i><sub>r</sub>, and <i>F</i><sub>0</sub> parameters are shown. a.u., arbitrary units.

Figure 3. Nitrogen Starvation Leads to Increased Chlororespiration.

(A) Steady state accumulation of the chlororespiratory enzymes PTOX2 and NDA2, probed with specific antibodies in WT-S24 (same samples as in Figure 1A) and <i>ΔpetB</i> strains subjected to nitrogen depletion for the indicated time points. A dilution series of the samples starved for 34 h is shown for the ease of quantification. Accumulation of the PSI subunit OEE2 provides a loading control.

(B) Abundance of PTOX2 or NDA2 in strains defective for the expression of the other enzyme, <i>nda2RNAi</i> or <i>ptox2</i>, respectively. The accumulation of diagnostic cytochrome <i>b</i><sub>6</sub><i>f</i> subunits was assessed in the same samples as that of the PSI subunit PsaA (as a loading control). A dilution series of the initial sample is shown for the ease of the comparison.

(C) Percentage of oxidation of the PQ pool in a cytochrome <i>b</i><sub>6</sub><i>f</i>-defective mutant, <i>ΔpetB</i>, or in a double mutant lacking both the cytochrome <i>b</i><sub>6</sub><i>f</i> complex and the terminal oxidase PTOX2, <i>ptox2</i> <i>ΔpetB</i>, in nitrogen-replete (gray curves) and nitrogen-starved (34 h; black curves) conditions. The PQ pool was first fully reduced by a saturating flash, and the number of electron acceptors available for PSIll was measured as a function of the duration of the subsequent dark period. The initial slope of the curve provides a measure of the rate of oxidation of the PQ pool.
prevented in the

of the catalytic ClpP subunit (Majeran et al., 2000). Here, we re-

visited these observations, now extended to the whole set of
cytochrome $b_6$ biogenesis factors and to Rubisco. We also
studied the $ftsh1-1$ mutant defective for FtsH activity because of
an $R_{122}C$ substitution altering an Arg finger essential for its
ATPase and protease activities (Karata et al., 1999). Both $ftsh1-1$
and $clpP-1$-AUU mutations were placed in a background expressing
tagged MCA1 by sexual crossing. The loss of cytochrome $b_6$
subunits was delayed in the $mH$ ($clpP$-AUU) mutant, in agree-
ment with our previous study (Majeran et al., 2000), and fully
prevented in the $mH$ $ftsh1-1$ mutant, as shown in Figure 6
(compared with WT-S24 in Figure 1A). CCB proteins behaved as
the cytochrome $b_6$ subunits, showing delayed degradation in
the $clpP$-AUU mutant and complete preservation in the $ftsh1-1$
mA1 was sensitive to both proteases, being markedly
stabilized by both mutations, so that its increase at a late stage
of nutrient stress was more visible than in the wild type. By
contrast, the soluble stromal protein Rubisco was insensitive to
the inactivation of the membrane-embedded protease FtsH, while
its loss was somewhat delayed in the mutant showing attenuated
expression of the soluble protease Clp. The loss of CCS1 and
CCS5 proved poorly sensitive to the altered activity of either Clp
or FtsH proteases, as expected from the topology of these pro-
teins that are little exposed to the stromal side of the thylakoid
membranes where both proteases are active.

Together, these results highlight the critical role of proteolysis
in the disappearance of cytochrome $b_6$ subunits and biogenesis
factors, which independently are targeted for degradation by
multiple proteases upon nitrogen starvation.

**The Degradations of Cytochrome $b_6$ Subunits, Cytochrome
$b_6$ Biogenesis Factors, and Rubisco Are Triggered by the
Same Signals**

The loss of cytochrome $b_6$ complex subunits is a regulated
process that is no longer observed when *C. reinhardtii* cells are
starved of nitrogen in the absence of acetate as a reduced carbon
source or when mitochondrial respiration is impaired (Bulté and
Wollman, 1992). To understand whether the losses of cytochrome
$b_6$ biogenesis factors and of Rubisco were similarly regulated, we
resorted to three distinct experimental conditions. First, we grew
*C. reinhardtii* WT-S24 and tmFH8 strains in phototrophic con-
ditions (i.e., in minimum medium), before subjecting them to
nitrogen starvation in the absence of acetate. Alternatively, we
prevented mitochondrial respiration by placing WT-S24 and
tmFH8 strains in anaerobic conditions during nitrogen starvation
in the presence of acetate. Finally, we deprived of nitrogen the
$dmu22$ mutant, which is defective for mitochondrial respiration
because of a deletion of the cob gene and a partial deletion of the
nd4 gene (Remacle et al., 2006). As shown on Figure 7, nei-
ther Rubisco nor cytochrome f, taken here as a typical cyto-
chrome $b_6$ complex subunit, nor the cytochrome $b_6$ biogenesis
factors, exemplified by CCB3, CCS5, and MCA1, decreased
significantly upon nitrogen starvation in these three experimental
conditions. Induction of LAO1 could still be detected in all cases
but was delayed and much weaker than when nitrogen starvation
was imposed in the presence of acetate and in aerobic con-
ditions, which suggests that the two phenomena are under
common control. Thus, when ATP production primarily depends
on photosynthesis, *C. reinhardtii* preserves the cytochrome $b_6$
subunits and biogenesis factors as well as Rubisco, even in the

![Figure 4. Cytochrome $b_6$ Biogenesis Factors Are Lost during Nitrogen Starvation.](image)

Accumulation of cytochrome $b_6$ biogenesis factors, assessed in the
same samples as in Figure 1A, using specific antibodies directed against
those proteins that are indicated between the two panels. Accumulation of
MCA1 and TCA1 was probed using antibodies directed against the
(NA for MCA1 and Flag for TCA1).

![Figure 5. Transcriptional Downregulation Is Not the Cause of the Loss of the Nucleus-Encoded Cytochrome $b_6$ Subunits and Biogenesis Factors.](image)

**A** Steady state accumulation of the petA mRNA during nitrogen star-
vation, probed by RNA gel blotting. Accumulation of the psbA mRNA is
shown as a loading control.

**B** Variations in transcript abundance, normalized to the initial value for
MCA1, CCB3, and PETC genes, assessed by quantitative RT-PCR.
Accumulation of the LAO1 transcript was normalized to the amount
reached after 4 h of starvation. a.u., arbitrary units.
Testing the Role of NO in the Loss of the Cytochrome \( b_{\mu} \) Complex

We then wondered which posttranslational modification might independently target the various cytochrome \( b_{\mu} \) subunits and biogenesis factors for proteolytic degradation. Nitrosylations frequently occur on metalloproteins, such as heme binding proteins. They represented a reasonable candidate to target cytochrome \( b_{\mu} \) subunits and biogenesis factors for degradation (see Discussion). To assess the possible involvement of NO in this process, we attempted to modify its concentration during nitrogen starvation and looked for possible changes in the kinetics of photosynthesis inactivation and loss in cytochrome \( b_{\mu} \) subunits, as studied by fluorescence and immunoblotting. We first used sodium nitroprusside (SNP) as a NO producer and 2-(4-carboxyphenyl)-4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (cPTIO) as a NO scavenger molecule. Since SNP slowly releases NO over a long time range (up to 10 h according to Ederli et al. [2009] and Mur et al. [2011]), both molecules were added 1 h after the onset of nitrogen starvation. Compared with the untreated control, \( \Phi_{\text{PSII}} \) decreased faster in cells treated with 1 mM SNP and more slowly in cells treated with 0.1 mM cPTIO (Figure 8A). The contrast between the two treatments was even larger in cells treated three times with SNP or cPTIO 1, 3, and 6 h after the beginning of nitrogen starvation (Supplemental Figures 5A and 5B). These changes in \( \Phi_{\text{PSII}} \) values corresponded to actual changes in the rate of degradation of cytochrome \( b_{\mu} \) and biogenesis factors, as shown in Figure 8B. At a given time point, the level of cytochrome \( f \), Subunit IV, or CCB3 level was lower and higher in SNP- or cPTIO-treated cells than in the control cultures (see quantifications for cytochrome \( f \) in Figure 8C). PSII was not affected by SNP and/or cPTIO addition, as judged from the maximum PSII quantum yield \( \left[ F_{\text{m}}/F_{\text{n}} = (F_{\text{m}} - F_{\text{n}})/F_{\text{n}} \right] \) and the D1 content (Figures 8A and 8B; Supplemental Figure 5A). We also note that Rubisco degradation showed some sensitivity to the addition of SNP and cPTIO at 10 h of nitrogen starvation but that their effect was no longer significant at later time points. That the faster degradation of the cytochrome \( b_{\mu} \) complex in the presence of SNP should be attributed to the release of NO rather than to other effects of the SNP molecule is supported by the antagonistic effect of cPTIO: When SNP and cPTIO were added simultaneously, the loss in cytochrome \( b_{\mu} \), instead of being faster, was even delayed when compared with the untreated control (Figure 8).

To further confirm that the effect of SNP was indeed due to the release of NO, we used S-nitrosoglutathione (GSNO) as an alternative NO donor. Compared with SNP, GSNO induces a much faster burst of NO within the first hour after its addition to a culture (Ederli et al., 2009; Mur et al., 2011), allowing a better temporal control of NO production. GSNO (0.1 mM) was therefore added to the starvation medium, either before any significant cytochrome \( b_{\mu} \) degradation or when this process had already started (i.e., 7 and 15 h after the onset of nitrogen depletion)
The accumulation of cytochrome b$_{6}$/f subunits and biogenesis factors, LAO1, Rubisco, and PTOX2, was studied as in Figure 1A in WT-S24 and tmFH8 strains deprived of nitrogen sources under low light, either in the absence of reduced carbon in the medium (−Ac; left panel) or in anaerobic conditions (−O$_{2}$; cultured with gentle shaking, 50 rpm, in sealed Erlenmeyer flasks; middle panel), or in the respiratory mutant dum22 grown under 80 μE m$^{-2}$ s$^{-1}$ (right panel). The antibodies used are directed against those proteins indicated on the left of the figure, and the accumulation of PsaA provides a loading control.

**C. reinhardtii Cells Starved for Nitrogen Produce NO**

To detect in situ endogenous NO production, we examined *C. reinhardtii* cells by confocal microscopy, after 1 h incubation with the NO-specific fluorescent probe 4-amino-5-methylamino-2',7'-difluoro-7-methoxy-4-methylfluorescein diacetate (DAF-FM DA). This permeant and nonfluorescent molecule enters the cell where it is esterified into the nonpermeant and weakly fluorescent DAF-FM molecule. Treatment with cPTIO (0.1 mM), at the same time point as GSNO, the decrease in Φ$_{PSII}$ observed during the nitrogen starvation was fully prevented (cells depleted of nitrogen for 7 h) or even reversed (cells depleted of nitrogen for 15 h). This suggests that the cytochrome b$_{6}$/f complex is first reversibly inactivated before being degraded. The simultaneous addition of both drugs fully reversed the action of GSNO (Figure 8D), demonstrating that its effect was indeed due to the release of NO.

When TAP-grown cultures were bubbled for 3 min with 90% N$_{2}$/10% NO, a strong green fluorescence signal was readily detected (Figure 9A) that was not restricted to the chloroplast compartment. The weaker green signal within the chloroplast may result either from a lower accumulation of NO and/or DAF DA in this compartment or, more likely, from the reabsorption of the green fluorescence by photosynthetic pigments. Strikingly, most WT-S24 cells starved for nitrogen sources during 20 h also displayed a strong green fluorescence signal (Figures 9A, right panel, 9B, and 10A) that we attribute to the presence of NO, as the signal was largely decreased when cPTIO (0.1 mM) was added to the starvation medium half an hour before the addition of DAF-FM DA (Figure 9B).

To get a statistical view of NO production over a large population, we conducted another starvation experiment in which cells were concentrated 20 times before confocal imaging performed over larger area (5 × 5 tiles). WT-S24 cultures grown in TAP, or cultures deprived of nitrogen for <12 h, almost completely lacked NO positive green fluorescent cells (Figure 10). The percentage of NO-positive cells then increased progressively with the duration of nitrogen starvation, reaching its maximum value of ~90% of the cells after 19 to 21 h of starvation, before decreasing slowly in the next hours (Figure 10B). To further test the relationships between nitrogen starvation, NO evolution, and the degradation of the cytochrome b$_{6}$/f complex, the same experiment was performed on a nitrogen-depleted culture of the nit1-137 mutant strain that lacks nitrate reductase (NaR) and does not lose the cytochrome b$_{6}$/f subunits and biogenesis factors (see below). That strain showed little detectable accumulation of NO over the time course of the experiment, with <15% of the cells exhibiting green fluorescence after 20 h of starvation (Figures 10A and 10B). In addition, when strain WT-S24 was subjected to nitrogen depletion in the absence of acetate (minimal medium), a condition that prevents the loss of the cytochrome b$_{6}$/f complex, almost no NO positive cells were observed after 20 h of starvation (Figure 10A) (i.e., at the time when the accumulation of NO is maximal in cultures starved of nitrogen in the presence of...
acetate). Last, the addition of 0.5 mM NH₄Cl together with the DAF-FM DA probe in a culture deprived of nitrogen for 19 h led to the disappearance of green fluorescence 1 h later (Figure 9B).

Nitrite Accumulation Increases Cytochrome $b_{6}f$ Complex Degradation

In an attempt to identify the source of NO produced upon nitrogen starvation, we investigated the possible contribution of nitrite (NO$_2^-$), a major source of NO through its reduction by a variety of pathways (reviewed in Gupta et al., 2011). Because nitrite reductase (NiR), by reducing nitrite to ammonium, should limit its availability for NO production, we compared the rate of degradation of cytochrome $b_{6}f$ complexes in mutants defective in NiR, when starved in the absence or presence of exogenously added nitrite. We compared the behavior of strains M3, carrying a deletion of the structural NII1 gene encoding NiR and M4, which, in addition, lacks the NTR2;2, NTR2;1, and NAR2 genes encoding components of the high-affinity nitrate/nitrite transporter HAN(i)T (Navarro et al., 2000; reviewed in Galvan and Fernández, 2001). As expected, strains M3 and M4 were unable to grow on nitrite or nitrate as their sole nitrogen source (Figure 11A). They lost the cytochrome $b_{6}f$ complex during nitrogen starvation, as revealed by their $F_{PSII}$ (Figure 11B) and cytochrome $f$ and subunit IV contents (Figure 11C), albeit to a lower extent and with delayed kinetics compared with our reference strain WT-S24. When the nitrogen-free medium was supplemented with 0.1 mM NaNO$_2$ 1, 3, and 5 h after starvation had started, the loss of the cytochrome $b_{6}f$ complex was accelerated in strain M3 (Figures 11B and 11C). By contrast, strain M4 proved insensitive to the addition of nitrite, in agreement with its lack of import ability (Figures 11B and 11C). In these strains, as in WT-S24, PSII activity (measured by...
Fv/Fm) remained unchanged during nitrogen starvation, whether nitrite was added or not. This result supports the view that NO, efficiently produced from nitrite (Sakihama et al., 2002), accelerates the loss of the cytochrome b6f complex.

Genetic Evidence for a Critical Role of Nitrite in the Loss of Cytochrome b6f Complexes

In a second series of experiments, we examined how the metabolism of nitrite in strains differing in their nitrate assimilation pathway correlated with the loss of the cytochrome b6f complex. We compared three groups of strains that were either wild type for nitrate/nitrite assimilation, defective for only nitrate assimilation, or defective for nitrate and nitrite assimilation (Figure 12A).

Our reference strain WT-S24 and all of the photosynthesis mutants used in this and previous studies to characterize the loss of cytochrome b6f complexes upon nitrogen starvation are derived from wild-type 137c. Like their 137c ancestor, they carry two mutations in genes required for nitrogen assimilation: nit1-137, a deleterious substitution that changes the axial ligand H542 of heme b5 to Q in the NIT1 structural gene encoding NaR, and nit2-124, an insertion of the TOC1 transposon into the first exon of the NIT2 gene (N. Tourasse, Y. Choquet, and O. Vallon, unpublished results), encoding the major nitrate/nitrite regulator (González-Ballester et al., 2004; Camargo et al., 2007). The NIT2 gene product is required for the expression of the NIT cluster that includes the gene for NaR, NIT1, together with genes for

![Figure 9.](image1)

**Figure 9.** The DAF-FM Signal Specifically Reflects the Accumulation of NO. Visualization of in vivo NO production by nitrogen-starved C. reinhardtii cells using confocal microscopy. Green and red fluorescence were due to the NO indicator DAF-FM and chlorophylls, respectively.

(A) Fluorescence pattern of isolated WT-S24 cells grown in TAP medium (TAP), grown in TAP medium and bubbled for 3 min with 10% NO (NO bubbled), or deprived of nitrogen for 20 h [-N(20 h)]. For a better view of the distribution of the green fluorescence in NO-bubbled cells, the same picture is shown on the right, with a lower gain during the acquisition of the green signal.

(B) Confocal images of cell populations grown in TAP (TAP) or deprived of nitrogen in the presence of acetate for 20 h (-N). Imaging was also performed on the same starved culture supplemented with either 0.1 mM cPTIO or 0.5 mM NH4Cl.

![Figure 10.](image2)

**Figure 10.** Upon Nitrogen Starvation, C. reinhardtii Cells Produce NO.

(A) Confocal images of cell populations from the WT-S24 strain grown in TAP (TAP), deprived of nitrogen in the presence of acetate for 20 h (-N), or deprived of nitrogen in the absence of acetate (MM-N) or from the nit1-137 strain deprived of nitrogen in the presence of acetate for 20 h. (B) Percentage of NO-positive cells over the time course of nitrogen starvation for the WT-S24 and the nit1-137 strains. For practical reasons, as indicated by the discontinuity of the plot, the first (0 to 18 h) and the second (20 to 25 h) parts represent independent starvation experiments. Error bars reflect, for each time point, the fraction of cells for which the DAF-FM fluorescence level is in between that of NO-negative and NO-positive cells.
nitrate/nitrite assimilation (NiR1, NTR2;1, NTR2;2, and NAR2) (Quesada et al., 1993, 1998; reviewed in Fernandez and Galvan, 2008). Therefore, a nit2-124 strain can grow neither on nitrite nor on nitrate as a nitrogen source, as is the case for a nit1-137 nit2-124 strain, exemplified here by WT-S24 (Figure 12A). These strains rapidly lose the cytochrome b$_6$f complex upon nitrogen starvation (as shown in Figures 1A, 1D, 12A, and 12B). Strains that are wild-type for nitrogen metabolism (i.e., growing on either nitrate or nitrite as a nitrogen source) (Figure 12A) also lose the cytochrome b$_6$f complex but to a lower extent and with delayed kinetics, as exemplified by wild-type 2 in Figures 12A and 12B. We conclude that in a nit2 genetic background, (1) the absence of NiR, a potent sink for nitrite, allows enhanced accumulation of nitrite that feeds the production of NO, resulting in the rapid degradation of the cytochrome b$_6$f complex; and (2) NaR is dispensable for NO production in the absence of NiR.

We then examined strains nit1-137 and nit4-104, which can grow on nitrite but not on nitrate because they lack only an active NaR, being respectively mutated in the NIT1 structural gene or in the CNX1E gene whose product is required for the biogenesis of the Molybdopterin cofactor (MoCo) (Chamizo-Ampudia et al., 2013). In contrast with the strains above, these two NaR-defective strains did not lose cytochrome b$_6$f complexes upon nitrogen starvation, as shown by the preserved abundance of cytochrome f, even over a longer time scale (Figure 11A). Accordingly, they showed a very limited decrease in Ψ$_{PSII}$ (Figure 12B). Similar results were obtained with other NaR-deficient strains, listed in Supplemental Table 1. Moreover, complementation of the nit1-305 mutant by the wild-type NIT1 gene (strain Tff5 in Kindle et al., 1989) restored active degradation of cytochrome b$_6$f complexes (Supplemental Figure 7). The nit1-137 (Figures 10A and 10B) and nit4-104 strains (data not shown) also produce very little NO when examined by confocal microscopy. Thus, three types of behavior can be observed: (1) no loss of cytochrome b$_6$f in NaR-deficient nit1, nit4, or nit7 mutants, (2) delayed loss in strains possessing both NaR and NiR, and (3) rapid loss in nit2 mutants lacking the whole nitrate/nitrite uptake and reduction pathways. In genetic crosses, these phenotypes consistently cosegregated with the corresponding genotypes, as detailed in Supplemental Tables 1 and 2. We conclude that the activity of NaR, although dispensable for NO production in the absence of NiR, is required when this nitrite sink is active. In the latter case, it helps keep the level of nitrite, formed at the expense of nitrate, high enough to sustain NO production.

**DISCUSSION**

**Nitrogen Starvation Remodels Thylakoid Membranes into a Matrix for Catabolic Processes**

When deprived of nitrogen, *C. reinhardtii* kept in mixotrophic conditions (in the presence of acetate) undergoes growth arrest after two rounds of cell division (Siersma and Chiang, 1971; Martin and
Goodenough, 1975) while losing its ability to fix carbon. This occurs through two coregulated processes: the selective depletion of Rubisco, the key enzyme of the Benson-Calvin cycle, a phenomenon that is also observed in sulfur-starved cultures of *C. reinhardtii* (Zhang et al., 2002), and the degradation of an essential component of the photosynthetic electron transfer chain, the cytochrome b$_6$f complex, required both for NADPH production through photosynthetic linear electron flow and ATP production through cyclic and linear electron flows. Thus, *C. reinhardtii* cells adapt their metabolism to nitrogen shortage by inhibiting their ability to reduce carbon. This regulation, ultimately aimed at maintaining balanced availability of nitrogen and carbon resources, does not result from an overflow/imbalance in the photoproduction of NADPH or ATP, as it is independent of the light regime, of the photosynthetic activity, and of state transitions. It is counteracted only by a block in mitochondrial respiration or by a drop in the intracellular flux of reduced carbon when acetate is omitted from the starvation medium, two conditions where the viability of *C. reinhardtii* rests exclusively on photosynthesis. Similarly, iron-limited *C. reinhardtii* cells maintain their photosynthetic apparatus when kept in phototrophic conditions, whereas photosynthesis decreases and respiration is prioritized in cells grown in the presence of acetate in the medium (Terauchi et al., 2010). Remarkably, *C. reinhardtii* has developed a conditional regulation of its photosynthetic properties to preserve its viability. During nitrogen starvation in mixotrophic conditions, the bioenergetic contribution of the thylakoid membranes changes dramatically since the decreased photosynthesis comes along with a marked increase in chlororespiration. The latter change was suggested about two decades ago on functional and molecular bases (Peltier and Schmidt, 1991), the latter of which turned out to be erroneous since mitochondrial cytochromes (Atteia et al., 1992) were mistaken for chlororespiration components. Here, we demonstrate that, in parallel with the block in photosynthetic electron transfer due to the loss of cytochrome b$_6$f complexes, the actual chlororespiratory enzymes NDA2 and PTOX2 markedly

**Figure 12.** The Kinetics and Extent of Cytochrome b$_6$f Complex Degradation Depends on the Genotype of the Strain with Regard to Nitrogen Metabolism.  
(A) Growth characteristics and accumulation of cytochrome f and LAO1 during nitrogen starvation of strains WT 4γ (*NIT1 NIT2*), nit1-137, nit4-104, nit2-124, and WT-S24 (nit1-137 nit2-124). Accumulation of the PSII subunit D1 is shown as a loading control. For nit1-137 NIT2 and nit4-104 NIT2 strains, the starvation was performed over 60 h to assess the stability of the cytochrome b$_6$f complex and the delayed and limited induction of LAO1.  
(B) Change in photosynthetic electron transfer (εPSII) in the same strains starved for nitrogen sources.
increased, leading to increased electron flux through chlororespiration, as we measured here by fluorescence techniques. However, the increase in chlororespiration is not mechanistically triggered by the degradation of the cytochrome \( b_6f \) complex since it was still observed in mutants lacking this complex or in respiratory mutants that do not lose the cytochrome \( b_6f \) complex. The increased chlororespiration has two major functional consequences. First, it contributes to counteract acetate assimilation and to limit starch/lipid storage by stimulating a chloroplast oxidative pathway. From the point of view of biofuel production, tuning down chlororespiration may be worth exploring in the future. The increase in PTOX2 also provides a safety valve, protecting PSII from low-light photoinhibition by plugging additional electron acceptors after the PQ pool. Oxygen thus becomes a genuine electron sink at low light in cytochrome \( b_6f \)-defective conditions (i.e., after 10 to 20 h of nitrogen deprivation). Yet, the chlororespiratory pathway is not efficient enough to prevent PSI and PSII photodestruction at higher light intensities, as documented in this study.

Cytochrome \( b_6f \) Biogenesis Factors: A Protein Network Specifically Sensitive to the Nitrogen Status

In earlier studies (Xie et al., 1998; Wostrikoff et al., 2001; Kuras et al., 2007; Raynaud et al., 2007; Loiselay et al., 2008), we identified several proteins that participate exclusively in the biogenesis of cytochrome \( b_6f \) complexes. Here, we show that most of them behaved as did the cytochrome \( b_6f \) subunits: They are degraded upon nitrogen starvation, and this degradation process is blocked when nitrogen-starved cells cannot perform mitochondrial respiration or are kept in photoautotrophic conditions. This concerted proteolysis is under the control of the FtsH protease with some specificity to the subunits: They are degraded active on the luminal side of the thylakoid membrane where the transmembrane CCS1 and CCS5 proteins protrude. That cytochrome \( b_6f \) or the COB proteins behave as genuine substrates for the FtsH protease upon nitrogen starvation is consistent with its role in the degradation of misassembled cytochrome \( b_6f \) complexes in nitrogen-replete conditions and in the degradation of CCB4 in the absence of CCB2 (Malnoë et al., 2014). In our attempts to identify a mechanism responsible for the degradation of such a diverse array of soluble and integral proteins, we were able to exclude a cascade hypothesis whereby the loss of one protein would trigger the degradation of the other cytochrome \( b_6f \) subunits or biogenesis factors, as was shown for the increased degradation of the various subunits of a photosynthetic protein complex in mutants lacking only one of its subunits (Wollman et al., 1999; Choquet and Vallon, 2000). As an alternative, we looked for posttranslational modification that would target these proteins for degradation.

NO Is Involved in the Remodeling of Photosynthesis during Nitrogen Starvation

The increase in PTOX2 upon nitrogen starvation was reminiscent of the behavior of the homologous alternative oxidase in mitochondria, which is triggered by NO-induced inhibition of cytochrome oxidase (Huang et al., 2002). In many biological systems, nitrosylation participates in signaling (reviewed in Astier et al., 2011, 2012) or induces protein degradation (Souza et al., 2000, Kim et al., 2004; Lee et al., 2008; Y. Wang et al., 2010; Wei et al., 2011; Tang et al., 2012; Jaba et al., 2013). Here, we reasoned that most of the cytochrome \( b_6f \) subunits and biogenesis factors belong to heme binding complexes or participate in the biogenesis of heme binding proteins, which are particularly prone to nitrosylation (Thomas et al., 2003; Angelo et al., 2008). Indeed, in a recent proteomic study, several cytochrome \( b_6f \) subunits were found to be nitrosylated in \( C.\ reinhardtii \) (Morisse et al., 2013).

In agreement with this hypothesis, confocal microscopy with an NO-sensitive fluorescence probe allowed us to detect NO production, but only in \( C.\ reinhardtii \) cells losing the cytochrome \( b_6f \) complex (i.e., starved for nitrogen in normoxic conditions and in the presence of acetate). By contrast, \( C.\ reinhardtii \) deprived of nitrogen in phototrophic or anoxic conditions or with a genetic background that did not lead to cytochrome \( b_6f \) degradation produced little, if any, NO upon nitrogen starvation. Furthermore, treatments increasing NO levels, such as addition of NO donors or nitrite in the starvation medium, increased the rate and extent of cytochrome \( b_6f \) complex degradation, whereas NO scavengers had the opposite effect.

Intracellular Nitrite Is the Most Likely Source of NO Produced during Nitrogen Starvation

How NO is generated in nitrogen-starved \( C.\ reinhardtii \) (i.e., in the absence of any external sources of nitrogen) remains to be elucidated. There are two well-identified intracellular sources of NO (reviewed in Besson-Bard et al., 2008; Wilson et al., 2008; Gupta et al., 2011). The first one is \( l \)-Arg, converted into NO and \( l \)-citrulline by the nitric oxide synthase (NOS). This is the main NO-producing pathway in animals, but its relevance in plants is still a matter of debate (reviewed in Zemojtel et al., 2006; Fröhlich and Durner, 2011). To date, a NOS homolog has been found in the green alga \( O.\ tauri \) (Foresi et al., 2010) but not in other algae nor in land plants. Furthermore \( l \)-Arg plays little role in NO production under nitrogen replete conditions, whether in \( C.\ reinhardtii \) (Sakihama et al., 2002) or in the closely related alga \( S.\ obliquus \) (Mallick et al., 2000b). Nevertheless, the NO-dependent repression of the nitrate assimilation pathway by ammonium is partially prevented upon addition of the \( l \)-Arg analog \( l \)-NAME, a known NOS inhibitor, suggesting some NOS-like activity in \( C.\ reinhardtii \) (de Montaigu et al., 2010). In any event, we could not test the effect of \( l \)-NAME on the loss of the cytochrome \( b_6f \) complex during nitrogen starvation because the drug was efficiently used as a source of nitrogen by \( C.\ reinhardtii \) (data not shown).

Reduction of nitrite thus seems to be the major source of NO in plants (Mallick et al., 1999, 2000a). NO is produced by a variety of pathways identified in several recent studies (reviewed in Gupta et al., 2011), competed by the reduction of nitrite to ammonium by NiR. These pathways are either associated with the plasma membrane (Stöhr et al., 2001; Eick and Stöhr, 2012) or localized in peroxisomes (Barroso et al., 1999; Corpas et al., 2001, 2009; Del Río, 2011), mitochondria (Tischner et al., 2004; Gupta et al., 2010; Gupta and Igamberdiev, 2011), chloroplasts (Jasid et al., 2006; Galatro et al., 2013; Tewari et al., 2013), or the cytosol. The cytosolic NaR appears to be critical for the nitrite-dependent
formation of NO since it provides the substrate to other nitrite-dependent NO-producing pathways by catalyzing the reduction of nitrate to nitrite, in addition to its intrinsic nitrite to NO reduction activity under hypoxic conditions (Dean and Harper, 1988; Yamasaki and Sakihama, 2000; reviewed in Meyer et al., 2005).

A number of our experiments support a predominant role of intracellular nitrite to NO production, as illustrated on the summary scheme shown in Figure 13. Addition of nitrite to NiR mutants was as efficient at accelerating the degradation of the cytochrome $b_{6f}$ complex as was the addition of an NO donor to our reference strain WT-S24. Furthermore, the kinetics and amplitude of the degradation of the cytochrome $b_{6f}$ complex correlate well with the predicted size of the nitrite pool in different genetic backgrounds. Nitrite accumulation should be very low in $nit1$ or $nit4$ mutants showing hampered nitrite production in the absence of NaR, while nitrite is still consumed by NiR. Accordingly, $nit1$ cells accumulate very little NO, as seen by confocal microscopy. Nitrite pools should increase in the wild type where, in spite of an active NiR, NaR is also active. In $nit2$ strains lacking both NiR and NaR activities, nitrite produced by NaR-independent pathways may accumulate. It is of note that in the normoxic conditions required to observe the degradation of the cytochrome $b_{6f}$ during nitrogen starvation, NO can be converted back to nitrite as a result of the chemistry of the GSNO:GSH system (Singh et al., 1996), as well as spontaneously by reacting with ambient O$_2$ or because of the NO oxidase activity of truncated hemoglobins. Among the 12 truncated hemoglobins found in C. reinhardtii (Hemschemeier et al., 2013a), THB2 is specifically induced more than 20-fold at the transcriptional level during nitrogen starvation (Sabeeha Merchant, personal communication). Thus, the status of intracellular nitrite and NO are intimately linked as part of a metabolic cycle and nitrite could be considered as a stable reservoir for NO production, at least in the absence of the NiR sink.

Based on our results, NaR activity is required in the presence of NiR to keep the concentration of nitrite high enough for NO production. Yet, the phenotype of $nit2$ cells shows that the NaR catalytic activity per se is not required for productive NO synthesis during nitrogen starvation, as was also observed in anoxic conditions (Hemschemeier et al., 2013a, 2013b). NaR activity appears also dispensable for nitrite-dependent NO synthesis in Scenedesmus (Mallick et al., 1999) but seems required for NO evolution from nitrite by C. reinhardtii cells grown in nitrogen-replete conditions (Sakihama et al., 2002). However, NO production by NaR-defective mutants has been documented during hypoxia (Hemschemeier et al., 2013a) in mastoparan-treated cells (Yordanova et al., 2010) or in iron-limited cultures exposed to CO (Liping et al., 2013). Together, these results point to the prevalence of alternative NO-producing pathways in C. reinhardtii, probably coupled to the increased catabolism of amino acids and nucleotides undergone by nitrogen-starved cells. For instance, in many organisms, including vascular plants, xanthine oxidase-dehydrogenase (XOR), aldehyde oxidase (AO), and sulfite oxidase (SO), all MoCo-requiring enzymes, can reduce nitrite to NO (Tewari et al., 2009; B.L. Wang et al., 2010; Maia and Moura, 2011), even under normoxia, when using NADH as a substrate (Li et al., 2004). In that respect, we noted that the induction of LAO1, which displays the same regulation pattern as the degradation of cytochrome $b_{6f}$ and Rubisco, as it is much delayed and limited in genetic or environmental contexts that prevent the degradation of these proteins, was even more delayed in strain $nit4$-104 lacking the four enzymes NaR, AO, XOR, and SO, because of the absence of MoCo, than in strain $nit1$-137 lacking NaR only. Interestingly, both XOR and AO are upregulated about 5-fold at the mRNA level during nitrogen starvation (Miller et al., 2010; Sabeeha Merchant, personal communication), probably to cope with the increased purine catabolism resulting from the extensive mobilization of RNA and chloroplast DNA.

**The Production of NO Is Part of a Signaling Pathway That Controls the Degradation of Cytochrome $b_{6f}$ and Biogenesis Factors**

At this stage, we cannot yet assess properly the respective contributions to the loss of the cytochrome $b_{6f}$ complex of (1) protein posttranslational modifications (nitrosylation) that would target them for degradation and (2) activation of a NO signal transduction pathway. NO has been involved in a number of signaling pathways in C. reinhardtii, including acclimation to anaerobiosis (Hemschemeier et al., 2013a), iron homeostasis (Liping et al., 2013), cell death (Yordanova et al., 2010), copper stress (Zhang et al., 2008), and regulation of nitrogen assimilation...
(De Montaigu et al., 2010; Sanz-Luque et al., 2013). Here, the coregulation of cytochrome b6f degradation and nucleus-encoded LAO1 induction upon nitrogen starvation, both being slowed in nit1 mutants and accelerated in nit2 mutants, suggests that NO acts as a signaling molecule at least for the transcriptional induction of LAO1. Actually, NO bubbling or SNP addition proved insufficient by themselves to trigger the degradation of the cytochrome b6f complex in anoxic or phototrophic conditions in absence of nitrogen (data not shown). These observations suggest the recruitment of other components in a signaling pathway, besides protein nitrosylation, to activate the degradation of cytochrome b6f complex and biogenesis factors, as is typical in many stress-induced responses. This signaling pathway may include the nitrogen starvation-induced transcription factor NITROGEN RESPONSE REGULATOR1 (Boyle et al., 2012) or the chloroplast ortholog of the PII protein (Hsieh et al., 1998; Ermiëlova et al., 2013), which in (cyanobacteria signals the nitrogen status under symbiotic conditions by α-ketoglutarate (reviewed in Ninfa and Jiang, 2005). It may also involve CYG11, one of the NO-responsive guanylate cyclases that were found in the nuclear genome of C. reinhardtii (De Montaigu et al., 2010), whose expression is increased 20-fold during nitrogen starvation (Sabeeha Merchant, personal communication). Last, the signaling pathway may include some adaptor proteins of proteases that recognize and bind both the substrates for degradation and the chaperone subunit of the protease (reviewed in Kirstein et al., 2009). For instance, in cyanobacteria, the NbA adaptor mediates the proteolytic degradation of phycobilisomes during nitrogen or sulfur starvation because of its double affinity for phycobilisome rods and for CipC (Collier and Grossman, 1994; Karrad et al., 2008).

This study shows how the extensive response of C. reinhardtii to N starvation is finely tuned by genetic and metabolic factors, some of which have been largely ignored thus far. As NO regulates photosynthesis shutdown that has deep implications for the intracellular allocation of carbon, it becomes critical to understand its signaling effects and to manipulate its production for better harnessing microalgae for biofuel production. A systematic study of nitrosylated proteins (focusing on cytochrome b6f subunits and biogenesis factors and Rubisco) in the experimental conditions or genetic contexts that do or do not lead to cytochrome b6f degradation upon nitrogen starvation will help identify the mechanisms that target proteins to proteolysis, while looking for changes in gene expression in the same conditions should provide clues to the signal transduction pathway.

METHODS

Strains, Media, Culture Conditions, and Chemicals

Wild-type and mutant strains (listed in Table 1) of Chlamydomonas reinhardtii were grown on a rotary shaker (120 rpm) in TAP medium, pH 7.2 (Harris, 1989) under continuous light (5 to 10 μE m−2 s−1), except for the dunn22 mutant that was grown in the same conditions but under 75 μE m−2 s−1. For nitrogen starvation experiments, cells pregrown either in TAP or in minimum medium (4.1 mM KH2PO4, 2.6 mM KH2PO4, 0.3 mM CaCl2, 0.4 mM MgSO4, and 1/1000 Huntner trace solution) without bubbling of ambient air nor extra CO2 up to the mid log phase (2 × 106 cells mL−1) were centrifuged at 3000g for 5 min, washed once in nitrogen-free medium (Harris, 1989), resuspended at 2 × 106 cells mL−1 in nitrogen-free medium (150 mL in a 500-mL Erlenmeyer), and kept on a rotary shaker with vigorous aeration (225 rpm) to ensure a good aeration. Strains m4 fsh1-1.2+ and m4 clpP-AUU231+ were generated by crossing, according to Harris (1989), the strain m4 mt+ (Boulois et al., 2011) with strains fsh1-1 mt+ (Malnoë et al., 2014) and clpP-AUU2 mt+ (Majeran et al., 2000) and selecting the desired progeny by appropriate screens. Strain ptxo2Δ (ΔpetB) was obtained by crossing the ptxo2 mt− strain (Houille-Vernes et al., 2011) with strain (ΔpetB) mt+ (Kuras and Wollman, 1994), cPTIO, SNP, and DAF-2 FM DA were purchased from Sigma-Aldrich, cPTIO was purchased from Invitrogen Life Technologies, and GSNO was obtained from BioVision. GSNO solutions were prepared extemporaneously, and GSNO concentration was determined spectrophotometrically using molar extinction coefficient of 920 M−1 cm−1 at 335 nm.

Protein Preparation, Separation, and Analysis

Protein isolation, separation, and immunoblot analyses were performed as described (Kuras and Wollman, 1994), except that, for detection of the CCB1, CCB2, and CSS5 factors, proteins were transferred onto polyvinylidene difluoride rather than onto nitrocellulose membranes. Cell extracts were loaded on an equal chlorophyll basis. At least three biological replicates were performed for each experiment. Proteins were detected by ECL. Primary antibodies, diluted 100,000-fold (antibodies against cytochrome f, LAO1, D1, LHClI, and PsaA), 50,000-fold (CF1s and RbcL), 10,000-fold (cytochrome b6, Rieske, cytochrome b6f subunit IV, and NDA2), 5000-fold (CCS5, PTOX2, and PETO2), 2500-fold (CCB3 and CCS1), or 1000-fold (CCB1,2,4) were revealed by an horseradish peroxidase-conjugated antibody against rabbit IgG (Promega). Antibodies against D1, PsaA, and Rubisco LS were purchased from Agrisera. Other antibodies were described previously: cytochrome b6f subunits (Kuras and Wollman, 1994), ATP synthase subunit β (Draper et al., 1992), CCBs (Kuras et al., 2007), CCS1 (Dreyfuss et al., 2003), CCS5 (Gabilly et al., 2010), PTOX2 (Houille-Vernes et al., 2011), and NDA2 (Desplats et al., 2009). TCA1-Flag and MCA1-Flag were detected by ECL using monoclonal antibodies against the tags anti-Flag M2 (Sigma-Aldrich), anti HA.11 (Covance), and horseradish peroxidase-conjugated antibody against mouse IgG (Promega). Protein accumulation (normalized to that of the D1 subunit of PSII as internal standard) was, when required, quantified from ChemiDoc XRS+ (Bio-Rad) scans of the membrane, using the ImageLab (v3.0) software. Cytochrome f accumulation, normalized to that of the CF1β subunit as an internal standard, was quantified from phosphor imager (Molecular Dynamics) scans of immunoblots revealed with 125I protein A, using the ImageQuant software, as described (Choquet et al., 2003).

RNA Isolation and Analysis

RNA extraction and RNA gel blot analysis were performed as described (Draper et al., 2002) with probes derived from coding sequences (Eberhard et al., 2002). When preparing the RNA templates for quantitative RT-PCR experiments, AIA was omitted from the lysis buffer. After extraction, 40 μg of RNA was resuspended in 200 μL of DNaseI buffer and treated at 37°C with DNase I (New England Biolabs; 25 units) before being further purified using the Qiagen RNaseasy MinElute Cleanup Kit according to the manufacturer’s instruction. Reverse transcription was done on 1 μg of total RNA with the SuperScript III first-strand synthesis SuperMix for quantitative RT-PCR kit (Invitrogen) according to the manufacturer’s protocol using the random primers included in the kit. Quantitative PCR was performed in a GeneRode 3000 apparatus (Corbet, Qiagen) in a final 25-μL reaction using the FastStart SYBR Green Master Mix (Roche Applied Science), 0.5 μL (out of 21 μL) of the reverse transcription reaction mixture, and the primers listed in Supplemental Table 3, with the following cycling parameters: 95°C for 10’ (95°C for 10’), 64°C for 15’ to 72°C for 20”) repeated 40 times. Fluorescence was recorded at the end of the elongation step. The specificity of PCR amplification was checked by...
a melting curve program (65 to 97°C with heating rate of 1°C per min and continuous fluorescence measurements). Absence of amplification from genomic DNA was checked by gel agarose electrophoresis. Results were analyzed using Rotor-Gene Q software (Qiagen), and data are expressed as relatives values with respect to the NAC2 mRNA level used as internal standard as this factor remains unaffected during nitrogen starvation (Raynaud et al., 2007).

Fluorescence Measurements
Fluorescence of liquid cultures, dark-adapted under strong agitation in open Erlenmeyer flasks for 30 min, was measured using a home-built fluorimeter.

Measurement of Reoxidation Kinetics of the PQ Pool
(ΔpetB) single mutant and ptox2 (ΔpetB) double mutant, starved for nitrogen for 34 h or grown in nitrogen-replete conditions, were collected by centrifugation and resuspended in 20 mM HEPES, pH 7.2, plus 20% Ficoll to avoid sedimentation. The in vivo rate of reoxidation of the plastoquinol pool in the dark was measured and calculated according to Houille-Vernes et al. (2011). In brief, the PSII photochemical rate and the variable part of fluorescence are linearly correlated. The area over the fluorescence rise curve represents the size of the pool of PSII electron acceptors. In mutants devoid of the cytochrome b6f complex, this area reflects the number of oxidized quinones before illumination (Bennoun, 1982). In the presence of DCMU, this area corresponds to one electron (transferred to Q0) and can be used for data normalization. Following the full reduction of the PQ pool by continuous illumination, measurements performed after increasing periods of darkness allow analysis of the kinetics of plastoquinol oxidation by PTOX (Bennoun, 2001).

Confocal Microscopy
Cells, deprived of nitrogen for the indicated times, were incubated 1 h in the presence of 5 mM DAF-FM DA, then washed and concentrated 20-fold by centrifugation in nitrogen-free medium and immediately imaged at room temperature with an inverted Zeiss LSM 710 laser scanning microscope equipped with a Plan Apochromat ×63/1.4 oil immersion objective and a motorized stage. Excitation was performed with a 488-nm argon laser (3% power), and emitted light was collected simultaneously on two different channels, between 493 and 599 nm and 647 and 797 nm, using a 568-nm dichroic mirror and a 63/1.4 oil immersion objective. Images were obtained by 5 × 5 tile scanning, and zooms were performed on representative individual cells. Images were collected and treated with the Zen 2011 software.

Accession Numbers
Sequence data from this article can be found in the GenBank/EMBL databases under the following accession numbers: cytochrome f, CA5A1422.1; cytochrome b6, CA5A1423; subunit IV, CA5A1424; petC, X76299.1; petO, AF222893; RbcL, A13091.1; AtpB, M13704.1; PsbA, X05845.1; PsaA, CAA25670; OEE2, M15187.1; LAO1, CRU78797; PTOX2, XP_001703466; NDA2, XP_001703643; CCS1, M15187.1; CCS5, GU362093; ccb1, EF190472; ccb2, EF190473; CCB3, EF190474; CCB4, EF190475; TCA1, EF503563.1; MCA1, AF330331.1; ClpP, L28803.1; Fsh1, XM_001690837; NAR, AF203033; and NIR, Y08937.1.

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

Supplemental Figure 1. The Induction of PTOX2 and NDA2 Is Independent of the Incident Light.

Supplemental Figure 2. The Redistribution of the Cytochrome b6f Complex during State Transitions Does Not Contribute to Its Loss during Nitrogen Deprivation.

Supplemental Figure 3. The Degradation of the Cytochrome b6f Biogenesis Factors Does Not Result from a Cascade of Proteolytic Events.

Supplemental Figure 4. The Degradation of Cytochrome b6f Complex Subunits and Biogenesis Factors Is Independent from the Photosynthetic Activity of the Cells.

Supplemental Figure 5. The Loss of the Cytochrome b6f Complex Is Accelerated upon Multiple Addition of SNP.

Supplemental Figure 6. Confocal Imaging of a Chlamydomonas Wild- Type Cell Grown in the Presence of Acetate but Not Preincubated with the DAF-FM Fluorochrome.

Supplemental Figure 7. Complementation of the nit1-305 Mutation Restores the Degradation of the Cytochrome b6f Complex.

Supplemental Table 1. Strains Used for the Crosses and Experiments Presented as Supplemental Data.

Supplemental Table 2. Genetic Analysis of the Determinism of the Cytochrome b6f Loss in Nitrate Assimilation Mutants.

Supplemental Table 3. Oligonucleotides Used in This Work.

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AUTHOR CONTRIBUTIONS

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### Supplemental Data

/content/suppl/2014/01/13/tpc.113.120121.DC1.html

### References

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Nitric Oxide–Triggered Remodeling of Chloroplast Bioenergetics and Thylakoid Proteins upon Nitrogen Starvation in Chlamydomonas reinhardtii

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