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Light History Influences the Response of the Marine Cyanobacterium *Synechococcus* sp. WH7803 to Oxidative Stress^{1[W][OA]}

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Marine *Synechococcus* undergo a wide range of environmental stressors, especially high and variable irradiance, which may induce oxidative stress through the generation of reactive oxygen species (ROS). While light and ROS could act synergistically on the impairment of photosynthesis, inducing photodamage and inhibiting photosystem II repair, acclimation to high irradiance is also thought to confer resistance to other stressors. To identify the respective roles of light and ROS in the photoinhibition process and detect a possible light-driven tolerance to oxidative stress, we compared the photophysiological and transcriptomic responses of *Synechococcus* sp. WH7803 acclimated to low light (LL) or high light (HL) to oxidative stress, induced by hydrogen peroxide (H_2O_2) or methylviologen. While photosynthetic activity was much more affected in HL than in LL cells, only HL cells were able to recover growth and photosynthesis after the addition of 25 μ M H₂O₂. Depending upon light conditions and H₂O₂ concentration, the latter oxidizing agent induced photosystem II inactivation through both direct damage to the reaction centers and inhibition of its repair cycle. Although the global transcriptome response appeared similar in LL and HL cells, some processes were specifically induced in HL cells that seemingly helped them withstand oxidative stress, including enhancement of photoprotection and ROS detoxification, repair of ROS-driven damage, and regulation of redox state. Detection of putative LexA binding sites allowed the identification of the putative LexA regulon, which was down-regulated in HL cells but up-regulated by oxidative stress under both growth irradiances.

Marine cyanobacteria must cope with a wide range of fluctuating environmental factors that, by affecting the cellular redox status, may lead to oxidative stress (Asada, 1994; Hirayama et al., 1995; Latifi et al., 2005, 2009; Schwarz and Forchhammer, 2005; Takahashi and Murata, 2008). For instance, absorption of solar radi-

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ation, especially UV wavelengths, by dissolved organic matter in aquatic environments leads to the photochemical production of various reactive transients, including reactive oxygen species (ROS; Mopper and Kieber, 2000). These compounds are partially reduced forms of oxygen, between atmospheric molecular oxygen and its fully reduced form, water, and constitute powerful oxidizing agents. Environmental stress usually induces an increase of ROS intracellular production in photosynthetic organisms, particularly when photosynthetic electron transport outpaces the rate of electron consumption during CO_2^1 fixation (Knox and Dodge, 1985; Latifi et al., 2009; Rastogi et al., 2010). This may occur when cells are exposed to excess light or UV radiation but also under nutrient and salt stress, which induce a slowing down of cell metabolism. For instance, Latifi et al. (2005) have shown that cells of Anabaena sp. PCC 7120 subjected to iron limitation exhibited a 100-fold increase in the amount of ROS compared with replete cells. Similarly, high salinity induces the release of hydrogen peroxide

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 (H_2O_2) from estuarine *Microcystis aeruginosa* cells (Ross et al., 2006). Thus, as environmental stress is intimately linked to oxidative stress, disentangling the direct effects of environmental stressors from the indirect effects of stress-induced ROS formation is not an easy task.

In cyanobacteria and other oxygenic phototrophs, the major source of ROS is the photosynthetic apparatus, although respiratory machinery also contributes to the generation of ROS compounds (Asada, 1999; Mittler, 2002; Ledford and Niyogi, 2005; Nishiyama et al., 2006; Latifi et al., 2009). Within PSI, transfer of electrons from reduced iron-sulfur centers to oxygen leads to the formation of superoxide radicals (O₂⁻) that undergo further dismutation into H₂O₂ and hydroxyl radicals (OH; Asada, 1999). PSII also contributes to ROS formation: at the PSII acceptor side, excessive reduction of electron acceptors, particularly during limitation of electron transport between PSII and PSI, may result in a reduction of molecular oxygen into O_2^{-} , which subsequently results in the formation of H_2O_2 and OH. At the PSII donor side, incomplete oxidation of water at the water-splitting manganese complex induces the formation of H_2O_2 , which can be further oxidized into O_2^{-} by manganese complex radicals or reduced into OH⁻ by free metals released by damaged PSII (Chen et al., 1992; Vass et al., 1992; Keren et al., 1997; Pospísil, 2009). Energy transfer from photoexcited pigments to oxygen in the light-harvesting antenna also leads to the formation of singlet oxygen (Knox and Dodge, 1985; Zolla and Rinalducci, 2002).

Excess ROS may severely affect the cell by damaging proteins, cleaving nucleic acids, peroxidizing unsaturated fatty acids, and inhibiting protein synthesis; this can ultimately lead to growth inhibition and cell death (Asada, 1999; Storz and Imlay, 1999; Ledford and Niyogi, 2005; Nishiyama et al., 2006). In particular, ROS have been reported to induce deleterious effects on phytoplankton by affecting cell membranes and photosynthesis (Lesser, 2006). Thus, the photosynthetic machinery is not only a major source of ROS but also constitutes a major target for these compounds. ROS are indeed thought to be involved in PSII photoinactivation, a process that has been intensively studied in freshwater cyanobacteria (Tichý and Vermaas, 1999; Nishiyama et al., 2001, 2004, 2006; Song et al., 2006; Kanesaki et al., 2007; Murata et al., 2007). Although PSII photoinactivation continually occurs under regular and stable growth conditions, it may increase under environmental stress conditions that induce an increase in the rate of ROS formation (Long et al., 1994; Hirayama et al., 1995; Asada, 1999; Latifi et al., 2005; Nishiyama et al., 2006; Takahashi and Murata, 2008). Furthermore, photoinhibition induced by ROS seems to be higher in cyanobacteria than in other phytoplanktonic organisms (Drabkova et al., 2007).

PSII photoinactivation is thought to occur primarily at the level of the oxygen-evolving complex, immediately followed by the destruction of the PSII core protein D1 (Hakala et al., 2005; Nishiyama et al., 2006). To maintain photosynthesis, photoinactivation is countered by a rapid repair cycle that includes the proteolytic removal of photodamaged D1 protein, the coordinated high rate of de novo D1 protein synthesis, and its insertion into PSII (Park et al., 1995; Tyystjärvi and Aro, 1996; Nixon et al., 2005; Ohnishi et al., 2005; Nishiyama et al., 2006). Thus, cells must modulate PSII repair cycle activity to avoid outrunning the repair by PSII photoinactivation, which would result in net photoinhibition of photosynthesis (Aro et al., 2005; Six et al., 2008). The role of ROS in PSII photoinactivation has been much debated over the past 15 years and remains controversial (Adir et al., 2003; Edelman and Mattoo, 2008; Latifi et al., 2009; Nixon et al., 2010). ROS have been suggested to be involved in the direct inactivation of the PSII reaction center through direct D1 degradation and by increasing D1 susceptibility to proteases (Vass et al., 1992; Aro et al., 1993; Miyao et al., 1995; Okada et al., 1996; Keren et al., 1997; Lupínková and Komenda, 2004; Hideg et al., 2007). Another nonexclusive hypothesis proposed that ROS primarily affect PSII by inhibiting the synthesis of D1 and therefore impairing the repair cycle (Nishiyama et al., 2004, 2006; Takahashi and Murata, 2008). Strengthening this idea, Kojima et al. (2007) have shown that the translation elongation factor EF-G is a primary target of H_2O_2 , thus suggesting that the synthesis of the D1 protein may be impaired at the elongation step of the protein translation machinery (Nishiyama et al., 2001, 2004, 2006).

Taken together, these data show the close interactions between light and ROS in photosynthetic organisms. On the one hand, the photosynthetic machinery constitutes a major source of ROS, with ROS generation particularly promoted when light absorption and the concomitant electron transport exceed the consumption of photochemically generated reductants. On the other hand, light and ROS seem to act synergistically in PSII photoinhibition, either through direct damage to PSII and/or by inhibiting repair. The influence of growth irradiance on the sensitivity to reactive oxygen stressors remains unclear. In particular, it is not well understood whether low light (LL)- and high light (HL)-acclimated cells, which show very different photophysiologies (Kana and Glibert, 1987a, 1987b; Kana et al., 1988; Moore et al., 1995; Six et al., 2004, 2005) and different tolerance to stress factors (MacDonald et al., 2003; Bouchard et al., 2005; Garczarek et al., 2008), are equally able to withstand oxidative stress. Cells acclimated to LL have a slow electron transport rate, presumably low rates of primary ROS generation, but would not be expected to maintain full induction of ROS detoxification paths. Cells acclimated to HL have a faster electron transport rate and presumably higher rates of primary ROS generation, but they are in parallel expected to maintain stronger induction of ROS detoxification paths. The net cellular response to additional ROS, therefore, depends upon the relative magnitudes of the underlying effects of primary cellular ROS generation and the counteracting effects of ROS detoxification.

In order to address this question, we used *Synechococcus* sp. WH7803, a physiologically well-characterized marine cyanobacterial isolate (Kana and Glibert, 1987a, 1987b; Toledo et al., 1999; Lindell and Post, 2001; Garczarek et al., 2008; Jia et al., 2010) for which the complete genome sequence (Dufresne et al., 2008) and a whole-genome microarray are available. We monitored PSII photophysiology and the global transcriptome response of LL- and HL-acclimated cells of this strain upon exposure to various concentrations of a direct reactive oxygen stressor, H_2O_2 , and a cell-dependent O_2^- -producing agent, methylviologen (MV).

RESULTS

Effect of H_2O_2 and MV on PSII Function of LL and HL Cells

Synechococcus sp. WH7803 cells acclimated to LL (18 $\mu mol~photons~m^{-2}~s^{-1})$ and HL (250 $\mu mol~photons$ m⁻² s⁻¹) are clearly in different initial states. In particular, the initial PSII activity, as indicated by the PSII maximal quantum yield (F_v/F_m) values before the application of any treatment, is lower in HL-acclimated cells than in LL-acclimated ones (Fig. 1). Incubation of LL- and HL-acclimated Synechococcus in the presence of different concentrations of H₂O₂ and MV for 2 h under their respective growth light irradiances indicated that the higher the concentration of oxidizing agent, the stronger the decrease of PSII activity (F_v/F_m) ; Fig. 1). LL and HL cells, however, showed quite distinct photoinactivation patterns in response to oxidizing stress. For LL-acclimated cells, increasing concentrations up to 1,250 μ M H₂O₂ induced a progressive PSII photoinactivation, but PSII of HL-grown cells was already completely inactivated at 250 μ M H₂O₂ (Fig. 1A). This suggests a synergistic effect of light and H_2O_2 on PSII photoinactivation. Concentrations of MV higher than approximately 25 μ M led to a stable PSII photoinactivation rate within 2 h, suggesting that 25 μ M MV is a saturating concentration, with no additional ROS production above this level in both LL and HL cells (Fig. 1B). The saturating level of 25 μ M MV induced a stress equivalent to exogenous application of about 125 μ M H_2O_2 , as at these concentrations both stressors induced a similar decrease of PSII quantum yield of about 20% for LL cells and 90% for HL cells.

Following this first experiment, two H_2O_2 concentrations inducing a comparable 50% PSII photoinactivation after 2 h were selected, 750 μ M H_2O_2 for LL cells and 25 μ M H_2O_2 for HL cells (Fig. 1A), to monitor the effect of oxidative stress on PSII activity over 3 d (Fig. 2A). For MV, since the saturating level was reached quickly and this oxidizing agent induced a very different photoinhibition level depending on the growth irradiance at which the culture was acclimated, comparable photoinactivation levels in LL and HL cells could not be reached within 2 h. Thus, two concentrations of MV, a subsaturating one (1 μ M) and a saturating one (50 μ M), were chosen to monitor their kinetic effects on PSII inactivation (Fig. 2B). As expected from the previous experiment, HL cells, under their growth irradiance, exhibited much faster and stronger photoinhibition than did LL cells in response to both H₂O₂ and MV. At both MV concentrations tested, HL cells reached full PSII inactivation about 2.5 h earlier than LL cells. However, all cultures submitted to MV eventually died, independently of their growth irradiance. This is probably due to the inability of cells to get rid of MV, which continued to produce ROS as long as the MV redox cycle was maintained and MV⁻⁺ was continuously reoxidized (Supplemental Fig. S1; Bus and Gibson, 1984).

The response to 25 μ M H₂O₂ differed tremendously between HL- and LL-acclimated cultures. While LL cells did not show any change in their PSII quantum yield for more than 10 h, HL cells exhibited fast and dramatic PSII inactivation under their growth irradiance once exposed to 25 μ M H₂O₂ (Fig. 2A). During the first 3 h, the rate of PSII inactivation was similar between LL cells incubated with 750 μ M H₂O₂ and HL cells incubated with 25 μ M H₂O₂. After this initial period, however, the evolution of PSII activity markedly differed between the two sets of cultures: while in LL cells subjected to 750 μ M H₂O₂, photoinactivation



Figure 1. Effect of photoacclimation on PSII photochemistry. $F_{\sqrt{F_m}}$ of *Synechococcus* sp. WH7803 cells acclimated to LL (black triangles) or HL (white squares) was measured 2 h after the addition of a range of concentrations of H₂O₂ (A) or MV (B). These data are based on three to five independent experiments and are expressed as means \pm sp.



Figure 2. PSII activity in response to H_2O_2 (A) and MV (B). The time course of changes in F_V/F_m of *Synechococcus* sp. WH7803 cells acclimated to LL (triangles) or HL (squares) was measured after the addition of 25 μ M (gray symbols) or 750 μ M (black symbols) H_2O_2 and 1 μ M (gray symbols) or 50 μ M (black symbols) MV. Control cultures without stress induction are indicated by white symbols. Broken vertical lines indicate changes of time scale on the *x* axis. These data are based on three to five independent experiments and are expressed as means \pm sp.

led to a total shutdown of PSII activity, HL cells exposed to 25 μ M H₂O₂ exhibited an increase of PSII function until full recovery after 3 d. Moreover, as measured by flow cytometry, the growth rate observed after this recovery was very similar to that observed before stress initiation (data not shown). Taken together, these data demonstrate the capacity of HL cells to recover after exposure to a low dose of H₂O₂ by completely detoxifying the cells from this oxidizing agent and restoring photosynthetic activity and most probably other affected cellular processes.

Effect of Light Acclimation and Electron Transport Rate on MV-Mediated ROS Production and PSII Inactivation

If it is correlated with photosynthetic electron flow at the PSI acceptor side, MV-mediated ROS production is expected to differ between LL- and HL-grown cells under their respective growth irradiance. To check this hypothesis, HL-acclimated cultures were either maintained under HL or shifted to LL conditions (inducing a sharp decrease in photosynthetic electron flow) 10 min before the addition of either subsaturating (1 μ M) or saturating (50 or 250 μ M) concentrations of MV (Fig. 3). For all MV concentrations tested, the shift to LL led to a much lower PSII inactivation after 2 h (Fig. 3A) and an interruption of H_2O_2 production, as assessed by the quasiconstancy of the fluorescence of scopoletin, a compound sensitive to H_2O_2 (Fig. 3B), compared with cells maintained under HL. This demonstrates that the MV-mediated PSII photoinactivation rate is highly dependent upon the photosynthetic electron transport rate. Surprisingly, the relative PSII photoinactivation in HL cells shifted to LL was only slightly lower than that of LL cells maintained in LL in the presence of MV (Fig. 3A), even though the H_2O_2 production was significantly higher in the LL cultures (Fig. 3B).

Relation between D1 Protein Turnover and ROS-Dependent Photoinactivation of PSII

In order to compare the PSII repair capacities of LL and HL cultures exposed to 25 μ M H₂O₂, we plotted



Figure 3. Effect of photosynthetic electron transport on the MV-mediated photoinhibition and H_2O_2 production. A, $F_{\sqrt{F_m}}$ was measured 2 h after the addition of 0, 1, 50, and 250 μ M MV (white, dashed, crossed, and dark bars, respectively) to *Synechococcus* sp. WH7803 cells acclimated to HL and maintained in HL (HL \rightarrow HL), acclimated to HL but shifted to LL 10 min before the stress initiation (HL \rightarrow LL), or acclimated to LL and maintained under LL (LL \rightarrow LL). The percentage of photoinactivation of stressed cells compared with untreated cells for the same light treatment is indicated above the bars. B, Scopoletin fluorescence was measured 0.5 h after the addition of MV in the same conditions as in A. The decrease of fluorescence, relative to its initial value, is due to the scopoletin degradation by H_2O_2 from the cells. All these data are based on three independent experiments and are expressed as means \pm sp.

the time-course variations of the relative D1 protein cell content (Fig. 4). While both LL and HL cells were able to increase their D1 protein content in response to oxidative stress during the first 8 h, only the HL cells sustained a high level of D1 protein over 2 d, thus enabling sufficient D1 protein turnover until complete recovery of PSII function. In contrast, PSII activity of LL cells dropped sharply and ceased after 30 h, suggesting that even a low dose of H_2O_2 provoked an irreversible impairment of the PSII repair cycle within about 1 d and likely subsequent cell death.

To assess the initial D1 repair capacities of HL and LL cells and the effect of H_2O_2 on the balance between PSII inactivation and repair, HL and LL *Synechococcus* cells were either maintained under their initial growth irradiance or transferred to dark and exposed to H_2O_2 and/or lincomycin, an inhibitor of protein synthesis. The results showed that no PSII damage was detectable after 2 h in LL-acclimated cells without the addition of any oxidant, either in the dark or under light, independently of the presence of lincomycin (Fig. 5A). In contrast, HL cells suffered a strong progressive light-dependent PSII inactivation in the presence of lincomycin (Fig. 5B), reaching 70% inactivation after 2 h compared with control (Ct) cultures without lincomycin. This demonstrates that, in contrast to LL



Figure 4. D1 protein content in response to 25 μ M H₂O₂. The D1 protein content (black symbols) in LL-acclimated (A) and HL-acclimated (B) *Synechococcus* sp. WH7803 cultures was measured by westernblot analysis after the addition of 25 μ M H₂O₂ and expressed as percentage of the initial D1 content. The evolution of the PSII quantum yield after the stress initiation, expressed as percentage of initial values, is indicated by white symbols. These data are based on three independent experiments and are expressed as means ± sp.

cells, PSII complexes from HL *Synechococcus* undergo considerable photoinactivation under their growth irradiance, which must be counteracted by the induction of a fast PSII repair cycle at a much higher rate than in LL-acclimated cells.

Second, the stronger effect on LL cells of 750 μ M H₂O₂ plus lincomycin compared with lincomycin alone (Fig. 5C) suggested that, at this concentration and under light exposure, H₂O₂ induces direct damage on PSII complexes. This effect is seemingly light dependent, since no significant decrease of the F_v/F_m occurred upon the addition of H₂O₂ plus lincomycin in the dark. A similar photoinactivation under light with or without lincomycin indicated that the D1 repair rate was too low to be measured, either because it was initially low, due to the lack of light-dependent photodamages, and/or because of an inhibitory effect of H_2O_2 . The absence of detectable light-driven damage in LL cells, which would necessitate an active repair (Fig. 5A), did not allow us to discriminate between these two hypotheses.

In HL cells, addition of 25 μ M H₂O₂ and lincomycin provoked a drop in F_v/F_m equal to the HL plus lincomycin only treatment, suggesting that this amount of oxidizing agent did not induce additional direct damage to PSII above that induced by HL alone, when repair was fully blocked (Fig. 5D). In contrast, the 25 μ M H₂O₂ treatment induced a modest decrease in PSII quantum yield compared with HL-Ct cells, suggesting some direct inhibitory effect of H₂O₂ on PSII repair. Notably, the inhibition was only partial at 25 μ M H₂O₂, since F_v/F_m did not drop to the level obtained with the lincomycin treatment to fully block PSII repair. This H₂O₂ effect on repair was confirmed by the absence of any significant effect of 25 μ M H₂O₂ alone in the dark, when no light-induced PSII damage occurred.

The stronger effect under light of 750 μ M H₂O₂ plus lincomycin compared with lincomycin alone indicated that, at this concentration, the oxidant also induced direct damage to PSII (Fig. 5D). Furthermore, 750 μ M H₂O₂ induced the same level of PSII photoinactivation with and without lincomycin, indicating that the HL cultures also underwent a complete inhibition of PSII repair at this oxidant concentration. Incubation of the culture in the dark further indicated that, in contrast to cells submitted to 25 μ M H₂O₂, HL cultures exposed to 750 μ M also underwent light-independent damage, since inactivation occurred with or without lincomycin in the dark.

Altogether, these results suggest that high concentrations of H_2O_2 affect photosynthetic activity by acting on PSII repair as well as by inducing both light-dependent and light-independent damage on PSII itself.

Global Transcriptomic Response of *Synechococcus* sp. WH7803 to Oxidative Stress

The global gene expression of *Synechococcus* sp. WH7803 was determined in response to H_2O_2 and



Figure 5. Effect of H_2O_2 on the balance between PSII damage and repair. To estimate the initial PSII repair capacity of *Synechococcus* sp. WH7803, cells acclimated to LL (A) and HL (B) were maintained under their initial growth irradiance (triangles and squares for LL and HL cells, respectively) or shifted to dark (upside down triangles and diamonds, respectively) in the presence (black symbols) or absence (white symbols) of lincomycin (Lin). F_V/F_m was then followed for 2 h. The effect of H_2O_2 on PSII damage and repair in LL (C) and HL (D) cells was estimated on cells maintained under their initial growth irradiance or shifted in the dark, by measuring the F_V/F_m 2 h after the addition of lincomycin (dashed bars) or water (open bars) and of 0 μ M (white bars), 25 μ M (light gray bars), or 750 μ M (dark gray bars) H_2O_2 . These data are based on three independent experiments and are expressed as means \pm sp.

MV doses leading to 50% PSII photoinactivation (note that the time needed to reach this state varies between treatments; Fig. 2). Microarray analyses were performed either by expressing the data with regard to the common reference hybridized on all arrays or by pairwise comparisons between nonstressed and stressed conditions (i.e. LL-Ct versus LL+MV, LL-Ct versus LL+H₂O₂, HL-Ct versus HL+MV, and HL-Ct versus $HL+H_2O_2$) or between HL and LL cultures (i.e. LL-Ct versus HL-Ct). Overall, 1,202 out of the 2,623 genes present on the array were significantly differentially expressed (false discovery rate [FDR] ≤ 0.05 using t test and/or LIMMA) in at least one of the pairwise comparisons. As expected, a higher number of genes were identified as differentially expressed in response to oxidative stress (450, 715, 530, and 585 genes in LL+H₂O₂ versus LL-Ct, LL+MV versus LL-Ct, HL+H₂O₂ versus HL-Ct, and HL+MV versus HL-Ct, respectively, when considering only the FDR cutoff) than between the two light acclimation conditions (121 genes). The complete set of gene expression data are available in Supplemental Table S1.

Principal component analysis (PCA; Fig. 6A) as well as hierarchical clustering dendrograms of the set of genes significantly differentially expressed in at least one of the pairwise comparisons (Fig. 6B) indicated good reproducibility among biological replicates used for microarray analyses, since all replicates grouped together except for the HL-stressed ones. This suggests a similar effect of H_2O_2 and MV on HL cells (Fig. 6). The most highly differentiated data sets were the oxidatively stressed cells versus Ct ones, independently of the reactive oxidizing agent used. The contribution of the stress factor seems to be dominant in PCA component 1, responsible for 35.4% of the variability of the data (Fig. 6A). The second major factor appears to be light acclimation, which segregated LL and HL data sets within both Ct and stressed conditions. This effect would be mainly represented by PCA component 2, responsible for 18.08% of the data variability. Furthermore, the greater distance between the HL and LL data sets in stressed conditions compared with Ct ones suggests that component 2 takes into account the synergistic effect of oxidative stress and HL acclimation.

In order to identify the proportion of genes responding to the different treatments, Venn diagrams were created using the same data sets (Fig. 7). These analyses showed that MV and H_2O_2 induced a similar effect on both LL and HL cells (Fig. 7A). Indeed, 82% of the genes affected by H_2O_2 in HL cells (434 out of 530 genes) were also affected by MV, and 74% (434 out of 585) were affected the opposite way around. Very similar results were obtained for LL-acclimated cells, 71% and 44%, respectively. Furthermore, among the genes common to MV- and H_2O_2 -induced stresses, **Figure 6.** Multivariate analyses of microarray data. PCA was performed (A) and hierarchical clustering dendrograms were drawn (B) using the R Stats package on a subset identified as significantly differentially expressed (FDR ≤ 0.05) in at least one of the pairwise comparisons. Experiments with LL- and HL-acclimated cells are indicated by colored triangles and white squares, respectively. Treatments using 25 μ M H₂O₂, 750 μ M H₂O₂, and 50 μ M MV are indicated by light blue, dark blue, and dark green symbols, respectively. Nontreated controls are in orange. In PCA, each replicate is indicated by one symbol, and percentages of variability accounted for by the two main components are shown near the axes.



100% showed concordance, meaning that no gene activated by one oxidizing agent was repressed by the other. This indicates that both oxidative treatments induced similar response mechanisms.

To be able to directly compare the response of LL and HL cells, genes responding to MV and/or H₂O₂ were pooled per light condition, and the resulting gene sets were called LL+Ox and HL+Ox. Pairwise comparison of LL+Ox versus LL-Ct and HL+Ox versus HL-Ct confirmed that, generally speaking, LL and HL cells responded similarly to oxidative stress. Indeed, Figure 7B shows that about half of the genes responding to oxidative stress in one light condition were also differentially expressed in the other (366 genes out of 847 for LL+Ox versus LL-Ct, 366 out of 681 for HL+Ox versus HL-Ct) and that the transcriptomic response of these genes was concordant in 93% of the cases (i.e. most genes were regulated in the same way by oxidant in LL- and HL-acclimated cells).

Additionally, these pooled sets were also compared with the effect of photoacclimation (HL-Ct versus LL-Ct) to check whether acclimation to HL and oxidative stress may lead to a common response. As shown in Figure 7C, only 42 genes (corresponding to about 5% of the LL+Ox versus LL-Ct) were differentially regulated in both LL+Ox versus LL-Ct and HL-Ct versus LL-Ct comparisons. Furthermore, 50% of these genes showed an opposite transcriptomic response (11 + 8 versus 15 + 8 genes). In contrast, although 61 genes were differentially regulated in both HL-Ox versus HL-Ct and HL-Ct versus LL-Ct sets, most of them (58 out of 61) were discordant, indicating a different expression behavior between LL and HL cells in our analyses. The full list of genes showing concordant or opposite transcriptomic patterns between the effects of acclimation to HL and ROS stress is available in Table I. Among discordant genes in HL cells, the few that are also discordant in LL cells were retrieved. Noteworthy, these common genes include *lexA* and *recA*, which play key roles in the control of DNA repair (Kuzminov, 1999; Butala et al., 2009; Kolowrat et al., 2010). The expression of these two genes was activated by oxidative stress but repressed by HL acclimation. Most of the other discordant genes encode ribosomal subunits or proteins without attributed function.

Identification of Functional Groups Responding to Oxidative Stress and Light Acclimation

Two-dimensional hierarchical clustering analysis allowed the identification of five gene clusters behaving similarly in the different treatments (Fig. 8): two clusters corresponding to transcripts mainly activated by ROS under LL (group A) or HL (group C) conditions only, a third cluster consisting of oxidantinduced genes under both light conditions (group B), and two clusters including transcripts mainly repressed by oxidative stress in HL cells (group D) or in both LL and HL cells (group E). These five clusters highlighted the behavior of coherent functional groups. The two most obvious ones included ribosomal and photosynthesis genes that were strongly downregulated by oxidants in both light conditions (group E), suggesting a decrease of photosynthetic and protein translation activities. Indeed, most of the genes encoding the subunits of the complexes driving these processes (ATP synthase, PSI and PSII, phycobilisomes, Calvin cycle, ribosomes) exhibited a similar response. Group E also includes numerous genes involved in phosphate uptake (pstB, pstSII, phnDE, etc.), storage (ppx, ppa), and regulation (phoB) as well as DNA repair genes such as *mutM* and most *rec* genes (except *recA*, located in group B). Another functional group with a characteristic behavior consists of genes involved in photoprotection, which are mainly induced by oxidative stress in LL and/or HL conditions (groups A-C). Among those, genes encoding HLinducible proteins are particularly well represented in group B, corresponding to transcripts induced in both light conditions, which also include the *ocp* gene coding for the so-called orange carotenoprotein (Kerfeld et al., 2005; Wilson et al., 2006; Kirilovsky,



Figure 7. Pairwise comparisons of microarray data sets. Venn diagrams were created with genes showing a statistically significant (FDR ≤ 0.05 ; see "Materials and Methods") differential expression in response to the different treatments. The number of genes differentially expressed in each data set is indicated near its representing symbol (see bottom right). The percentage concordance is related to genes that were significantly differentially expressed between the two compared conditions (represented by overlapping symbols) and that are additionally regulated in the same way (up- or down-regulation) in both conditions. A, Comparison of the transcriptomic responses to H₂O₂ and MV treatments for both LL- and HL-acclimated cells. B, Comparison of the pooled responses to oxidative stress (+/- oxidizing agent; i.e. genes responding to H₂O₂ and/or MV) between LL- and HL-acclimated cells. C, Comparison of the pooled responses to oxidative stress (+/- reactive oxygen stressors) in LL (top signs/values) and HL (bottom signs/values) cells and of the genes differentially expressed in HL- compared with LL-acclimated conditions. Concordant and discordant genes (i.e. common genes regulated in opposite ways between two conditions) from this diagram are listed in Table I.

2007). In contrast, genes involved in the synthesis of other carotenoids (*crtDEHQR* genes), including zeaxanthin, are located in group C, containing genes induced by ROS in HL cells only. As expected, chaperones and proteases encoding genes (*clpBCPX*, *dnaJK*, *groLS*, etc.) were induced by ROS and were also mainly represented in groups A to C.

DISCUSSION

Are HL Acclimation and Oxidative Stress Cumulative?

The physiological experiments performed in this study revealed that under their growth irradiance, HL-acclimated cells were more affected than LL cells by H_2O_2 stress with regard to PSII photoinactivation, and this was verified across the whole range of H_2O_2 concentrations tested (Figs. 1 and 2). A possible hypothesis explaining this interesting observation could be that cells grown at 250 μ mol photons m⁻² s⁻¹ were already facing a stress situation, so that they could not handle an additional stress. However, marine *Synechococcus* exhibit a large physiological

flexibility that allows them to grow under a wide range of irradiance in laboratory conditions (Kana and Glibert, 1987a, 1987b; Six et al., 2004, 2005), with the maximum growth rates usually observed between 200 to 300 μ mol photons m⁻² s⁻¹ (Kana and Glibert, 1987a; Moore et al., 1995; Six et al., 2004). Thus, the HL irradiance used in this study should not represent highly stressful growth conditions. Furthermore, microarray analyses revealed that only 121 genes were differentially expressed in the HL-Ct versus LL-Ct pairwise comparison, while four to six times more genes responded to the addition of oxidizing agents. None of the 121 genes seemed to be involved in ROS defense or detoxification mechanisms, except for a ferredoxin-encoding gene ($petF_{1580}$) and a peroxiredoxin-encoding gene $(prxQ_{1236})$, which were both significantly up-regulated in HL compared with LL cells. Interestingly, lexA and recA genes, which are involved in the SOS response (Butala et al., 2009; Kolowrat et al., 2010), were down-regulated in HL cells, supporting the interpretation that those cells were not under HL stress, which would require the activation of this DNA repair system.

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| Open Pooding Frame Identifier | Droduct |
|--|---|
| Open keading rrame identifier | Product |
| Concordant genes | |
| Up-regulated by ROS in LL cells and by HL acclimation | |
| 0203 | Putative potassium channel, VIC family |
| 0343 | Putative 3-methyladenine DNA glycosylase |
| 0514 | Predicted HAD superfamily phosphatase |
| 0791 | HII family hydrolase protein |
| 094/ | Conserved hypothetical protein |
| 1068 | Uncharacterized conserved membrane protein |
| 1501 | Hypothetical protein |
| 2334 Deven regulated by BOC in LL cells and by LU certinestices | Conserved hypothetical memorane protein |
| 1020 2182 | Conserved hypothetical proteins |
| 1029, 2102 | Hypothetical proteins |
| 1540 | Lincharacterized conserved membrane protein |
| 1732 | Sulfate ion transporter |
| 1796 | Uncharacterized conserved secreted protein pili subunit superfamily |
| 2068 | $\beta_{\rm C}$ Clycosidase (chitinase-like), family 18 |
| 2167 | HlvD-like secretion protein superfamily related to AcrA |
| galF 0242 | LIDP-GIc 4-enimerase |
| mneY 0.0494 | Putative phycobilin C-PE II lyase |
| nshA_0784 | PSII protein D1 |
| $r_{DOD} = 2501$ | Alternative RNA polymerase σ factor σ -70 family |
| 15 Svn WH7803i235 s415 208 Svn WH7803i519 s480 3i | 09 Svn WH7803i1818 a 350 |
| Up-regulated by ROS in HL cells and by HL acclimation | 55_5)1_111 00511010_q550 |
| 0947 | Conserved hypothetical protein |
| metK 0513 | S-Adenosyl-Met synthetase |
| tmk 2244 | Thymidylate kinase |
| Down-regulated by ROS in HL cells and by HL acclimation* | |
| None | |
| Discordant genes | |
| Up-regulated by ROS in LL cells and down-regulated by HL acc | limation* |
| 0095, 0096, 0934, 1953 | Hypothetical proteins |
| 1500 | Conserved hypothetical protein |
| 1797, 2532 | Uncharacterized conserved secreted protein |
| 1799 | Uncharacterized conserved secreted protein, pili subunit superfamily |
| 2116 | Uncharacterized membrane protein |
| lexA_1680 | SOS-response transcriptional repressor LexA |
| recA_0439 | Recombinase A |
| Down-regulated by ROS in LL cells and up-regulated by HL acc | limation |
| 0347 | Conserved hypothetical protein |
| 1329 | 30S ribosomal protein S2 |
| 2236 | Porin, P stress induced |
| petF_1580 | Ferredoxin |
| ppiB_1251 | Peptidyl-prolyl cis-trans-isomerase |
| psb29_1213 | PSII biogenesis protein Psb29 |
| rpIE_0422, rpmF_1115 | 50S ribosomal proteins L5 and L32 |
| Up-regulated by ROS in HL cells and down-regulated by HL ac | climation* |
| 0095, 0096, 0934, 1953 | Hypothetical proteins |
| 0543 | LysM-repeat protein |
| 0739, 1797 | Uncharacterized conserved secreted proteins |
| 0971 | Possible multisubunit Na'/H' antiporter, MnhD subunit |
| | Uncharacterized conserved membrane protein |
| 0/85, 0940, 1497, 1500, 2182 | Conserved hypothetical proteins |
| 1/32 | Suitate ion transporter |
| 1/96, 1/98, 1/99 | Uncharacterized conserved secreted proteins, pill subunit superfamily |
| 2116 | Uncharacterized membrane protein |
| 2208 | iron-suitur oxidoreductase |
| 2329 | para-Aminobenzoate synthase component II |
| | (Table continues on following page.) |

Table I. List of genes showing concordant or opposite transcriptomic patterns between the effects of acclimation to HL and the effects of oxidative stress in LL- and HL-acclimated cells, as represented in Figure 7C

Asterisks indicate genes down-regulated by HL acclimation or up-regulated by LL acclimation.

| Open Reading Frame Identifier | Product SulP-type bicarbonate transporter | | | | | |
|---|--|--|--|--|--|--|
| bicA_0719 | | | | | | |
| ctaD_1872 | Cytochrome <i>c</i> oxidase subunit I | | | | | |
| hemD_2221 | Úroporphyrinogen-III synthase | | | | | |
| lexA_1680 | SOS-response transcriptional repressor LexA | | | | | |
| mpeY_0494 | Putative phycobilin:C-PE II lyase | | | | | |
| recA_0439 | Recombinase A | | | | | |
| trpB_2299 | Trp synthase β -chain | | | | | |
| Down-regulated by ROS in HL cells and up-regulate | d by HL acclimation | | | | | |
| 1068 | Uncharacterized conserved membrane protein | | | | | |
| 1115 | 50S ribosomal protein L32 | | | | | |
| 0347, 1312, 1313, 1379, 1423, 2521 | Conserved hypothetical proteins | | | | | |
| 2002 | Probable 30S ribosomal protein PSRP-3 (Ycf65-like protein) | | | | | |
| 2198 | Inositol monophosphatase family protein | | | | | |
| 2257 | RNA-binding protein, RRM domain | | | | | |
| 2494 | Conserved hypothetical protein specific to cyanobacteria | | | | | |
| amt_0297 | Ammonium transporter | | | | | |
| focA_2493 | FNT formate/nitrate transporter family protein | | | | | |
| glnAI_1347 | Gln synthetase, Glu ammonia ligase | | | | | |
| mraY_2522 | Phospho-N-acetylmuramoyl-pentapeptide-transferase | | | | | |
| nusA_2077 | Transcription elongation factor NusA | | | | | |
| pdxJ_1371 | Pyridoxal phosphate biosynthetic protein PdxJ | | | | | |
| petF_1580 | Ferredoxin | | | | | |
| ppiB_1251 | Peptidyl-prolyl cis-trans-isomerase | | | | | |
| rp1E_0422 | 50S ribosomal proteins L5, L15, L19, L23 | | | | | |
| rpsB_1329 | 30S ribosomal proteins S2, S5, S14, S16 | | | | | |
| pstSII_1045 | ABC transporter, substrate-binding protein, phosphate | | | | | |
| 325_Syn_WH7803i1973_q360 | | | | | | |

Why Are the HL Cells More Sensitive Than LL Cells to MV-Mediated Oxidative Stress?

MV is a very low potential redox compound that primarily accepts electrons from the F_A and F_B ironsulfur clusters of PSI centers at the expense of ferredoxin reduction (Supplemental Fig. S1; Fujii et al., 1990). By reacting very quickly with molecular oxygen, the reduced MV⁺ radical is spontaneously reoxidized back to MV with the formation of the membraneimpermeable superoxide anion. The latter compound undergoes further dismutation to form H_2O_2 , which can readily pass through biological membranes (Bus and Gibson, 1984; Goldstein et al., 2002). PSI electron transport includes both linear electron flow (LEF) from PSII through PSI to NADP⁺ and cyclic electron flow (CEF) from the PSI acceptor side back to the cytochrome $b_6 f$ complex (Supplemental Fig. S1; Bendall and Manasse, 1995; Joliot and Joliot, 2002). By reacting with any electrons coming from PSI, MV inhibits CEF and diverts LEF (Yu et al., 1993; Herbert et al., 1995) toward ROS production (Jia et al., 2008; Fan et al., 2009). In the presence of MV, the ROS production in a cell is thus mostly dependent upon the absolute number of electrons passing through PSI complexes per time unit, which results from two parameters that are both strongly influenced by growth irradiance: the electron transport rate at PSI complexes and the number of PSI complexes per cell. LEF is thus a critical parameter when considering the differential response of LL and HL cells to MV-mediated oxidative stress.

At their growth irradiance, HL cells mediate much faster LEF than do LL cells, thereby producing more ROS per PSI. For PSI content, previous studies have shown that in cyanobacteria, acclimation to high irradiance is accompanied by a strong decrease of the thylakoid surface and chlorophyll *a* content per cell (Kana and Glibert, 1987a; Moore et al., 1995; Six et al., 2004). In cyanobacteria, chlorophyll *a* is almost exclusively bound to reaction centers (especially to PSI complexes); therefore, HL cells must contain fewer photosystems than LL cells. Thus, HL cells exhibit a high PSI electron transport rate favoring a large MVmediated ROS production rate, but a lower photosystem cell content and therefore a lower number of potential ROS formation sites, than LL cells.

Addition of MV concentrations higher than 25 μ M induced a saturation of PSII photoinactivation in both LL and HL cells (Fig. 1B), most likely originating from a saturation in the ROS production rate from PSI centers. Such a saturating effect has previously been observed in the freshwater *Synechococcus* sp. PCC 7942, albeit at a somewhat higher MV concentration (100 μ M; Roncel et al., 1988). While the MV concentration leading to the saturation of PSII photoinactivation rate was similar between LL and HL cells, the remaining PSII quantum yield at the saturating MV concentration was, in contrast, almost 10-fold lower in HL cells than in LL cells (Fig. 1B). If the number of PSI complexes per cell was a major factor influencing ROS production, a severe PSII photoinactivation would have been



Figure 8. Two-way cluster analysis diagram of gene expression data. This analysis is based on 1,548 genes corresponding to all genes present on the array but microarray controls, small RNA oligonucleotides, as well as unknown and hypothetical proteinencoding genes. Each row in the diagram represents a gene, and each column represents a light acclimation or a stress condition. The dendrogram at the top is similar to Figure 6, except that each column was obtained from the mean of four biological replicates performed for each condition. The color saturation represents differences in gene expression across the samples: red indicates higher than median expression (black), and green indicates lower than median expression. The color intensity indicates the degree of gene regulation. The boxes to the right represent gene clusters A to E and show some of the genes present in these clusters and belonging to specific functional categories. Genes showing a statistically significant differential expression in at least one condition (Student's *t* test and/or LIMMA with FDR ≤ 0.05) are indicated in blue.

expected in LL cells, given their high PSI cell content. Since this is not the case, this rather suggests that the LEF rate is the major factor controlling the amplitude of the MV-induced oxidative stress. Furthermore, when HL cells were suddenly shifted to LL conditions, lowering LEF without affecting the PSI cell content, the production of H_2O_2 almost stopped (Fig. 3B), with a consequent slowing down of PSII photoinactivation compared with the cells maintained in HL conditions (Fig. 3A). This clearly highlights the importance of LEF in the amplitude of MV-induced oxidative stress in HL cells. Still, it is worth noting that this only partially explains the higher sensitivity of HL cells to

oxidative stress. First, despite the sharp reduction in MV-dependent ROS production, the HL cells shifted to LL still exhibited a 25% photoinhibition (Fig. 3). Moreover, even though this stronger PSII inactivation could, at least in part, be due to a higher rate of endogenous ROS formation in the presence of MV, this justification does not hold true for H_2O_2 , since the same oxidant concentration, added exogenously, also produces a much stronger photoinhibition in HL cells than in LL ones (Fig. 1A). Thus, besides inducing a higher MV-dependent ROS production, HL acclimation also seems to result in a higher sensitivity to ROS-driven damage.

Effect of ROS on PSII Damage and Repair

The PSII repair cycle is a crucial process in the response of photosynthetic organisms to various stressful conditions (Park et al., 1995; Nishiyama et al., 2006; Six et al., 2007a). Several hypotheses have been proposed to explain the detrimental effect of ROS on PSII function, either through direct damage to PSII reaction centers with consequent degradation of the D1 protein (Hideg et al., 2007; Vass and Cser, 2009) or through effects on the de novo D1 protein synthesis through ROS inhibition of protein translation (Nishiyama et al., 2004; Kojima et al., 2007; Takahashi and Murata, 2008). Our results suggest that H₂O₂ induces both direct PSII damage and inhibition of PSII repair, with the respective importance of these processes being dependent upon both photoacclimation and oxidant concentration.

In HL cells, a treatment with lincomycin and 25 μ M H_2O_2 induced the same level of photoinactivation as the lincomycin-only treatment. This shows that at this concentration and under this growth irradiance, H_2O_2 does not induce damage beyond that observed when the PSII repair cycle is inhibited (Fig. 5D). Therefore, the moderate decrease of PSII function in the presence of 25 μ M H₂O₂ and the absence of lincomycin suggests that H₂O₂, at this concentration and under this growth irradiance, acts by partially inhibiting PSII repair rather than by directly damaging PSII. Despite the inhibitory effect of H₂O₂ in these conditions, a strong induction of D1 synthesis occurred right after stress initiation (Fig. 4). This de novo synthesis of D1 was not sufficient to maintain PSII quantum yield, indicating that most of these D1 proteins were not included in functional PSII centers (Garczarek et al., 2008).

While PSII repair cycle inhibition was partial at 25 μ M, it seemed to be complete in the presence of 750 μ M H_2O_2 , since this oxidant had the same effect on HL cells with and without lincomycin (Fig. 5D). Furthermore, this H₂O₂ concentration also induced strong direct PSII damage, since the level of photoinactivation was about twice higher than when the repair cycle was blocked by lincomycin alone. These ROS-driven damages seemed mostly light independent, since the photoinactivation in the presence of 750 μ M H₂O₂ alone was similar in the dark. Such damages could not be restricted to PSII only but could be attributed to a more global degradation of cellular components by ROS (Imlay, 2003), including translation machinery (Kojima et al., 2007). Thus, at high H₂O₂ concentrations, the decrease of PSII quantum yield in HL cells reflected both an impairment of the PSII repair cycle and direct damage to PSII complexes.

In contrast to HL cells, LL cells initially showed a nearly constant D1 pool in the presence of $25 \,\mu\text{M}\,\text{H}_2\text{O}_2$, which appeared sufficient to maintain PSII quantum yield at its initial level during 6 h (Figs. 2 and 4). This suggests that $25 \,\mu\text{M}\,\text{H}_2\text{O}_2$ did not provoke additional PSII damage beyond the very low photoinactivation rate at this light level, which was below detection.

However, an effect leading to PSII damage was clearly noted at 750 μ M H₂O₂ (Fig. 5C). In contrast to HL cells, these damages were light dependent, as almost no damage occurred when H_2O_2 was added in the dark (Fig. 5, C and D). A possible explanation for the light dependence of these damages relies on the formation of reactive OH through oxidation by H₂O₂ of reduced metals, such as the iron or manganese atoms located in reaction center II (Imlay, 2003). Indeed, PSII nonheme iron, one of the prosthetic groups of the D1 protein, has been reported to be oxidized by H₂O₂ and is thought to be involved in a specific OH-induced cleavage of this protein (Diner and Petrouleas, 1987; Lupínková and Komenda, 2004). Such reactions are expected to be light dependent, since the redox state of the nonheme iron depends on photosynthetic electron transfer (Ishikita and Knapp, 2005). Similarly, at the PSII donor side, the redox status of the manganese cluster, which can also be oxidized by H₂O₂ (Lupínková and Komenda, 2004), is also dependent on PSII electron transport. In darkness, the oxidized state of such PSII-bound metals might prevent a reaction with H₂O₂ and therefore the resulting formation of OH⁻ leading to D1 protein cleavage.

Interestingly, a cumulative effect of light and oxidative stress on PSII function does not agree with the results obtained on Synechocystis sp. PCC 6803, for which oxidative stress induced by H₂O₂ did not accentuate light-induced damage but only affected the rate of PSII repair (Allakhverdiev and Murata, 2004). It should be noted, however, that there are major experimental differences between those results and our study. Indeed, the Synechocystis experiments were mainly performed by monitoring the recovery of PSII activity after cultures were briefly exposed to very HL conditions (250–2,000 μ mol photons m⁻² s⁻¹), inducing strong PSII photoinactivation that the cells were only able to withstand for a short time. In contrast, in our study, Synechococcus sp. WH7803 cells were fully acclimated to the different growth irradiances. Furthermore, a large range of H_2O_2 concentrations was tested, as the dose proved to be important to differentiate between ROS-induced direct damage and inhibition of repair. For instance, the effect of H₂O₂ on PSII repair activity could only be detected at 25 μ M and on HL cells (i.e. when some light-mediated but no ROSmediated damages were generated). Although our approach did not allow the assessment of the PSII repair cycle when no detectable photodamage had occurred, it had the advantage of taking into account the differential damage resistance/repair capacities of these cells that cannot be appreciated through a light shift. For instance, one could expect that HL cells, which possess a higher ratio of D1:2 to D1:1 protein isoforms than LL cells (Garczarek et al., 2008), also possess a higher PSII resistance to photoinactivation (Campbell et al., 1995, 1998; Tichý et al., 2003). The fact that HL cells, in contrast, were even more sensitive to photoinhibition indicates that oxidative stress has a strong effect on direct PSII damage independently of the dominant D1 isoform present in the PSII. Altogether, these data suggest that the cumulative effect of light- and ROS-driven damage in HL cells and the resulting stronger dependence on D1 repair are probably responsible for the higher PSII sensitivity of those cells to oxidative stress. For cells shifted from HL to darkness, we observed a moderate, progressive decline in F_v/F_m (Fig. 5B). Addition of lincomycin prevented this decline, which therefore depends upon protein synthesis in the dark. This decline in F_v/F_m in the dark resulted mostly from an increase in the F0 level of fluorescence in cells shifted from HL to darkness (Supplemental Fig. S2). Similar increases in F0 occurred in the absence or presence of H₂O₂.

Transcriptome Response to Oxidative Stress

Effects of oxidative stress on PSII do not alone determine the survival capacities of the cells. Indeed, while HL cells were able to recover from a $25 \,\mu\text{M}\,\text{H}_2\text{O}_2$ stress, LL cells eventually died, even though their photosynthetic activity seemed initially less affected (Fig. 1). We performed transcriptome analyses to observe the global response of the cells and to understand why HL cells were capable of better resistance and recovery than LL cells when facing oxidative stress.

Microarray analyses showed that H₂O₂ altered the expression of functional groups of genes (e.g. photosynthesis, NADPH dehydrogenase, ribosomal proteins, some chaperones) in the marine Synechococcus sp. WH7803, in much the same way as what has been observed in the freshwater Synechocystis sp. PCC 6803 in response to approximately millimolar H_2O_2 concentrations (Li et al., 2004; Houot et al., 2007). Moreover, global transcriptomic data clearly indicated that H_2O_2 and MV induced similar transcriptomic responses in both LL and HL Synechococcus sp. WH7803 cells (Figs. 6 and 7A). Pairwise comparison between pooled genes responding to H₂O₂ or MV resulted in 93% concordance between LL and HL cells, meaning that most genes differentially expressed in both light conditions responded similarly in HL and LL cells (Fig. 7B). Hierarchical clustering analyses (Fig. 8), however, allowed highlighting some differences associated with one or the other light condition.

The photosynthetic genes are among the most affected by oxidative stress in both LL and HL cells (Fig. 8; Supplemental Table S1), including the four *psbA* gene copies encoding the D1 protein of the PSII core in *Synechococcus* sp. WH7803. At the transcriptomic level, microarray data showed that the two transcripts making up most of the *psbA* mRNA pool in this strain (Garczarek et al., 2008) are inversely regulated in response to oxidative stress, indicating an exchange between D1:1 (encoded by *psbA_0784*) and D1:2 (encoded by *psbA_0790*) isoforms in both HL and LL cells. The resulting D1 pool would then be dominated by the D1:2 isoform, which is thought to provide resistance to photoinhibition (Krupa et al., 1991; Clarke et al., 1993a, 1993b; Campbell et al., 1995, 1998; Tichý et al., 2003). Accordingly, ftsH_1216, the ROS-induced direct ortholog of Synechocystis sp. PCC 6803 slr0228 gene (Li et al., 2004; Houot et al., 2007), encoding the FtsH2 protease involved in the clearance of damaged D1 proteins from inactivated PSII (Komenda et al., 2006, 2010), belongs to the cluster of genes mainly activated in response to ROS in Synechococcus sp. WH7803 (cluster B; Fig. 8). It is worth noting that, as previously described, the D1 pool in the PSII of HL cells is dominated by the D1:2 isoform (Garczarek et al., 2008). HL cells would then be better prepared to sustain ROSdriven photoinhibition not only thanks to the initial dominance of the transcript coding for the D1:2 isoform but also by completing the isoform switch in response to additional stress. Most other photosynthetic genes belong to some of the main gene clusters inhibited by oxidative stress (clusters D and E; Fig. 8). This includes most of the ATP synthase, PSI and PSII subunits, as well as the genes involved in phycobilisome synthesis and the Calvin cycle, suggesting a decrease in energy supply in response to stress.

Although most of the photosynthetic genes were not significantly differentially affected by oxidative stress in LL and HL cells, some differences in gene induction gave us hints to explain the survival of HL cells in oxidative stress conditions that were lethal for LL cells. This includes a number of genes involved in photoprotection mechanisms, such as the ocp gene (Kerfeld et al., 2005; Wilson et al., 2006; Kirilovsky, 2007), whose induction was significantly higher in HL cells than in LL cells (Supplemental Table S1). In addition, the crtDEHQR genes involved in carotenoid synthesis (Masamoto et al., 1998; Klassen, 2010) mainly belong to the cluster C genes more activated in HL cells than in LL cells (Fig. 8), with *crtR*, a gene involved in the conversion of β -carotene into zeaxanthin, being clearly the most up-regulated. Carotenoids are thought to dissipate energy from photosensitized chlorophyll or from singlet oxygen and may have intrinsic antioxidant properties (Edge et al., 1997). In particular, zeaxanthin-deficient mutants of Synechocystis sp. PCC 6803 and Synechococcus sp. PCC 7002 are more sensitive than their respective wild types to HL and oxidative stress, and it has been shown that such mutants indeed accumulate ROS (Schäfer et al., 2005; Zhu et al., 2010). The genes differentially regulated by oxidative stress also include the high-light-inducible proteins encoding genes, three of which are induced at higher levels in HL cells than in LL cells (*hli_0937*, *hli_0797*, *hli* 1441). HL-inducible proteins have been shown to be involved in protecting cells against photodamage, either by direct or indirect dissipation of excess absorbed light energy (Montané and Kloppstech, 2000; He et al., 2001; Havaux et al., 2003) or by binding and storage of free chlorophylls (Funk and Vermaas, 1999; Vavilin et al., 2007; Kufryk et al., 2008).

Among the genes clearly induced by oxidants in HL cells (cluster C; Fig. 8), those encoding the NADPH dehydrogenase subunits are highly represented. This

complex is involved in a variety of cellular processes, including respiratory electron flow, CEF around PSI, and indirect control of the redox state of the plastoquinone pool, as well as in CO₂ uptake (Cooley and Vermaas, 2001; Ogawa and Mi, 2007; Battchikova et al., 2010). The active induction of *ndh* genes would allow maintaining energy production of HL cells under oxidative stress. Notably, a direct effect of NDHdependent CEF in alleviating photooxidative stress has also been suggested in tobacco (*Nicotiana tabacum*) chloroplast by compensating the stromal overreduction that induces the formation of ROS (Wang et al., 2006).

Tolerance of the deleterious effects of ROS may result from the activity of a number of mechanisms that maintain ROS at a harmless level or that actively repair ROS-induced damage. ROS-scavenging enzymes such as catalases, peroxidases, and superoxide dismutases, as well as antioxidants such as glutathione, tocopherol, and ascorbic acid, are involved in the maintenance of low levels of ROS (Noctor and Foyer, 1998; Storz and Imlay, 1999; Mittler, 2002; Masip et al., 2006). Such antioxidant activities involved in ROS scavenging and redox homeostasis have been reported in freshwater cyanobacteria such as Synechocystis sp. PCC 6803 and Synechococcus sp. PCC 7942 (Regelsberger et al., 2002; Yousef et al., 2003; Kobayashi et al., 2004; Hosoya-Matsuda et al., 2005; Kanesaki et al., 2007). Most of the genes involved in antioxidant activities have also been reported in the genome of Synechococcus sp. WH7803, suggesting that a similar detoxification systems occurs in this marine cyanobacterium (Scanlan et al., 2009). Among these genes, some, like the catalase/peroxidase-encoding gene *katG*, were induced in response to oxidative stress in both LL and HL cells, but others showed an obvious difference between HL and LL. While the expression of the cytosolic (iron [Fe]) superoxide dismutase (Fe-SOD)-encoding gene *sodB* was down-regulated in both LL and HL cells submitted to oxidative stress, the response of the periplasmic (copper/zinc [Cu/Zn]) superoxide dismutase (Cu/Zn-SOD) gene sodC was up-regulated in HL cells only (Supplemental Table S1; Langford et al., 2002; Culotta et al., 2006). In contrast, in Synechocystis sp. PCC 6803, which only carries the *sodB* gene, the latter seemed to be induced in response to H₂O₂ (Li et al., 2004; Houot et al., 2007). In Synechococcus sp. PCC 7942, the cytosolic Fe-SOD does not protect the cells from ROS generated within the thylakoid membrane (Thomas et al., 1998), suggesting that other SODs might confer such a protection. Thus, the increase in periplasmic Cu/Zn-SOD activity would confer a resistance to the light-driven ROS production in HL cells. Another gene involved in ROS detoxification, dpsA, encoding a DNA-binding protein, was also up-regulated in response to oxidative stress in HL cells only. DpsA has been shown to be involved in the detoxification of Fe²⁺ ions and H₂O₂ and to more specifically protect DNA from oxidative damage (Chiancone and Ceci, 2010). In Synechococcus sp. PCC 7942, while an insoluble DpsA fraction would bind DNA, a soluble faction, localized into the thylakoid membranes, would allow maintaining metal homeostasis of the photosynthetic apparatus (Durham and Bullerjahn, 2002). Thus, as the Cu/Zn-SOD, DpsA would confer a better resistance of HL cells to the ROS generated within the thylakoid membrane. The genes *ftrC* and ftrV, encoding subunits of the ferredoxin-thioredoxin reductase (FTR) complex, are clearly differentially regulated in response to oxidative stress between LL and HL cells: both genes are down-regulated in LL cells but not in HL cells, for which *ftrC* is up-regulated. The FTR complex links PSI redox status with the regulation of numerous photosynthetic processes and is required for the growth of plants that are submitted to oxidative stress (Buchanan and Balmer, 2005; Schürmann and Buchanan, 2008). Thus, the activation of the FTR system in HL cells, while it is inhibited in LL cells, may partially prevent HL cells from cellular damage due to reactive oxygen stressors by means of a better retrocontrol of the redox status of the cells.

A number of chaperone- and protease-encoding genes were also affected by ROS in only one of the light acclimation conditions. Among these, the three genes encoding the GroEL/GroES system (groS, groL1, *groL2*) and *htpG* were particularly activated in LL cells. These two major chaperone systems in bacteria are known to be activated in response to cold stress, high irradiance, and MV in Synechococcus sp. PCC 7942 (Hossain and Nakamoto, 2002, 2003; Sato et al., 2010). In contrast, *clpP* genes mostly responded to oxidative stress in HL cells, with three being activated (*clpP2*, clpP4, clpP3), while the clpP1 gene was repressed in both LL and HL cells (Fig. 8; Supplemental Table S1). The expression of these genes may also be related to *clpX*, which encodes a regulatory ATPase/chaperone interacting with ClpP and was surprisingly inhibited by ROS. Indeed, despite the fact that *clpX* and *clpP* are typically induced by heat, high salt, oxidation, and Glc deprivation in bacteria, such an inhibitory effect by H_2O_2 on *clpX* expression has already been observed in Synechococcus sp. PCC 7942 (Schelin et al., 2002) and may be linked to a possible exchange of Clp proteolytic core complexes in response to stress (Stanne et al., 2007; A. Clarke, personal communication). Finally, *degQ*, which encodes a periplasmic protease known to confer some stress resistance (Barker et al., 2006), was also somewhat more induced in HL cells than in LL cells.

An element that could contribute to the better ROS resistance of HL cells is the capacity to induce an SOS response, which is involved in protecting the cells from the damaging effects of increased mutation rates. Although some authors have suggested that cyanobacteria could lack an SOS response similar to that of *Escherichia coli* (Domain et al., 2004; Michel, 2005; Patterson-Fortin et al., 2006), it seems that *Synechococcus* sp. WH7803, like *Prochlorococcus* (Kolowrat et al., 2010), could well possess such an inducible pathway involved in DNA repair. The SOS response is under

the control of two regulatory components, RecA and LexA, the latter being a transcriptional regulator of SOS genes, including itself. The consensus sequence motif for the LexA-binding site in cyanobacteria has been proposed to be RGTACNNNDGTWCB (Mazón et al., 2004). Among the 43 genes belonging to the LexA regulon in E. coli (Courcelle et al., 2001), 10 seem to be present in the Synechococcus sp. WH7803 genome, namely lexA, recAN, umuCD, uvrAD, ssb, and ruvAB, but only *lexA*, *recA*, and *umuD* show potential LexAbinding sites in their regulatory regions, independently on their orientation (Table II). In order to identify other putative SOS genes, Synechococcus sp. WH7803 intergenic regions were screened with the consensus motif RRTACNNNDGTWYB, which integrates one degenerated position from the LexA box of the lexA gene. Six other putative LexA-binding sites have been detected downstream of the wcaG, dprA (or smf), WH7803_0096, WH7803_0814, WH7803_0843, and WH7803_1500 genes (Table II). These findings differ slightly from those of Li et al. (2010), who, by using another cyanobacterial consensus, failed to detect significant sites in the regulatory regions of lexA and umuCD, leading them to conclude that the SOS response regulation might be attenuated in Synechococ*cus* sp. WH7803. According to our analysis, all genes from the LexA regulon (i.e. that carry a putative LexA box in their regulatory region) were very similarly regulated in response to HL acclimation and to oxidative stress, strengthening the hypothesis that LexA may indeed regulate these genes and that the gene products with unknown functions might be involved in the SOS response. In contrast, genes that are thought to be part of the LexA regulon in *E. coli* but that do not have a LexA-binding motif in their promoter region in *Synechococcus* sp. WH7803 are not similarly regulated and are most probably not under the control of LexA. Using the WebLogo tool (Crooks et al., 2004), the nine putative binding sites found in Synechococcus sp. WH7803 allowed us to define the consensus of the LexA box in this strain, which seems to be degenerated from the palindromic TACAN₂TGTA consensus (Supplemental Fig. S3). Interestingly, the expression of lexA was more induced in HL cells than in LL cells, suggesting that the SOS response may indeed play a role in the better resistance of HL cells to oxidative stress. Although it might appear surprising that both repressor-encoding (lexA) and derepressor-encoding (recA) genes are simultaneously induced, the constant production of LexA would allow its reaccumulation to repress SOS genes once DNA damage has been repaired (Michel, 2005). At last, the expression of *uvrA*,

| Gene | Putative LexA-Binding Site | Position from ATG ^a | Log ₂ Fold Change from Microarray Data ^b | | | | C | | |
|-------------------|----------------------------------|--------------------------------------|--|---|-----------------|---|---------------------|------------------------------|---|
| | | | HL-Ct/ LL-Ct | LL+H ₂ O ₂ / LL-Ct | LL+MV/ LL-Ct | HL+H ₂ O ₂ / HL-Ct | HL+MV/ HL-Ct | Gene Cluster ^c | Gene Product |
| Genes wit | h a putative LexA-bindin | g site in th | eir upstre | am interger | ic region | | | | |
| lexA | GGTACACATGTATT | -21 | -0.1 | 0.9 | (0.2) | 1.4 | 1.4 | В | SOS-response transcriptional repressor |
| recA | AGTACAGATGTACG | -137 | -1.1 | 1.9 | 1.0 | 1.1 | 1.4 | В | Recombinase A |
| umuD | AGTACAGATGTTCT | -40 | (-0.3) | 2.6 | 1.2 | 1.2 | 1.3 | В | Subunits of DNA polymerase |
| umuC ^d | | | (-0.3) | 1.6 | 0.6 | (0.4) | 0.6 | В | for DNA repair |
| dprA | AGTACACATGTATT | +1 | -0.5 | 0.6 | (0.2) | 0.6 | 0.8 | В | Putative DNA-protecting protein |
| wcaG | AGTACAAACGTATT | -8 | (-0.3) | (-0.1) | 0.5 | 0.7 | 0.9 | В | Nucleoside-diphosphate-sugar epimerase |
| 0096^{e} | AATACAGTTGTATT | -88 | -2.1 | 2.2 | (0.6) | 2.9 | 3.2 | ND | Hypothetical protein |
| 0814 ^e | GGTACGCCTGTTCT | -96 | (-0.6) | 1.7 | 1.4 | 1.9 | 2.3 | ND | Conserved hypothetical protein |
| 0843 ^e | AGTACAGATGTACT | -20 | -0.7 | 1.6 | 1.8 | 1.4 | 1.5 | ND | Conserved hypothetical protein |
| 1500 ^e | AGTACAGATGTACT | -13 | -1.5 | 1.5 | (0.4) | 2.2 | 2.0 | ND | Conserved hypothetical protein |
| Consens | sus | | | | | | | | |
| | RRTACRNNYGTWYK | | | | | | | | |
| Genes wit | h no LexA-binding site fo | ound but w | hose orth | ologs are th | nought to l | pe part of the | e <i>E. coli</i> Le | xA regulo | n |
| recN | | | (-0.4) | (-0.2) | (-0.5) | (0.1) | (-0.3) | D | ATPase involved in DNA repair |
| ruvB | | | (-0.3) | (-0.2) | (0.7) | 0.9 | (0.4) | В | Holliday ATP-dependent DNA helicase |
| ssb | | | -0.8 | (-0.1) | (0.5) | 1.0 | 1.0 | А | Single-stranded DNA-binding protein |
| uvrA | | | (0.1) | (0.1) | -0.7 | -1.2 | -1.2 | D | Excinuclease ATPase subunit |
| uvrD | | | (-0.4) | (-0.4) | (-0.1) | (-0.4) | (-0.2) | А | DNA-dependent ATPase I and helicase II |

^aPositions have been calculated between the first nucleotide of the start codon and the middle of the putative binding site. ^bValues in parentheses are not statistically significant (i.e. with FDR > 0.05). ^cSee hierachical clustering analysis in Figure 8. ND, Not determined (hypothetical proteins were excluded from the analysis). ^dumuD and umuC genes are likely organized in an operon. ^eThe complete open reading frame number is SynWH7803_XXXX.

which was clearly differently regulated compared with all other SOS genes, was strongly repressed in response to oxidative stress in HL cells, and that may be due to the fact that *uvrA* is only expressed in very early stages of the SOS response (Kuzminov, 1999; Michel, 2005).

CONCLUSION

Our data show that the susceptibility of Synechococcus cells to oxidative stress varies depending on culture light history. HL cells are more immediately affected by oxidant exposure than are LL cells, but HL cells then show better capacity to ultimately acclimate and recover from low concentrations of H_2O_2 . By testing a range of oxidant concentrations and treatment durations on LL and HL cultures, we show that, depending upon the oxidant concentration, PSII damage observed in oxidant-stressed HL cells results both from cumulative effects of direct ROS damage to PSII and ROS inhibition of the rapid PSII repair needed to counter the high photoinactivation rate. These findings largely reconcile earlier findings that ROS directly inactivates PSII with other studies showing ROS inhibition of PSII repair. Even though the active PSII repair cycle of HL cells is initially inhibited by oxidative stress, it nonetheless allowed them to fully recover when H_2O_2 exposure did not exceed 25 μ M. Although the oxidative stress induced a similar global transcriptomic response in HL and LL cells, in HL cells we detected the induction of other cellular processes, including more efficient photoprotection mechanisms, some ROS detoxification, and a better redox/energy homeostasis, that most likely helped prevent the death of HL cells. Thus, in HL cells, physiological changes induced by ROS were reversible when the intracellular level of ROS decreased, while in LL cells, similar ROS treatment led to cell death. More generally, identification of the factors that set the threshold at which a cell makes the transition from successful acclimation/resistance to oxidative stress-induced cell death will be of major interest to understanding the cumulative effect of environmental stress conditions.

MATERIALS AND METHODS

Culture and Oxidative Stress Conditions

Synechococcus sp. WH7803 (Roscoff Culture Collection strain RCC752) was grown at 22°C in 0.2 μ M filtered PCR-S11 medium (Rippka et al., 2000) supplemented with 1 mM NaNO₃. Cultures were acclimated during many generations to continuous LL (18 μ mol photons m⁻² s⁻¹) and HL (250 μ mol photons m⁻² s⁻¹) provided by Sylvania Daylight 58W/154 fluorescent bulbs. Growth rates for this strain grown at several irradiances have been published elsewhere (Kana and Glibert, 1987a; Garczarek et al., 2008). For all stress experiments performed in this study, exponentially growing cultures (1–3 × 10⁷ cells mL⁻¹) were split into subcultures and incubated in the presence of different concentrations (see "Results") of oxidizing agent under various light conditions: the subcultures were either set back to their growth light conditions, or shifted to lower light to investigate the influence of photosynthetic electron flux on ROS production, or shifted to dark to differentiate direct ROS-

induced damage from light-driven damage. The oxidative stress induced by H_2O_2 was compared with that provoked by the herbicide MV (Sigma Aldrich). H_2O_2 stock solutions were titrated before each experiment by A_{240} (Nanodrop ND-1000; NanoDrop Technologies) using a molar extinction coefficient of 43.6 M^{-1} cm⁻¹ (Hildebraunt and Roots, 1975) and adjusted at 100 mM.

Synechococcus sp. WH7803 growth was monitored by flow cytometry using a FACS Canto flow cytometer (Becton Dickinson Biosciences) as described previously (Marie et al., 1997, 1999) and by phycoerythrin fluorescence emission maximum (excitation at 530 nm) using an LS-50B spectrofluorometer (Perkin-Elmer).

PSII Quantum Yield and Repair

PSII fluorescence quantum yield (F_v/F_m) was measured using a pulse amplitude-modulated fluorometer (PhytoPAM; Walz) as described previously (Six et al., 2007b; Garczarek et al., 2008).

In order to measure PSII damage and repair, 500 μ g mL⁻¹ lincomycin (Sigma-Aldrich), an inhibitor of protein synthesis, was added or not to the culture immediately before submitting the cells to oxidative stress, following a procedure similar to the one described by Six et al. (2007a). When cultures were shifted from one light condition to another, cells were shifted to the new light condition 10 min prior to the addition of oxidizing agent, immediately after the lincomycin.

Measurement of H₂O₂ Production during MV Stress

H₂O₂ production by cells upon MV-induced oxidative stress was monitored by the decrease of scopoletin fluorescence (Patterson and Myers, 1973; Tichý and Vermaas, 1999), as this molecule is degraded by H₂O₂ in the presence of horseradish peroxidase. Cells (0.5–1 × 10⁸) from exponentially growing cultures were pelleted (6,000g for 10 min; Eppendorf 5424R), resuspended in 3 mL of 25 mM HEPES (pH 7.0), and incubated in a cuvette for 30 min under their initial HL or LL acclimation conditions. After the addition of 1.33 μM scopoletin (Sigma-Aldrich) and 25 units mL⁻¹ horseradish peroxidase (Sigma-Aldrich), the cells where incubated for 10 min under their initial irradiance or shifted to LL irradiance, and 0, 50, or 250 μM MV was then added. After 30 min, the scopoletin fluorescence emission was monitored at 460 nm (excitation at 350 nm) using an LS-50B spectrofluorometer (Perkin-Elmer). After subtracting the dye-free background signal, fluorescence was expressed as a percentage of the initial value.

D1 Protein Quantification by Immunoblotting

A culture volume of 50 mL was centrifuged in the presence of the nonionic surfactant Pluronic F-68 (10 mg L⁻¹; Sigma-Aldrich) to avoid cell adhesion at the surface of the tubes, and the pellet was flash frozen in liquid nitrogen and stored at -80° C until analysis. Total protein assays, PAGE, protein transfer onto a polyvinylidene fluoride membrane, immunoreaction, and chemoluminescence detection were performed as described by Garczarek et al. (2008). D1 protein amounts were expressed as a percentage of the initial conditions.

RNA Sampling and Extraction

For the transcriptomic analyses, *Synechococcus* sp. WH7803 cultures were submitted to oxidative stress by the addition of H_2O_2 or MV and harvested when PSII quantum yield fell to half of the initial value. For H_2O_2 experiments, this level of PSII photoinactivation was reached 2 h after submitting LL and HL cultures to 750 μ M and 25 μ M, respectively (Fig. 2). Because of the large divergence in dose and kinetics responses to MV between LL and HL cells, it was not possible to find MV concentrations leading to 50% decrease of quantum yield at the same time for both light acclimations. Thus, array analyses for MV were performed on HL and LL cultures incubated at the same MV concentration (50 μ M) but harvested once PSII quantum yield was halved (i.e. after 1 and 3.5 h of stress, respectively; Fig. 2).

For every experiment, a 300-mL volume of stressed culture was harvested by centrifugation (10,000g for 7 min; Eppendorf 5417R) in the presence of the nonionic surfactant Pluronic F-68 (10 mg L⁻¹; Sigma-Aldrich). Pellets were then flash frozen in liquid nitrogen and stored at -80° C until analysis. Four independent biological replicates of the experiments were carried out.

Total RNA was extracted with acid pH-guanidinium thiocyanate/phenol/ chloroform (TRIzol reagent; Invitrogen) following a modified procedure from Millican and Bird (1998) and purified on a column (Qiagen RNeasy Mini Kit). Briefly, frozen cells were resuspended in 500 μ L of TRIzol (Invitrogen) and submitted to three cycles of 2 min of heating at 65°C and flash freezing in liquid nitrogen. Following 10 min of incubation at 65°C with vortexing, the clear lysate was vigorously mixed with 100 μ L of chloroform and incubated at room temperature for 5 min in microtubes containing an interphase gel barrier (Phase Lock Gel Heavy; 5 PRIME). After centrifugation at 9,000g for 15 min, the upper aqueous fraction was collected and the total RNA was purified following the Qiagen RNeasy Mini Kit procedure. Two consecutive DNase treatments (Qiagen RNase-Free DNase Set) were performed on the RNeasy column during the purification step, as recommended by the manufacturer. RNA was retrieved by two consecutive elutions with 30 μ L of RNA-free water (Ambion), and its quality was checked by capillary electrophoresis on a Bioanalyzer 2100 using the Procayote total RNA nano chips (Agilent).

Microarray Analysis

Array Design, cDNA Labeling, and Hybridization

Microarray experiments were performed using a homemade array targeting 2,497 protein-coding genes out of the 2,586 genes identified so far in the *Synechococcus* sp. WH7803 genome (Dufresne et al., 2008) as well as 126 potential small RNAs (W.R. Hess, personal communication). The 60-mer oligonucleotides were designed and synthesized by Eurogentec, resuspended at a final concentration of 20 μ M into a 1× spotting buffer (Schott Nexterion), and spotted in duplicate on Schott Nexterion slides using facilities of the Rennes transcriptomic platform.

cDNA synthesis from 5 μ g of total RNA and indirect CyDye cDNA labeling were performed using the ChipShot Indirect Labeling and Clean-Up System (Promega) following the manufacturer's procedure. After resuspension, labeled cDNA was quantified using a NanoDrop 1000 spectrophotometer (Thermo Scientific) and vacuum concentrated to use equal amounts of Cy3- and Cy5-labeled probes (20–100 pmol) for hybridization. All hybridizations were performed on four independent biological replicates. A pool of RNA from all samples investigated in this study was used as a reference sample. This allowed us to maximize the number of spots with a significant signal. Furthermore, in order to minimize bias due to differential dye bleaching or unequal incorporation of the Cy3 and Cy5 dyes during reverse transcription reactions, two out of the four replicate samples for each condition were hybridized in dye-swap experiments.

Printed slides were rinsed and blocked by 5 min of incubation in 0.1% Triton X-100 followed by 2 min in 0.0037% HCl, 10 min in 100 mM KCl, and finally 15 min at 55°C in a blocking solution (0.1% SDS, 0.1 M Tris, pH 9, and 0.03% ethanolamine). The slides were then prehybridized during 60 min at 55°C in prehybridization buffer (150 μ M bovine serum albumin, 3.5% SSC, and 0.7% SDS). Labeled cDNA was denatured at 90°C for 3 min and immediately spun down and kept at 55°C until loading onto the slide. The slides were then covered with a coverslip (Corning), placed in a hybridization chamber (Telechem), and hybridized for 17 h in a water bath at 55°C. The washing consisted in steps of 5 min at 55°C in 4× SSC, 5 min in the same buffer at room temperature, 1 min in 0.2× SSC at room temperature, and finally 1 min in 0.1× SSC at room temperature. Immediately after washing, slides were briefly rinsed with distilled water before being dried by centrifugation at 300g for 90 s. Scanning was performed immediately after this step.

Image Acquisition and Statistical Analysis

Scanning of the arrays was performed using a Genepix 4000A scanner (Molecular Devices). Photomultiplier gain at 570 nm (Cy3) and 660 nm (Cy5) was adjusted automatically using a threshold saturation of 0.001%, as implemented in Genepix 6.0 (Molecular Devices). Addressing and segmentation of spots were automatically detected and manually corrected, and intensities were quantified using the Genepix 6.0 software. All microarray experiments were MIAME compliant, and raw data were deposited under accession number E-MTAB-681 at the ArrayExpress database of the EMBL-European Bioinformatics Institute (http://www.ebi.ac.uk/microarray-as/ae/).

Data treatments were done using custom-designed scripts written under an R environment (R Development Core Team, 2009). Normalization of the Cy3 and Cy5 signal intensities within arrays was performed by Loess normalization (Yang and Thorne, 2003) followed by quantile normalization between arrays, as implemented in the Bioconductor package LIMMA (Smyth and Speed, 2003; Gentleman et al., 2004). The value of each sampling time point is an average of the technical replicates on a slide. Student's *t* test, linear modeling features, and empirical Bayes test statistics of the LIMMA package (Smyth, 2004) were used to perform pairwise comparison of the stress induced by either H₂O₂ or MV on both LL and HL cultures (i.e. LL-Ct versus LL+MV, LL-Ct versus LL+H₂O₂, HL-Ct versus HL+MV, and HL-Ct versus HL+H₂O₂) as well as comparing the steady-state acclimation to different light conditions (i.e. LL-Ct versus HL-Ct). To take into account multiple testing, *P* values were adjusted using the Benjamini and Hochberg algorithm (Thissen et al., 2002). We particularly focused our attention on genes that were statistically significant using Student's *t* test and/or LIMMA with FDR lower than or equal to 0.05, also taking into account a fold change (FC) cutoff between two conditions [log₂(FC) < -1 or log₂(FC) > 1].

Finally, hierarchical clustering analyses (Bolstad et al., 2004) and PCA (Raychaudhuri et al., 2000) were performed to investigate the technical and biological reproducibility of our results and highlight the main groups of genes, excluding Ct oligonucleotides, that shared similar expression patterns over multiple conditions. The clustering was performed using the hclust function from the R Stats package (R Development Core Team, 2009), using the clustering method "ward" and a Pearson correlation, and PCA analysis was carried out using the FactoMineR package (Husson et al., 2008). Analyses were performed on a subset of genes corresponding to the genes significantly differentially expressed (FDR \leq 0.05 using *t* test and/or LIMMA) in at least one of the pairwise comparisons (1,202 genes left), except for two-dimensional (genes versus arrays) hierarchical clustering analyses, where no FDR restriction had been applied but where oligonucleotides targeting unknown proteins, hypothetical proteins, and small RNAs were removed (1,548 genes left).

Validation by Real-Time Quantitative PCR

To validate microarray data, real-time quantitative PCR (qPCR) was performed on 10 Synechococcus sp. WH7803 genes. This set includes genes that were either differentially expressed in microarray analyses or representative of key processes, including ROS detoxification processes and PSII turnover. Gene-specific primers for both reverse transcription and qPCR were designed using PrimerExpress software version 2.0 (Applied Biosystems; Supplemental Table S2), and the linearity of the cDNA content to C_T (for cycle at threshold) ratio was checked for every set of primers within the dilution range used. Reverse transcription was carried out on 100 ng of RNA using SuperScriptII reverse transcriptase (Gibco-BRL) as described previously (Six et al., 2007b). qPCR was performed on the 1:100 diluted cDNA obtained, using the DNA Engine/Chromo4 Real Time PCR-Detector (Bio-Rad) and using absolute SYBR Green ROX Mix (Abgene), as described previously (Garczarek et al., 2008), in the presence of 200 nm primers, except for the sodB primers, which were used at 500 nm. The relative expression of each gene between two conditions was calculated using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen, 2001), using the *rnpB* gene as an internal standard (Mary and Vaulot, 2003). All genes and pairwise comparisons tested showed a similar response (up- or down-regulation) in qPCR and microarray experiments (Supplemental Table S2: Pearson correlation coefficient of 0.85).

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number NC_009481.

Supplemental Data

- The following materials are available in the online version of this article.
- Supplemental Figure S1. Effect of MV on ROS production through photosynthetic electron transport.
- Supplemental Figure S2. Effect of dark adaptation on PSII basal fluorescence in HL cells.
- Supplemental Figure S3. Graphical representation of the predicted LexA motif identified in *Synechococcus* sp. WH7803.
- Supplemental Table S1. Complete set of gene expression data as measured by microarray analyses.
- Supplemental Table S2. Validation of microarray data using qPCR analyses on selected genes from *Synechococcus* sp. WH7803.

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