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Redox crisis underlies conditional light–dark lethality in cyanobacterial mutants that lack the circadian regulator, RpaA

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Cyanobacteria evolved a robust circadian clock, which has a profound influence on fitness and metabolism under daily light–dark (LD) cycles. In the model cyanobacterium *Synechococcus elongatus* PCC 7942, a functional clock is not required for diurnal growth, but mutants defective for the response regulator that mediates transcriptional rhythms in the wild-type, regulator of phycobilisome association A (RpaA), cannot be cultured under LD conditions. We found that *rpaA*-null mutants are inviable after several hours in the dark and compared the metabolomes of wild-type and *rpaA*-null strains to identify the source of lethality. Here, we show that the wild-type metabolome is very stable throughout the night, and this stability is lost in the absence of RpaA. Additionally, an *rpaA* mutant accumulates excessive reactive oxygen species (ROS) during the day and is unable to clear it during the night. The *rpaA*-null metabolome indicates that these cells are reductant-starved in the dark, likely because enzymes of the primary nighttime NADPH-producing pathway are direct targets of RpaA. Because NADPH is required for processes that detoxify ROS, conditional LD lethality likely results from inability of the mutant to activate reductant-requiring pathways that detoxify ROS when photosynthesis is not active. We identified second-site mutations and growth conditions that suppress LD lethality in the mutant background that support these conclusions. These results provide a mechanistic explanation as to why *rpaA*-null mutants die in the dark, further connect the clock to metabolism under diurnal growth, and indicate that RpaA likely has important unidentified functions during the day.

cyanobacteria | metabolism | circadian clock | metabolomics | diurnal

Cyanobacteria are both key agents of global carbon and nitrogen cycles and promising platforms for renewable chemicals, fuels, and nutraceuticals (1–3). Understanding the control mechanisms that govern the flow of carbon and nitrogen through these organisms is crucial for predicting their behavior in natural environments as well as for improving engineering strategies. Although the basic pathways for carbon and nitrogen metabolism, and their regulation, are well understood in heterotrophic bacteria, cyanobacteria exhibit important deviations in these core metabolic pathways (4–7). Additionally, metabolic control mechanisms in cyanobacteria evolved to be compatible with photoautotrophic metabolism and the dramatic shifts that are imposed on those pathways by predictable daily light–dark (LD) cycles. Examples include enzymatic activity that responds to light-dependent cellular redox changes (8–11); the preference for NADPH, the reductant produced by the photochemical reactions, over NADH by many biosynthetic enzymes (12, 13); and a circadian clock that drives 24-h transcriptional rhythms in most genes (14–16).

A daily LD cycle presents a strong metabolic driver for the photosynthetic cyanobacteria, but a circadian clock also imposes daily cycles in transcription and redox regulatory systems (17, 18). Circadian measurements historically have been performed

in constant light (LL) conditions to distinguish internal circadian regulation from that which is environmentally driven (16, 17). However, diurnal physiology in a natural environment must integrate the two sources of regulation. We recently showed that the circadian clock regulates carbon metabolism in *Synechococcus elongatus* PCC 7942 as cells transition from the dark into the light during diurnal growth (19). Specifically, in the morning the clock represses the activity of the conserved circadian transcriptional regulator of phycobilisome association A (RpaA), which normally activates nighttime metabolic processes (19, 20). This action suppresses primary metabolic processes in the morning, allowing carbon to flow toward secondary metabolic processes when light energy is not limiting (19).

The circadian clock in *S. elongatus* comprises a core oscillator formed by the proteins KaiA, KaiB, and KaiC (16). The oscillator relays timing information to the SasA–RpaA two-component output pathway, in which RpaA is a transcription factor that binds 170 known downstream gene targets (20, 21). RpaA was first identified in another cyanobacterium, *Synechocystis* sp. strain PCC 6803, as an OmpR-type response regulator that influences the ratio of light energy transfer from light-harvesting phycobilisomes to photosystem I (PSI) vs. PSII (22). Thus, it is not surprising that RpaA affects core energy-producing pathways when cells are exposed to light. RpaA protein activity is directly controlled by the

Significance

The evolution of photosynthetic cyanobacteria under 24-h cycles of light and darkness selected for a robust circadian clock. Understanding how cyanobacteria integrate circadian clock signals with natural light–dark cycles to control metabolism is critical, because these organisms are central to global carbon cycling and hold promise for development of renewable energy. Here we assess how the circadian transcription factor regulator of phycobilisome association A (RpaA) influences metabolism as a cyanobacterium goes through a light-to-dark transition. The data show that RpaA plays a key role in maintaining metabolic stability during the night period. Additionally, RpaA is important in controlling redox balance, which in turn is very important for regulating metabolism at night.

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The authors declare no conflict of interest.

Dedicated to the memory of Dr. David B. Knaff.

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circadian oscillator, which represses the activity of RpaA in the late night/early morning, and subsequently relieves repression throughout the day such that RpaA reaches its peak activity at dusk (19, 20, 23). This temporal activity pattern, and the fact that RpaA directly binds and activates nighttime metabolism genes (20, 24), suggests that it also plays an important role in metabolic control at night. Although the transcriptional targets of RpaA have been identified, and it is clear that LD conditions are deleterious to *rpaA*-null mutants (25), the metabolic and physiological changes that attenuate growth under LD conditions have not been explored.

Under LD conditions, *S. elongatus* performs photosynthesis and carbon fixation during the day via the Calvin–Benson cycle, with excess fixed carbon stored as the branched chain glucose polymer glycogen (19). As cells enter a dark period, glycogen is rapidly degraded via the oxidative pentose phosphate pathway (OPPP), which serves as the primary source of energy and reducing power (NADPH) at night (26, 27). The OPPP shares many reactions with the Calvin–Benson cycle, and the transition from photosynthetic to oxidative metabolism occurs through both transcriptional and redox-regulated steps (28–30). Strict control of cellular redox via the NADPH/NADP⁺ ratio is a common and important mechanism across plant and cyanobacterial species (28). Additionally, RpaA transcriptionally activates genes that code for sugar catabolic and OPPP enzymes at the end of the day, before entering the dark, including *glgP* (glycogen phosphorylase), *gap1* (glyceraldehyde-3-phosphate dehydrogenase 1), *opcA* (OxPP cycle protein A), and the OPPP rate-limiting enzyme *zwf* (glucose-6-phosphate dehydrogenase) (20). In *rpaA* mutants, glycogen degradation is strongly attenuated, which reflects an inability to activate these sugar catabolic pathways (19).

In this study, we investigated whether the LD growth defect in a *rpaA*-null mutant is attributable to specific misregulation of metabolism and physiology as *S. elongatus* transitions into darkness and over the night period. We initially addressed the viability of an *rpaA*-null mutant (hereafter $\Delta rpaA$) over a 12-h dark period. Subsequently, we used untargeted metabolic profiling to investigate how loss of RpaA affects the abundance of primary metabolites at time points after cells enter the dark. Finally, we identified both second-site mutations and physiological growth conditions that suppress LD lethality in the $\Delta rpaA$ mutant and correlated these data with metabolomics, gene expression, and measurements of global oxidative stress. We present a model in which RpaA acts as a critical transcriptional activator of reductant-producing pathways and show that its activity is important to maintain strict metabolic stability at night. This work shows that, even in cyanobacteria that do not carry out obvious nighttime programs such as nitrogen fixation, carbon catabolism and reductant production at night are crucial for homeostasis. Thus, metabolic quiescence is not sustainable under diurnal growth conditions.

Results

Darkness Initiates Pigmentation Changes and Rapid Cell Death in the $\Delta rpaA$ Mutant. Although it is known that $\Delta rpaA$ strains do not grow under LD conditions (19, 25), the nature of the defect has not been characterized. We initially examined changes in cell viability and whole-cell absorbance as cultures entered the dark. WT and $\Delta rpaA$ cultures were sampled immediately before entry into the dark (0 h) and at intervals thereafter. Before dark exposure, the $\Delta rpaA$ strain had significantly elevated absorbance at 440 and 680 nm relative to WT (Fig. 1A), indicating an increase in chlorophyll absorbance. Although this study did not directly address changes in photosynthesis, a similar relative increase in chlorophyll was observed in *Synechocystis* sp. strain PCC 6803 $\Delta rpaA$ mutants (22). The broad differences in pigmentation at 0 h between WT and $\Delta rpaA$ (Fig. 1A) may indicate that the *S. elongatus* $\Delta rpaA$ mutant has altered energy-transfer kinetics

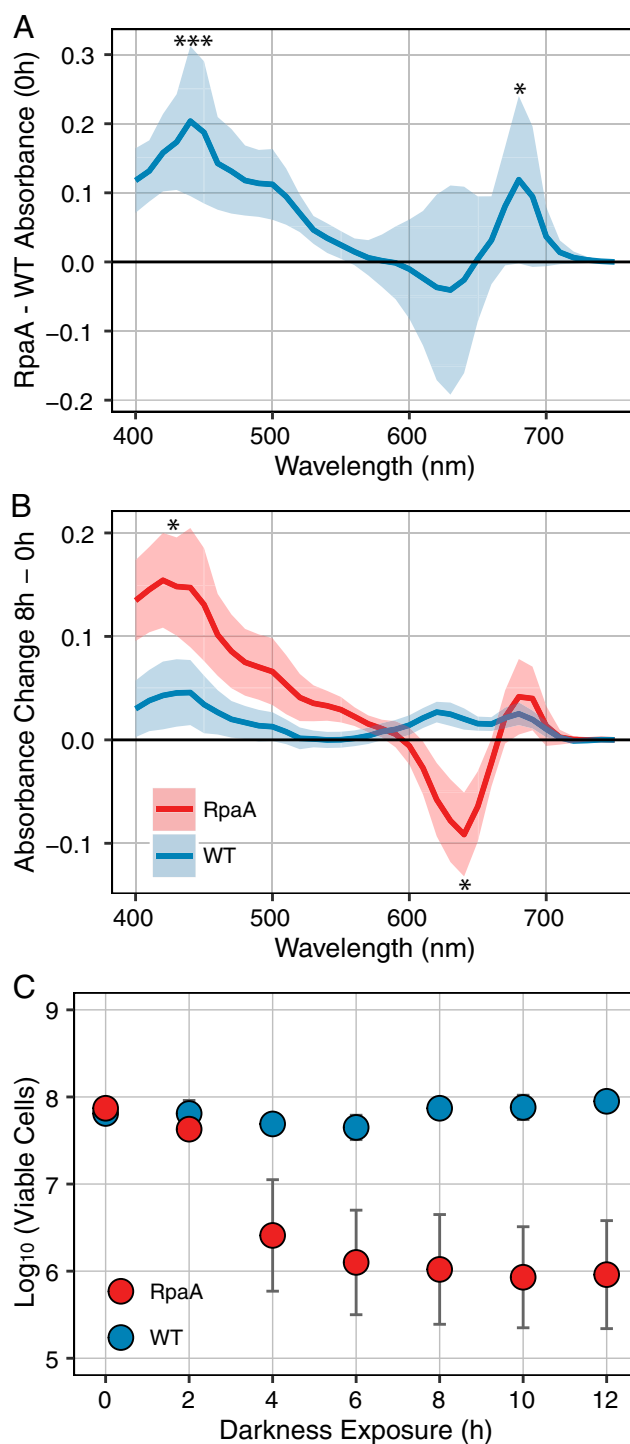


Fig. 1. Absorbance and viability data from WT and the $\Delta rpaA$ mutant. (A) Mean absorbance of WT subtracted from the $\Delta rpaA$ mutant at 0 h, immediately before entering darkness. Shaded area indicates SD of mean, significance of difference between WT and $\Delta rpaA$ calculated by Student's *t* test ($n = 8$). * $P < 0.05$; *** $P < 0.001$. (B) Change in absorbance of WT and $\Delta rpaA$ from 0 h immediately before entering darkness to 8 h of dark exposure. Shaded area indicates SD of mean, significance of difference between 0 and 8 h for each strain calculated by Student's *t* test ($n = 8$). * $P < 0.05$. (C) Mean viable cells counted at time points after WT and the $\Delta rpaA$ mutant entered the dark. Error bars indicate SEM. Significance was calculated by using Student's *t* test ($n = 4$).

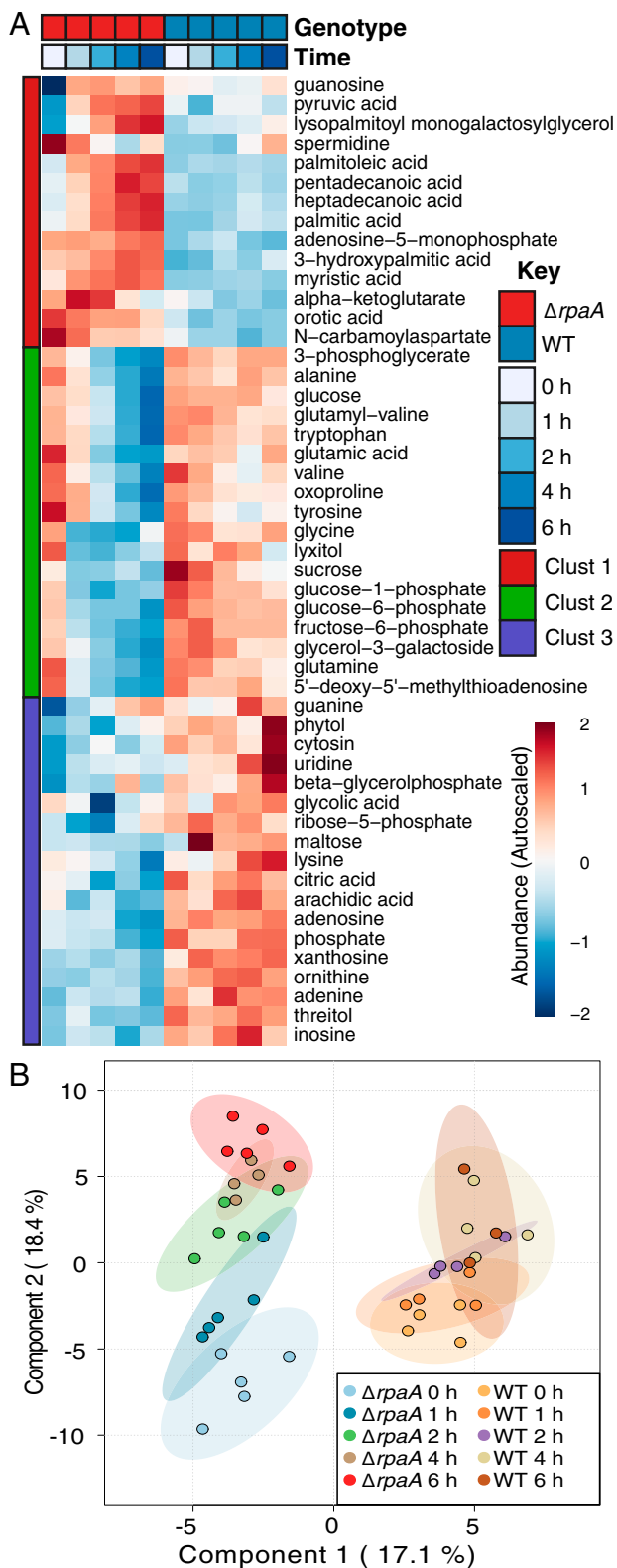


Fig. 2. Summary of metabolic changes in WT and $\Delta rpaA$. (A) Heatmap showing the autoscaled abundances of all metabolites where a significant difference was detected between WT and $\Delta rpaA$ over the time course as analyzed by two-way ANOVA and Tukey's honest significant difference ($n = 4$ for WT; $n = 5$ for $\Delta rpaA$; $P < 0.05$). Autoscaling represents a Z-score difference from the mean value of the metabolite across all time points. (B) Plot of PLS-DA components 1 and 2 for all metabolomics samples. Components 1 and 2 account for 35.5% of the variance in the dataset and are significant

between phycobilisomes, PSI, and PSII, as is true for *Synechocystis* sp. strain PCC 6803. Beginning 1–2 h after entering the dark and reaching a maximal change at 8 h after dark exposure, $\Delta rpaA$ had a significant decrease in absorbance at 630 nm and a further increase in absorbance at 440 nm, whereas WT showed no significant change in its absorbance spectrum (Fig. 1B and Fig. S14). The decrease in absorbance at 630 nm indicates a loss of phycobilisome-specific pigmentation, which is a well-characterized response to stress and macronutrient deprivation (31).

In parallel, separate samples were removed and plated under LL conditions to assess viable cell counts (colony-forming units) at each time point. Samples taken immediately before dark exposure (0 h) showed similar numbers of viable cells in the two strains (Fig. 1C). After 4 h of dark exposure, a large reproducible decrease in cell number was evident for the $\Delta rpaA$ strain, with no corresponding decrease for WT (Fig. 1C); by 8 h of darkness, only ~1% of $\Delta rpaA$ cells were viable (Fig. 1C). Optical density measurements of the sampled cultures also showed that the $\Delta rpaA$ strain did not resume growth during a following light period, even when transitioned back to LL growth conditions (Fig. S1B). These data indicate that $\Delta rpaA$ cells die soon after entering the dark. Together, the speed of the cell death response, the rapid changes in pigment absorbance, and an inability to regain viability in LL support an active mechanism that drives cell death in LD, as opposed to simple failure to thrive under LD conditions.

Temporal Metabolic Changes in the $\Delta rpaA$ Mutant. Previous data showing attenuated glycogen degradation in the $\Delta rpaA$ mutant (19) and RpaA transcriptional regulation of carbon catabolic pathways (20), and our observation of active nutrient deprivation-like bleaching (Fig. 1B), suggest that broad changes in central carbon metabolism likely occur in $\Delta rpaA$ cells after a light-to-dark transition. To characterize metabolic changes, we applied untargeted gas chromatography time-of-flight mass spectrometry (GC-TOF-MS) to samples collected from photobioreactors under LD (12-h light:12-h dark) growth conditions. Samples were collected directly before the dark onset (0 h) and at 1, 2, 4, and 6 h thereafter. A total of 114 known compounds were identified and measured (Dataset S1). Dramatic differences in primary carbon metabolites and metabolites that require NADPH for their biosynthesis were found between WT and $\Delta rpaA$ samples, suggesting that RpaA-mediated reductant production via the OPPP at night is critically important to keep metabolite levels stable. Additionally, the detection of stress-associated metabolites before dark exposure suggests that RpaA also plays a role in mitigating cellular stress during the day.

Elevated polyamines in $\Delta rpaA$ before entering dark indicate stress. Before entering the dark (0 h), polyamines were highly elevated in $\Delta rpaA$ with spermidine and putrescine showing a 48.8- and 4.5-fold increase relative to WT, respectively (Fig. 2A and Fig. S2). Correspondingly, ornithine, which is a known precursor for polyamines (32), was one-third less abundant in $\Delta rpaA$, suggesting mobilization of carbon toward polyamines. Accumulation of polyamines is a known general stress response in cyanobacteria (33), and the observed differences indicate that $\Delta rpaA$ cells may be stressed even before they enter the dark. Thus, in addition to nighttime functions, RpaA may have other, less understood, functions during the day.

WT maintains strict metabolic stability at night, which is lost in $\Delta rpaA$. Over the first half of the dark period, a large number of additional metabolite differences rapidly formed between WT and the $\Delta rpaA$ mutant. Using two-way ANOVA, we identified 50 compounds

predictors of class membership (Materials and Methods). Ellipses indicate the 95% confidence interval (CI) for each sample grouping ($n = 4$ for WT; $n = 5$ for $\Delta rpaA$; 114 metabolites per sample).

with significant differences in abundance patterns over the time course (Fig. 2A and Dataset S1). We also applied a multivariate modeling method, partial least-squares discriminate analysis (PLS-DA), to visualize and statistically test the overall similarity of sample groups, as well as determine which metabolites were more associated with differences in genotype or time (Fig. 2B).

One of the most striking observations from both analysis methods was the overall stability of metabolite levels in WT relative to $\Delta rpaA$ (Fig. 2). We had expected that WT would show significant metabolic changes downstream of the OPPP, because cyanobacteria have significant flux through glycogen degradation and the OPPP at night (19, 34, 35). However, WT maintained a stable metabolic profile, whereas $\Delta rpaA$ exhibited broad metabolic changes. Metabolites in the first and second clusters of the heatmap (Fig. 2A) showed large increases and decreases, respectively, over time in $\Delta rpaA$. In WT, the corresponding metabolites showed very gradual or no change in abundance (Fig. 2A). This effect was also pronounced in the plot of PLS-DA components 1 and 2 (Fig. 2B). Component 1 discriminated well between the two sample genotypes, and component 2 discriminated based on sampling time (Fig. 2B). The $\Delta rpaA$ mutant shows a clear separation across component 2, with temporally close samples more similar to each

other than temporally distant samples (Fig. 2B). This pattern was absent in WT samples, indicating that these samples were globally similar over the time course (Fig. 2B). Thus, ANOVA and PLS-DA both indicated that WT cells maintain a high level of metabolic stability in the early night period, which is lost in $\Delta rpaA$ mutants.

Metabolic changes in $\Delta rpaA$ indicate OPPP depression and NADPH deficit. Metabolites connected to the OPPP and those that require NADPH for biosynthesis rapidly decrease in $\Delta rpaA$ after dark exposure. The OPPP-connected compounds sucrose, glucose-1-phosphate (G1P), glucose-6-phosphate (G6P), and fructose-6-phosphate (F6P) all showed a precipitous drop in abundance as soon as $\Delta rpaA$ entered the dark (Figs. 2A and 3A). WT also showed decreases in these metabolites over time, which was expected as glycogen stores are used, but the decrease was much more gradual (Fig. 2A). However, WT, unlike $\Delta rpaA$, showed an increase in F6P at the 1-h time point after dark onset. F6P is a known indicator of OPPP activation in cyanobacteria (28), and its stark decrease in $\Delta rpaA$ cells is consistent with highly attenuated OPPP activity (Fig. 2A).

A primary sink of NADPH in cyanobacteria is amino acid biosynthesis, and many amino acids showed strong decreases in the $\Delta rpaA$ mutant (Fig. 2A). The primary nitrogen donors to

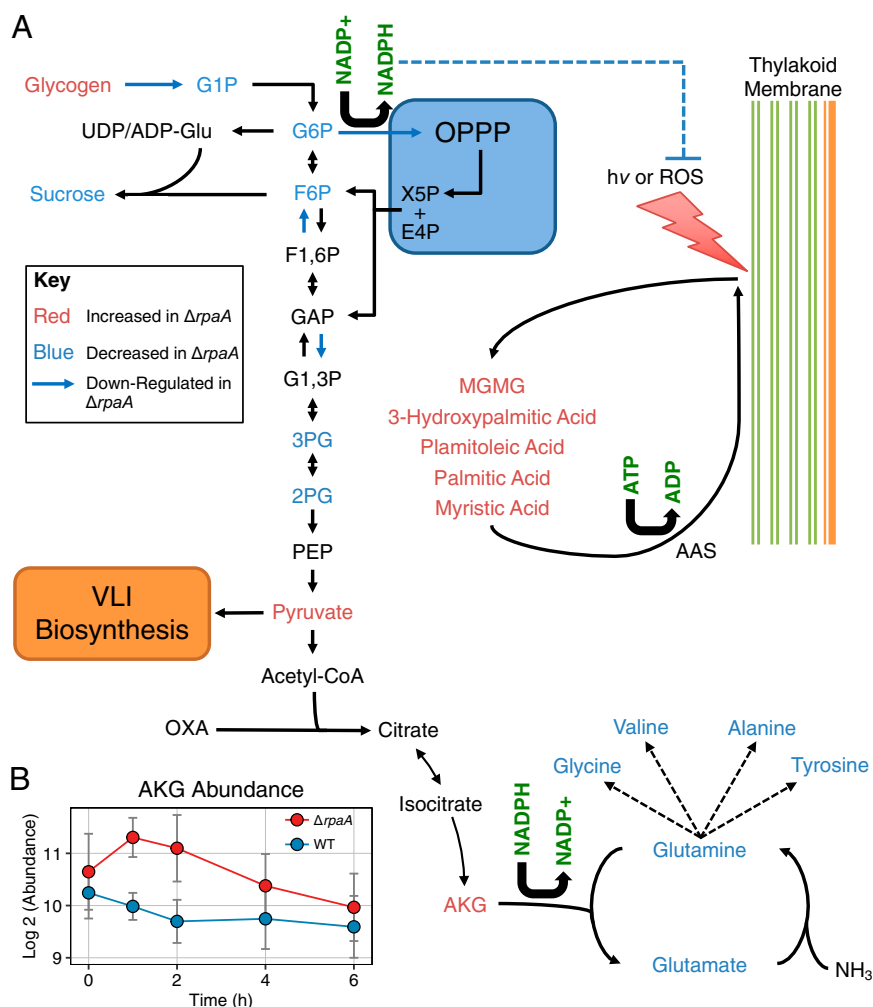


Fig. 3. Metabolic changes in the context of central carbon and nitrogen metabolism. (A) Diagram of relevant reactions in central carbon and nitrogen metabolism. Also included is a model of lipid recycling from photosynthetic membranes (right side). Metabolites are colored based on whether they were elevated (red text) or decreased (blue text) in $\Delta rpaA$ relative to WT at some point during the 6-h time course. Data for RpaA gene regulation were taken from Markson et al. (20). Dotted lines indicate that metabolites are linked, but details are not displayed. (B) Plot of AKG abundance in $\Delta rpaA$ and WT across the metabolic time course. Error bars indicate SD ($n = 4$ for WT; $n = 5$ for $\Delta rpaA$).

amino acid synthesis, glutamine and glutamate, dropped in abundance rapidly when $\Delta rpaA$ cells entered the dark (Figs. 2A and 3A). In turn, $\Delta rpaA$ showed a corresponding increase in α -ketoglutarate (AKG), the precursor metabolite for nitrogen assimilation (Fig. 3B). The conversion of AKG into glutamate in cyanobacteria is catalyzed by ferredoxin-dependent enzymes that use NADPH exclusively in their oxidation/reduction cycle (12, 13). A decrease in amino acid pools with a corresponding increase in AKG is consistent with an NADPH deficit in $\Delta rpaA$ cells. Additionally, elevated AKG can activate a nitrogen-starvation transcriptional response in cyanobacteria (36, 37), and although these cells were not nitrogen-starved, we found that elevated AKG levels in $\Delta rpaA$ were accompanied by this transcriptional response (Fig. S3 and *SI Text*).

Fatty acid accumulation in $\Delta rpaA$ likely causes damage and may result from redox imbalance. Intracellular accumulation of free fatty acids (FFAs) is uncommon in cyanobacteria, because they are activated and recycled into membrane lipids in an ATP-dependent reaction catalyzed by acyl-ACP synthetase (AAS) (38). However, in $\Delta rpaA$, we observed a large increase in the abundance of palmitic acid, palmitoleic acid, and myristic acid at the 4- to 6-h time points after dark exposure (Figs. 2A and 3A). The accumulation of intracellular FFAs in *S. elongatus* is generally toxic and directly causes damage to photosystem complexes, potentially exacerbating redox stress (39, 40). We saw that the accumulation of FFAs in $\Delta rpaA$ temporally coincided with the observed decrease in $\Delta rpaA$ cell viability (Figs. 1C and 2A), which is consistent with accumulation of these compounds contributing to the LD lethality phenotype.

Fatty acid recycling in *S. elongatus* increases dramatically under high light, and functional AAS is important to maintain cell viability under these conditions (40). In the $\Delta rpaA$ mutant, the presence of elevated lysopalmitoyl monogalactosylglycerol and 3-hydroxypalmitic acid indicated that active membrane remodeling and fatty acid recycling are taking place (Figs. 2A and 3A). Although the mechanism that drives membrane remodeling and subsequent lipid accumulation in cyanobacteria is still unclear, de novo synthesis of lipids would be unlikely in $\Delta rpaA$ cells that are reductant-poor. We hypothesize that regulation of membrane lipid turnover and FFA accumulation may be responsive to a change in cellular redox state. This hypothesis is consistent with membrane remodeling occurring under both high light (40) and in $\Delta rpaA$ cells that lack sufficient reductant (NADPH) to control cellular redox state in the dark. We posit that, although both WT and $\Delta rpaA$ may respond to oxidative stress by activation of lipid recycling, only the mutant reaches a triggering threshold of activation under moderate-light growth and does so during the night period, when AAS activity may be limited by ATP availability.

Interventions That Suppress the $\Delta rpaA$ LD Lethality Phenotype Support Reductant Imbalance as a Cause of Cell Death.

Suppression of $\Delta rpaA$ LD lethality by second-site mutagenesis. Older cultures of $\Delta rpaA$ mutants accumulate cells with the ability to grow under normally restrictive LD conditions (Fig. S4A). These clones still maintain fully segregated deletions at the *rpaA* locus; thus, it was surmised that they have accumulated compensatory changes at secondary genetic loci. To investigate the types of mutations that could suppress $\Delta rpaA$ LD lethality, we mutagenized freshly constructed, and still LD-sensitive, $\Delta rpaA$ mutant cells with ethyl methanesulfonate (EMS). Both EMS-exposed and unexposed $\Delta rpaA$ samples were then incubated under a restrictive LD growth condition. Hundreds of colonies with a wide degree of coloration and morphology appeared exclusively on the plate containing EMS-exposed $\Delta rpaA$ cells (Fig. S4B). We isolated 20 colonies, confirmed that all maintained fully segregated deletions at the *rpaA* locus (Fig. S4C), and performed full genome resequencing on each. Comparison of the mutagenized

genomes to both a WT control and the $\Delta rpaA$ parent strain revealed a total of 63 single nucleotide changes across all strains with an average of 3.15 ± 1.2 new mutations per strain. Subsequently, we filtered the mutations (*Materials and Methods*) and identified a subset of 56 that we categorized as “high confidence for biological effect” (Dataset S2).

Of the mutations, 47% occurred in genes that code for metabolic enzymes (Dataset S2). Pathways that synthesize and use amino acids were significantly enriched in these mutations, including valine, leucine, and isoleucine (VLI) biosynthesis [false discovery rate (FDR) = $2.8e-4$]; aminoacyl-tRNA biosynthesis (FDR = 0.01); and global amino acid biosynthesis (FDR = $2.8e-4$)

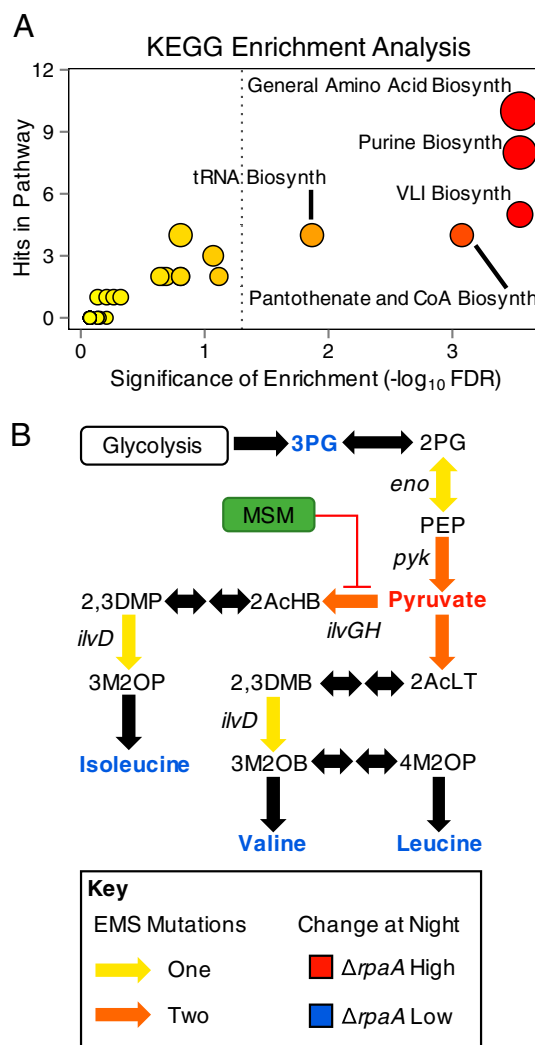


Fig. 4. Summary of enriched KEGG functional categories identified by suppressor mutations and metabolic pathway topology represented by mutated genes. (A) Plot of KEGG metabolic categories that were enriched in the gene set of suppressor mutations. The x axis indicates the number of times a specific KEGG pathway was matched to genes in the set; dots scale from small to large with increasing number of matches, and color of dots scales from yellow to red with increasing significance. Significance was calculated by using the binomial distribution, corrected for multiple testing using the method of Benjamini-Hochberg, and significance cutoff is indicated with a gray dotted line (FDR < 0.05). (B) Subpathway diagram of VLI biosynthesis indicating locations of $\Delta rpaA$ suppressor mutations and average abundance of compounds in the $\Delta rpaA$ strain relative to WT over the metabolomics time course. Genes were named for reactions where a suppressing mutation was identified, and colors are detailed in the key.

(Fig. 4A and Dataset S2). Additionally, multiple independent strains carried mutations affecting the same biochemical step in some of these pathways (Fig. 4B). These mutations affecting the same biochemical step included independent point mutations in *ilvG* and *ilvH* (acetolactate synthase catalytic and regulatory subunits), the rate-limiting enzyme complex in VLI biosynthesis (41), and two in *pyk* (pyruvate kinase), which produces pyruvate, the substrate of *ilvGH* (Fig. 4B) (42). Although mutations were found that could affect a number of metabolic pathways (Dataset S2), the high concentration of mutations in pathways that produce and consume amino acids was interesting, considering the large decreases observed for these compounds in $\Delta rpaA$ at night (Figs. 2A and 4B).

Blocking VLI biosynthesis suppresses LD lethality in $\Delta rpaA$. Because each suppressed $\Delta rpaA$ strain carries multiple EMS generated mutations, we targeted the VLI biosynthetic pathway to test whether manipulation of a single pathway identified through EMS mutagenesis is sufficient for the suppression of LD lethality in $\Delta rpaA$. VLI biosynthesis was chosen because a known herbicide, metsulfuron methyl (MSM), specifically and potently inhibits IlvGH (6, 43), and two suppressor mutations mapped to genes coding for this complex (Fig. 4B). Suppression of LD sensitivity with MSM, if successful, would also imply that the point mutations identified have a negative impact on complex activity.

Treatment of $\Delta rpaA$ with MSM under LD growth suppressed cell death of $\Delta rpaA$ on solid (100 nM MSM) and in liquid (25 μ M MSM) media (Fig. 5A and Fig. S5A). Additionally, although all MSM-treated cultures showed a change in pigmentation (Fig.

5A), we confirmed that when $\Delta rpaA$ entered the dark, treated cells no longer showed phycobilisome-specific pigment bleaching or the activation of a nitrogen-deprivation transcriptional response (Fig. 5B and Fig. S3). Therefore, inhibition of the VLI biosynthetic pathway is sufficient to rescue $\Delta rpaA$ cells from LD lethality, as well as inhibit specific phenotypes associated with cell death in the dark. Additionally, the fact that MSM exerts its suppressive effect by *ilvGH* inhibition suggests that the point mutations identified in *ilvG* and *ilvH* reduce native enzyme complex activity.

Suppression of VLI biosynthesis lowers phycobilisome content during the daytime. $\Delta rpaA$ strains treated with MSM and those that carry VLI pathway mutations had a strong yellow color (Fig. 5A and Fig. S5B). Absorbance scans taken before genomic DNA extraction for sequencing revealed that these mutants had a decreased phycobilisome-to-chlorophyll ratio (630/680 nm) relative to WT (Table S1). To investigate whether pigmentation changes occur during repression of VLI biosynthesis, we incubated WT and $\Delta rpaA$ with 25 μ M MSM in the light for 12 h and compared whole-cell absorbance spectra of treated and untreated cells. Cultures exposed to MSM visibly appeared more yellow. WT treated with MSM had decreases in all three pigment absorption maxima: 440, 630, and 680 nm, with the phycobilisome peak at 630 nm showing the largest difference (Fig. 5C). MSM-treated $\Delta rpaA$ samples also had a significant reduction in absorbance at the 630-nm phycobilisome absorbance peak (Fig. 5C). These results show that treatment with MSM reduces phycobilisome content in both WT and $\Delta rpaA$. Because phycobilisomes are the

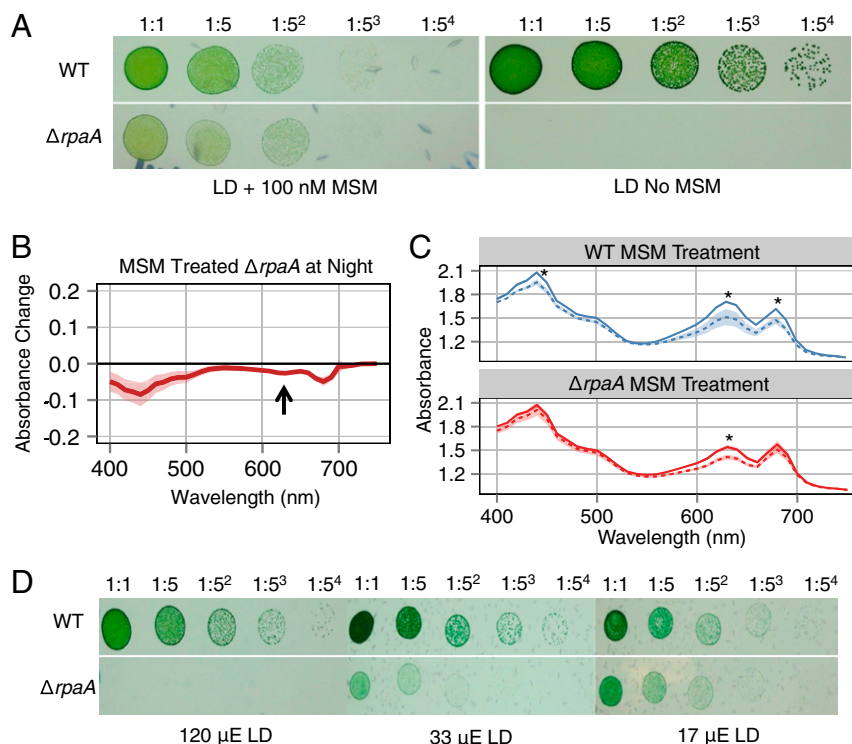


Fig. 5. Summary of data on MSM- and light intensity-mediated suppression of the $\Delta rpaA$ LD lethality phenotype. (A) Representative photo of dilution series of WT and $\Delta rpaA$ cells treated (Left) and not treated (Right) with 100 nM MSM ($n = 6$ biological replications of experiment). Pictured samples were grown in an LD cycle with a light intensity of 120 μ E·m⁻²·s⁻¹. (B) Difference in absorbance of $\Delta rpaA$ cells treated with 25 μ M MSM between 0 and 8 h after dark exposure. Shaded region indicates SD of mean, and black arrow points to absorbance at 630 nm highlighting no significant change. Significance was calculated by using Student's *t* test for absorbance at 440, 630, and 680 nm, with no significant change observed ($n = 3$). (C) Mean absorbance values of WT and $\Delta rpaA$ untreated (solid line) and treated (dotted line) with 25 μ M MSM after 12 h in the light. Shaded area indicates SEM. MSM-exposed cells show significantly lower absorption values at 440, 630, and 680 nm, as calculated by a one-sided Student's *t* test ($n = 3$ for MSM treated samples and $n = 4$ for untreated samples). * $P < 0.05$. (D) Representative photo of dilution series of WT and $\Delta rpaA$ cells grown in an LD cycle with decreasing daytime light intensity (indicated below each image; $n = 2$).

Synechocystis PCC 6803 that show proper NADPH balance is important for appropriate diurnal regulation of metabolic processes (52), as well as data indicating that the fitness advantage conferred by the *S. elongatus* circadian clock is more likely due to the ability to anticipate a coming dark period rather than a morning period (53).

One of the surprising findings is that events during the day ultimately affect death of the $\Delta rpaA$ mutant in the dark. This outcome was not anticipated because the level of light that is lethal under LD growth was well tolerated by $\Delta rpaA$ cells when they are grown under LL (Fig. S6A). Thus, it was originally hypothesized that events exclusively occurring at night were driving cell death. However, the original identification of RpaA derived from its effect on the association of phycobilisomes with photosystem reaction centers in *Synechocystis* sp. strain PCC 6803. In that strain, *rpaA*-null mutants have an increased efficiency of energy transfer to PSII relative to PSI (22). Our pigment absorbance data collected at 0 h support the hypothesis that RpaA also affects photosynthetic parameters in *S. elongatus* (Fig. 1A). If energy transfer to PSII relative to PSI is also increased in $\Delta rpaA$ *S. elongatus* cells, this alteration could both serve as a source of redox stress via excess excitation energy in PSII as well as a decrease the cells' ability to produce NADPH via PSI-driven reduction. Regardless of the source, the ROS and metabolomics data both indicate that once cells reach the light-to-dark transition, the $\Delta rpaA$ mutant is already under a great deal of cellular redox stress (Figs. 1A, 2A, and 6 and Fig. S2) (33, 46).

We propose that high redox stress generated from photosynthetic activity is a critical component of the $\Delta rpaA$ LD lethality phenotype. Manipulations that reduce light energy absorbed by cells, including reduction of growth light intensity (Fig. 5D) and treatment with MSM (Fig. 5A), rescue $\Delta rpaA$ from death under LD conditions. These results are consistent with overstimulation of photosynthetic pathways as a source of ROS. Although we focused on the EMS suppressor mutations that affect the VLI biosynthetic pathway, via treatment with MSM, strains that carry other mutations in amino acid and aminoacyl-tRNA biosynthesis exhibited similarly depressed phycobilisome absorbance (Fig. S5B and Table S1). As some of the most abundant proteins in cyanobacteria (54), phycobilisome levels are affected significantly by amino acid limitation. Cells with reduced phycobilisome content would collect less light energy and generate less ROS during the day period. Additionally, amino acid biosynthetic pathways consume large amounts of NADPH, and partially blocking these pathways may serve to preserve the limited NADPH pools present in $\Delta rpaA$ at night, which are needed for ROS-scavenging pathways.

The viability of the $\Delta rpaA$ mutant in LL (Fig. S6A), at a light intensity that generates high ROS, suggests that NADPH produced via photosynthesis can drive the ROS detoxification mechanisms necessary to maintain viability while $\Delta rpaA$ cells are in the light; however, when the $\Delta rpaA$ mutant enters the dark under redox stress, it lacks a source of NADPH because of an attenuated ability to degrade glycogen and activate the OPPP (19, 20). Consistent with an inability to activate the OPPP in the dark, $\Delta rpaA$ cells rapidly deplete soluble sugars connected to this pathway (Fig. 2A). Because glycolysis is still active in $\Delta rpaA$ and the OPPP functions as a cycle at night, recycling its inputs (34, 35), we saw a significant increase only in metabolites downstream of glycolysis in the $\Delta rpaA$ mutant, including pyruvate and AKG (Figs. 2A and 3A). Although glycolysis can produce some NADH as reducing power, NADH is a poor electron source for ROS-detoxifying processes in *S. elongatus* (46). AKG buildup normally results in its conversion to glutamine and glutamate via nitrogen assimilation (12, 13). The rapid depletion of glutamine, glutamate, and many other amino acid species in the $\Delta rpaA$ mutant, concurrent with AKG elevation (Figs. 2A and 3B), is consistent with an NADPH deficit that precludes cells from performing

AKG to amino acid biosynthesis. Additionally, the activation of a nitrogen-deprivation transcriptional response (Fig. S3) and phycobilisome degradation (Fig. 1B) indicate that $\Delta rpaA$ cells accumulate AKG to a level that is perceived as C/N imbalance. The EMS mutations in amino acid biosynthetic and utilization pathways slow the mobilization of carbon toward amino acids and lower NADPH consumption, allowing $\Delta rpaA$ to more easily achieve homeostasis with limited NADPH pools. This hypothesis is consistent with the fact that $\Delta rpaA$ cells treated with MSM no longer show nitrogen-deprivation transcriptional or bleaching responses at night (Fig. 5B and Fig. S3).

Protein redox modifications drive important metabolic shifts in cyanobacteria and plant chloroplasts (10, 28, 55, 56). Recent work has shown that redox modifications are pervasive across all metabolic pathways in cyanobacteria and that LD transitions drive global changes in the oxidation state of redox-modified proteins (8, 9, 49). Because de novo transcription is limited to the early night period in *S. elongatus* (57), redox modifications on metabolic enzymes likely play a major role in modulating enzymatic activity and dictating metabolic flux over the dark period. Some of the metabolic changes in the $\Delta rpaA$ mutant after the termination of ROS detoxification may be driven by an inability to further modulate the protein redox state. Specifically, there is evidence that accumulation of lipids can be driven by redox changes and may contribute directly to LD lethality (40, 58).

Lipids are particularly sensitive to oxidative stress, and their turnover is important because oxidized lipid species can further perpetuate oxidative damage (59). Activation of lipid recycling by high light in *S. elongatus* is consistent with a redox-driven mechanism to signal this process. The increase of lysopalmitoyl monogalactosylglycerol and 3-hydroxypalmitic acid in $\Delta rpaA$ indicates that a lipid-recycling response has been activated at night (43). We propose that ROS detoxification terminates in the $\Delta rpaA$ mutant when the limited NADPH pool is exhausted, and subsequent redox stress activates the membrane recycling process (46). We propose that the $\Delta rpaA$ mutant, but not WT, has reached an ROS threshold that would trigger lipid recycling by the end of the day. Although both WT and $\Delta rpaA$ cells should properly express the AAS protein required to activate lipids for recycling (20), AAS is ATP-dependent, and its functionality at night in cyanobacteria would likely be impaired (39). Thus, the activation of lipid recycling in $\Delta rpaA$ occurs when ATP levels are dropping in the dark, and AAS may be unable to deal with FAA load. Indeed, we observed bleaching (Fig. 1B), AKG elevation (Figs. 2A and 3B), fatty acid accumulation (Fig. 2A), and the start of cell death (Fig. 1C) around the time that ROS detoxification terminated (Fig. 6). Thus, the primary mechanisms directly driving cell death likely occur between the 2- to 6-h period after entering the dark, and these mechanisms are likely the result of a strong redox imbalance.

The daytime functions of RpaA are unexpected and should be further explored. In particular, the pigmentation changes during LL growth of the $\Delta rpaA$ mutant (Fig. 1A) suggest that core photosynthetic parameters are altered. Metrics such as photosystem efficiency, capacity, and oxygen evolution in *S. elongatus* have not been explored in the context of circadian rhythms. Additionally, it is interesting that ROS levels in the $\Delta rpaA$ mutant consistently increase at ~6 h after a transition from darkness into light (Fig. 6), because this mutant lacks a clock output mechanism and does not exhibit transcriptional rhythms (20, 21). This timing may represent a point where ROS scavenging resources are naturally exhausted, or it may indicate that other known rhythmic processes, such as 24-h peroxiredoxin rhythms, may be important for modulating ROS (18, 60). The participation of peroxiredoxins in ROS modulation would be consistent with their roles in oxidative stress-mediated signaling (61). This observation hints that Kai-mediated and other rhythmic processes may interact to control the cellular redox state. Overall, the integration of light conditions,

circadian rhythms, and the cellular redox state in the control of cyanobacterial metabolism will be of crucial importance to advance the engineering and understanding of cyanobacteria growing under natural diurnal conditions.

Materials and Methods

The full description of experimental techniques is provided in *SI Materials and Methods*.

Cyanobacterial Strains, Media, and Culture Conditions. All $\Delta rpaA$ mutants were constructed by transformation in a WT *S. elongatus* background with plasmid pAM4420 (25) and were validated by PCR. For all experiments, precultures were first prepared in 100 mL of fresh BG-11 medium as in Diamond et al. (19).

For metabolomics experiments, precultures were used to inoculate Phenometrics ePBR photobioreactors (Version 1.1; Phenometrics Inc.) at an initial density of $OD_{750} = 0.1$ in 400 mL of BG-11 medium without antibiotics. Temperature was maintained at 30 °C; filtered (0.2 μm) air was sparged at a rate of 50 mL/min; and light intensity was either 150 or 500 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ provided from the top of the culture while lights were on. After inoculation, cultures were grown at a constant light intensity of 150 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ until $OD_{750} = 0.3$, then maintained turbidostatically at this density for the duration of the experiment. In the metabolomics experiment, WT circadian rhythms were entrained by growth in a 12:12 LD at a light intensity of 150 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ cycle for 1 d and subsequently at a light intensity of 500 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ for 2 d before release into experimental conditions and sampling. The $\Delta rpaA$ strains were maintained in constant light at the same intensities as for the WT strain before the sampling procedure (Fig. S7).

For absorbance scanning, viable cell counts, quantitative reverse transcription-polymerase chain reaction (qRT-PCR), MSM-treatment absorbance measurements, and oxidative stress measurements precultures were used to inoculate the photobioreactors at an initial density of $OD_{750} = 0.2$ in 400 mL of BG-11 medium without antibiotics. Temperature, airflow rate, and light intensity settings were the same as above. For these experiments, both WT and $\Delta rpaA$ mutants were maintained at a constant light intensity of 150 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ for 1 d. Subsequently, both strains were subjected to growth in a 12:12 LD cycle at a light intensity of 150 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ for 2 d. Light intensity was then increased to 500 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ over the final 12:12 LD period, during which sampling took place.

For viable cell plating, 200 μL of the indicated sample was serially diluted 1:5 in fresh BG-11 medium without antibiotics five times. For LD sensitivity testing, samples were first all diluted to an $OD_{750} = 0.2$, and the same dilution scheme was then followed. Subsequently, 4 μL of each sample was spotted onto solid BG-11 plates without antibiotics. Plates were incubated at 30 °C and 150 $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ constant light for 5–6 d. For LD sensitivity testing, samples were plated in duplicate, with one set incubated in constant light for 5–6 d, and a second set under a 12:12 LD cycle for 6–8 d.

Metabolomics and Data Analysis. Strains for metabolomics were grown in Photobioreactors and sampled ($n = 5$ at each time point and for each genotype) as described above (Fig. S7). Metabolite extraction and GC-TOF-MS were conducted by the West Coast Metabolomics Center (WCMC) at the University of California, Davis identically to the methods used in Diamond et al. (19) and Fiehn et al. (62, 63).

Raw metabolite abundance data for known metabolites (Dataset S1) were analyzed by using a combination of the online analysis platform MetaboAnalyst (Version 3.0) (64) and the statistical package R (65). Principal component analysis (PCA) was applied to \log_2 -normalized and autoscaled data to detect outlying samples. Based on PCA, replicate A of the WT sample was removed from the dataset before statistical analysis (Fig. S8A). In addition, WT replicate C time T6 and $\Delta rpaA$ replicate C time T4 were removed before analysis because of problems during sample extraction reported by the WCMC. Statistical analysis of metabolomics data are detailed in *SI Materials and Methods*. Further details and statistical methods are provided in *SI Text*.

Mutagenesis and Identification of $\Delta rpaA$ Suppressing Mutations. EMS mutagenesis of fresh $\Delta rpaA$ mutant cyanobacterial strains was carried out as in Kondo et al. (66). Absorbance scans were taken of all cultures, as detailed in *SI Materials and Methods*, and genomic DNA was extracted by using standard methods (67). Before sequencing genomic DNA, the disruption of *rpaA* in all strains was verified by PCR (Fig. S4C). Genomic library preparation for Illumina short-read sequencing was performed by using the NEBNext DNA library preparation kit (NEB, catalog no. E6040S/L) with NEXTflex barcoded cDNA adaptors (BIOO Scientific, catalog no. 514104). Samples were run on an Illumina HiSeq 2500 DNA sequencer at the University of California, Berkeley QB3 Genomics Sequencing Laboratory. Sequencing runs resulted in 50-bp reads with a median coverage depth of 45.7 \times per sample over the *S. elongatus* genome. Reads from all sequencing methods were mapped against the *S. elongatus* genome (GenBank accession no. NC_007604) and the large plasmid pANL (GenBank accession no. AF441790), and polymorphisms were called by using the program breseq (68). Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis of mutated genes was conducted by using a custom-written R script and the metabolic categories in Dataset S2. Statistical overrepresentation was determined by using the binomial test, and *P* values were corrected by using the method of Benjamini and Hochberg (69).

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