An unexpected role for leucyl aminopeptidase in UV tolerance revealed by a genome-wide fitness assessment in a model cyanobacterium

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UV radiation (UVR) has significant physiological effects on organisms living at or near the Earth’s surface, yet the full suite of genes required for fitness of a photosynthetic organism in a UVR-rich environment remains unknown. This study reports a genome-wide fitness assessment of the genes that affect UVR tolerance under environmentally relevant UVR dosages in the model cyanobacterium Synechococcus elongatus PCC 7942. Our results highlight the importance of specific genes that encode proteins involved in DNA repair, glutathione synthesis, and the assembly and maintenance of photosystem II, as well as genes that encode hypothetical proteins and others without an obvious connection to canonical methods of UVR tolerance. Disruption of a gene that encodes a leucyl aminopeptidase (LAP) conferred the greatest UVR-specific decrease in fitness. Enzymatic assays demonstrated a strong pH-dependent affinity of the LAP for the dipeptide cysteinyl-glycine, suggesting an involvement in glutathione catabolism as a function of night-time cytosolic pH level. A low differential expression of the LAP gene under acute UVR exposure suggests that its relative importance would be overlooked in transcript-dependent screens. Subsequent experiments revealed a similar UVR-sensitivity phenotype in LAP knockouts of other organisms, indicating conservation of the functional role of LAPs in UVR tolerance.

UV radiation | cyanobacteria | fitness | leucyl aminopeptidase | RB-TnSeq

UV radiation (UVR, 300 to 400 nm) is the most photochemically reactive band of the solar spectrum and can have deleterious effects on the physiology of organisms that live on Earth’s surface or in the near-surface region of lakes and oceans. While photosynthetic organisms primarily rely on photosynthetically available radiation (PAR, 400 to 700 nm) and near-infrared radiation for photosynthesis, they are not immune to the harmful effects of UVR that are concurrent with PAR. These effects can occur at biologically relevant levels in the water column to depths of up to 20 m (1, 2). In the era of anthropogenically induced climate change, defined by changes to ice melt, aerosol concentrations, cloud cover, the ozone layer, and variations in water-column mixed layer depths, phytoplankton production and community structure will respond directly to related changes in UVR dosage, although these changes remain difficult to predict (3). Natural populations of phytoplankton will respond differently to changes in UVR stress; for example, many large-celled microbes can efficiently utilize sunscreen molecules as a photoprotective mechanism and are often less affected by high light irradiances and UVB (290 to 320 nm), while picoplankton typically experience more damage (4, 5).

Types of damage induced by UVR in phytoplankton have long been understood, such as the formation of reactive oxygen species (ROS), direct damage to DNA and proteins, and the inhibition of photosynthesis (6, 7). The general mechanisms of tolerance are thus considered to be DNA repair, protein repair and recycling, and the synthesis of antioxidants, UV-absorbing sunscreen, and photoprotective molecules, as well as avoidance of UVR via motility and mat formation (7). Most studies on UVR tolerance in phytoplankton have targeted specific processes in these categories that were previously shown to confer resistance, using transcript-dependent methodologies such as RNA Sequencing (RNA-Seq) to detect UVR-regulated genes as a proxy for gene relevance. While useful, screens of this nature do not provide direct measures of gene-specific fitness impacts. Additionally, there is a necessity to provide realistic UVR spectra in these studies, as previous research has demonstrated that the harmful effects of UVR on phytoplankton are highly wavelength dependent; wavelengths in the UVB region inhibit photosynthetic activity exponentially more than longer wavelengths in the UVA region, and direct damage to DNA and proteins occurs as a result of UVB exposure (7, 8). A dearth of genome-wide fitness screens for UVR tolerance in phytoplankton, under realistic conditions, suggests that mechanisms and genes important for UVR tolerance remain to be discovered.

Significance

Cyanobacteria account for ~25% of global primary production and dominate vast regions of the ocean, as well as soil crusts. While they rely upon light availability for photosynthesis, they are simultaneously exposed to UV radiation (UVR). Damage induced by UVR decreases photosynthetic rates, suggesting that previous calculations of global primary production may be overestimated. Despite the strong impact of UVR on phytoplankton, there has not yet been a genome-wide fitness assessment to identify genes that are important for UVR tolerance in a photosynthetic organism under environmentally relevant UVR dosages. We report genes that are most critical for UVR tolerance in a cyanobacterium and the function of a conserved leucyl aminopeptidase whose disruption results in loss of fitness under UVR.
Random barcode transposon site sequencing (RB-TnSeq) is a methodology that has proven to be particularly useful for elucidating genes that are most important to sustain a variety of conditions and processes in the model cyanobacterium *Synechococcus elongatus* PCC 7942 (9–12). In this study we leveraged RB-TnSeq to elucidate the gene set that is most critical in *S. elongatus* for tolerating UVR stress under a light cycle that resembles natural UVR dosage and short-wavelength spectral quality. Our data highlight the importance of a leucyl aminopeptidase with cysteinyl-glycinase activity that was previously overlooked and support the involvement of many canonical processes in UVR tolerance.

**Results**

**Physiological Response and Transcriptomic Landscape of Wild-Type *S. elongatus* Grown under a Light:Dark Cycle with Three UVR Dosages.** A suite of experiments was conducted to assess how *S. elongatus* is affected by three environmentally relevant UVR dosages. Wild-type (WT) cultures of *S. elongatus* grown to exponential phase and transferred to quartz photobioreactors were exposed to a 12-h light:12-h dark cycle with the addition of UVR for the central 6 h of the light period at one of three dosages levels: low UVR (LUV), medium UVR (MUV), or high UVR (HUV), or the PAR-only control (Fig. 1A). Growth rates were equivalent among the LUV, MUV, and PAR-only conditions throughout the experimental period. However, cells in the HUV condition grew more slowly than the PAR-only control during an initial 53-h phase, after which all growth rates were equivalent (experimental h 53 to 104) (Fig. 1B). ROS measured in cultures over the course of the light cycle illustrated a distinct pattern of ROS formation and dissipation (Fig. 1C). At the end of the 6-h UVR exposure period, ROS was higher in all UVR samples relative to the PAR-only control (P value LUV < 0.05; MUV < 0.01; and HUV < 0.001). At the end of the following 6 h of white light, ROS levels in the UVR treatments dropped below the PAR-only treatment. No ROS was detected the next morning following 12 h of darkness, implying that cells were able to fully scavenge all ROS and/or the ROS had fully reacted with cellular targets. Surprisingly, after the 6-h white-light period at virtual dawn, there was significantly greater ROS detected in the UVR samples, despite no UVR exposure for this period. It is possible that an accumulation of damaged DNA and proteins from the UVR exposure of the prior light cycle stimulated enhanced photosynthetic activity, resulting in elevated ROS. Whole cell absorbance spectra normalized to 675 nm showed pronounced variations in the ratio of the 630-nm to the 675-nm peak, indicating greater absorption by phycobiliproteins (PBPs) in PAR-only relative to UVR treatments and suggesting either the down-regulation of PBPs or destruction of PBPs under UVR, which is consistent with prior studies if we assume that there was little change in the chlorophyll concentration (Fig. 1D) (13).

Differential expression of genes during the UVR period was analyzed 15 min and 2 h after the onset of the UVR period following three light:dark cycles under the HUV condition using RNA-Seq. After 15 min of UVR exposure, 28 genes were up-regulated (log2 fold change > 2; adjusted P value < 0.05), while 18 were down-regulated (Fig. 2). Transcripts strongly down-regulated upon 15 min of UVR exposure primarily encode proteins related to photosystem I subunits and phycobiliprotein-containing light-harvesting complexes. Transcripts most up-regulated upon 15 min of UVR exposure encode proteins involved in light quenching, the synthesis of photosystem II D1 protein isomers PsbA2/PsbA3; protein turnover; carotenoid cycling; transcription; and replication, recombination, and repair. After 2 h of UVR exposure, 41 genes were differentially up-regulated, while 37 were down-regulated. Strongly down-regulated genes relate to photosystems I and II, energy conversion, phycobiliprotein-containing light harvesting complexes, and translation. Of the transcripts belonging to the “translation” Clusters of Orthologous Groups (COG) (14) functional group, those showing strong down-regulation included the *rpl*, *rpm*, and *rps* gene families. A complete list of the differential expression of transcripts is provided in Dataset S1.

**Assessment of Genome-Wide Fitness under UVR Using RB-TnSeq.** A fresh culture of the RB-TnSeq mutant library of *S. elongatus* PCC 7942, recovered from a frozen archive following a standardized protocol (9), was entrained in a 12-h light:12-h dark cycle for 48 h and inoculated into quartz tubes; cultures were incubated with or without the addition of a 6-h LUV, MUV, or HUV exposure period to a 12-h light:12-h dark regimen. The cultures were diluted to an OD750 of 0.05 halfway through the experimental period to minimize the attenuation of incident light. After five to six generations of growth the library was harvested and the barcodes were sequenced (9). Based on the relative number of barcode sequence reads per gene under each UVR condition to the PAR-only condition, we were able to quantify the relative fitness contribution of each gene under each UVR dosage level (Dataset S2).

* S. elongatus PCC 7942 has 2,723 genes in its 2.7-Mbp genome, of which 718 were previously determined to be essential (9); thus, those mutants are absent from the library and were not sampled in this assay. The essentiality status of all genes that were differentially expressed after acute UVR exposure is available in Dataset S1. Our experiment revealed that 5 of the nonessential genes, when disrupted, conferred a strong decrease in fitness specifically under the HUV condition relative to the PAR-only condition (log2 fold < −1.0; false discovery rate [FDR] adjusted P value < 0.05), and mutations in 14 additional genes led to a moderate decrease in fitness (log2 fold < −0.5; FDR adjusted P value < 0.05) (Fig. 3). The top 5 mutant loci that strongly decreased fitness are *Synpcc7942_1190*, *Synpcc7942_0112*, *Synpcc7942_1945*, *Synpcc7942_1679*, and *Synpcc7942_1616*. These genes are currently annotated as encoding a leucyl aminopeptidase (LAP), a deoxyribodipyrimidine photolyase, an excinuclease ABC subunit C, a photosystem II reaction center W protein, and a hypothetical protein, respectively. A strong increase in fitness was calculated only for mutants with insertions in 3 genes: *Synpcc7942_1628* (hypothetical protein), *Synpcc7942_0319* (hemolysin A), and *Synpcc7942_1933* (isopentenyl pyrophosphate isomerase). However, closer examination of the data revealed that the positive values in these cases resulted from exceptionally poor representation of these barcodes in the PAR-only sample rather than enhanced growth in PAR + HUV, indicating an amelioration of the mutant defect under UVR. The same was true for other mutant loci that were identified as moderately improving fitness.

Four of the top five genes revealed by the RB-TnSeq screen to be important under HUV exhibited a fitness response that is linearly correlated with the UVR dosage incident on the mutant pools. For example, mutants defective for *Synpcc7942_1190* had a decrease in log2 fold fitness values under HUV, MUV, and LUV of −2.1, −1.6, −0.9 (P < 0.01, and FDR < 0.01; t test), respectively. Only *Synpcc7942_1616* exhibited a dosage-independent response (HUV: −1.0, MUV: −1.1, and LUV: −0.8), suggesting that the mechanism of UVR tolerance conferred by this gene is saturated even at lower dosages of UVR.
The genes whose loss confers moderate to strong fitness decreases were classified and subsequently binned into functional groupings based on merged COG and cyanobacterial clusters of orthologous groups of proteins (CyOG) (15) databases. One is classified as being involved in coenzyme transport and metabolism (gshB); one in energy production and conversion (Synpcc7942_0567); one in nucleotide transport and metabolism (cppB); one in posttranslational modification and protein turnover (ppiB); three in replication, recombination, and repair (psbA, psbG, and psbI); one in translation (miaA); and eight were unclassified by the database (Synpcc7942_1190, Synpcc7942_1616, Synpcc7942_0037, Synpcc7942_2194, Synpcc7942_B2641, Synpcc7942_1812, Synpcc7942_B2643, and Synpcc7942_1511).

**LAP Knockout Phenotype Verification and Complementation.**

The greatest decrease in fitness observed under our HUV condition was conferred by mutations in Synpcc7942_1190, a homolog of the *Escherichia coli* pepA gene (UniProt accession no. P68767), called lap in *Synechocystis* sp. PCC 6803 (16), which encodes a putative leucyl aminopeptidase. We assessed the phenotype of a knockout mutant of *S. elongatus* Synpcc7942_1190, hereafter called lap, and performed kinetic assays of the LAP protein. To further examine the UV-sensitive phenotype identified by the RB-TnSeq screen, a fully segregated insertion mutant of *lap* with a kanamycin-resistance marker was generated (*V*lap) and grown in coculture with a WT strain containing a spectinomycin–streptomycin selective media, and colony forming units (CFUs) were used to calculate growth rates (Fig. 4). There was no statistical difference in growth rate under the PAR-only condition; however, growth of the *V* lap mutant was significantly inhibited under HUV relative to the WT (*P < 0.05; Fig. 4 A and B). To ensure that the phenotype of the *V* lap mutant is not the result of a second-site mutation elsewhere in the genome, the *V* lap mutant was complemented with a segment of DNA extending from 243 bp upstream to 71 bp downstream of SynPCC7942_1190 inserted into neutral site 1 of the genome (*V* lap::*lap*) (17). When spotted and grown on BG-11 agar, the WT phenotype was restored in the *V* lap::*lap* strain (Fig. 5 A). To assess whether the *V* lap mutant experienced enhanced sensitivity due to the dark period of the light–dark cycle in combination with prior UVR exposure, an identical experiment was conducted between the *V* lap mutant and the WT under constant light, with the UVR lamp active during the same 6-h exposure period as in previous studies. While the CFU circumference of the *V* lap mutant was reduced relative to the WT, the UV-sensitive phenotype, as assessed by reduced CFU viability in the 12-h light:12-h dark + UVR condition, was alleviated. This result indicates that the lethality of a *lap* insertional mutant relative to the WT when exposed to UVR is enhanced under the ecologically relevant condition of a subsequent period of darkness (Fig. 5 B).

**Recombinant Expression and Purification of SynPCC7942_1190 in *E. coli* and Characterization of Enzymatic Properties.**

SynPCC7942_1190 is predicted to be of the M17 family of proteases and function as an LAP; however, prior research has shown...
that some LAPs have Cys-Gly hydrolase activity. Given that Cys-Gly is a metabolite produced during the two-step catabolism of glutathione, and glutathione synthase was identified as an important enzyme for fitness under UVR, we investigated the enzymatic properties of SynPCC7942_1190. The recombinant protein was overexpressed in E. coli, and the purified 51-kDa protein migrated as a single band of the expected size on a sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gel (Fig. 6A). The substrate specificity of SynPCC7942_1190 was then assessed against Cys-Gly and L-leucine-p-nitroanilide (L-leu-p-NA) at pH 7.5. Using Cys-Gly as a substrate, a $K_m$ of 1.21 mM was observed, whereas experiments with L-leu-p-NA yielded a $K_m$ of 7.44 mM (Fig. 6B). No activity was detected against various other substrates including cystine (Cys-Cys), glycylglycine (Gly-Gly) or glutathione (Glu-Cys-Gly), suggesting specificity and preferential Cys-Gly affinity for SynPCC492_1190 at pH 7.5 despite exhibiting leucyl aminopeptidase activity (Fig. 6B).

Given that the M17 family of proteases are metalloproteases, the purified protein was tested to determine metal requirements for Cys-Gly activity. In the presence of ethylenediaminetetraacetic acid (EDTA), no activity was observed (Fig. 6C). Maximal activity was observed with the addition of Mn$^{2+}$, and a small degree of activity was observed with the addition of Co$^{2+}$. No activity was observed with the addition of Ca$^{2+}$, Zn$^{2+}$, or Mg$^{2+}$.

To assess the effect of pH on the activity of the enzyme, both Cys-Gly and L-leu-p-NA were tested as substrates across a pH gradient from 6.5 to 9.5 (Fig. 6D). Maximal enzymatic activity against Cys-Gly was observed at pH 7.5, while maximal activity against L-leu-p-NA occurred at pH 8.0. The enzyme was not as sensitive to pH changes against L-leu-p-NA when compared to Cys-Gly, similar to results published previously for an LAP from the pathogenic bacterium Treponema denticola (18).

**Synpcc7942_1190 Is Conserved across Ecologically Relevant Phytoplankton and Knockout Phenotype Is Conserved across Prokaryotic Species.** Proteins with strong sequence similarity to Synpcc7942_1190 were identified using BlastP (19) against the genomes of select ecologically relevant phytoplankton that are present in surface waters, such as Fragilariastris, Prochlorococcus marinus, Pseudonitzschia multiserial, Synechococcus sp. CCMP2592, Synechococcus sp. WH7803, and Trichodesmium erythraeum. In each homologous protein, conservation of amino acids at the seven residues canonically involved in M17 family metal binding and catalysis were identified (SI Appendix; Fig. S1). Additionally, all 97 genomes from marine and brackish Prochlorococcus, Cyanobium, and Synechoccus isolates available on the Cyanobase database v2.1 (20) encode a homologous LAP.

To test whether the UV-sensitive phenotype of the LAP-deficient mutant is observable among diverse species, deletion mutants of genes that encode the characteristic M17 family peptidase domains found in the S. elongatus LAP were generated in model organisms Anabaena sp. PCC 7120 (ΔRS03160) and E. coli (ΔpapA). The E. coli LAP gene chosen for this study was previously described to confer cysteinyl-glycinase activity in E. coli K-12 (21). Relative to the WT strain, the Anabaena sp. PCC 7120 ΔRS03160 mutants produced fewer CFUs when grown under UVR in a 12-h light:12-h dark cycle relative to white light only, indicating conservation of the UV-sensitive phenotype across species of cyanobacteria (Fig. 7A). Similarly, the E. coli ΔpapA mutant resulted in reduced colony forming units when grown under UVR relative to white light only and the respective WT controls (Fig. 7B), indicating the conservation of the UVR-sensitivity phenotype of LAP knockouts broadly across prokaryotic genera. Additionally, to test whether the lap gene from Anabaena sp. PCC 7120 (RS03160) could be used to complement the Vlap strain of S. elongatus, RS03160 was cloned under the regulation of a Ptra promoter and introduced into neutral site 1 of the chromosome in the Vlap strain of S. elongatus. When spotted and grown on BG-11 agar, the WT phenotype was restored in the Vlap:RS03160 strain, corroborating conservation of the UVR tolerance mechanism in a leucyl aminopeptidase across cyanobacterial species (Fig. 7C).

**Discussion**

This work has identified an unexpected protein, the aminopeptidase LAP, encoded by Synpcc7942_1190, as the strongest determinant of fitness under all UVR dosage conditions. This protein has a greater impact on UVR fitness than previously described proteins that strongly contribute to UVR tolerance through DNA repair, maintenance of the photosynthetic apparatus, and antioxidant synthesis (22, 23).

![Volcano plot highlighting genes whose loss have moderate to strong fitness effects under HUV. Colors indicate whether loss of a gene causes an estimated fitness decrease (red) or increase (blue). A horizontal line indicates the P value cutoff threshold, and dotted vertical lines indicate the cutoff thresholds for moderate and strong fitness effects (−1, −0.5, 0.5, and 1). Bold text highlights genes with the strongest fitness effects (−<1 or >1).](https://doi.org/10.1073/pnas.2211789119)
LAPs preferentially catalyze the hydrolysis of leucine residues at the N terminus of peptides and proteins and are often thought of as housekeeping enzymes. In addition to this functionality, LAPs have distinct and complex roles in diverse organisms ranging from bacteria to higher eukaryotes. For example, LAPs are involved in oxidative lens aging in the cattle species Bos taurus (24, 25), serve as a defense protein in the tomato plant Lycopersicon esculentum (26, 27), are a central component of H₂S production, which may be important for pathogenesis in the pathogenic bacterium T. denticola (18), and aid the decay of particulate organic matter in marine systems (28). No LAPs have been previously implicated as a critical factor in UVR tolerance.

This study demonstrated that the S. elongatus LAP is capable of the hydrolysis of cysteine-glycine in addition to the expected activity against dipeptides containing leucine moieties. Several aminopeptidases from E. coli K-12, including PepA, PepB, PepD, and PepN, have been shown to confer cysteinyl-glycinase activity (21), and LAP cysteinyl-glycinase activity has also been reported in LAP purified from bovine lens, Arabidopsis thaliana, rat liver, and T. denticola (18, 25, 29, 30).

While the LAP demonstrated broad activity against L-Leu-p-NA across the pH gradient tested, there was no Cys-Gly hydrolase activity detected at pH levels at or above 8.5. Previous measurements of the cytosolic pH of S. elongatus have identified an increase in cytosolic pH from 7.3 ± 0.2 in the dark to 8.4 ± 0.1 in the light, due to H⁺ pumping from the cytosol into the thylakoid lumen triggered by photosynthetic activity (31). The results presented here suggest that, while the LAP may be functionally active as a leucyl aminopeptidase across all intracellular pH levels, it may preferentially function as a cysteinyl-glycinase when the cell is exposed to darkness. A similar phenomenon of pH dependence has been demonstrated for ribulose bisphosphate carboxylase/oxygenase (RuBisCO) and has been suggested for the regulation of chaperone function during light–dark cycles in S. elongatus (31, 32).

During the catabolism of glutathione (GSH), GSH is initially cleaved into glutamate and Cys-Gly by a gamma-glutamyltransferase (GGT), and Cys-Gly is subsequently degraded into glycine and cysteine by a cysteinyl-glycinase. Previous work in Synechocystis sp. PCC 6803, as well as Emiliania huxleyi and Thalassiosira pseudonana, have demonstrated that the onset of darkness triggers glutathione catabolism (33, 34). The affinity of the S. elongatus LAP toward cysteinyl-glycine under pH levels that are representative of the cytosolic pH during darkness likely facilitates the complete catabolism of glutathione during the dark period. The improved survival in UVR of the Vlap mutant under constant light relative to a light–dark cycle provides further evidence that this enzyme's relevance to UVR tolerance is most pronounced during the dark period and reduced cytosolic pH levels.

There are several arguments for importance of a complete glutathione catabolism pathway in S. elongatus. Cysteine appears to be the main limiting amino acid for glutathione synthesis, in addition to being an important source of sulfur for other thiols and general cell metabolism in cyanobacteria (35–37). Additionally, it has been proposed that Cys-Gly has important redox properties and can be toxic in high quantities. This toxicity is likely due to the prooxidant nature of Cys-Gly, as it is a more reactive thiol than GSH, capable of starting iron redox-cycling processes that produce ROS and subsequent oxidative reactions (38). In the absence of a complete or efficient catabolism cycle, there is potential for the buildup of the dipeptide Cys-Gly within the cell, which results in a paucity of free cysteine and glycine. Abundant Cys-Gly would simultaneously create a prooxidant environment and a lack of Cys substrate to reallocate toward glutathione-dependent mechanisms of tolerance of oxidative stress conditions. It is possible that the severe fitness defect under UVR upon loss of this gene is due to the
enzyme’s dual role as a housekeeping leucyl aminopeptidase during the day, and a cysteinyl-glycinase at night. Previous screens of microorganisms have not implicated a cysteinyl-glycinase or leucyl aminopeptidase as being major factors in UVR tolerance (39, 40). The lack of differential expression of lap upon acute UVR exposure may explain this oversight in screens that have relied upon transcriptomic data (Fig. 2). Furthermore, the survival of lap insertional mutants under the constant light + UVR condition suggests that the strong UV-sensitive phenotype in S. elongatus would require subsequent exposure to a dark period to be detected. Thus, this phenotype may be overlooked in screens that focused on acute UVR exposure to photosynthetic organisms, or UVR exposure under constant light (39, 41).

Cyanobacteria such as P. marinus and marine Synechococcus spp., which dominate vast oceanic areas, and other ecologically relevant phytoplankton such as the diazotroph T. erythraeum, Bacillariophyceae such as F. cylindrus and the domoic acid producing diatom P. multistriata, and the dinoflagellate photosymbiont of corals Symbiodinium microadriaticum, all harbor LAPs with strong sequence similarity to Synpcc7942_1190 and identical amino acids at the seven residues canonically involved in M17 family metal binding and catalysis (SI Appendix, Fig. S1). This conservation of functional residues suggests that LAPs may support UVR stress tolerance in key primary producers residing in upper mixed layers of marine systems. The conservation of the UV-sensitive phenotype of LAP knockout strains in E. coli and Anabaena sp. PCC 7120 support the proposal that LAPs serve a UVR tolerance role in a diversity of organisms. It is possible that the E. coli LAP tested in this study may not be subject to the same pH dependent mechanism as the phototrophic S. elongatus LAPs yet is nevertheless important under UVR stress.

The only gene other than Synpcc7942_1190 that confers moderate to strong fitness and has an obvious connection to the antioxidant cycling functional family is gshB, encoding glutathione synthase. GSH is canonically synthesized through two sequential adenosine triphosphate (ATP)-dependent steps, catalyzed by Glu-Cys ligase (gshA) and gshB; however, no homolog of gshA has been identified in S. elongatus PCC 7942. When GSH is oxidized (GSSG), it can be recycled back to GSH via glutathione reductase (gor; Synpcc7942_0842); however, mutation of gor had no more fitness cost under UVR stress than in white light alone. GSH is thought to be the major antioxidant present in cyanobacteria, playing a central role in ROS scavenging by providing electrons to various GSH-dependent enzymes (42). ΔgshB strains of Synechocystis sp. PCC 6803 have severe growth retardation in the presence of H2O2, as well as other forms of ROS, at concentrations that do not affect the WT or complemented strains, indicating that GSH is crucial for protection against diverse ROS species (33). Glutathione levels can be modulated by both light spectra and intensity in cyanobacteria (33, 43). Our results confirm the critical nature of GSH synthesis in S. elongatus under UVR and suggest that a complete catabolic cycle of GSH is necessary under natural conditions that involve UVR and a light–dark transition in S. elongatus as well as Anabaena sp. PCC 7120 and other microorganisms.

Fig. 6. Kinetic assays of recombinant Synpcc7942_1190 Lap protein reveal preferential affinity for cysteiny1-glycine. (A) SDS-PAGE gel of the 51-kDa recombinant Synpcc7942_1190 Lap protein. (B) Activity of the recombinant protein against various substrates, at pH 7.5 and 8.0. (C) Activity of the recombinant protein against cysteinyl-glycine in the presence of metal cofactors. (D) Activity of the recombinant protein against cysteinyl-glycine and L-leucine-p-nitroanilide over a pH gradient. Average intracellular pH levels of S. elongatus in the dark and light are overlaid, per Mangan et al. (31). (E) Samples containing 5 μg/mL of the recombinant protein were assayed for cysteinyl-glycine or L-leucine-p-nitroanilide at different substrate concentrations at pH 7.5. Km values are overlaid vertically as red dotted lines. Data points are mean (SD); n = 3.
The two genes from the DNA repair functional family that strongly affect fitness under UVR are \( phr \), encoding a deoxyribodipyrimidine type I photolyase, and \( uvr \), encoding excinuclease ABC subunit C. Deoxyribodipyrimidine photolyases catalyze the light-dependent repair of cyclobutyl pyrimidine dimers that form between adjacent DNA bases. Cyanobacteria from high-light biotopes tend to possess \( Phr \), whereas those from low-light biotopes may not (22, 44). \( Phr \) is important for resistance against UVC radiation in \( S. elongatus \), however, its enzymatic activity requires a light period following UVR exposure to drive catalysis, as would be experienced by an organism in the natural environment (45). Our attempt to recreate a natural midday UVR exposure environment in the laboratory detected \( phr \) as critical for fitness. The UvrABC endonuclease enzyme complex proteins are encoded in all analyzed cyanobacterial genomes, supporting a core role in DNA repair (46). In \( E. coli \), UvrA initiates contact with damaged DNA, transfers the DNA to UvrB, and UvrC creates incisions around the damaged site. Subsequently DNA helicase II releases the incised nucleotide (47). Interestingly, in this study, \( uvr \) was the only gene of the UvrABC complex to display a strong fitness effect (\( uvrA \) 0.15 ± 0.03 \( P < 0.05 \), \( uvrB -0.03 ± 0.08 \), \( pcrA -0.05 ± 0.23 \), under HUV). This result suggests redundancy and interchangeability of the functions of UvrA and UvrB.

The results of this screen suggest that the photosynthetic functional family genes \( psb27, psb28-1 \), and \( psb27 \) are fundamental for tolerating UVR stress in a cyanobacterium, likely by promoting the assembly and/or repair of PSII. \( Psb27 \) and \( Psb28-1 \) are needed for optimal growth under intermittent high-light/dark conditions, and \( Psb28-1 \) appears to be important for the synthesis of chlorophylls and apoproteins of chlorophyll-binding proteins (48–53). While two paralogs of \( Psb28 \) are present in \( S. elongatus \) (\( Psb28-1 \) and \( Psb28-2 \)), \( Psb28-2 \) mutants conferred no observable fitness change under UVR stress in our screen, corroborating previous reports that \( Psb28-2 \) may serve an alternative function (54). \( Psb \) mutants have light-sensitive phenotypes due to improper assembly of PSII cores and deregulated electron flow, in addition to an accumulation of \( Psb27- \) and \( Psb28-2 \)-containing PSII intermediates (53, 55, 56). Here, we identify these three photosystem genes as being of importance for UVR tolerance.

None of the genes identified as critical for fitness under UVR stress by the RB-TnSeq experiment, with the exception of \( Synpcc7942_1511 \) (fitness estimate −0.5), was differentially expressed following an 15-min exposure to the HUV period (SI Appendix, Fig. S2). This highlights that studies focusing on differential expression at discrete timepoints may overlook many genes that are important for UVR tolerance. Three genes important for UVR fitness were down-regulated after 2 h of HUV exposure: \( Psb28-1 \), \( Psb27 \), and \( Synpcc7942_1616 \). This decrease in expression was observed in tandem with a global down-regulation of genes involved with photosystems I and II and phycobilin-containing light-harvesting complexes. This down-regulation by UVR may decrease photosynthetic activity, with subsequent up-regulation to be expected late in the dark period in anticipation of dawn. Notably, the hypothetical protein encoded by \( Synpcc7942_1616 \) previously has been detected in PSI and PSI pulldown experiments, which when taken with the results of the RB-TnSeq experiment, suggests a potentially overlooked and important role of \( Synpcc7942_1616 \) in association with photosystem complexes (57).

The results of this study define the gene set that is critical for UVR tolerance under environmentally relevant UVR conditions in a model photosynthetic organism; however, it should be noted that in comparison to natural sunlight incident on Earth’s surface, the spectral quality of the UV lamp used in this study is enhanced in the UVB relative to UVA, weighting the fitness effects observed toward the impact of short-wavelength UVR. It is likely that essential genes, such as superoxide dismutase, which are not assayed with the RB-TnSeq library, are also important for fitness under UVR stress, and that relevant genes with redundant homologs may have escaped detection. While the importance of such genes should not be overlooked, the utilization of an RB-TnSeq library in \( S. elongatus \) under realistic UVR conditions has corroborated the importance of several canonical gene functional families to UVR tolerance, such as DNA repair, photosystem maintenance, and glutathione synthesis, and highlighted the key role of a previously overlooked leucyl aminopeptidase with cysteinyl-glycine activity, which may be conserved among many ecologically relevant phytoplankton.

**Materials and Methods**

**Growth Conditions and Initial Growth Experiments.** \( S. elongatus \) PCC 7942, stored in our laboratory as AMC66, (\( S. elongatus \)) was grown to an initial OD\(_{750} \) of 0.6, and 50 mL of culture was inoculated in triplicate into quartz bio-reactors (100 mL, outer diameter 30 mm, inner diameter 25 mm, length 200 mm). The bioreactors were suspended above white LED panels with diffusion screens at an incident intensity of 160 \( \mu \)mol photons-m\(^{-2}\)-s\(^{-1} \) at the bottom of the bioreactors. Two Q-Lab UVA 340 lamps (QUV-UVA 340, Q Lab) were positioned perpendicular to the white LED panels. The spectral output of the Q-Lab UVA 340 lamp resembles the spectrum of sunlight incident on Earth’s surface from ∼290 to 360 nm, albeit lacking in relative intensity from ∼365 to 400 nm (Dataset S3). The tubes were positioned at various distances from the UV lamps, in triplicate, resulting in the following UVR conditions: high UVR (HUV, 1 mW·cm\(^{-2} \)), medium UVR (MUV, 0.6 mW·cm\(^{-2} \)), and low UVR (LUV, 0.4 mW·cm\(^{-2} \)). UVR levels were measured with a Solarmeter Model
5.0 Standard UVA-B meter. A control group, PAR alone, was created by wrapping a single layer of Edmund Polyester High-Pass UV Filter Sheet (<10% transmission below 390 nm) around three bioreactors positioned at the same distance from the UVR source as the LUV condition. All bioreactors were bubbled with ambient air filtered through a 0.2 µm filter and subjected to a 12-h light:dark regime, with activation of the UV lamp during the central 6 h of the light period to mimic highest UVR exposure during the midday solar zenith. OD_{750} was continually monitored, and cultures were diluted upon reaching an OD_{750} of ~0.3 to maintain the optical thinness of the cultures, limiting the maximum attenuation of UVR across the 25-mm reactor to less than 30%.

**Growth Rates and Absorption Spectra.** S. elongatus was grown to an initial OD_{750} of 0.6, and 50 mL of culture was inoculated in triplicate into quartz bioreactors at an OD_{750} of 0.025. Growth rates were determined by measuring OD_{750} daily. Additionally, absorption spectra from 300 to 800 nm were recorded on a Tcean M200 plate reader, with BG-11 media used as a blank, and corrected to 0 at 750 nm. The spectra were normalized to the local maximum at 675 nm.

**ROS.** Samples were collected for analysis of ROS following the completion of three light:dark cycles. The fluorescent marker 2',7'-dichlorodihydrofluorescein diacetate (H_{2}DCFDA) was used following the methods described in Rastogi et al. (58). A total of 2.5 µL of 2 mM H_{2}DCFDA was added to 1 mL of sample for a final concentration of 5 µM and incubated at 30°C for 30 min. The fluorescence was then quantified at an excitation of 480 nm and an emission of 520 nm on a Tcean M200 plate reader with the gain manually set to 120. Fluorescence data were normalized to OD_{750} for each sample, and untreated-sample background fluorescence was then subtracted from treated-sample fluorescence values.

**RB-TnSeq Library Growth and Sampling.** A 1-mL aliquot of an S. elongatus PCC 7942 RB-TnSeq library, previously archived at ~80°C, was quickly thawed for 2 min at 37°C, resuspended in two flasks of 100 mL BG-11 with kanamycic (Km), and incubated at 30°C for 1 h at 30 µmol photons·m^{-2}·s^{-1} without shaking (9). The culture flasks were then transferred to 70 µmol photons·m^{-2}·s^{-1} on an orbital shaker. The cultures were subjected to a 12-h light:12-h dark regime for entrainment for 2 d, until the OD_{750} reached 0.25. The cultures were then combined and diluted to an OD_{750} of 0.025, and four replicates of 15 mL were spun down at 4,500 × g and frozen at ~80°C as time 0 samples to determine the population baseline. Aliquots of 50 mL of the RB-TnSeq outgrowth were then transferred to 12 quartz tubes and grown in identical conditions to the preliminary growth experiments for five generations. A total of 15 mL of culture from each tube was subsequently collected by centrifugation at 4,500 × g, and DNA was extracted from the pellet using a phenol-chloroform extraction (17).

**Barseq.** The method for Barseq analysis was conducted as previously described (59). In brief, amplification of barcodes from extracted DNA was conducted using 1 of 96 indexed forward primers for multiplexing, Barseq_P2_ITXX, and a common reverse primer, Barseq_P1. PCR was performed in a final volume of 50 µL using Q5 DNA polymerase, Q5 GC Enhancer (New England Biolabs), and the following thermocycler conditions: 1) 98°C for 4 min; 2) 25 cycles of 30 s at 98°C, 30 s at 55°C, and 30 s at 72°C; and a final extension at 72°C for 5 min. A total of 10 µL of each PCR product was then combined, purified with a DNA Clean and Concentrator Kit (Zymo Research), and quantified using a NanoDrop 2000 Spectrophotometer (Thermo Scientific). The pooled and purified product was then sequenced using Illumina HiSeq4000 SR75 by the IG5 Genomics Center at the University of California San Diego.

**Fitness Calculations.** Fitness values were calculated using a previously described R script (10). In brief, the number of reads for each sample was used as a normalizing factor between samples. Any barcode falling outside or within the peripheral 20% of a gene coding sequence was ignored. Of the remaining barcodes, any genes represented by fewer than three barcodes were removed. For each barcode in each sample, a pseudocount of one was added to the number of reads, divided by the total number of reads for the sample, and the sample-normalized number of reads was log_{2} transformed. Any gene without at least 15 T0 reads was also removed. For each gene, maximum likelihood was used to fit a pair of nested linear mixed effects models to the sample- and read-normalized log_{2} transformed counts. Genes with significant fitness differences between the UVR condition and PAR only control were identified by comparing the difference in the −2-log likelihoods of the models to a χ² distribution with one degree of freedom, estimating a P value, accounting for multiple testing by the FDR method of Benjamin-Hochberg (60), and selecting those genes with adjusted P values less than 0.02. We took the concept of C_{H} - C_{PAR} to be the estimated UVR-specific fitness effect of knocking out a given gene. Mutants that had fitness effects below a confidence threshold (FDR adjusted P value <0.05) were not included. Mutants with gene disruptions conferring a fitness effect of ≥1 or ≤−1 were classified as having a strong fitness effect, whereas mutants with gene disruptions conferring a fitness effect of 0.5 or ≤−0.5 were classified as having a moderate fitness effect. Fitness scores of >0.5 or <−0.05 were classified as having negligible fitness effects.

**RNA-Seq Sampling and Library Preparation.** S. elongatus was grown under the HUV and PAR conditions described for the initial growth and RB-TnSeq experiments for 3 d prior to sampling 15 min and 2 h after the onset of the UVR period. Triplicate 10-mL samples at each timepoint were collected, and 2 mL of an ethanol-phenol stop solution was added on ice. Samples were centrifuged at 4,500 × g for 10 min at 4°C, decanted, and the cell pellets were stored at −80°C. Total RNA was isolated and purified using the Zymo Research Quick-RNA Fungal/Bacterial MicroPrep Kit from frozen cell pellets previously harvested according to the manufacturer’s protocols. Ribosomal RNA was removed from 1 µg total RNA with the use of a QiAseq FastSelect - 5S/16S/23S kit (Qiagen). The resulting RNA-subtracted RNA preparations were made into libraries with the KAPA RNA HyperPrep kit incorporating short Y adapters and barcoded PCR primers. The libraries were quantified with a fluorescent assay (dsDNA AccuGreen quantitation kit, Biotium) and checked for proper size distribution and average size with a TapeStation (D1000 Tape, Agilent). Library pools were then assembled and a 1× SPRI bead cleanup was performed to remove traces of carryover PCR primers. The final library pool was quantified and run on an Illumina NovaSeq6000 with a PESO run configuration. The sequencing depth ranged from 8.8 to 12.8 million reads per sample.

**RNA-Seq Differential Expression Analysis.** Differential expression analysis was conducted using the following R packages: R/amr (R package version 1.30.0), GenomineDb (R package version 1.14.0), GenomicFeatures (61), GenomicAlignments, GenomicRanges (61), and DESeq2 (62). The workflow was followed as described by Love et al. (62) and the differential expression was performed using DESeq2. Differentially expressed genes between the HUV and PAR condition were selected for an adjusted P value <0.05 (Benjamini-Hochberg correction for multiple testing) and an absolute log_{2} of fold change equal to or greater than 2.

**Construction of Mutants and Complemented Strains.** The plasmid for the inserional mutation of Synpcc7942_1190 (Vlap) was taken from the unigenic set (JGS-24-F-1). The plasmid for complementation of the Vlap mutant strain with the native lap gene and promoter (Vlap-lap) was created as follows: pAM5217, the destination vector for the gene for complementation, was generated via seamless assembly with a GeneArt Seamless Cloning and Assembly Kit (Thermo Fisher Scientific) of individual parts pCVD020 (NS1-Tc, pAM4836), pCVD002 (SpSm, pAM4818), and pCVD015 (ccdB-Swal), available from the Cyanovector system (63). pAM5217 was then digested with Swal (NEB), and a Gibson assembly (NEB) between the linearized product and the PCR product of the region 243 bp upstream to 71 bp downstream of SynPCC7942_1190 was performed, forming plasmid pAM5816 (Table 1). The SynPCC7942_1190 PCR product was generated using the primer pair 5217 to 1190-F and 5217 to 1190-R using standard protocols (SI Appendix, Table S1). The lap mutant strain was then transformed with the pAM5816 plasmid.

The plasmid for complementation of the lap mutant with the LAP derived from Anabaena sp. PCC 7120 (pAM5835) was created by linearizing pAM2991 (Addgene plasmid #40248) (64) with EcoRI and BamHI, and performing a Gibson assembly (NEB) between the linearized backbone and the PCR product of PCC7120DELTAR_RS03160 (WP_010994414.1) using the primer pair 2991-A7120-LAP-R using standard protocols (SI Appendix, Table S1).

To generate the plasmid for the markerless deletion of a native LAP (PCC7120DELTAR_RS03160) in Anabaena sp. PCC 7120, the oligos A7120_gRNA-LAP-F and A7120_gRNA-LAP-R were annealed to produce a guide RNA template and were cloned into the Aar site of the CRISPR/cpf1 containing plasmid pAM5572 (11) by Golden Gate cloning (65). The guide RNA-containing plasmid was digested with KpnI-HF (NEB) and dephosphorylated with Antarctic
phosphatase, and the linearized backbone was gel purified. Homologous repair templates upstream and downstream of the target PCC7120DELTAR303160 gene were generated by PCR with the primer pairs AT120-LAP-U-F/AT120-LAP-U-R and AT120-LAP-D-F/AT120-LAP-D-R and cloned into the linearized backbone using a three-part Gibson Assembly (NEB) (SI Appendix, Table S1). The final plasmid product (pAM5799) was verified by sequencing.

Transformation of S. elongatus was achieved using standard protocols (17). Genotyping of S. elongatus was performed using colony PCR with Taq DNA Polymerase (Millipore Sigma) with sequence to encode a Strep-tag on the N terminus of the protein product. An additional primer pair (pet28-linear-streptag-F and pet28-linear-streptag-R) was used to amplify the PCR product of PCC7120DELTAR303160 (Sp, Sm) and native promoter (Sp, Sm, Tc) This study.

Table 1. List of plasmids used in this study

| Plasmid name | Description | Source |
|--------------|-------------|--------|
| UGS-24-F-1   | Unigene set plasmid for insertional mutation of Synpcc7942_1190 | (69)   |
| pCVD002      | SpR, SmR gene carried on a CYANO-VECTOR device (Sp, Sm, Ap) | (63)   |
| pCVD015      | Counter selectable ccdB-based cloning cassette carried on a CYANOVECTOR donor plasmid (Ap) | (63)   |
| pCVD020      | S. elongatus NS1 carried on a CYANO-VECTOR donor plasmid (Ap, Tc) | (63)   |
| pAM2991      | One-step cloning vector for overexpression with Ptrc promoter (Sm, Sm) | (64)   |
| pAM5217      | Cloning vector generated by seamless assembly of pCVD002, pCVD015, and pCVD020 (Sp, Sm, Tc) | This study |
| pAM5572      | RSF1010-based broad host range plasmid for genome editing using Cpf1/CRISPR technology (Sp, Sm) | (11)   |
| pAM5799      | pAM5572 with gRNA spacer sequence targeting alr0236, and homology regions flanking PCC7120DELTAR303160 (Sp, Sm) | This study |
| pAM5816      | Plasmid for chromosomal integration of Synpcc7942_1190 and native promoter (Sp, Sm) | This study |
| pAM5835      | pAM2991 with PCR product of PCC7120DELTAR303160 (Sp, Sm) | This study |
| pAM5836      | Synpcc7942_1190 cloned into a pet-28b(+) vector backbone with Strep-tag appended on 3' end | This study |

LAP protein was incubated at 5 µg/mL for 20 min at 37 °C and terminated with 5% trichloroacetic acid (TCA). The reaction product L-cysteine was measured as described by Gaitonde (67). L-cysteine concentrations were determined by absorbance at 560 nm, after subtracting a blank.

Leucine aminopeptidase activity was determined using L-Leu-p-Na (Millipore Sigma) as a substrate (68). The standard reaction buffer for these reactions was 50 mM Tris-HCl (pH 7.5 or 8.0), 2 mM L-Leu-p-Na, and 0.2 mM MnCl2, unless another pH, substrate concentration, or cation is indicated. The purified LAP protein was incubated at 5 µg/mL for 20 min at 37 °C and terminat

Table S1). The final plasmid product (pAM5799) was verified by sequencing.

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Recombinant Expression and Purification of SynPCC7942_1190 from E. coli.

Two primers (pet28-linear-streptag-F and pet28-linear-streptag-R; SI Appendix, Table S1) were used to linearize a commercially available pET-28b(+) vector (Millipore Sigma) with sequence to encode a Strep-tag on the N terminus of the protein product. An additional primer pair (pet28-1190-streptag-F and pet28-1190-streptag-R; SI Appendix, Table S1) was used to amplify Synpcc7942_1190 with appropriate overhangs for subsequent Gibson Assembly (NEB). The resulting plasmid (pAM5836) was used to transform E. coli DH5α cells for plasmid generation and sequence confirmation. The plasmid was subsequently moved into E. coli BL21 (DE3).

Plasmid-bearing E. coli BL21 (DE3) strains were grown to OD600 of 0.5 with 50 mg/mL kanamycin before overnight induction of protein expression at room temperature by the addition of 200 µM isopropyl β-D-1-thiogalactopyranoside. Cells were then collected by centrifugation and the pellet was resuspended in 50 mL of Streptactin wash buffer (50 mM Tris-HCl, pH 8.0, 150 mM NaCl, 5% glycerol). The resuspended cells were disrupted on ice by French press, and the lysate was clarified by centrifugation at 27,000 × g (Sorvall SS-34 rotor) at 4 °C for 30 min. Recombinant protein was purified using a Streptactin XT Superflow resin column (IBA Lifesciences). The column was washed with 20× the column volume of Streptactin wash buffer, followed by protein elution with same buffer containing 50 mM biotin, and collected in six elution aliquots. The elution aliquots were run on SDS-PAGE to assess purity, and the central three elution aliquots were pooled.

Enzymatic Assays. The standard reaction buffer for all cysteinyl-glycinase assays was 50 mM Tris-HCl (pH 7.5 or 8.0), 2 mM Cys-Gly, and 0.2 mM MnCl2, unless another pH, substrate concentration, or cation is indicated. The purified LAP protein was incubated at 5 µg/mL for 20 min at 37 °C and terminated with 5% trichloroacetic acid (TCA). The reaction product L-cysteine was measured as described by Gaitonde (67). L-cysteine concentrations were determined by absorbance at 560 nm, after subtracting a blank.

Leucine aminopeptidase activity was determined using L-Leu-p-Na (Millipore Sigma) as a substrate (68). The standard reaction buffer for these reactions was 50 mM Tris-HCl (pH 7.5 or 8.0), 2 mM L-Leu-p-Na, and 0.2 mM MnCl2, unless another pH, substrate concentration, or cation is indicated. The purified LAP protein was incubated at 5 µg/mL for 20 min at 37 °C and terminated with 5% TCA. The concentration of the reaction product was determined by absorbance at 405 nm, after subtracting a blank. All assays were performed in triplicate.

LAP Sequence Alignment. Proteins with sequence similarity to Synpcc7942_1190 were identified using Blastp (19) against the genomes of Anabaena sp. PCC 7120 (accession WP_010994414.1), F. cylindrus (accession OEU09452.1), P. marinus (accession WP_158467072.1), P. multistriata (accession VEU35739.1), Symbiodinium sp. CCMC2592 (accession CAE724717.1), Synecococcus sp. WH7803 (accession ASG62.1), and T. erythraeum (accession Q11A96.1). A multiple alignment of the amino acid sequences was conducted with Geneious Alignment in Geneious Prime version 2022.0.2. Additionally, Synpcc7942_1190 was queried against the Cyanorak database v2.1 (20).

Data, Materials, and Software Availability. RNA-Seq and RB-TnSeq data generated in this study have been deposited in the NCBI Sequence Reads Archive (SRA) database, https://www.ncbi.nlm.nih.gov/sra (Bio-Project identifier PRJNA854269) (70).

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