The moderating role of population succession in the adaptive responses of *Synechococcus* assemblages: evidence from light intensity simulation experiment

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Abstract

*Synechococcus* is one of the most abundant photoautotrophic picoplankton in the marine ecosystem. However, it is not clear how *Synechococcus* assemblages respond to light intensity variation in a genus group. Here, enriched *Synechococcus* assemblages from *in situ* coastal seawater were subjected to light intensity simulation experiments in a range of 9–243 μmol(photon) m⁻² s⁻¹. Characteristics concerning physiology, genomics, and metatranscriptomics were analyzed. Physiologically, the fitting model predicted photosynthesis indications and pigment contents increased with different trends following the light intensity. Genomic sequencing demonstrated that both the phylogenetic and phenotypic compositions of *Synechococcus* assemblage exhibited population succession. Especially, the proportion of *Synechococcus* pigment type 2 was changed significantly. In metatranscriptomics, most genes were downregulated in the high-light intensity group, while photosynthesis-related genes were entirely upregulated. The high upregulation of photosynthesis-related genes, such as *psbO*, *psbA*, *apcB*, and *cpcB*, corresponded to the succession of *Synechococcus* genotype and was responsible for the physiological shift in response to light intensity.

Keywords: enrichment culture; genomics; irradiance; metatranscriptomics; picocyanobacteria.

Highlights

- Both phylogenetic and phenotypic compositions of *Synechococcus* showed alterations
- Pigment contents and photosynthesis indicators increased with light intensity at different trends
- Regulation of photosynthetic genes was associated with the succession of genotypes

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Abbreviations: ANOVA – analysis of variance; AP – allophycocyanin; CA – carbonic anhydrase; Chl – chlorophyll; DEGs – differentially expressed genes; FDR – false discovery rate; GO – Gene Ontology; KEGG – Kyoto Encyclopedia of Genes and Genomes; LI – light intensity; LSM – least-square method; OTU – operational taxonomic unit; PE – phycoerythrin; PEC – phycoerythrocyanin; PC – phycocyanin; PCR – polymerase chain reaction; PT – pigment type; RDA – redundancy analysis; VIF – variance inflation factor; WGCNA – weighted gene co-expression network analysis.
Introduction

Marine Synechococcus, a polyphyletic group of picoplanktonic cyanobacteria, is one of the most abundant photosynthetic autotrophs in the oceans worldwide (Sohn et al. 2016). As an important marine picoplankton with global distribution, Synechococcus plays a significant role in the carbon flux and primary production in oceanic waters (Guidi et al. 2016). Estimates show that Synechococcus contributes 8 Gt of C to the net primary production per year, which is responsible for 16.7% of the marine net primary production, exceeding the production of other phytoplankton species (Flombaum et al. 2013).

In response to global climate change, the distribution of Synechococcus has changed considerably both in spatial and vertical scales. For example, a massive northward shift of species was observed with an increase in temperature. Indeed, Synechococcus has been recorded far beyond the polar circle (Paulsen et al. 2016); an expansion of its dominant area in the marine ecosystem is predicted (Flombaum et al. 2013). Light is considered an effective factor that determines the vertical distribution of Synechococcus in natural water bodies (Everroad and Wood 2012). The turbidity caused by suspended particulate matters, which has increased with an intensification of human activities (van der Hout et al. 2017), and the occurrence of algal blooms, such as the red tide due to eutrophication (Zohdi and Abbaspour 2019), have altered the light environment, including both light quality and light intensity (LI). Studies have shown that the abundance of Synechococcus is maximum in the surface mixed layer and declines with depth (Rocap et al. 2003, Scanlan et al. 2009). Synechococcus is typically less abundant below 100 m; however, it is found at depths of 200–600 m and even deeper during deep mixing events (Lindell and Post 1995, DuRand et al. 2001). The high physiological plasticity of Synechococcus indicates that its distribution may increase in the future. Hence, investigation of deep adaptation mechanisms in response to changes in environmental conditions is required.

Studies regarding the photosynthetic adaptive strategies of phytoplankton are important in the field of marine biology and physiology (Halsey and Jones 2015). Until now, researchers have conducted a series of in situ studies or pure culture-based physiological analyses to understand the intrinsic adaptation mechanisms in response to LI variation. Field measurements of chlorophyll fluorescence revealed that picophytoplankton in high-light and low-nutrient open sea preserved PSII activity by diverting electrons from the PSI complex-mediated carbon assimilation to oxygen (Mackey et al. 2008). Meanwhile, it maintained high photosynthetic efficiency and simultaneously prevent photodamage via a propyl gallate-sensitive oxidase associated with the photosynthetic electron transport chain. Marine Synechococcus lineages have also evolved ecotype differentiation in response to light, as observed in isolated culture experiments and clone library data (Ahlgren and Rocap 2006). Furthermore, proteomic research on light responses, ranging from optimized low light to optimized high light, revealed that the proteins of Synechococcus strains involved in photosynthetic electron transport, carbon fixation, and oxidative stress protection (superoxide dismutases) varied. The unique combinations of light responses in different strains resulted in distinct physiologically phenotypes that may result in Synechococcus niche division in the ocean (Mackey et al. 2017).

Culture experiments based on purified Synechococcus isolates are the most common methods used for investigating the adaptation mechanisms of phytoplankton under different environmental stresses (Śliwinska-Wilewska et al. 2020). However, Synechococcus is a genus with rich phylogenetic and phenotypic diversity, and pure cultures of only rare strains have been successfully established so far (Grébert et al. 2018, Xia et al. 2019). In addition, the succession of community and reciprocal or competitive interactions between strains are neglected to a considerable extent in pure culture.

Hence, in this study, we conducted a controlled simulation experiment to analyze response to LI using enriched culture samples of Synechococcus obtained from in situ coastal water and investigated the variation characteristics at physiological, genomics, and transcriptomics levels. The specific aims are as follows: (1) to detect the variation in physiological parameters, including growth rate, photosynthetic efficiency, and pigment content of Synechococcus assemblages in the LI-controlled experiment; (2) to determine the phylogenetic and phenogenetic community successions from in situ coastal water to enriched culture under LI gradient; (3) to explore the mechanisms underlying gene regulation of Synechococcus group in response to LI variation; (4) to construct the relationship among physiology, genomic structure, and gene regulation of Synechococcus assemblages for LI adaption.

Materials and methods

Sampling: Field sampling was conducted in the BHB10 station of the Bohai Bay on Research Vessel Kexue III on 24 August 2018 (Fig. 1S, supplement). Bohai Bay, a semi-closed bay, is located in the mid-latitude monsoon area. Its geohydrological conditions, along with the large amounts of pollutants discharged by the inflowing rivers, make it one of the most polluted coastal areas in China (Gao and Chen 2012). The BHB10 station is located at the center of the Bohai Bay, which is adjacent to the Tianjin Port that experiences frequent occurrence of harmful algal blooms (Liu et al. 2019). Two parallel original seawaters, B1 and B2, were collected at BHB10. To detect the in situ composition of Synechococcus assemblages, 1,000 ml of seawater was filtered on 0.22-μm polycarbonate filters (Millipore Co., Bedford, MA, USA). Triplicate membranes from each sample were placed in sterile cryopreservation tubes and flash-frozen in liquid nitrogen. Upon returning to the laboratory, the samples were stored at −80°C for subsequent molecular experiments. For the simulation experiment, 100 ml of seawater was collected in a sterile
culture bottle containing 50 ml of SNAX medium with actinomycin solution (final concentration 0.05 mg ml⁻¹) to exclude eukaryotes (Deng et al. 2014, Hunter-Cevera et al. 2016). After returning to the laboratory, the samples were enrich-cultured for subsequent light-controlled experiments.

**Culture experiments**: The two enriched culture samples, B1 and B2, were inoculated with the same initial cell density in conical flasks containing the SNAX medium. The simulation experiment was performed in ZQZY-BGF8 combined light shaker incubators (Shanghai Zhichu Instruments Co., Ltd., China). The four LI were set as low (L), middle-low (ML), middle-high (MH), and high (H) LI gradients using 9, 27, 81, and 243 μmol(photon) m⁻² s⁻¹, respectively, in three parallel samples. The light and dark cycle was set at 12:12 h and the temperature was maintained at 21°C. Physiological parameters were determined after cultivating for 15 d. All samples were collected in the end using 0.22 μm-polycarbonate membranes (Millipore Co., Bedford, MA, USA) for genomic and transcriptomic analysis.

**Physiological parameters**: During experimental periods, 1.40 ml of culture liquid of each flask was taken in a sterile tube every day, followed by the addition of polyoxymethylene (the final concentration was 0.5%) for recording cell density using flow cytometry (Nowack et al. 2015). The growth rate of each sample was calculated over the experimental period from the daily cell density using linear regression models (Jodłowska and Śliwińska 2014). The net photosynthetic rate (net oxygen growth rate) was monitored using an oxygen electrode (Hansatech, King's Lynn, UK) (Ueno et al. 2020). Carbonic anhydrase (CA, EC 4.2.1.1) activity was determined from the time required for the pH to decrease from 8.1 to 7.3 after ultrasonic extraction in 20 mM barbiturate buffered saline (Hu et al. 2010). Chlorophyll (Chl) a content was determined from the optical densities at 647 and 664 nm after extraction with 90% acetone (Wagner et al. 2016). Contents of phycobiliproteins, including phycocerythin (PE), phycocyanin (PC), and allophycocyanin (AP), were determined from the optical densities at 565, 620, and 650 nm (UV-VIS spectrophotometer TU-1810, Persee, China), respectively, after ultrasonic extraction in 100 mM phosphate-buffered saline (Mittal et al. 2017).

**Genome sequencing and analysis**: DNA was extracted from the filter membranes using the Fast DNA spin kit (MP BIO, USA) according to the manufacturer's protocol. Polymerase chain reaction (PCR)-mediated amplification of DNA was performed using the thermostyler GeneAmp® PCR system 9700 (Perkin Elmer, USA) in triplicate following previous methods (Li et al. 2019). *rpoC1* was amplified using the primer pair *rpoC1*-39F (5-GGNATNNTNNTGGAGCGGTYG-3) and *rpoC1*-462R (5-CGYGRCGCTTGRCTAGCTT-3) (Xia et al. 2019). Then, the purified *rpoC1* gene products were sequenced at the Majorbio Co., Ltd. (Shanghai, China) using the PE300 Illumina MiSeq sequencing platform. The *cpeBA* operon sequences were amplified using the primer pair *peBF* (5-GACCTACATCGCWCTGGGYG-3) and *peAR* (5-CMACAACGRCARGCGTAGTT-3) (Xia et al. 2018). The purified products of the *cpeBA* operon were sequenced at the Shanghai Biozoon Biological Technology Co., Ltd. (Shanghai, China) using the PE250 Illumina MiSeq sequencing platform.

Raw FASTQ data were quality-controlled, paired, chimera-filtered, and redundancy eliminated for downstream analysis (Caporaso et al. 2010). Operational taxonomic unit (OTU) clustering was performed on the nonrepetitive sequences considering 97% similarity and the representative OTU sequences were obtained. The representative sequence of each OTU was identified using BLAST search (Altschul et al. 1997). Sequences that do not belong to the phylum Cyanobacteria were deleted from the results of MegaBlast. The most similar reference sequences were retrieved from the NCBI database [reference sequence information of *rpoC1* is listed in Table 1S (supplement), and that of the *cpeBA* operon is listed in Table 2S (supplement)]. The sequence that accounted for more than 1% in each sample was aligned with reference sequences using ClustalW according to codon structure, and the Maximum Likelihood phylogenetic tree was constructed with MEGA 7 using the best model GTR+G+I for nucleotide evolution with 200 bootstraps (Xia et al. 2018). Phylogenetic and phenotypic information of *Synechococcus* strains was derived from published literature (Grébert et al. 2018) and the Roscoff Culture Collection (https://roscoff-culture-collection.org/strains/shortlists/taxonomic-groups/marine-synechococcus).

**Metatranscriptomic sequencing and analysis**: Metatranscriptome analysis was performed to investigate the effect of LI on gene regulation. The physiological parameters of *Synechococcus* assemblage showed a nonlinear increase from 9 to 243 μmol(photon) m⁻² s⁻¹; hence, we used the samples cultured under L and H for analysis. Total RNA was extracted from the filtered membranes using TRIzol® reagent according to the manufacturer's instructions and genomic DNA was removed using DNase I (TaKara, Otsu, Japan). Then, *rRNA* was removed using the Ribo-Zero rRNA removal kits per the manufacturer's instructions (Illumina). Optimized high-quality RNA samples were used for downstream experiments. Metatranscriptome libraries were prepared using TruSeq™ RNA sample preparation kit from Illumina (San Diego, CA, USA). Metatranscriptome sequencing was performed by Shanghai Biozoon Biotechnology Co., Ltd. using the Illumina Novaseq 6000 instrument.

The raw paired-end reads were trimmed and quality-controlled using Trimmomatic v. 0.36 (Bolger et al. 2014). High-quality reads were then aligned to the SILVA SSU (16S/18S) and SILVA LSU (23S/28S) databases to remove *rRNA*-related reads using the SortMeRNA software (Kopylova et al. 2012). Then, clean data from all samples were used for assembly with MegaHit (Li et al. 2016). All the genes were predicted using METAProdigal (Hyatt et al. 2012). The nonredundant gene catalog was
constructed with 95% identity and 90% coverage using CD-HIT (Fu et al. 2012). To identify the proteins that showed the highest sequence similarity with the given transcripts and retrieve their functional annotations, all genes were searched against the NCBI protein non-redundant, String, and Kyoto Encyclopedia of Genes and Genomes (KEGG) databases using BLASTp with E-values less than $1.0 \times 10^{-5}$. Metabolic pathway analysis was performed using KEGG (Kanehisa and Goto 2000). Salmon was used for calculating the expression level of each transcript, and RSEM was used to quantify gene and isoform abundance (Li and Dewey 2011, Roberts and Pachter 2013). EdgeR in R was utilized for differential expression analysis (Robinson et al. 2010). The differentially expressed genes (DEGs) with logarithmic fold change in expression $> 2$ and false discovery rate (FDR) $< 0.05$ were selected for Gene Ontology (GO) functional enrichment and KEGG pathway analysis (Klopfenstein et al. 2018). In total, 678 genes, the expression level of which was $\geq 10$ in at least one sample, were selected to perform weighted gene co-expression network analysis (WGCNA) using an R package (Langfelder and Horvath 2008). Gephi and Cytoscape were used to visualize results from the co-expression networks. Hubba genes of each module were determined using Clusterine Coefficient of CytoHubba, a Cytoscape plugin (Chin et al. 2014).

**Statistical analysis:** Package basicTrendline in R was used for fitting each physiological parameter with the corresponding LI using three models: linear, logarithmic curve, and power curve (Mei et al. 2018). Package agricolae in R was used for one-way analysis of variance (ANOVA) with post hoc test (LSD) (de Mendiburu 2014). Package vegan in R was used for calculating redundancy analysis (RDA) between physiological parameters and community composition (Oksanen et al. 2013). The least-square method (LSM) was used to further calculate the linear relationship between physiological parameters and the percentage of OTU. Packages psych and pheatmap in R were used for sketching correlation heat map between gene expression as well as physiological parameters and Synechococcus composition.

**Data availability:** The sequence data of this study were deposited in the sequence read archive of NCBI. A bio-project associated with this study was applied for and processed in NCBI with the accession number PRJNA680823. All raw sequencing data were stored under this bio-project.

**Results**

**Physiological responses to LI:** The physiological responses of B1 and B2 to LI showed similar trends (Fig. 2S, supplement). All parameters increased with LI. The highest growth rate, net photosynthetic rate, CA activity, PC content, and AP content all appeared in B2-H, reaching 0.45 $d^{-1}$, 1.42 mmol $L^{-1} s^{-1}$, $1.81 \times 10^{-2}$ EU $ml^{-1}$, $3.77 \times 10^{-2}$ mg $ml^{-1}$, and $1.45 \times 10^{-2}$ mg $ml^{-1}$, respectively. The Chl $a$ and PE contents were maximal in B1-H, which were $1.06 \times 10^{-3}$ mg $ml^{-1}$ and $1.12 \times 10^{-2}$ mg $ml^{-1}$, respectively. Most PE, PC, and AP contents under L were below the detection limit. In B1, the CA activity under MH ($1.64 \times 10^{-2}$ EU $ml^{-1}$) was slightly higher than that under H ($1.58 \times 10^{-2}$ EU $ml^{-1}$) but did not differ significantly.

The nonlinear power curve model was always the best model for fitting each parameter with its LI (Table 3S, supplement). The power values, $b$, were less than 0 in models of growth rate, net photosynthetic rate, and CA activity; hence, the maximal value of these physiological parameters was dependent on the constant $c$ (Fig. 1). The values of $b$ in models of Chl $a$, PE, PC, and AP content were between 0 and 1, and consequently, these physiological parameters continued to increase with LI without attaining any maximal value. The $p$-value of each model was less than 0.01, indicating the dependability of this nonlinear relationship between each parameter and LI.

**Changes in genomic composition in response to LI:** On average, rpoC1 generated 21174 sequences with coverage of more than 0.99 in each sample (Table 4S, supplement). In total, 16 OTUs were detected with relative abundances accounting for more than 1% in each sample, among which 11 OTUs were from in situ samples, which decreased to 5 OTUs in cultured samples (Fig. 2A). These OTUs occupied 97.8% of the total sequences on average. All three Synechococcus subtypes, S5.1, S5.2, and S5.3 were detected, of which S5.1 was further divided into two clades: VIII and IX. Both samples of in situ seawater, B1-I and B2-I, were dominated by S5.1 clade IX, followed by S5.3 and S5.2. Clade VIII was the least abundant in the in situ water samples. Compared to those of the in situ samples, the Chao index and Shannon index of the cultured samples decreased to 86.3 and 85.4%, respectively. The Synechococcus assemblage in all culture samples was composed of two OTUs of clade VIII, which contributed to 87.4% (B1-MH) and 99.9% (B2-H) sequence abundance. OTU 2380, most dominant in B2-MH and all culture samples of B1, occupied more than 82.8% of the sequences. OTU31 was the most abundant in B2-H, reaching 97.5% abundance. In the remaining samples B2-ML and B2-L, each of these two OTUs accounted for about half of the sequences.

The cpeBA operon generated 40,825 sequences on average, with coverage of more than 0.99 in each sample (Table 4S). Thirty-three OTUs were detected using the cpeBA operon, accounting for more than 1% in each sample, among which 22 OTUs were from in situ samples; this decreased to 11 OTUs in the cultured samples (Fig. 2B), which occupied 82.90% of total sequences on average and included three PTs: PT2, PT3a, and PT3c/PT3d. A similar trend in both in situ and cultured samples was observed – the proportion of PT2 was higher than that of PT3a and PT3c/PT3d. Compared to those of the in situ samples, the Chao index and Shannon index of the cultured samples decreased to 28.9 and 41.8%, respectively. The proportion of PT2 increased two-fold in the samples after culture. In general, OTU1 (30.4–40.1%) and OTU2 (32.2–40.0%) accounted for a large bulk of the Synechococcus assemblage composition, while the other OTUs were rarely detected.
Significant differences in PT2 percentage between B1-MH, B1-ML, and B2-L were observed (Fig. 2D).

The VIF value indicated strong autocorrelation between contents of PE, PC, and AP, leaving only PC for eliminating the effect of collinearity in calculating RDA. The permutation test for all constrained eigenvalues in the model of RDA between physiological parameters and OTUs based on rpoC1 showed statistical significance (Fig. 3A). Three physiological parameters, net photosynthetic rate ($R^2 = 0.34$, $p<0.01$), Chl $a$ content ($R^2 = 0.37$, $p<0.01$), and PC content ($R^2 = 0.24$, $p<0.05$), significantly correlated with the community. Noticeably, the relationship between two main OTUs, OTU31 and OTU2380, and the parameters were opposite. Further fitting calculation showed a significant linear relationship between the percentage of OTU31 and PC content (Fig. 3S, supplement). However, relationships between most of the other OTUs and physiological parameters were not simply linear. Comparably, the permutation test for all constrained eigenvalues showed that the $p$-value of the RDA model between physiological parameters and OTUs based on cpeBA operon was 0.14, which was > 0.05, and hence did not reach statistical significance (Fig. 3B). Therefore, Spearman’s correlation was calculated to investigate the interaction between phenotypes of *Synechococcus* and physiological parameters (Fig. 4S, supplement). We found that OTU2, OTU3, OTU6, and OTU8, which belonged to different PTs, showed significant correlations with all physiological parameters. OTU15 and OTU54 correlated with Chl $a$ content. Notably, the relationship between OTUs belonging to the same PT and physiological parameters could differ considerably.

**Metatranscriptional response to LI:** The metatranscriptome sequencing of each sample generated about 20 GB of data. In total, 7,145 genes were differentially expressed between the H group (combined B1-H and B2-H) and the L group (combined B1-L and B2-L). Among them, the expression of 1,839 genes was significantly more in group H than in group L (hereafter, upregulated), and 5,306 genes were expressed significantly less in group H than in the L group (hereafter, downregulated) (Fig. 4A). In total, 3,252 differentially expressed genes were annotated in 81 KEGG pathways, which could be classified into six functions of KEGG level 1 and 34 functions of KEGG level 2 (Fig. 4B). Among the 50 genes with the highest expression level, eight genes were related to photosynthesis, and all of them were
The genes associated with photosynthesis were mostly classified in two KEGG pathways, ko00195 and ko00196 (Fig. 5). One hundred and twenty-three genes were annotated in the pathway map of ko00195 with 27 unique functions, involving PSI, PSII, cytochrome b/f complex, photosynthetic electron transport, and F-type ATPase. Forty-five of these genes were upregulated and 78 were downregulated. In addition to the genes encoding the F-type ATPase, all other matched genes were upregulated.

Sixteen genes were annotated in the pathway map of ko00196 with 12 unique functions, involving regulation of AP, PC/phycoerythrocyanin (PEC), and PE; all these genes were upregulated.

Five significant Spearman correlations between gene expression associated with photosynthesis and physiological parameters were detected (Fig. 6A). All the correlations were strongly negative (R^2 < 0.6). The expression levels of psaB, psbO, and psbV were significantly associated with percentages of OTU31 and
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OTU2380 (both OTUs belong to Synechococcus clade VIII) in the sequence of rpoC1 (Fig. 6B). The expression of 13 types of genes correlated with the percentages of three OTUs, OTU108, OTU218, and OTU36 in the sequence of rpoC1 (Fig. 6B). The expression of 13 types of genes correlated with the percentages of three OTUs, OTU108, OTU218, and OTU36 in the sequence of rpoC1 (Fig. 6B).
OTU108 and OTU218 belonged to *Synechococcus* PT2, while OTU36 belonged to *Synechococcus* PT3a.

**Gene co-expression network analysis:** Seven gene modules were identified from the results of WGCNA (Fig. 7; Fig. 5S, supplement). Among them, modules 1, 5, and 6 clustered together based on the module eigengenes of modules and were separated from others (Fig. 5S,B). The topological feature of co-expression networks constructed based on each module is listed in Table 5S (supplement). Module 7 occupied more than 44% of 678 genes with expression level ≥ 10 in at least one sample. Modules 1 and 2 accounted for 27 and 16% genes, respectively, and the other four modules added up to account for 13% genes. After annotation using the

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**Fig. 5.** Pathway maps of entry ko00195 and ko00196 associated with photosynthesis annotated using the *KEGG* database. The red box marked genes significantly upregulated in group H (high-light intensity) compared with those in group L (low-light intensity). The yellow box represents the presence of both upregulated and downregulated genes.

**Fig. 6.** Spearman correlation heat map between photosynthesis associated gene expression and physiological parameters (*A*), operational taxonomic units (OTUs) accounting for more than 1% abundance based on *rpoC1* (*B*) and *cpeBA* operon (*C*). The color legends on the right represent the value of the Spearman correlation coefficient. *– p* < 0.05, ** –* p < 0.01. AP – allophycocyanin; CA – carbonic anhydrase; PC – phycocyanin; PE – phycoerythrin.
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KEGG database, more than 53% of genes were assigned to at least one function. Three hubba genes of networks were annotated in pathway ko03010, one was annotated in ko01100, while the others could not be assigned to any pathways. In total, 19 genes targeting KEGG pathways ko00195 and ko00196 were found, of which 79% occurred in module 1.

**Discussion**

**Photosynthetic capacity increased but efficiency decreased in high LI.** The power values of growth rate, net photosynthetic rate, and CA activity were less than 0, indicating the appearance of their predictable maximums with the increase in LI in the power curve model. This indicated that 0.46 d^{-1}, 2.45 mmol L^{-1} s^{-1}, and 0.60 \times 10^{-4} EU ml^{-1} were probably their upper limits, respectively, even when the LI increased beyond H. Comparatively, the power values of Chl a, PE, PC, and AP content were between 0 and 1, and hence these parameters continuously increased with LI, although the increase rate might be slow. CA plays a key role in catalyzing the critical first step of C_4 photosynthesis and the hydration of CO_2 to bicarbonate (Jensen et al. 2019), and the value of net photosynthetic rate reflected the ability to produce oxygen. The continuous increase in photosynthetic pigment content with stagnation in the abilities to absorb carbon dioxide and produce oxygen indicated that the photosynthetic efficiency per unit pigments might decrease with a further increase in LI. Furthermore, the content of pigment-proteins in the enrichment culture of *Synechococcus* assemblage might be lower than that of pure culture. The PC and APC contents of *Synechococcus* strain PCC 6715 (PT1) were determined; it showed that under continuous white light, the PC and APC contents were 8 \times 10^{-2} and 2 \times 10^{-2} mg mL^{-1}, which were about eight times higher than our results (Klepacz-Smółka et al. 2020).

**Alteration of Synechococcus assemblages from in situ water to laboratory culture:** Enrichment culture is closer to the natural state of *Synechococcus* assemblages than pure culture. However, after cultivation, the change in population composition and relative abundance was still large. Phylogenetically, marine *Synechococcus* formed a well-defined clade termed cluster 5, which was divided into three subclusters: S5.1, S5.2, and S5.3; each subcluster can
be further divided into various clades (Fuller et al. 2003). Generally, Synechococcus S5.1 is the most abundant and widespread subcluster in the ocean worldwide; in particular, its clades I, II, III, and IV dominate most sea waters (Zwirglmaier et al. 2008). After enrichment culture, Synechococcus assemblages of B1-I and B2-I were dominant in clade IX of S5.1. This clade was also observed in the adjacent Yellow Sea and the East China Sea but is rarely observed in other areas of the sea (Choi and Noh 2009, Wang et al. 2021). S5.2, occupying more than 15% of the site, is globally distributed in estuaries and brackish waters (Jing et al. 2009). The abundance of S5.3 is usually low (Ahlgren and Rocap 2012), and it was almost twice as much as S5.2 in this study. It is not clear whether this phenomenon is due to the unique niche occupation pattern of Synechococcus formed by the conditions of the semi-enclosed Bohai Sea. For further exploration, samples should be collected and cultured from more stations and different depths in the future research. Interestingly, the original dominant assemblages, including clade IX, S5.2, and S5.3, almost disappeared after enrichment culture. On the contrary, clade VIII, which accounts for less than 1% of the population in situ, was found to be the main population. Clade VIII is not a common Synechococcus lineage. Physiologically, some strains of clade VIII lack PE and cannot grow on nitrate (Fuller et al. 2003), and its distribution may be related to salinity, as it is found in hypersaline pools (Huang et al. 2012).

Synechococcus strains possess rich phenotypic diversity (Everroad and Wood 2012). In particular, the differences in evolutionary history between the major light-harvesting pigment PE and the core genome results in diverse PTs (Grébert et al. 2018). Some lineages are preferentially associated with a specific PT, such as most strains in clades I–IV and CRD1, while others exhibit a wide range of adaptability of PTs. As mentioned above, clade VIII normally lacks PE and is considered as PT1 (Fuller et al. 2003). However, according to the sequencing result based on the cpeBA operon, Synechococcus PT2 containing both PC and PE-I is the spectral phenotype corresponding to clade VIII. This can be attributed to the changing LI of the cultivating environment, which represents a selective agent that promotes the evolution of phenotypes adapted to different light environments, probably via mutation, positive directional selection, and horizontal gene transfer (Stomp et al. 2004).

Upregulation of genes related to photosynthesis: Metatranscriptomics is a powerful tool for capturing gene expression patterns in microbial communities without previous assumptions regarding the ongoing activities or dominant taxa (Frias-Lopez et al. 2008). Metatranscriptome research is still limited by difficulties in the construction of gene sets. In cases of prokaryotic polycistronic structure and low abundance of some genes, the lengths of the contigs are not sufficient for assembly, resulting in mapping that is far from satisfactory. In addition, deviations in results from the transcriptional data can be resolved only for individual genes with extremely high expression. To circumvent the above limitations, a mixed sample was used for metagenomic sequencing to obtain more comprehensive and accurate DNA-based metagenomic sequencing (Gilbert et al. 2008, Poretsky et al. 2010). The elimination of symbiotic species in the culture experiment of Synechococcus, even in the pure culture, is difficult (Zheng et al. 2020). Hence, the result of metatranscriptomic analysis may contain genes from these symbiotic species. However, Synechococcus is the only photoautotrophic organism in the culture system; hence, we believed that the regulation of genes related to photosynthesis was mostly because of the Synechococcus assemblage. Among the 7,145 differentially expressed genes, the downregulated genes occupied about 74% of the total, which is more than the proportion of upregulated genes. But interestingly, after annotation using the KEGG database, all matched genes were significantly upregulated in pathway map ko00190 and ko00196 associated with the photosynthetic system, except for genes encoding F-type ATPase. Proteins or subunits of F-type ATPase are associated with the hydrolysis of a nucleoside triphosphate and electron transport and are also enriched in other pathways overlapping with ko00190 and ko01100 (Kanazawa et al. 1981). Therefore, under high LI, the metabolic activities associated with photosynthesis were enhanced in Synechococcus assemblages. This may inhibit other metabolic activities, although whether these repressed metabolic activities are due to Synechococcus or other symbiotic species remains unknown. Upregulation of genes related to photosynthesis did not necessarily correlate with the increase in photosynthetic efficiency. The interactions between gene expression and physiological parameters showed a negative correlation. A gene encoding the photosystem II protein, Psb27, was upregulated, which is a key protein for efficient repair of photodamaged PSII (Chen et al. 2006). An increase in its expression indicated that the PSII had already undergone a certain extent of damage under the LI of H [243 μmol(photon) m⁻² s⁻¹].

Synergistic effect of population succession and gene regulation in photosynthetic adaptation: Population succession in Synechococcus assemblage is involved in its light adaptation, which was largely neglected in studies using pure culture. At the OTU level, both the phylogenetic and phenotypic composition of Synechococcus assemblage varied with the LI. In particular, in terms of phenotype, significant differences in the proportions of PTs between samples cultured under different LI were detected. Although RDA cannot be used to investigate the interaction between Synechococcus phenotypic composition and physiological parameters, Spearman correlations were observed between them. It is noteworthy that in addition to OTUs belonging to different PT, the relationship between OTUs belonging to the same PT and physiological parameters may differ, and may even be opposite. Therefore, the use of only one strain for studying physiological characteristics will lead to overgeneralization. Moreover, the relationship between the expression of some genes and physiological parameters, as well as some OTUs, demonstrated that gene regulation
is also an important part of adaptation. Population succession and gene regulation are jointly responsible for photosynthetic adaptation with LI change. Compared to our results, photosynthetic parameters of purified Synechococcus isolates appear to vary more strongly with LI change in culture experiments in previous studies (Jodlowska and Śliwińska 2014, Śliwińska-Wilczewska et al. 2020). Such differences in the variation trends might be because of the specific characteristics of the selected isolates in purified experiments. In the screening of Synechococcus isolate, the adaptation from natural light to culture light is the key process, which may result in a similar response to LI among different strains (Nowack et al. 2015). However, changes in community diversity and self-regulation in population structure in enrichment experiments lead to influences from the environment that tend to be moderating and steady. Whether there is an effect of the quorum sensing in assemblages needs further experiments.

Potential genes involved in adapted regulation of light intensity: The enormity and complexity of the metatranscriptomic data render exploration of relationships between phenotypes and genes difficult. In fact, in addition to the two known pathways (ko00195 and ko00196), numerous other genes are involved in the regulation. WGCNA revealed that 678 expressed genes were transformed into seven co-expression modules. Among them, module 1 collected more than 79% of genes annotated in KEGG pathways ko00195 and ko00196. In total, 52.7% of genes in module 1 could be annotated using the KEGG database, of which 56.3% belonged to ko03010 (ribosome). The hubba gene of module 1, RP-L4, encoding the large subunit ribosomal protein L4, was also in this pathway, which indicated that the regulation in response to light intensity activated the translation of various proteins. Previous proteomics research has shown that regulation of Synechococcus proteins involved in photosynthetic electron transport, carbon fixation, oxidative stress protection, and iron and nitrogen metabolism varied among Synechococcus strains that responded to an increase in LI (Mackey et al. 2017). rpoC1, a single copy housekeeping gene targeting the DNA-directed RNA polymerase beta subunit, is used to study the phylogenetic diversity of Synechococcus (Mühling et al. 2006). Consistent with the results of genomic sequencing, rpoC1 was not differentially expressed. Furthermore, other genes encoding the RNA polymerase subunits alpha (rpoA) and beta (rpoB) were found to be downregulated. The key function of rpoA is RNA polymerase assembly (Lee et al. 2012), while rpoB and rpoC1 may collaborate to bind DNA and RNA or synthesize RNA (Sutherland and Murakami 2018). Therefore, it is reasonable to speculate that certain changes occur in the process of transcribing the genetic information stored in DNA to RNA under a gradient of light intensity. In addition, many upregulated genes were clustered in module 1, the functions of which were associated with metabolic pathways (such as bsmB, ndhC, and gsmt), biosynthesis of secondary metabolites (chlL and pbsA1), carbon metabolism (aceE and prcL), and ABC transporters (psbB). The functions of these genes could be closely interconnected to regulate adaptation under the stress of fluctuating light intensity.

Conclusions: The characteristics of Synechococcus assemblage in terms of physiology, genomics, and metatranscriptomics were revealed in the LI simulation experiment. We found that: (1) Physiological parameters exhibited positive responses to the LI in the studied scale. However, the model of the power curve predicted that the growth rate, net photosynthetic rate, and CA activity would reach their maximum value, while the contents of Chl a, PE, PC, and AP would keep increasing with further increase in LI beyond 243 μmol(photon) m⁻² s⁻¹. (2) Compositions of Synechococcus assemblages derived from rpoC1 and cpeBA operon sequences shifted with LI change. Synechococcus PT2 was the most affected genotype. Such genomic succession could explain variations in physiological parameters, especially in the content of pigment proteins. (3) 74% of genes in metatranscriptome were downregulated in the H group. However, genes related to photosynthesis, most annotated in two KEGG pathways of ko00195 and ko00196, were commonly upregulated. Regulation of these genes was closely associated with the succession of genotypes and physiological shifts.

Based on the results, it is reasonable to infer that Synechococcus transfers energy to photosynthesis-related genes by reducing the expression of other metabolic activity-related genes, increasing photosynthetic parameters under high light intensity. Self-regulation in Synechococcus assemblages may lead to a moderate and steady response to the light intensity. However, whether there is an effect of the quorum sensing needs further experiments. Further studies will focus on investigating the expression and function of related genes using gene silencing assays to reveal the complete and accurate regulation strategies under conditions of varying LI. Mesoscale and large-scale field experiments should be conducted to better simulate the in situ Synechococcus assemblages and their adaptive strategies should be discussed.

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