Effects of a Type I RM System on Gene Expression and Glycogen Catabolism in Synechocystis sp. PCC 6803

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Increasing evidence has shown that DNA methylation is involved in gene regulation in prokaryotes. However, there have been very limited reports about the role of DNA methylation in regulation of gene expression and physiological functions in cyanobacteria. In Synechocystis sp. PCC 6803, four genes on the plasmid pSYSX are predicted to encode the type I restriction-methylation system, slr6095 and slr6096 for the M subunit, slr6097 for the S subunit and slr6102 for the R subunit. Compared to the wild type, slr6095, slr6096, and slr6097 mutants lacked the GG\textsuperscript{m6}AN\textsubscript{7}TTGG/CCA\textsuperscript{m6}AN\textsubscript{7}TCC methylation in genomic DNA. Transcriptomic analysis indicated that 171 genes were reproducibly up- or down-regulated in all three mutants relative to the wild type. The changed expression of some genes, including sll1356 for glycogen phosphorylase (GlgP), was associated with the loss of GG\textsuperscript{m6}AN\textsubscript{7}TTGG/CCA\textsuperscript{m6}AN\textsubscript{7}TCC methylation in the coding regions or the upstream non-coding sequences. Inactivation of slr6095, slr6096, or slr6097 increased the expression of sll1356 and the GlgP activity but lowered the glycogen content. These results indicated that the DNA methylation by a type I RM system could alter the expression of certain genes and physiological functions in Synechocystis sp. PCC 6803.

Keywords: Synechocystis, type I RM system, gene regulation, glgP, glycogen catabolism

INTRODUCTION

DNA methylation as an epigenetic signal is found in both eukaryotes and prokaryotes. In bacteria, a methyl group from S-adenosyl-L-methionine (AdoMet) is transferred to cytosine or adenine, resulting in N\textsuperscript{4}-methyl-cytosine, C\textsuperscript{5}-methyl-cytosine, or N\textsuperscript{6}-methyl-adenine on DNA (Sánchez-Romero and Casadesús, 2020). Because methyl groups located in the major groove of double helix alter protein-DNA interactions and DNA curvature, DNA methylation may protect DNA from
cleavage by restriction enzymes and regulate gene expression, cell cycle, chromosome replication, phase variation, virulence, and mismatch repair (Sánchez-Romero and Casadesús, 2020). DNA methylation is catalyzed by DNA methyltransferases (MTases), which include those of restriction-modification (RM) systems and orphan methylylases (such as Dam and Dcm in Escherichia coli; Blow et al., 2016).

Cyanobacteria are a group of prokaryotes that perform ogenic photosynthesis (Castenholz, 2001). Restriction-methylation systems have been predicted in genomes of most cyanobacteria. In addition, some orphan methylylases were identified. In Anabaena sp. PCC 7120, a heterocyst-forming filamentous cyanobacterium, there are four predicted orphan MTases, denoted as Dmt A to D, in addition to the MTases of RM systems (Matveyev et al., 2001). These orphan MTases modify DNA at GATC (DmtA), GGCC (DmtB), CGATCG (DmtC), or RCCGGY (DmtD). In Synechocystis sp. PCC 6803 (hereafter Synechocystis 6803), a unicellular cyanobacterium, there are at least five different methylation activities toward specific DNA sequences: M.Ssp6803I for m3CGATCG (Scharnagl et al., 1998), M.Ssp6803II for Gm4CC, M.Ssp6803III for Gm6ATC, M.Ssp6803IV for GA4AGGC, and M.Ssp6803V for the bipartite motif GGm6AN7TTGG/CCAm6AN7TCC (Hagemann et al., 2018). DNA methylation may play roles in cellular and physiological functions. For an example, the GGm4CC methylation is involved in regulation of gene expression, fine-tuning of DNA replication and DNA repair in Synechocystis sp. PCC 6803 (Gärtner et al., 2019).

In cyanobacteria, glycogen is the most important storage polysaccharide. It is an α-1,4 linked α-1,6 branched glucose polymer. The synthesis of glycogen depends on ADP glucose pyrophosphorylase, glycogen synthase, and a branching enzyme (Preiss, 1984), whereas the degradation of glycogen depends on glycogen phosphorylase (GlgP), the debranching enzyme, α-1,4-glucanotransferase, and maltodextrin phosphorylase (Ball and Morell, 2003). GlgP is the key enzyme involved in glycogen digestion. In Synechocystis 6803, there are two GlgP genes, glgP1 (sll1356), and glgP2 (slr1367), which are functionally diverged (Fu and Xu, 2006) and differently regulated (Osnai et al., 2014). Under autotrophic growth conditions, glgP1 plays a more important role in glycogen digestion (Fu and Xu, 2006). When we analyzed the previously reported DNA methylase M.Ssp6803V (slr6095), we realized that it is probably not an orphan methylase as proposed, because slr6095 (N-terminal portion of M subunit, for methylation), slr6096 (C-terminal portion of M subunit), slr6097 (S subunit, for recognition specificity), and slr6102 (R subunit, for restriction) are predicted to encode a type I RM system. In this type of RM system, M and S subunits are required for the restriction activity as well (Murray, 2000), therefore their encoding genes can be inactivated in the presence of a functional R subunit gene. We generated mutants for slr6095, slr6096, and slr6097, and analyzed differences in genome-wide gene expression and DNA methylation between mutants and the wild type strain. Inactivation of these genes consistently abolished DNA methylation at GGm6AN7TTGG/CCAm6AN7TCC motifs in the genome and indirectly altered the expression of many genes and the glycogen catabolism in Synechocystis 6803.

MATERIALS AND METHODS

Strains and Culture Conditions

Synechocystis sp. PCC 6803 and derivatives were cultured in BG11 in flasks on a rotary shaker at 30°C under continuous illumination of 30 µE m−2 s−1. Kanamycin (25 µg mL−1) was added as needed.

DNA Preparation

Genomic DNA was extracted using the cetyltrimethylammonium bromide (CTAB) method (Murray and Thompson, 1980) with modifications. The cyanobacterial cells were broken by grinding in liquid nitrogen and suspended in CTAB solution. The broken cell suspension was treated with proteinase K (100 µg mL−1) and β-mercaptoethanol (2%, v/v) at 65°C for 1 h and centrifuged at 17,226 g for 30 min. The supernatant was extracted with phenol:chloroform:isoamyl alcohol (25:24:1), followed by chloroform:isoamyl alcohol (24:1). Nucleic acids were precipitated with isopropanol, washed with 70% ethanol, dissolved in TE buffer. After removal of RNA with RNase A (100 µg mL−1), DNA was extracted with phenol and chloroform, precipitated with ethanol and dissolved in TE buffer. The quality of DNA was assessed by agarose-gel electrophoresis and OD_{260}/OD_{280} (1.8~2.0).

RNA Preparation

Total RNA was extracted using Trizol reagent (Invitrogen, Carlsbad, CA, United States) according to the manufacturer’s instructions. After removal of DNA with DNase RQ1 (Promega, Madison, WI, United States), the sample was extracted with Trizol reagent. RNA was precipitated with isopropanol, washed with 70% ethanol, followed by chloroform:isoamyl alcohol (24:1). Nucleic acids were precipitated with isopropanol, washed with 70% ethanol, dissolved in nuclease-free ddH2O. RNA integrity was verified on Bioanalyzer (Agilent, Santa Clara, CA, United States) using RNA 6000 nano chip, and the RNA integrity numbers (RIN) were shown to be >6.5.

Construction of Plasmids and Generation of Mutants

Molecular manipulations were performed according to manufacturers’ instructions. Clones of PCR products were confirmed by sequencing. Construction of plasmids and primers used are described in Supplementary Table S1 in the Supplementary Material.

Using overlap PCR, an XhoI site was introduced into slr6095, slr6096, and slr6097, and the DNA fragment containing one of these genes, interrupted by the XhoI site, was cloned into pMD18-T, then a kanamycin-resistance (Km) cassette was inserted into the XhoI site, producing plasmids pHB6515, pHB6520, and pHB6517. These plasmids were used to inactivate slr6095, slr6096, and slr6097 in Synechocystis 6803.
Transformation of *Synechocystis* 6803 was performed as described by Williams (1988). Briefly, 100 µl of cell suspension was mixed with DNA (10 µg ml⁻¹) and incubated for 4 to 6 h at 30°C in the light of 30 µE m⁻² s⁻¹, spread onto membrane filters resting on BG11 agar plates and allowed to grow for 20 h. The filters were then transferred to plates supplemented with glucose (5 mM) and kanamycin (25 µg ml⁻¹). Transformants usually appeared within 4 days. After streaking on plates, the transformants were cultured in liquid BG11 with kanamycin. The complete segregation of mutants was confirmed by PCR. *Synechocystis* strains and primers used are described in Supplementary Table S1 in the Supplementary Material.

**SMRT (Single Molecule Real Time) Sequencing and DNA Methylation Analysis**

DNA methylation was analyzed by whole-genome SMRT sequencing. Briefly, libraries with 10 kb inserts were prepared according to the PacBio Sequel Microbiological sequencing protocol, and the genome DNA was sequenced with a PacBio Sequel platform. DNA methylation was then analyzed with the SMRT raw data using the SMRT LINK 6.0.0 software. Base modifications and motifs were analyzed using P_modificationDetection and P_motifFinder module from the SMRT Analysis software. Because SMRT sequencing is not suitable for m⁵C methylation (Murray et al., 2012), we collected the data for m⁶A and m⁴C only (consistent between two biological repeats). Raw sequence read data has been deposited in the NCBI Sequence Read Archive (SRA) with the study identifier SRP256040.

**RNA Sequencing and Differential Expression Analysis**

*Synechocystis* 6803 and mutants were cultured in BG11 to an OD₇₃₀ of 1.0 at 30°C in the light of 30 µE m⁻² s⁻¹. Total RNA was extracted using TRIzol reagent (Invitrogen, Carlsbad, CA, United States). DNA was removed with RNase-free DNase I (Promega, Madison, WI, United States). RNA was precipitated with isopropanol and stored at 70% ethanol. Strand-specific RNA-seq libraries were prepared using the Illumina small RNA sample preparation kit. RNA sequencing was carried out using the HiSeq 2500 sequencing instrument (Illumina, San Diego, CA, United States) to generate paired-end reads with a length of 150 bp. Reads were aligned against the Cyanobase genome sequence for *Synechocystis* 6803 using TopHat (Trapnell et al., 2009). Differential gene expression analysis was performed by using the DESeq package (Anders and Huber, 2010). Fold changes of >2 or <0.5 with P-value of <0.01 were defined as differentially expressed. Data are means ± SD produced from 3 biological replicates. Raw sequence read data has been deposited in the NCBI SRA with the study identifier SRP258660.

**Reverse-Transcriptase Quantitative Polymerase Chain Reaction (RT-qPCR)**

Reverse transcription and the RT-qPCR analysis were performed according to Gao and Xu (2012). *rnpB* (RNase P subunit B; Vioque, 1992) was used as the internal control. PCR primers (indicated with “RT” in name) are listed in Supplementary Table S1. Data are means ± SD produced from 3 technical repeats.

**Determination of Glycogen Contents and Glycogen Phosphorylase Activity**

*Synechocystis* 6803 and mutants were cultured in BG11 to an OD₇₃₀ of 1.0. Glycogen contents and GlgP activities were determined according to previously described (Fu and Xu, 2006). Data are means ± SD produced from 3 biological replicates.

**Prediction of Gene Functions**

The type I RM system was predicted according to the REBASE⁴ and the Cyanobase⁵. Structural domains were predicted using SMART analysis⁶. Similarity searches were performed using Blast⁷. Multiple sequence alignment was performed using the ClustalW².

**RESULTS**

**Gene Cluster Analysis and Mutant Construction**

*slr6095* is located in a gene cluster on the plasmid pSYSX (Figure 1). According to the REBASE and the Cyanobase, *slr6095* and *slr6096* are predicted to encode two portions of the M subunit of a type I RM system, *slr6097* encodes the S subunit, *slr6102* the R subunit. Alignment with N6-Mtases from *Rhizobiales bacterium* and *Thiorhodovibrio* sp. 970 shows that Slr6095 and Slr6096 are similar to the N-terminal and C-terminal portions of the M subunit (for methylation), respectively (Supplementary Figure S1). Slr6097 is predicted to contain a methylase-S domain and shows high similarity to the S subunit HsdS (for sequence recognition, GenBank accession no. AFWS0200001.1) of a predicted type I RM system in *Thiorhodovibrio* sp. 970 (69% similarity; e-value, 4e-108). In the same gene cluster, Slr6012 contains an HSDR_N domain and is similar to the R subunit HsdR (for restriction) in *Thiorhodovibrio* sp. 970 (59% similarity; e-value, 0.0). We tentatively call these proteins M. *Ssp6803V*, M. *Ssp6803V-c* (*n* and *c*, N-terminal, or C-terminal portion), S. *Ssp6803V*, and R. *Ssp6803V*.

To investigate the functions of *slr6095*, *slr6096*, and *slr6097*, we generated mutant strains for each of them (Figure 2A). A Km² cassette was inserted into the target genes via

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1. https://www.pacb.com/support/software-downloads/
2. http://genome.microbedb.jp/cyanobase
3. http://rebase.neb.com/rebase/rebase.html
4. http://genome.microbedb.jp/cyanobase
5. http://smart.embl-heidelberg.de
6. https://blast.ncbi.nlm.nih.gov/Blast.cgi
7. https://www.genome.jp/tools-bin/clustalw
FIGURE 1 | Genes encoding the type I restriction-modification system on the plasmid pSYSX in Synechocystis 6803.

FIGURE 2 | Inactivation of genes in Synechocystis 6803. (A) Schematic diagram showing insertion of target genes with the Km\textsuperscript{r} cassette. (B) Agarose electrophoresis of PCR products for examination of mutants.

homologous double crossover, and the complete segregation of mutants was confirmed with PCR (Figure 2B). Because M and S subunits of a type I RM system are involved in the restriction activity, the complete segregation of the three mutants does not imply that the predicted R subunit (Slr6102) is not functional.

Methylome and Transcriptome Analyses
We examined the whole-genome methylation landscape using SMRT sequencing technology. This approach is modestly sensitive to m\textsuperscript{5}C methylation but detects m\textsuperscript{6}A and m\textsuperscript{4}C with high accuracy and sensitivity. As shown in Table 1, in the wild-type genome there are methylations at motifs G\textsuperscript{m6}ATC, G\textsuperscript{m4}CC, G\textsuperscript{m6}AGGC, and G\textsuperscript{m6}AN\textsuperscript{7}TTGG/CCA\textsuperscript{m6}AN\textsuperscript{7}TCC. Unlike the wild type, mutant strains slr6095::Km\textsuperscript{r}, slr6096::Km\textsuperscript{r}, and slr6097::Km\textsuperscript{r} showed no bipartite methylation at G\textsuperscript{m6}AN\textsuperscript{7}TTGG/CCA\textsuperscript{m6}AN\textsuperscript{7}TCC (hereafter Ssp6803V methylation). This is consistent with the predicted functions of the three genes. We also noticed that the G\textsuperscript{m4}CC methylation decreased in slr6096::Km\textsuperscript{r} and slr6097::Km\textsuperscript{r} in comparison to the wild type level. This could be due to an indirect effect of the two genes on the efficiency of G\textsuperscript{m4}CC methylation.

DNA methylation may alter gene expression. We performed RNA-seq analyses to compare gene expression in the wild type and mutants (Supplementary Tables S3–S5). Relative to gene expression in the wild type, 143 genes were up-regulated and 160 genes down-regulated in slr6095::Km\textsuperscript{r}, 137 genes up-regulated, and 172 genes down-regulated in slr6096::Km\textsuperscript{r}, 109 genes up-regulated, and 150 genes down-regulated in slr6097::Km\textsuperscript{r}. There are 73 genes up-regulated and 98 genes down-regulated in all three mutants (Table 2). These 171 genes (73 + 98) are involved in photosynthesis, ATP synthesis, pigment syntheses, carbon metabolism, nutrient transport, stress tolerance, signal transduction, and gene regulation, etc.

To understand the effects of DNA methylation on gene expression in Synechocystis 6803, we attempted to find out the relationship between DNA methylation and gene expression. The Ssp6803V methylation sites are distributed in the encoding
sequences of 869 genes (23.74% of all in the genome) and the sequences (1 kb) upstream of 270 genes (7.38% of all). Of the 171 genes with changed expression, 75 have no recognition sequence for the Ssp6803V methylation, 40 have the motifs only in their encoding regions (Table 2 and Supplementary Table S2). It is not possible for these genes to be directly regulated by the DNA methylation. However, there are some genes, such as slr1286 (transcriptional regulator), slr1883 (argJ), slr1926 (hypothetical protein), and slr0093 (dnaJ), whose Ssp6803V methylation sites may overlap regulatory sequences (see transcription start sites and methylation positions in Table 2). The expression of these genes could be modulated by methylation due to altered interactions between regulatory proteins and recognition sites.

**Effect of DNA Methylation on Glycogen Catabolism**

Among genes that showed changed expression in the mutants relative to the wild type, slr0653 encodes the principal RNA polymerase sigma factor (SigA), slr1356 (glgP1) encodes GlgP. Both were upregulated in the mutants. RT-qPCR analysis confirmed the increased expression of sigA and glgP1 in three mutants; glgP2, however, showed no or slight change in mRNA level (Figure 3).

Because GlgP1 plays an important role in glycogen turnover (Fu and Xu, 2006), we further measured the GlgP activities and total glycogen contents in these strains under photoautotrophic conditions. As shown in Figure 4A, all three mutants showed higher GlgP activities than the wild type. Higher GlgP activity implies more rapid digestion of glycogen. Consistently, the mutants showed lower glycogen contents than the wild type (Figure 4B).

Some genes involved in photosynthesis (phycobilisome, photosystem I, photosystem II, and CO2 concentrating mechanism), electron transfer (petC2, petF), or ATP synthesis were downregulated in the mutants (Table 2), but petE and rbcX (Rubisco chaperonin) were significantly upregulated. Some heat shock protein/molecular chaperone genes were also downregulated in the mutants (Table 2). The mutants showed no or very slight change in growth at 30°C but reduced growth at 40°C compared to the wild type (Supplementary Figure S2 and data no shown).

**DISCUSSION**

In Synechocystis 6803, the role of GGm4CC methylation in regulation of gene expression has been reported (Gärtner et al., 2019). Now we show that the GGm4AN7TTGG/CAM6AN7TCC methylation, catalyzed by the type I RM system Ssp6803V, is also involved in regulation/modulation of gene expression.

Genes slr6095-slr6096-slr6097 and slr6102 on the plasmid pSYSX are predicted to encode a type I RM system, but the ancestor M subunit gene split into slr6095 and slr6096 in the evolution. Unlike a type II RM system, in which the restriction activity is only dependent on the restriction enzyme, cleavage of the target DNA by a type I RM system is dependent on S, R, and M subunits (Murray, 2000). Therefore, successful inactivation of genes for M and S subunits does not imply that the R subunit gene is not functional. In the RNA-seq data, the mRNA abundance of slr6102 is similar to slr6095-slr6096-slr6097. It is possible that this type I RM system is fully functional.

In Synechocystis 6803 mutants slr6095::Km1, slr6096::Km1, and slr6097::Km1, the Ssp6803V methylation was completely...
**TABLE 2** | Ssp6803V methylation and differential expression*.

| Gene     | Annotation                                      | TSS site** | Methylation position | Ratio of gene expression |
|----------|-------------------------------------------------|------------|-----------------------|--------------------------|
| slr0170  | DnaK protein 2, heat shock protein 70           | −20        | +6                    | 0.4151 0.0026 0.2577 0.0000 0.3237 0.0007 |
| slr0199  | Plastocyanin                                    | −98        | None                  | 37.0240 0.0000 44.7207 0.0000 48.7945 0.0000 |
| slr0254  | Phytene dehydrogenase Rieske iron-sulfur component | −49        | +20, +918, +1253     | 0.1844 0.0005 0.1729 0.0007 0.1610 0.0003 |
| slr0416  | GroEL-2                                         | Not ident. | −298                  | 0.2612 0.0003 0.1348 0.0000 0.1730 0.0000 |
| slr1031  | CO2 concentrating mechanism protein CcmM        | Not ident. | +1191                 | 0.3955 0.0031 0.3169 0.0009 0.3209 0.0027 |
| slr1281  | Photosystem II protein PsbZ                     | −282       | None                  | 0.2460 0.0037 0.2027 0.0007 0.1737 0.0004 |
| slr1286  | Transcriptional regulator                       | −27        | −257                  | 3.7930 0.0002 4.3482 0.0002 4.1379 0.0004 |
| slr1325  | ATP synthase delta chain of CF1                 | Not ident. | None                  | 0.2789 0.0005 0.3369 0.0051 0.2163 0.0004 |
| slr1327  | ATP synthase gamma chain                        | Not ident. | −642                  | 0.3708 0.0043 0.3381 0.0031 0.3022 0.0045 |
| slr1356  | Glycogen phosphorylase                          | −14        | +2153, +2285          | 2.8489 0.0010 3.2206 0.0007 3.6499 0.0005 |
| slr1382  | Ferredoxin, PetF-like protein                    | −10        | None                  | 0.4459 0.0059 0.3637 0.0038 0.3825 0.0042 |
| slr1514  | 16.6 kDa small heat shock protein               | −41        | None                  | 0.0890 0.0000 0.0800 0.0000 0.0491 0.0000 |
| slr1579  | Phycobilisome rod linker polypeptide            | Not ident. | None                  | 0.2747 0.0003 0.1525 0.0000 0.1419 0.0000 |
| slr1580  | Phycobilisome rod linker polypeptide            | Not ident. | None                  | 0.3043 0.0004 0.1843 0.0000 0.1535 0.0000 |
| slr1883  | Arginine biosynthesis bifunctional protein ArgJ | −28        | −223, −716            | 0.2984 0.0001 0.3149 0.0013 0.2580 0.0002 |
| slr1926  | Hypothetical protein                            | −81        | −660, −336, −227      | 14.0133 0.0000 15.4892 0.0002 12.8917 0.0001 |
| slr0011  | Rubisco chaperonoin                              | Not ident. | None                  | 4.2534 0.0000 3.4429 0.0049 3.9258 0.0000 |
| slr0056  | Chlorophyll a synthase                          | Not ident. | +724                  | 0.2575 0.0001 0.2858 0.0007 0.3234 0.0028 |
| slr0093  | DnaJ protein, heat shock protein 40             | −108       | −193                  | 0.1370 0.0000 0.1181 0.0000 0.1099 0.0000 |
| slr0653  | Principal RNA polymerase sigma factor SigA      | −223       | +29, +845             | 4.7465 0.0000 4.5298 0.0000 4.2149 0.0000 |
| slr0895  | Transcriptional regulator                       | Not ident. | None                  | 0.1921 0.0004 0.1458 0.0006 0.1657 0.0001 |
| slr1185  | Cytochrome b6-type complex alternative iron-sulfur subunit | Not ident. | None                  | 0.3490 0.0053 0.2335 0.0010 0.2333 0.0048 |
| slr1265  | Two-component sensor histidine kinase           | Not ident. | None                  | 0.3257 0.0001 0.2269 0.0001 0.1969 0.0000 |
| slr1291  | NADH dehydrogenase subunit 4                    | Not ident. | −337, +428            | 0.2032 0.0000 0.1512 0.0000 0.1420 0.0000 |
| slr1330  | ATP synthase epsilon chain of CF1               | Not ident. | −680                  | 0.2507 0.0000 0.3063 0.0025 0.2177 0.0005 |
| slr1655  | Photosystem I subunit XI                        | Not ident. | None                  | 0.1807 0.0002 0.1193 0.0000 0.1053 0.0000 |
| ssl2598  | Photosystem II protein PsbH                     | −37        | None                  | 0.1785 0.0001 0.1469 0.0000 0.1617 0.0002 |

* This table includes some of the differentially expressed genes, whereas Supplementary Table S2 includes a full list of the genes. **not ident., not identified.
eliminated. Downstream of slr6097 is sll6098, which is oriented 3' to 3' relative to slr6097 (Figure 1). Therefore, the phenotype of slr6097::Km could not be attributed to a polar effect. Insertion of slr6095 and slr6096 may have polar effects on the downstream gene(s), but these two and slr6097 are very closely related in function, their phenotypes are supposed to be similar to each other. The very similar phenotypes of the three mutants excluded the possibility that a second mutation might have caused the elimination of DNA methylation and physiological effects, if any.

To analyze the effects of Ssp6803V methylation on gene expression in Synechocystis 6803, we compared the mRNA expression profiles of the wild type and mutants and identified 171 genes that showed up- or down-regulated expression in all three mutants. Most of these genes are apparently indirectly regulated by the Ssp6803V methylation (Table 2). glgP1 is one of those upregulated genes (Table 2 and Figure 3), suggesting that the type I RM system may tune glycogen catabolism via gene regulation. Differences in GlgP activities and glycogen contents between the wild type and mutants (Figure 4) confirmed such a physiological effect of the Ssp6803 methylation. Apparently, the DNA methylation reduces the expression of GlgP1 for glycogen turnover and enhances the storage of glycogen in Synechocystis 6803. The mutants also showed reduced growth at 40°C. This phenotype may reflect an integrated effect of changed gene expression (such as expression of genes for photosynthesis, stress responses, etc.) on the high temperature tolerance. In addition, the changed expression of heat shock protein genes (Table 2) could affect salt and oxidative stress tolerance (Asadulghani et al., 2004; Sakthivel et al., 2009; Kim et al., 2014) as well.

The impact of type I RM systems on gene expression has been reported in other bacteria (De Ste Croix et al., 2017). For example, in Streptococcus pneumoniae, a human pathogen, the virulence was shown to be regulated by a type I RM system (SpnD39III) with phase variable recognition specificities (Manso et al., 2014). SpnD39III variants showed distinct genome methylation patterns and gene expression profiles. However, there are no SpnD39III sites in promoters or regulatory regions of those differentially expressed genes. Similarly, no Ssp6803V sites can be found upstream of most genes differentially expressed in the wild type and methylation mutants. Possibly, addition of methyl groups to target sequences can alter protein-DNA interactions that are involved in maintaining regional DNA conformation, therefore indirectly modulates the expression of glgP1 and many other genes. The underlying mechanism for type I RM systems to impact gene expression in bacteria should be a further question to be tackled in the future.
DATA AVAILABILITY STATEMENT

Raw sequence read data for DNA methylation analysis has been deposited in the NCBI SRA with the study identifier SRP256040 (runs: SRR11528407-SRR11528414). Raw sequence read data for RNA-seq has been deposited in the NCBI SRA with the study identifier SRP258660 (runs: SRR11614230-SRR11614241).

AUTHOR CONTRIBUTIONS

XX designed and supervised the project. DW performed the experiments and bioinformatic analyses. YW performed bioinformatic analyses. DW, and YW analyzed the data. DW and XX wrote the manuscript. XX edited the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.01258/full#supplementary-material

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