Salt-Tolerant *Synechococcus elongatus* UTEX 2973 Obtained via Engineering of Heterologous Synthesis of Compatible Solute Glucosylglycerol

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The recently isolated cyanobacterium *Synechococcus elongatus* UTEX 2973 (Syn2973) is characterized by a faster growth rate and greater tolerance to high temperature and high light, making it a good candidate chassis for autotrophic photosynthetic microbial cell factories. However, Syn2973 is sensitive to salt stress, making it urgently important to improve the salt tolerance of Syn2973 for future biotechnological applications. Glucosylglycerol, a compatible solute, plays an important role in resisting salt stress in moderate and marine halotolerant cyanobacteria. In this study, the salt tolerance of Syn2973 was successfully improved by introducing the glucosylglycerol (GG) biosynthetic pathway (OD750 improved by 24% at 60 h). In addition, the salt tolerance of Syn2973 was further enhanced by overexpressing the rate-limiting step of glycerol-3-phosphate dehydrogenase and downregulating the gene rfbA, which encodes UDP glucose pyrophosphorylase. Taken together, these results indicate that the growth of the end-point strain M-2522-GgpPS-drfbA was improved by 62% compared with the control strain M-pSI-pSII at 60 h under treatment with 0.5 M NaCl. Finally, a comparative metabolomic analysis between strains M-pSI-pSII and M-2522-GgpPS-drfbA was performed to characterize the carbon flux in the engineered M-2522-GgpPS-drfbA strain, and the results showed that more carbon flux was redirected from ADP-GLC to GG synthesis. This study provides important engineering strategies to improve salt tolerance and GG production in Syn2973 in the future.

**Keywords:** cyanobacteria, GG, metabolomic analysis, salt tolerance, small RNA regulation

**INTRODUCTION**

Cyanobacteria have attracted significant attention recently for the production of green fuels and chemicals due to their ability to absorb sunlight and CO2 as sole energy and carbon sources, respectively (Gao et al., 2012, 2016a,b; Luan and Lu, 2018; Matson and Atsumi, 2018). A recently isolated cyanobacterium, *Synechococcus elongatus* UTEX 2973 (hereafter Syn2973), is characterized...
by a faster growth rate and more tolerance to high temperature and high light (Mueller et al., 2017; Ungerer et al., 2018b). The doubling time of Syn2973 can reach 1.9 h in BG11 medium at 41°C under continuous 500 µmol photons m⁻² s⁻¹ white light with 3% CO₂ (Yu et al., 2015), which is very similar to that of the heterotrophic yeast Saccharomyces cerevisiae (1.67 h) (Herskowitz, 1988). However, as a freshwater cyanobacterium, Syn2973 is very sensitive to salt stress, and its growth was significantly inhibited by high concentrations of salt (Song et al., 2016; Cui et al., 2020b). Meanwhile, large-scale cultivation of cyanobacteria with seawater is necessary for industrial biotechnological applications (Silkina et al., 2019; Cui et al., 2020a). It is therefore urgent to improve the salt tolerance of Syn2973 for the future application of this promising chassis for green fuels and chemical production.

Acclimation to high external salinities involves the accumulation of compatible solutes and the active export of ions (Hagemann, 2011). Compatible solutes are a diverse class of low-molecular-weight compounds that accumulate at high intracellular concentrations in response to hyperosmotic or salt stress (Da Costa et al., 1998). Glucosylglycerol (GG) is a potent compatible solute consisting of glucose and glycerol linked by a glycosidic bond (Tan et al., 2016) and has a wide range of beneficial functions as a health food material, therapeutic agent, cosmetic ingredient, and enzyme stabilizer (Sawangwan et al., 2010; Luley-Goedl and Nidetzky, 2011). The marine cyanobacterium Synechococcus sp. PCC 7002 (hereafter Syn7002) and moderate halotolerant cyanobacterium Synechocystis sp. PCC 6803 (hereafter Syn6803) could accumulate GG to resist salt stress (Hagemann, 2011). The synthesis of GG has been elucidated in Syn6803, which depends strictly on ADP-glucose and consists of two steps involved in two enzymes [GG-phosphate-synthase (GGPS) and GG-phosphate-phosphatase (GGPP)] (Hagemann and Erdmann, 1994). In addition, the regulatory mechanisms involved in the synthesis of GG have mainly been investigated in Synechocystis (Kirsch et al., 2019). However, GG was not synthesized natively in Syn2973; instead, sucrose was synthesized as the only compatible solute under salt stress (Song et al., 2016). Heterologous synthesis of compatible solutes in salt-sensitive strains has been considered a key strategy for improving the salt tolerance of hosts (Waditee-Sirisattha et al., 2012; Singh et al., 2013). For example, the introduction of glycine methylation genes (ApGSMT-DMT) associated with glycine betaine synthesis successfully improved the salt tolerance of Anabaena sp. PCC 7120, and the growth of Anabaena sp. The PCC 7120-expressing ApGSMT-DMT gene cluster was improved by ~70% under conditions with 0.12 M NaCl (Waditee-Sirisattha et al., 2012). It is thus interesting to explore the effect of heterologous GG synthesis on the salt tolerance of Syn2973. Notably, Syn2973 is known to undergo higher carbon flux into the sugar phosphate pathway and accumulates greater amounts of glycogen and sucrose (Abernathy et al., 2017; Ungerer et al., 2018a), which might also be beneficial for the heterologous synthesis of GG to improve salt tolerance.

In this study, multiple efforts were made to improve the salt tolerance of Syn2973 by achieving highly efficient biosynthesis of GG (Figure 1). Specifically, GG biosynthesis was first established in Syn2973 upon introducing genes encoding GgpS and GgpP. Second, the rate-limiting step in the process of converting DHAP to G3P was identified and strengthened by overexpressing genes from different sources. Third, redirection of carbon flux toward GG production by a small RNA (sRNA) regulatory tool was performed. Finally, the salt tolerance of different mutants was tested under different culture conditions. Together, these integrated efforts improved the salt tolerance of Syn2973 by 62% and represent the first attempt to increase salt stress tolerance by de novo biosynthesis of GG in Syn2973.

MATERIALS AND METHODS

Bacterial Growth Conditions

The wild-type Syn2973 and engineered strains were grown on BG11 medium (pH 7.5) under a light intensity of approximately 200 µmol photons m⁻² s⁻¹ or 500 µmol photons m⁻² s⁻¹ in a shaking incubator at 130 rpm at 37°C (HNYS-202T, Honour, Tianjin, China) (Li et al., 2018). CO₂ from the air was used as the only carbon source. Appropriate antibiotics, 20 µg/ml spectinomycin or chloramycin (Solarbio, Beijing, China), were added to maintain the stability of the Syn2973 engineered strains. Inoculum and 0.1 M NaCl BG11 medium were mixed with 1 M NaCl BG11 medium and adjusted to a certain NaCl concentration. Cell optical density was measured by a spectrophotometer (UV-1750, Shimadzu, Kyoto, Japan) at 750 nm. Escherichia coli DH5α was grown on LB agar plates or in LB liquid medium in an incubator at 37°C or shaking incubator at 200 rpm supplemented with 50 µg/ml spectinomycin or 200 µg/ml ampicillin (Solarbio, Beijing, China).

For GG production, prior to production experiments, mutants were inoculated in BG11 medium containing appropriate antibiotics and grown photoautotrophically. For the production experiments, mutant cultures were incubated in fresh BG11 medium without antibiotics under different NaCl concentrations at the initial OD₇₅₀ of 0.1 and cultivated photoautotrophically.

Construction of Strains and Plasmids

The strains and plasmids used in this study are listed in Table 1. Among them, E. coli DH5α was used for vector construction and amplification. For gene overexpression, the integrative vector pSI with a spectinomycin-resistant cassette and pSII with a chloramycin-resistant cassette were used (Li et al., 2018). Primers for gene amplification are listed

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**Abbreviations:** AcCoA, acetyl-CoA; AGK, α-ketoglutarate; ADP-GLC, adenosine diphosphoglucose; ATP, adenosine triphosphate; CIT, citrate; D, day; DHAP, dihydroxyacetone phosphate; E4P, erythrose-4-phosphate; F6P, fructose-6-phosphate; FBP, fructose-1,6-diphosphate; G6P, glucose-6-phosphate; G1P, glucose-1-phosphate; G3P, glyceraldehyde-3-phosphate; GPP, glyceraldehyde-3-phosphate; PEP, phosphoenolpyruvate; Ru5P, ribulose 5-phosphate; RUBP, ribulose 1,5-bisphosphate; SRP, ribose 5-phosphate; S7P, sedoheptulose-7-phosphate; SUC, succinate; Suc-6-P, sucrose-6-phosphate; UDP-GLC, uridine diphosphate glucose; UTP, uridine-5′(1-thiotriphosphate); X3P, xylose-5-phosphate.
Briefly, E. coli HB101 harboring pRL443 and pRL623 and E. coli DH5α harboring the target plasmid were cultured overnight and then transferred into fresh LB medium with 50 μg/ml spectinomycin or 200 μg/ml ampicillin (Solarbio, Beijing, China) at a 1:50 ratio. When cells grew to exponential phase (OD₆₀₀ ≈ 0.5), 2 ml of each E. coli strain was washed twice with fresh (LB) medium to remove all antibiotics, resuspended in 0.1 ml of LB medium, then phosphorylated by T4 Polynucleotide Kinase (Thermo Fisher Scientific Inc., CA, United States). Then, the gene products were ligated into pSI/pSII (linearized by PCR) by blunt end connection. All constructs were validated by colony PCR and confirmed by Sanger sequencing. The constructed plasmid was finally transferred into Syn2973 according to a method reported previously (Li et al., 2018). In this study, the gene products were ligated into pSI/pSII (linearized by PCR) by blunt end connection. All constructs were validated by colony PCR and confirmed by Sanger sequencing. The constructed plasmid was finally transferred into Syn2973 according to a method reported previously (Li et al., 2018).

All primers were synthesized by GENEWIZ Inc. (Suzhou, China). The target genes were amplified by Phanta Super-Fidelity DNA Polymerase (Vazyme Biotech, Nanjing, China), purified by a Cycle Pure Kit (Omega Bio-tek, Nanjing, China), and then phosphorylated by T4 Polynucleotide Kinase (Thermo Fisher Scientific Inc., CA, United States). Then, the gene products were ligated into pSI/pSII (linearized by PCR) by blunt end connection. All constructs were validated by colony PCR and confirmed by Sanger sequencing. The constructed plasmid was finally transferred into Syn2973 according to a method reported previously (Li et al., 2018). In this study, the gene products were ligated into pSI/pSII (linearized by PCR) by blunt end connection. All constructs were validated by colony PCR and confirmed by Sanger sequencing. The constructed plasmid was finally transferred into Syn2973 according to a method reported previously (Li et al., 2018).
were validated by colony PCR (Supplementary Figure 1) and Sanger sequencing.

**GG and Sucrose Extraction and Measurement**

Based on the methods of a previous study (with minor modifications) (Tan et al., 2015), samples (1 OD) of mid-exponential cultures under salt stress were centrifuged at 12,000 rpm for 5 min. Harvested cells were suspended in the same volume (0.8 ml) of 80% ethanol, incubated at 65°C for 4 h, and centrifuged at 12,000 rpm for 5 min; 0.2 ml of supernatant solution was taken for analysis, and internal standard 13C-sorbitol with a final certain concentration of 62.5 mg/L was added. The solvents were removed using a vacuum concentration system (ZLS-1, Her-exi, Hunan, China). For GC-MS analysis, each sample was further derivatized in a two-step process as previously published (Cui et al., 2016). The derivatized samples were analyzed by GCMS using an Agilent 5975 MSD/7890 instrument (Agilent Corp., Santa Clara, CA, United States). The column was a HP-5MS (Restek, Bellefonte, PA, United States). The oven temperature was initially held at 80°C for 2 min and reached 280°C at 10°C per min, then held at 280°C for 5 min. The GG production per OD was calculated by internal standard method. The dry cell weight corresponded to OD<sub>750</sub> nm by the regression equation \( y = 0.9x + 4E-5 \) (\( r^2 = 0.9998, p < 0.05 \)), where \( y \) is the dry cell weight (mg) and \( x \) is the absorbance of the cell suspension at 750 nm.

**LC-MS-Based Targeted Metabolomics Analysis**

Samples (8 ml) of mid-exponential cultures (48 h) at an OD<sub>750</sub> of 1 ± 0.1 were rapidly harvested by centrifugation at 8000 × g for 8 min at 25°C (Eppendorf 5430R, Hamburg, Germany). The extraction of metabolites was carried out as previously published with slight modifications (Bennette et al., 2011; Wang et al., 2014).<sup>13</sup>C<sub>3</sub>,<sup>15</sup>N-alanine (Cambridge Isotope Laboratories, Inc., Andover, MA, United States) was added as the internal standard to correct for variation due to sample extraction and injection. Briefly, approximately eight OD<sub>750</sub> cells were added to 900 µl of the solution containing methanol/H<sub>2</sub>O (8:2, v/v) and then frozen-thawed three times. Samples were centrifuged at 15,000 × g for 5 min at 4°C. The supernatant was extracted, and the sediment was subjected to the above extraction process. The supernatants were mixed, and the solvents were removed using a vacuum concentration system (ZLS-1, Her-exi, Hunan, China). For LC-MS analysis, each dried sample was dissolved in 100 µl of purified water.

LC-MS analysis was conducted on an Agilent 1260 series binary HPLC system (Agilent Technologies, Santa Clara, CA, United States) using an XBridge Amide column (150 × 2.1 mm, 3.5 µm; Waters, Milford, MA, United States) coupled to an Agilent 6410 550 triple quadrupole mass analyzer equipped with an electrospray ionization source (ESI). Multiple reaction monitoring (MRM) mode was used for scanning according to Cui et al. (2020b). All of the peaks were integrated by Qualitative Analysis B.06.00 software and Xcalibur (version 2.1) (Niu et al., 2015). Sucrose was derivatized and analyzed on an Agilent 5975B/6890N GC-MS instrument (Agilent Technologies, Santa Clara, CA, United States) as previously described (Cui et al., 2016). The metabolite data were normalized to internal standards and cell numbers. Glycogen extraction and determination were performed according to the method described by Song et al. (2016).

**Quantitative RT-PCR Analysis**

Total RNA extraction of samples was achieved through a miRNeasy Mini Kit (Qiagen, Valencia, CA, United States) following the protocols described by Sun et al. (2017). In addition, DNase treatment after RNA preparation to ensure degradation of copurified gDNA and 500 ng of total RNA were subjected to cDNA synthesis using a ReverTra Aid First Strand cDNA Synthesis Kit following the manufacturer’s protocol (Thermo Fisher Scientific Inc., CA, United States). Then, 1 µl of each dilution was used as a template for the qRT-PCR described previously (Sun et al., 2018). Three technical replicates were performed for each condition. Data analysis was carried out using StepOnePlus analytical software (Applied Biosystems, CA, United States) and the 2<sup>−ΔΔCt</sup> method (Livak and Schmittgen, 2001).

**Statistical Analysis**

In this study, each experiment was performed with three biological replicates. All data were reported as the means ± standard deviations. A statistical t-test model was applied for the comparative analysis, and a p-value of less than 0.05 was considered statistically significant.

**RESULTS**

**Improved Salt Tolerance of Syn2973 by Heterologous Expression of the GG Biosynthetic Pathway**

To improve the salt tolerance of Syn2973, the synthetic pathway of the compatible solute GG was introduced into Syn2973. First, the ggps and ggpp genes were cloned from Syn6803 onto plasmid pSI with a strong P<sub>trc</sub> promoter. Then, the resulting plasmid was introduced into Syn2973, generating strain M-GgpPS. Compared with the control strain M-pSI, which only harbored the empty plasmid, the growth of strain M-GgpPS showed no difference under 0 M NaCl addition but was improved by approximately 24% under 0.4 M NaCl at a cultivation time point of 60 h (Figure 2A). In addition, a time course of GG production of strain M-GgpPS was determined under 0.4 M NaCl (Supplementary Figure 2), and the chromatogram analysis of GG is shown in Supplementary Figure 3. The results showed that GG production reached 51 mg/g under 0.4 M NaCl at a cultivation time point of 60 h. Meanwhile, no GG production was detected under normal conditions without salt stress (Figure 2B). Early studies showed that the GgpS protein is activated and catalyzes the synthesis of α-glucosylglycerol-phosphate in Syn6803 once under certain salt concentrations (Novak et al., 2011). Moreover, a recent study showed that NaCl
stress was required to initiate αGG synthesis in *Corynebacterium glutamicum* cells, suggesting that the mechanism of osmosensing of GGPS by salt-dependent protein–nucleic acid interactions might be conserved even when the coding gene of the enzyme is transferred into a heterologous host (Ronneke et al., 2018). Consistently, our results showed that upon NaCl-induced stress, GgpS was activated in Syn2973, similar to Syn6803 (Tan et al., 2015), leading to the synthesis of GG. Furthermore, the synthesis of GG slightly enhanced the salt tolerance of Syn2973 (Figure 2A) under the tested condition. Next, it is necessary to improve GG synthesis to further enhance the salt tolerance of Syn2973.

### Screening the Rate-Limiting Steps of Precursor Supply for GG Synthesis

Metabolomics, which involves the measurement of various cellular metabolites, is indispensable for understanding the cellular status, rate-limiting steps of metabolic flow, and productivity of metabolites. Metabolomics based on gas chromatography (GC), liquid chromatography (LC), and capillary chromatography with mass spectrometry (MS) have been performed to analyze cyanobacterial cells for the production of ethanol (Yoshikawa et al., 2017), succinic acid (Osanai et al., 2015), PHB (Arisaka et al., 2019), and others. To determine the possible regulatory mechanism relevant to GG biosynthesis in Syn2973, a better understanding of the intracellular metabolism in the GG-producing strain is required. Thus, the metabolites of strains M-pSI and M-GgpPS cultivated under 0.4 M NaCl conditions were extracted and subjected to LC-MS-based targeted metabolomics analysis. The constructed central metabolic pathways in Syn2973 are shown in Figure 3. Notably, the metabolomic results showed that the precursor metabolite of GG synthesis, G3P, was decreased by 26.02% in strain M-GgpPS compared with strain M-pSI, indicating that the synthesis of G3P might be a limiting step and could be further strengthened to improve GG biosynthesis. Interestingly, the metabolites that were involved in sucrose and glycogen metabolism, such as ADP-GLC and UDP-GLC, in strain M-GgpPS were increased by 2.31- and 1.44-fold compared with those of the control strain M-pSI, respectively. To further evaluate the effect of GG production on sucrose and glycogen metabolism, the contents of sucrose and glycogen were also measured, and the results showed that the content of sucrose was significantly decreased by approximately 50%, while no significant change in the glycogen content was observed in strain M-GgpPS. The glycogen synthase (encoded by gene *glnA*) incorporates glucose monomers into the growing 1,4-α-linked glucose polymer. UDP glucose pyrophosphorylase (encoded by gene *rfbA*) catalyzes the production of UDP-GLC from G1P and UTP. In addition, the sucrose phosphate synthase (SPS) and sucrose phosphophatase (SPP) encoded by gene *Syn0808* catalyzes the synthesis of sucrose from UDP-GLC and F6P. Previous studies have shown that this enzyme reaction is the limiting step of sucrose synthesis in cyanobacteria (Du et al., 2013; Lin et al., 2020). To further explore the effects of GG synthesis on sucrose and glycogen metabolism, qRT-PCR was performed to determine the mRNA abundance of three target genes (*glgA*, *rfbA*, and *Syn0808*) of strains M-pSI and M-GgpPS under salt stress. As illustrated in Supplementary Figure 4, the transcriptional levels of *glgA* and *rfbA* were increased by 39 and 62%, respectively, while the transcriptional level of *Syn0808* was decreased by 42% compared with that of strain M-pSI. In addition, the metabolites R5P and E4P associated with the pentose phosphate pathway were decreased by 13.8 and 20.2%, respectively. These results suggested that more carbon flux might be redirected though sucrose and GG metabolism in the GG-producing strain M-GgpPS, while decreased carbon flux was directed to the pentose phosphate pathway. We next sought to enhance GG synthesis in Syn2973 by improving G3P contents and regulating ADP-GLC flux toward GG biosynthesis and away from glycogen and sucrose biosynthesis.

### Enhancing GG Synthesis by Overexpressing the Glycerol-3-Phosphate Dehydrogenase Gene

To further improve GG synthesis, heterologous genes that encoded glyceraldehyde-3-phosphate dehydrogenase (GlpD) were overexpressed in Syn2973 to enhance G3P supply and GG synthesis. In detail, four genes, *sll1085* encoding GlpD (glycerol-3-phosphate dehydrogenase) from Syn6803, *sbr1755* encoding GpsA (NAD⁺-dependent glycerol-3-phosphate dehydrogenase) from Syn6803, Synpcp7942-2522 from Syn7942, and YDL022W (*glpd1*) from *S. cerevisiae*, were cloned into the plasmid...
pSI-ggpPS with a strong Pcep560 promoter and introduced into the wild-type Syn2973 strain. The strain M-Glpd1-GgpPS overexpressing glpd1 could not survive under salt stress (data not shown), indicating that glpd1 from S. cerevisiae might be toxic to Syn2973 under salt stress, although it had a positive effect on the G3P supply for lipid accumulation in Syn6803 (Wang et al., 2016). Then, the total GG production of strains M-1085-GgpPS, M-1175-GgpPS, and M-2522-GgpPS reached 65, 62, and 70 mg/g at a cultivation time point of 60 h, respectively, improving 32, 26, and 42% compared with strain M-GgpPS, respectively (Figure 4A).

**Knockdown of the Competing Pathway Using Hfq-MicC to Improve GG Production**

The glycogen, sucrose, and pentose phosphate pathways are the major carbon competition pathways for GG synthesis (Figure 1). However, knockout of genes in essential pathways might cause severe growth inhibition. An sRNA regulatory tool that introduces an exogenous Hfq chaperone and MicC scaffold with a binding sequence (Hfq-MicC) was developed in Syn6803, achieving the accurate regulation of endogenous and exogenous genes (Sun et al., 2018). Recently, the utilization of the Hfq-MicC tool was also established and tested in Syn2973, suggesting that this tool was effective in Syn2973 and demonstrated the essential role of Hfq (Li et al., 2018). To drive more carbon flux from the competing pathways to GG biosynthesis, the sRNA regulatory tool above was employed, and the M744-00310-encoding gene pgl related to the pentose phosphate pathway, the M744-05875-encoding gene rfbA related to sucrose synthesis, and the M744-03630-encoding gene glgA related to glycogen synthesis were selected as target genes. The strains M-2522-GgpPS-dpgl, M-2522-GgpPS-dglgA, and M-2522-GgpPS-drfbA were constructed, targeting the pgl, rfbA, and glgA genes, respectively (Table 1). qRT-PCR assays were carried out to determine the mRNA abundances of three target genes in the mutant strains. As illustrated in Supplementary Figure 5, the transcriptional levels of pgl, rfbA, and glgA were decreased by 52, 45, and 38%, respectively, compared to those in the control strain M-2522-GgpPS-pSII. In addition, the GG production of strains M-2522-GgpPS-pSII, M-2522-GgpPS-dpgl, M-2522-GgpPS-dglgA, and M-2522-GgpPS-drfbA achieved 68, 66, 64, and 80 mg/g, respectively, at a cultivation time point of 60 h (Figure 4B). Compared with the control strain M-2522-GgpPS-pSII, GG synthesis in strains M-2522-GgpPS-dpgl and M-2522-GgpPS-dglgA was slightly decreased, while GG production in strain M-2522-GgpPS-drfbA was increased by 17%. The GG productivity was up to 1.3 mg/g/h, which represented an increase of 56.8% compared with the initial strain M-GgpPS. Together, these results demonstrated that among these three tested genes, downregulating the gene rfbA produced the best enhancement of GG production.

**Comparative Targeted Metabolomics Analysis of Strains M-2522-GgpPS-drfbA and M-pSI-pSII**

To further explore the mechanisms relevant to high GG biosynthesis, the metabolites of strain M-2522-GgpPS-drfbA and the control strain M-pSI-pSII under 0.4 M NaCl and 200 μmol photons m⁻² s⁻¹ conditions were extracted and subjected to LCMS-based targeted metabolomics analysis. As shown in Figure 5, the G3P content in strain M-2522-GgpPS-drfbA was increased by threefold compared to the control strain M-pSI-pSII, suggesting that it was effective for G3P and GG accumulation by overexpressing the gene glpD. The UDP-GLC in strain M-2522-GgpPS-drfbA was decreased by 24%, while sucrose was further decreased by 60% compared with the control strain M-pSI-pSII, suggesting that more carbon flux might be redirected from sucrose metabolism to GG synthesis. The key metabolites of the gluconeogenic pathway in strain M-2522-GgpPS-drfbA, such as FBP and F6P, were decreased by 11 and
Figure 4 | (A) GG synthesis of strains M-GgpPS, M-1085-GgpPS, M-1175-GgpPS, and M-2522-GgpPS; (B) GG synthesis of strains M-2522-GgpPS-pSII, M-2522-GgpPS-dpgl, M-2522-GgpPS-drfbA, and M-2522-GgpPS-dglgA. Statistical analysis was conducted as described in the text, as statistical significance indicated by **p < 0.01; *p < 0.05.

Figure 5 | Targeted metabolomics analysis of strain M-2522-GgpPS-drfbA and M-pSI-pSII. (A) Comparison of the metabolites in central metabolic pathway between strain M-2522-GgpPS-drfbA and the control strain M-pSI-pSII, red: increased, blue: decreased; (B) heatmaps of metabolomics profiles in strains M-2522-GgpPS-drfbA and M-pSI-pSII. Each colored cell on the heatmap corresponds to a concentration value. The higher the concentration, the darker the color (red represents the increase, and blue represents the decrease).

12%, respectively. However, GAP as the precursor to G3P was increased by 25% compared with the control strain M-pSI-pSII. In addition, the metabolites 3PG, PEP and PYR in the glycolytic pathway were increased by 6-, 1.9-, and 2.4-fold compared with the control strain M-pSI-pSII, respectively, suggesting that more carbon might be redirected to the TCA cycle, which is necessary for biomass accumulation.

GG Synthesis for the Salt Tolerance of Syn2973
Glucosylglycerol, a compatible solute, plays an important role in moderate and halotolerant cyanobacterial strains with salt tolerance (Klahn and Hagemann, 2011). To test the salt tolerance of different GG-producing strains, growth curves of strains were measured under added 0.5 M NaCl. The results showed that the OD_{750} values of strains M-GgpPS and M-2522-GgpPS were increased by 27 and 52% compared with the control strain M-pSI at a cultivation time point of 60 h, respectively. The OD_{750} value of strain M-2522-GgpPS-drfbA was improved by 62% compared with that of the control strain M-pSI-pSII (Figure 6). These results indicated that the salt tolerance of Syn2973 was enhanced in association with the increase in GG synthesis in Syn2973.

In addition, the growth of strain M-2522-GgpPS-drfbA and control strain M-pSI-pSII was also measured under high light (500 µmol photons m^{-2} s^{-1}) and salt stress conditions, and the growth of strain M-2522-GgpPS-drfbA was improved by threefold compared with the control strain M-pSI-pSII under high light and 0.4 M NaCl (Supplementary Figure 6). However, neither of the strains survived under high light and 0.5 M NaCl, suggesting that further improvement is still necessary for growth under high-light and high-salt conditions.
DISCUSSION

Syn2973 has a faster growth rate and is more tolerant to high temperature and high light, making it a good candidate chassis for autotrophic photosynthetic microbial cell factories. However, Syn2973 is sensitive to salt stress. In this study, the salt tolerance of Syn2973 was preliminarily improved by heterologously expressing ggpP and ggpS. Overexpressing the glpD gene and downregulating the rfbA gene further improved GG production and enhanced the salt tolerance of Syn2973. Under 0.5 M NaCl, the OD_{750} value of strain M-2522-GgpPS-drfbA was improved by 62% compared with that of the control strain M-pSI-pSII. A comparative metabolomic analysis between strains M-pSI-pSII and M-2522-GgpPS-drfbA showed that more carbon flux was redirected from ADP-GLC to GG synthesis. In addition, more carbon was directed to the TCA cycle for cell growth compared with the control strain M-pSI-pSII.

Glucosylglycerol synthesis was achieved in Syn2973 under salt stress conditions, and GG production reached 51 mg/g under salt stress in strain M-GgpPS. The GG production in wild-type Synechocystis was up to 100 mg/L (Tan et al., 2015), suggesting that the accumulation level of GG was relatively lower than that in Synechocystis. Reasons for the limited accumulation of GG in Syn2973 cells are not entirely clear but could be speculated, such as insufficient precursor supply. To confirm this hypothesis, targeted metabolomics analysis was conducted, and the results showed that the G3P content was decreased in strain M-GgpPS, suggesting that it was a limiting step for GG synthesis. G3P is a key precursor for triacylglycerol synthesis (Dulermo and Nicaud, 2011; Wang et al., 2016). A previous study showed that lipid production was enhanced by overexpressing the glpD gene (Liu et al., 2016). In addition, the production of 1,3-propanediol was increased by heterologous overexpression of the glpD gene (Liu et al., 2016). It is interesting to explore the relationship between GG synthesis and G3P supply. Moreover, ADP-GLC and UDP-GLC in strain M-GgpPS were increased, and pentose phosphate pathways, such as E4P and R5P, were decreased, suggesting that there might be more flux through the sucrase phosphate pathway due to the introduction of GG synthesis. The sucrose content decreased, which might be due to GG synthesis driving part of the flux. The qRT-PCR assay showed that expression of the sps gene was downregulated and that of rfbA was upregulated. The sps gene in Syn2973 was the limiting step for sucrose synthesis (Lin et al., 2020), suggesting that some carbon might accumulate in UDP-GLC. Cyanobacterial glycogen can be utilized as a sugar feedstock by microbes for ethanol fermentation (Aikawa et al., 2013) and has significant roles in tolerance to nitrogen, salt, or oxidative stress. Previous studies showed that sucrose and GG could be enhanced by blocking glycogen synthesis (Miao et al., 2003; Xu et al., 2013). The glycogen content did not show a significant change in strain M-GgpPS. It is necessary to enhance GG synthesis by improving G3P contents and directing ADP-GLC flux toward GG and away from glycogen and sucrose biosynthesis in future research.

By performing overexpression of glpD genes from different sources, we found that glpD from Syn7942 demonstrated the best effect in terms of improved GG synthesis, suggesting that it was effective for improving GG production by increasing the G3P supply. Surprisingly, GG production by strain M-2522-GgpPS-dglgA with glgA downregulation was decreased. A recent study showed that downregulation of glycogen has little effect on sucrose production (Lin et al., 2020). Another study suggested that reducing glycogen content could not increase sucrose content in Syn7942 (Ducat et al., 2012). Meanwhile, a previous study indicated a close link between the glycogen pool and compatible solute synthesis in Syn7942 and strongly suggested that intracellular glycogen serves, at least partially, as a carbon pool supporting sucrose synthesis rather than competing with sucrose synthesis (Qiao et al., 2018). A previous study also showed that overexpression of genes associated with glycogen synthesis, such as ADP-glucose pyrophosphorylase (GlgC), could enhance sucrose production (Lin et al., 2020). More recently, a study showed that trehalase production was increased by overexpressing glgC and isoaomylase-type debranching enzyme (glgX) (associated with glycogen degradation) (Choi and Woo, 2020). It is thus worth further exploring how to increase glycogen metabolism for GG synthesis in the future. In addition, GG synthesis in strain M-2522-GgpPS-dglg with downregulation of pgl was slightly decreased. The reason might be associated with the roles of the pgl gene in the pentose phosphate pathway, which was associated with CO₂ fixation. The GG synthesis of strain M-2522-GgpPS-drfbA was increased by downregulating the rfbA gene, suggesting that more carbon flux was redirected from sucrose to GG synthesis.
To further decipher the possible mechanisms involved in GG synthesis and its regulation, targeted metabolomics analysis was also performed for strain M-2522-GgpPS-drfbA and the control strain M-pSI-pSII. The G3P in strain M-2522-GgpPS-drfbA was increased, suggesting that overexpressing glpD was effective for G3P accumulation. In addition, sucrose as a compatible solute in strain M-2522-GgpPS-drfbA was further decreased, which was closely related to the limited improvement of salt tolerance. The key metabolites of the gluconeogenic pathway in strain M-2522-GgpPS-drfbA, such as FBP, F6P, and GAP, were decreased. A previous study showed that fructose 1,6-bisphosphate aldolase (FBA) was associated with glycolysis, gluconeogenesis, and the Calvin cycle (Zgiby et al., 2000), which is known to be associated with salt stress (Patipong et al., 2019). It is interesting to check whether improving FBA activity could further redirect more carbon flux to the sugar synthesis pathway to further improve GG production. The metabolites 3PG, PEP, and PYR in the glycolytic pathway were increased compared with the control strain M-pSI-pSII, which was consistent with the better growth phenotype of strain M-2522-GgpPS-drfbA. It is expected that more carbon flow can be driven into the TCA cycle to maintain cell growth in the early stage, and more carbon flow can be redirected to the gluconeogenic pathway to promote GG synthesis by induced sRNA interference in the later stage.

Glucosylglycerol represents the primary compatible solute for salt resistance in many marine cyanobacteria (Kirsch et al., 2019). To validate whether the GG accumulation level was associated with the salt tolerance of Syn2973, different mutant strains were chosen to evaluate their growth under salt stress. The results suggested that the salt tolerance of mutants increased with increasing GG production ability, indicating that GG production was beneficial for Syn2973 under salt stress. Together, these results indicated that the heterologous synthetic compatible solutes could function as compatible solutes for resisting salt stress. In addition, the strain M-2522-GgpPS-drfbA and control strain M-pSI-pSII were tested under high-light culture conditions. The growth of strain M-2522-GgpPS-drfbA was far better than that of control strain M-pSI-pSII under high light and 0.4 M NaCl; however, neither of the strains survived under conditions of high light and 0.5 M NaCl, suggesting that high light could inhibit the growth of Syn2973 under salt stress. It is speculated that several mechanisms might be involved: (i) excess light energy could break the balance between energy supply and consumption of cyanobacteria, leading to the intracellular accumulation of reactive oxygen species (ROS) (Muramatsu and Hihara, 2012); meanwhile, high salt stress could also induce ROS accumulation (Swapnil et al., 2017). The accumulated ROS further disrupted key cellular components, seriously restricting cell viability; (ii) salt stress inhibited photosystems II and I in cyanobacteria (Allakhverdiev and Murata, 2008), which could interfere with photosynthesis under high light; and (iii) a previous study showed that energy metabolites such as ATP and NADPH might be necessary contributors to the rapid growth of Syn2973 under high-light conditions (Ungerer et al., 2018b). However, energy metabolites are also directly associated with Na+ excretion from the cytoplasm, such as Na+ H+ antiporters that utilize the energy of the transmembrane proton gradient for Na+ exclusion (Cui et al., 2020a). Therefore, energy metabolites might be urgent for cell growth under high-light and high-salt-stress conditions. In this study, we preliminarily tested the heterologous synthesis of GG and its effect on salt tolerance of Syn2973 under low CO2 concentration from the air, the same condition for both engineered strain and wild-type, which could tell the preliminary difference. It also worth evaluating the detailed performance of the engineered strains and wild-type employing other conditions or methods, such as under carbon-sufficient cultivation condition, or a spot assay on agar plates for salt tolerance monitoring in future research. Together, the study provides important engineering strategies to further improve salt tolerance and GG production of Syn2973 in the near future, and further work is still necessary to engineer Syn2973 to reach high salt and high light tolerance for future cultivation applications.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

LC and WZ designed the research. JC and TS performed the major experiments and wrote the draft manuscript. JC, TS, LC, and WZ analyzed the data and drafted and revised the manuscript. All authors contributed to the article and approved the submitted version.

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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.2021.650217/full#supplementary-material

**Supplementary Figure 1** | Identification of correct strains by colony PCR. (A) Amplification of the fragment Prc-sll1566-Tbrcl from strains M-pSII (control strain) and M-GgpPS; (B) amplification of the fragment Prc-sll0746-Prc-sll1566-Tbrcl from strains M-pSII and M-2522-GgpPS; (C) amplification of the fragment sll0746-Tbrcl from strains M-pSII-pSII (control strain) and M-2522-GgpPS-drfbA; M, marker.
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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