Influence of culture conditions on extracellular polysaccharide production and the activities of enzymes involved in the polysaccharide synthesis of *Nostoc flagelliforme*

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Cultures conditions significantly influence extracellular polysaccharide (EPS) production of *Nostoc flagelliforme*, however, the key enzyme controlling EPS synthesis has not been fully explored yet. The influence of different culture conditions including light quality, carbon source and nitrogen source on EPS production of *N. flagelliforme* and activities of EPS synthesis enzymes was investigated. Three experimental groups produced higher amounts of EPS than the control group, including the carbon source group with 1.26 g L\(^{-1}\) NaHCO\(_3\), the nitrogen source group with 0 g L\(^{-1}\) NaNO\(_2\) and the light quality group with blue light. Activities of seven related enzymes phosphoglucone isomerase (PGI), fructose-1,6-bisphosphatase (FBPase), UDP-glucose pyrophosphorylase (UGPase), UDP-galactose-4-epimerase (UGE), UDP-glucose dehydrogenase (UGDH), phosphomannose isomerase (PMI), and phosphofructokinase (PFK) were significantly influenced by culture conditions. Partial least-squares analysis and correlation analysis methods were used to analyze the relationship between the activities of these enzymes and EPS production, and a correlation between the production of EPS and the activities of PGI, PMI, FBPase, UGDH, and UGPase was found under different culture conditions. Subsequent analysis of the transcription level of genes encoding the five enzymes showed genes *pgi* and *fbp1* in three experimental groups were significantly up regulated. The results revealed PGI and FBPase might be important enzymes positively influencing the biosynthesis of *N. flagelliforme* EPS. The findings would be helpful to further understand the pathway of EPS biosynthesis aimed to improve the EPS production of *N. flagelliforme*.

**Introduction**

The cyanobacteria extracellular polysaccharide can be divided into two main forms: the capsule polysaccharide (CPS) fraction bound to the cell surface and the extracellular polysaccharide (EPS) fraction released into the surrounding environment.\(^1\)\(^,\)\(^2\) The EPS shows advantages over the polysaccharides from plants or microalgae, given its characteristic easy recovery from the culture medium and its physicochemical properties.\(^3\)\(^,\)\(^4\) The future looks bright for its potential applications in the food, pharmaceutical and environmental industries, consequently, the interest toward EPS has significantly increased in recent years.\(^5\)\(^,\)\(^6\)

*Nostoc flagelliforme* is a species of edible terrestrial filamentous cyanobacteria, distributed throughout arid and semi-arid areas.\(^6\)\(^,\)\(^9\) The properties of *N. flagelliforme* EPS, including anti-virus, antioxidant and anti-tumor, have been intensively investigated.\(^10\)\(^,\)\(^11\) Considerable research efforts have been devoted to find the optimum culture medium and culture condition to obtain high EPS production yields.\(^12\)\(^,\)\(^13\)\(^–\)\(^17\) Light quality, carbon and nitrogen sources have been reported as important factors controlling *N. flagelliforme* EPS synthesis.\(^13\)\(^,\)\(^17\)\(^–\)\(^21\) As a group of photoautotrophic microorganisms, several studies reported that light quality especially red-light and blue-light could promote the accumulation of *N. flagelliforme* EPS.\(^17\)\(^,\)\(^20\)\(^–\)\(^25\) Most of cyanobacteria have efficient carbon concentration mechanisms (CCM) which could transport and accumulate inorganic carbon actively (Ci; HCO\(_3\)-, and CO\(_2\))\(^26\)\(^,\)\(^28\) and could utilize very different forms of nitrogen, including combined nitrogen source or atmospheric nitrogen,\(^22\)\(^,\)\(^29\)\(^–\)\(^31\) which was helpful for increasing EPS production.

In previous studies, various complex EPS biosynthetic mechanisms were investigated and described in many bacterial, fungal and microalgae species, but the necessary pathways and enzymes involved in typical EPS biosynthesis, such as UDP-
N. agelliforme, the key enzyme that controls EPS production in cyanobacterial EPS was summarized (Fig. 1). In the complex pathway, a simplification was subsequently analyzed for key type and nitrogen concentration on EPS production and activities in different conditions. Furthermore, expression level of genes encoding different light quality, NaHCO$_3$ concentration, nitrogen source, and light quality were evaluated in submerged fermentation of N. agelliforme. The enzyme extract method is modified by the method of Looijesteijn et al. and Tamoi et al. The N. flagelliforme cells were harvested by centrifugation at 10 000 × g for 10 min at 4 °C, washed twice with cell extract buffer pre-cooled at 4 °C, which was 20 mM phosphate buffer containing 50 mM NaCl, 10 mM MgCl$_2$, and 1 mM dithiothreitol at pH 6.5. The cells pellet resuspended in the extract buffer were disrupted ultrasonically at 20 kHz for 12 min on ice using an ultrasonic processor (SCIENTZ-HD sonicator, Ningbo Scientz Biotechnology Co., Ltd., China). Cell debris was removed by centrifuged at 12 000 × g for 15 min at 4 °C, and the resulting supernatant (cell extract) was used for the enzyme activity assay. The protein content of the cell extract was determined by the method of Bradford.

Enzyme extract

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Enzyme assays

Activity of EPS synthesis enzymes were measured spectrophotometrically by kinetic assay in a final volume of 250 μL at 24 °C. Reactions were initiated by adding 30 μL cell extract to various reaction mixtures depending on the enzyme. The enzyme activity was determined by monitoring the formation or disappearance of NAD(P)H by measuring the absorbance at 340 nm (ε$_{340}$ = 6220 M$^{-1}$ cm$^{-1}$), and specific activities were expressed in μmol NAD(P)H (ng protein)$^{-1}$ min$^{-1}$. The blank for each enzyme was the same reaction condition with the same volume of cell extract that had been heated at 100 °C for 5 min. In all of the assays, the reaction velocity was linearly proportional to the amount of cell extract. The reaction rate (Δabsorbance/min) calculated from the slope of linear plots ($R^2 = 0.95–1.0$) from 0–3 min or 0–5 min that was normalized to the control. The enzyme activities were calculated using the following equations:

\[
\text{Enzyme activity (U, μmol min$^{-1}$)} = (\Delta A_{340}/\Delta t) \times V_1 \times D (V_2 \times \varepsilon_{340} \times d)
\]
Specific activities = enzyme activity/protein content

where $\Delta A_{340}/\Delta t$ is the reaction rate, $V_i$ is the final reaction volume, $V_o$ is the cell extract volume, $D$ is the dilution ratio, $e_{340}$ is extinction coefficients of NAD(P)H and NADH, $d$ is colorimetric cup light path.

Phosphoglucom isomerase (PGI)\textsuperscript{37} reaction mixture contained 50 mM potassium phosphate buffer (pH 6.8), 5 mM MgCl\textsubscript{2}, 4 U glucose-6-phosphate dehydrogenase, 0.4 mM NADP\textsuperscript{+}, 10 mM fructose-6-phosphate.

Fructose-1,6-bisphosphatase (FBPase)\textsuperscript{37} reaction mixture contained 100 mM Tris-HCl buffer (pH 8.0), 0.4 mM NADP\textsuperscript{+}, 20 mM MgCl\textsubscript{2}, 20 mM DTT, 0.5 U phosphoglucom isomerase, 0.5 U glucose-6-phosphate dehydrogenase, 2 mM fructose-1,6-bisphosphate.

UDP-glucose pyrophosphorylase (UGPase)\textsuperscript{49} reaction mixture contained 50 mM Tris-HCl buffer (pH 7.8), 0.4 mM UDP-glucose, 14 mM MgCl\textsubscript{2}, 4 U UDP-glucose-6-phosphate.

UDP-galactose-4-epimerase (UGE)\textsuperscript{37} reaction mixture contained 100 mM Tris-HCl buffer (pH 7.5), 5 mM MgCl\textsubscript{2}, 0.015 U UDP-glucose dehydrogenase, 0.5 mM NAD, 0.2 mM UDP-galactose.

UDP-glucose dehydrogenase (UGDH)\textsuperscript{39} reaction mixture contained 100 mM Tris-HCl buffer (pH 7.5), 5 mM UDP-glucose, 1 mM dithiothreitol, 1 mM MgCl\textsubscript{2}, 1 mM NADP\textsuperscript{+}.

Phosphomannose isomerase (PMI)\textsuperscript{39} reaction mixture contained 50 mM MOPS buffer (pH 7.0), 1 mM CoCl\textsubscript{2}, 1 mM NADP\textsuperscript{+}, 10 mM mannose-6-phosphate, 4 U phosphoglucom isomerase, 4 U glucose-6-phosphate dehydrogenase.

Phosphofructokinase (PFK)\textsuperscript{39,37} reaction mixture contained 50 mM Tris-HCl buffer (pH 7.5), 5 mM MgCl\textsubscript{2}, 50 mM KCl, 1.25 mM ATP, 0.15 mM NADH, 4.5 U aldolase, 18 U triose-phosphate isomerase, 6.2 U glycerol-3-phosphate dehydrogenase, 5 mM fructose-6-phosphate.

### Total RNA extraction

Total RNA of \textit{N. flagelliforme} cells was prepared using MiniBEST Universal RNA Extraction Kit (TaKaRa, Japan) according to the manufacturer’s instruction. For cDNA synthesis of total RNA was transcribed with the PrimeScript\textsuperscript{TM} RT reagent Kit (TaKaRa, Japan) according to the manufacturer’s instruction. Total RNA was quantified through the absorbance spectrum at wave-lengths 260 nm and 280 nm using UV-mini1246 spectrophotometer (SHIMADZU, Japan). Total RNA solution was loaded on 2% agarose gel to assess the integrity of total RNA bands. The DNA maker was DL2000 maker (TaKaRa, Japan).

### Primer designing and real-time quantitative PCR analysis

Expression levels of \textit{pgi}, \textit{pmi}, \textit{fbp1}, \textit{fbp2}, \textit{ugp} and \textit{ugdh} (encoding enzymes PGI, PMI, FBPase, UGPase and UGDH, respectively) were analyzed by qRT-PCR using SYBR® Premix Ex Taq\textsuperscript{TM}(Tli RNaseH Plus) Kit (TaKaRa, Japan) according to the manufacturer’s instructions with Bio-Rad IQ5 (Bio-Rad, American). Primers used for qRT-PCR were listed in Table 1. GAPDH was used as the reference gene to normalize the expression levels of different genes. For each gene, the reference sample (the control group) was defined as 1.0, and the $2^{-\Delta\Delta C\text{t}}$ method was applied to calculate other gene expression level which was expressed as the fold of the reference sample.\textsuperscript{2,3} Negative controls (reactions without cDNA template) were included to detect the presence of any DNA contamination. All the reactions of these samples were done in triplicate to determine the accuracy and repeatability of the qRT-PCR reaction.

### Data analysis and statistical analysis

Experimental data in this study was obtained from three replicates for each treatment, and values were shown as mean ± standard deviation. Supervised partial least-squares (PLS) analysis was used to verify the relative enzymes which were significant response to the EPS production for different culture conditions. Supervised PLS was performed by SIMCA 11.5 software (Umetrics, Umea, Sweden). Statistical significance of correlations between enzyme activities and EPS production was based on correlation test, which were analyzed based on data obtained under various culture conditions by SPSS statistics vision 2.0. Correlation coefficient ($R^2$ > 0.5) and significant effects ($P < 0.05$) were considered statistically significant.

### Results

#### Effects of culture conditions on EPS production

In order to optimize the culture condition with maximum production of \textit{N. flagelliforme} EPS, the effects of carbon source, nitrogen source, and light quality on EPS production were studied. As shown in Fig. 2A, the productions of U-EPS were similar and high under both blue-light and red-light condition, which were 48.01 ± 3.70 and 45.84 ± 0.68 mg g\textsuperscript{-1} DCW, respectively. The U-EPS productions under blue-light and red-light condition were 1.52 and 1.45 times compared to that of white-light. Because of the highest biomass, the T-EPS obtained the highest production (37.32 ± 1.75 mg L\textsuperscript{-1}) under blue-light, which was 1.78 times to that of white-light.

The carbon sources used for submerged fermentation of \textit{N. flagelliforme} were NaHCO\textsubscript{3} with different initial concentrations,
and the profiles of EPS production were showed in Fig. 2B. The highest T-EPS (28.38 ± 3.19 mg L^{-1}) and U-EPS (40.93 ± 4.70 mg g^{-1} DCW) production were both observed in the medium containing 1.26 g L^{-1} NaHCO₃, which were 4.27 and 3.75 times compared to that of the medium containing 0 g L^{-1} NaHCO₃ respectively.

The effects of nitrogen source type and initial concentration of NaNO₃ on EPS production by N. flagelliforme were shown in Fig. 2C and D, respectively. Using EPS production as maker, we tested three nitrogen source types (Fig. 2C), including NaNO₃, NH₄Cl and CO(NH₂)₂. It showed that NaNO₃ was the best nitrogen source for accumulating EPS (29.27 ± 0.37 mg L^{-1} T-EPS and 42.87 ± 1.52 mg g^{-1} DCW U-EPS) among these three nitrogen source types. The low level of EPS production was caused by adding NH₄Cl and CO(NH₂)₂ as nitrogen source, especially in the medium containing NH₄Cl as solo nitrogen source with few EPS production (1.32 ± 0.27 mg L^{-1} T-EPS and 3.17 ± 0.81 mg g^{-1} DCW U-EPS) detected, although NH₄⁺ was considered to be the preferred reduced N form for cyanobacteria uptake in general.³⁰ As shown in Fig. 2D, the changes of initial NaNO₃ concentration had different effects on EPS production. The highest EPS production (37.95 ± 1.87 mg L^{-1} T-EPS and 67.73 ± 3.98 mg g^{-1} DCW U-EPS) was observed in the media without exogenous addition of NaNO₃, and EPS production was reduced when initial NaNO₃ concentration was increased.

### Effects of culture conditions on enzyme activities involved in N. flagelliforme EPS synthesis

Activities of seven enzymes were evaluated under different culture conditions (Fig. 3), which were involved in N. flagelliforme EPS synthesis. Fig. 3A showed the profiles of the activity levels of seven enzymes measured under light quality condition. The activities of PGI, PMI, UGDH, UGE and UGPase were significantly improved under blue-light and red-light condition. The order of activity of FBPase was red-light > white-light > blue-light condition. As can be seen from Fig. 3B, it was interesting that for the five enzymes, the lowest activity was obtained at 1.26 g L^{-1} NaHCO₃ and then their activities were increased at 2.94 g L^{-1} NaHCO₃, which was an opposite trend with EPS production. However, FBPase activity was increased with the increase of NaHCO₃ concentration.

For different nitrogen source types, the activities of PGI, PMI, UGDH, UGPase in Fig. 3C had similar trend with EPS production, which was declined after adding NH₄Cl and CO(NH₂)₂ as nitrogen source, except FBPase and PFK. For different initial NaNO₃ concentrations (Fig. 3D), increased concentration of NaNO₃ leaded to the reduction of enzyme activity, which showed similar trend with EPS production, except UGDH. Therefore, the activity of PGI, PMI, and UGPase had the same trend with the changes of EPS production regardless of nitrogen source type and the initial NaNO₃ concentration used.
Relationship between activities of enzymes and EPS production under differing culture conditions

To study relationship between activities of enzymes and EPS production under differing culture conditions and avoid the impact of collinearity, multivariate statistical analysis was performed by using supervised partial least-squares (PLS) analysis and correlation analysis. The PLS was used to validate the differences between the different groups. The t1/t2 scatter plot used to explain the applicability of the PLS Method. The score plot displayed the space formed by the two predictive components as first predictive component ($t_1$) and second predictive component ($t_2$), respectively. These models were all well-constructed and possessed excellent fit and satisfactory predictive capability (Table 2). The first two components accounted for >95% of the total variance for each culture conditions. In the PLS score plots, the confidence interval is defined by Hotelling’s T2 ellipse (95% confidence interval), and observations outside the confidence ellipse are considered outliers.

Variable Importance Plot (VIP) is one of the important plots used to explain or determine the fitting effect of PLS model and VIP coefficients reflect the contribution of each variables to the PLS models. In this study, VIP values ≥ 0.5 were considered as an index indicating that these were possible key enzymes which had contribution to EPS production.

Three groups clearly separated in each profiles of PLS score plots (Fig. 4A and D and Fig. 5A and D). These results showed that the activities of enzymes involved in EPS biosynthesis of N.

Table 2  Statistical data from PLS analysis under different culture conditions

| Culture condition     | $R^2_X$ | $R^2_Y$ | $Q^2$  | $t_1$ | $t_2$ | $t_1 + t_2$ |
|-----------------------|---------|---------|--------|-------|-------|-------------|
| Light quality         | 0.985   | 0.971   | 0.958  | 72.43%| 26.03%| 98.46%      |
| NaHCO$_3$ concentration| 0.991   | 0.947   | 0.913  | 85.48%| 13.65%| 99.13%      |
| Nitrogen source type  | 0.972   | 0.993   | 0.990  | 86.48%| 10.74%| 97.22%      |
| NaNO$_3$ concentration| 0.993   | 0.998   | 0.997  | 85.48%| 13.78%| 99.26%      |

Fig. 3  Effects of culture conditions on enzyme activities involved in N. flagelliforme EPS synthesis. (A) Light quality; (B) NaHCO$_3$ concentration; (C) nitrogen source type; (D) NaNO$_3$ concentration. * indicates $p < 0.05$ and ** indicates $p < 0.01$. 

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flagelliforme differed significantly. PLS loading plots and VIP were used to analyze the contribution of each enzyme to the EPS production. The results from Fig. 4B and C showed that PGI, PMI, UGDH and UGPase were possible key enzymes affecting EPS production under light quality condition. As shown in Fig. 4E and F, PGI, UGDH, FBPass, UGPase and PMI might be
the potential major drivers of EPS accumulation under carbon conditions. The order of enzyme contribution was UGDH, UGPase, PGI, FBPase, PFK and PMI under various nitrogen source conditions (Fig. 5B and C), while PGI, UGPase, PFK and PMI might be the possible key enzymes in the medium containing different concentration of NaNO₃ (Fig. 5E and F).

The correlation test was performed by correlation coefficient ($R^2 > 0.5$) and the results of linear correlation analysis showed in Table 3. The result of correlation analysis between T-EPS and enzyme activities was in accord with that between U-EPS and enzyme activities under different culture conditions, except light quality condition. PGI and UGDH showed positive correlation with both T-EPS and U-EPS under light quality condition. PGI, UGDH, UGPase, PFK and PMI were negatively associated with EPS (T-EPS and U-EPS) under carbon source condition. The effect of UGDH and UGPase was positive on EPS, while that of FBPase was negative under various nitrogen source conditions. The positive correlations were observed between five enzymes (PGI, FBPase, UGPase, PFK and PMI) and EPS production when different concentrations of NaNO₃ were added.

Combining the results of PLS analysis and correlation analysis, by choosing the enzymes showing significant correlation with EPS production under all tested conditions, PGI, PMI, UGDH, UGPase and FBPase were preliminarily selected as the potential important enzymes influencing EPS production.

### Table 3  Correlation analysis between activities of enzymes and EPS production under differing culture conditions

| Culture condition | EPS type | Related enzyme | PGI | FBPase | PFK | PMI | UGDH | UGE | UGPase |
|------------------|---------|----------------|-----|--------|-----|-----|------|-----|--------|
| Light quality    | T-EPS   | ++/-            | +/− | +/−    | ++  | +   | +/−  | +   | +/−    |
|                  | U-EPS   | ++/-            | +/− | −      | ++  | +   | +/−  | +   | +/−    |
| NaHCO₃ concentration | T-EPS | −/−            | −/− | −      | −/− | −   | −/−  | −   | −/−    |
|                  | U-EPS   | −/−            | −/− | −      | −/− | −   | −/−  | −   | −/−    |
| Nitrogen source type | T-EPS | +/−            | +/− | −      | +/− | +   | +/−  | +   | +/−    |
|                  | U-EPS   | +/−            | +/− | −      | +/− | +   | +/−  | +   | +/−    |
| NaNO₃ concentration | T-EPS | ++            | ++  | ++     | ++  | +   | +/−  | +   | ++     |
|                  | U-EPS   | ++            | ++  | ++     | ++  | +   | +/−  | +   | ++     |

**Note:** ++: significant positive, *p* < 0.01; +: positive, *p* < 0.05; −: negative, *p* < 0.05; −−: significant negative, *p* < 0.01; +/−: uncorrelated with the addition of different NaNO₃ concentrations.

### Transcription levels of genes encoding key EPS synthesis enzymes

To evaluate the effects of culture conditions on expression of key enzymes, qRT-PCR was used to analyze the transcription levels of genes including \( pgi \) (encoding PGI), \( pmi \) (encoding PMI), \( fbp1 \) and \( fbp2 \) (encoding FBPase), \( ugdh \) (encoding UGDH) and \( ugp1 \) (encoding UGPase) in control group and three experimental groups with high-yield production of EPS consisting of carbon source group with 1.26 g L⁻¹ NaHCO₃, nitrogen source group with 0 g L⁻¹ NaNO₃, and light quality group with blue-light.

The relative quantitative results of the above genes under four culture conditions were shown in Fig. 6. The mRNA levels were normalized against GAPDH level of control group. In three experimental groups, \( pgi \) and \( fbp1 \) genes were highly expressed compared with that of control group, while the transcription levels of other four genes were decreased in three experimental groups.

### Discussion

In recent two decades, various efforts have been devoted to increase \( N. \) flagelliforme EPS production, which mostly focus on optimization of culture conditions. So far, little information is available about the effects of culture conditions on activities of enzymes involved in EPS production. Therefore, in this study, the possible relationship between \( N. \) flagelliforme EPS production and enzymes activities related to EPS biosynthesis was evaluated.

In this study, the results showed that the EPS production of \( N. \) flagelliforme was significantly influenced by culture conditions, which were consistent with previous reports. In the PLS analysis, PGI was the only enzyme with VIP value > 1 under all four culture conditions. The results of correlation analysis showed that PGI was significantly correlated with EPS production, except under the nitrogen source condition. PGI catalyzes the reversible isomerization of glucopyranose-6-phosphate and fructofuranose-6-phosphate playing a key role in glycolysis and gluconeogenesis pathway. The highest EPS production was also associated with the highest activity of...
PGI under four culture conditions, except under the condition of adding different concentration of NaHCO₃. The gene pgi encoding PGI was highly expressed in three experimental groups verified the results of correlation analysis, referring that pgi might be one important gene influencing EPS production under different culture conditions.

FBPase can catalyze the conversion of fructose-1,6-diphosphate into fructose-6-phosphate, which occurs in two essential steps: one involved in gluconeogenesis and the other in Calvin cycle. The transcription analysis of genes encoding FBPase revealed that two genes (fbp1 and fbp2) encoding two forms of FBPase (FBPase-I and FBPase-II) were detected in N. flagelliforme, which were mainly involved in Calvin cycle and involved in gluconeogenesis respectively. Interestingly, the activity of FBPase was significantly enhanced under red-light, and was obviously increased with increasing initial NaHCO₃ concentration and decreasing NaNO₃ concentration, but the FBPase activity was more closely associated with EPS production under nitrogen condition, which might indicate different regulation mechanisms were involved in different culture conditions.

UGPase was involved in the production of UDP-glucose, which was a key precursor to polysaccharide synthesis. UGDH was required for the synthesis of the precursor UDP-glucuronic acid, which was an important intermediate in polysaccharide synthesis pathway. In this study, both partial least-squares analysis and correlation analysis showed the activities of UGPase and UGDH had strong correlation with EPS production under most of culture conditions. However, the activities were not all increased in experimental groups with higher EPS production. In addition to that, the transcription levels of corresponding genes ugdh and uppG were decreased. Therefore it might be deducted that UGPase and UGDH were not directly associated with EPS yield. In our previous study, it was found that culture conditions showed significant influences on monosaccharide composition of EPS, and UGPase and UGDH were correlated with monosaccharide ratio under different culture conditions. Thus it might be speculated that changes of UGPase and UGDH activities induced by culture conditions had greater influence on EPS monosaccharide composition than yield.

In many cyanobacteria, PMI possess mannose-6-phosphate isomerase activity and nucleotidyltransferase activity which can catalyze the reversible isomerization of fructose-6-phosphate from central metabolism into mannose-6-phosphate and forming GDP-mannose, respectively. The correlation between the activity of PMI and the amount of EPS produced was observed, indicating that EPS production might be affected by activity of PMI. However, in contrast to the increased activity, down-regulation of gene pmi expression level was detected. The reason for discrepancy between encoding gene transcription and activity levels remains unclear and needs to be further investigation.

In addition, UGE, the Leloir pathway key enzyme catalyzing the interconversion of UDP-galactose and UDP-glucose, was not significantly changed, except light quality condition, which seemed to be not closely associated with EPS synthesis. PFK was the first committed step of glycolysis catalyzing the phosphorylation of fructose-6-phosphate to fructose 1,6-bisphosphate by ATP, which seemed not to play an important role in controlling N. flagelliforme EPS production.

In this study, PGI and FBPase were found to be more important enzymes positively influencing the EPS biosynthesis than the other five enzymes, however, through comparing the relationship between enzyme activities and EPS production under four kinds of culture conditions, the results showed some discrepancies that the activities of enzymes had positive effects on EPS production under some culture conditions but had negative effects under other culture conditions. It might be due to the complex EPS biosynthesis process and regulation mechanism. Different influence patterns were found under different culture conditions indicating the utilization of distinct regulation mechanism for different culture conditions, which might explain the discrepancies, and further indicate the existence of coordinated regulation of several enzymes in the EPS production, which still requires further investigation.

**Conclusions**

The effects of culture conditions on N. flagelliforme EPS production and related enzymes were investigated in the study. Culture conditions showed significant influences on EPS production, and related enzyme activities. Both PLS analysis and correlation analysis were used to analyze the relationship between activities of seven enzymes and EPS production. The results showed that the activities of PGI, PMI, FBPase, UGDH and UGPase were correlated with EPS production under different culture conditions. Subsequent analysis of the transcription level of genes encoding the five enzymes showed pgi and fbp1 in three experimental groups was significantly up regulated, which implied PGI and FBPase might be the important enzymes positively influencing EPS production of N. flagelliforme. The findings would improve the understanding of EPS synthesis process in N. flagelliforme.

**Conflicts of interest**

There are no conflicts of interest to declare.

**Abbreviations**

dTDP-Rha
dTDP-rhamnose
F6P
Fructose-6-phosphate
FBP
Fructose-1,6-bisphosphatase
FBPase
Fructose-1,6-bisphosphate
G1P
Glucose-1-phosphate
G6P
Glucose-6-phosphate
M6P Mannose-6-phosphate
PFK Phosphofructokinase
PGI Phosphoglucone isomerase
PMI Phosphomannose isomerase
T-EPS Total EPS
UDP-Gal UDP-galactose
UDP-Glu UDP-glucose
U-EPS Unit quality EPS
UGDH UDP-glucose dehydrogenase
UGE UDP-galactose-4-epimerase
UGPase UDP-glucose pyrophosphorylase

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