Synechocystis PCC 6803 overexpressing RuBisCO grow faster with increased photosynthesis

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ABSTRACT

The ribulose-1,5-bisphosphate (RuBP) oxygenation reaction catalyzed by Ribulose-1,5-bisphosphate carboxylase/oxygenase (RuBisCO) is competing with carboxylation, being negative for both energy and carbon balances in photoautotrophic organisms. This makes RuBisCO one of the bottlenecks for oxygenic photosynthesis and carbon fixation. In this study, RuBisCO was overexpressed in the unicellular cyanobacterium Synechocystis PCC 6803. Relative RuBisCO levels in the engineered strains FL50 and FL52 increased 2.1 times and 1.4 times, respectively, and both strains showed increased growth, photosynthesis and in vitro RuBisCO activity. The oxygen evolution rate increased by 54% and 42% on per chlorophyll basis, while the in vitro RuBisCO activity increased by 52% and 8.6%, respectively. The overexpressed RuBisCO were tagged with a FLAG tag, in strain FL50 on the N terminus of the large subunit while in strain FL52 on the C terminus of the small subunit. The presence of a FLAG tag enhanced transcription of the genes encoding RuBisCO, and, with high possibility, also enhanced the initiation of translation or stability of the enzyme. However, when using a streptavidin-binding tag II (strep-tag II), we did not observe a similar effect. Tagged RuBisCO offers an opportunity for further studying RuBisCO expression and stability. Increased levels of RuBisCO can further improve photosynthesis and growth in the cyanobacterium Synechocystis PCC 6803 under certain growth conditions.

1. Background

Ribulose-1,5-bisphosphate carboxylase/oxygenase (RuBisCO, EC 4.1.1.39) has attracted intensive research interest as an abundant, low efficiency but critical enzyme (Spreitzer and Salvucci, 2002) since it was firstly purified from spinach in 1947 (Wildman and Bonner, 1947). Presently, food supply becomes an outstanding social problem as world population increases rapidly. Optimizing RuBisCO performance is one of the main strategies to increase crop yield to meet the high food requirement (Durall and Lindblad, 2015; Parry et al., 2013).

RuBisCO evolved about 3.8 billion years ago. Until now, three types of RuBisCO (Form I, II, and III) and one RuBisCO like protein (RLP, also called form IV) which lacks the carboxylation ability have been demonstrated (Tabita et al., 2008). Form I hexadecameric RuBisCO exists in most plants, algae, cyanobacteria and proteobacteria. It has eight large subunits forming the catalytic core (4L2) and four small subunits capping on the top and the bottom. L2 is anti-parallel and the catalytic site is formed by N terminus of one large subunit and C terminus of the neighbor large subunit, altogether 8 catalytic sites in one holoenzyme (Tabita et al., 2008). Significant efforts have aimed to increase RuBisCO performance. Unfortunately, only limited successes have been achieved so far due to the turnoff between CO2 affinity and catalytic ratio (Carmo-Silva et al., 2015; Durall and Lindblad, 2015). RuBisCO is an abundant protein in leaves consisting of almost half of the soluble proteins. This makes it difficult to further increase RuBisCO amount in the cells. The complex regulatory system of RuBisCO does not only make homologous expression difficult, but also make heterologous expression challenging (Parry et al., 2013). Overexpressing Synechococcus elongatus PCC 6301 rbcLS in Synechococcus elongatus PCC 7942 resulted in cells with higher in vitro RuBisCO activity and twofold isobutyraldehyde production whereas oxygen evolution remained the same as the strains without additional rbcLS expression (Atsumi et al., 2009). This is a successful example of heterologous expression of RuBisCO even though the Synechococcus elongatus PCC 6301 genome is almost identical to the Synechococcus elongatus PCC 7942 genome, except for a 188.6 kb inversion (Sugita et al., 2007). In addition, positive effects on the free fatty acid production in Synechococcus sp. PCC 7002 have been reported by heterologous expression of RuBisCO from Synechococcus elongates PCC 7942 (Ruffing, 2014). In addition, an increased RuBisCO activity in engi-
neered cells of Synechococcus elongates PCC 7942 containing heterologous rbcLS has been observed (Iwaki et al., 2006). These positive examples of heterologous overexpression of RuBisCO may benefit from avoiding post-transcriptional regulations of the introduced, foreign RuBisCO by the native machinery (Ruffing, 2014).

Cyanobacterial and algae RuBisCO are reported to have higher efficiency compared to plant RuBisCO and are regarded as alternatives to increase crop yield (Whitney et al., 2011). Unfortunately, there is no breakthrough report on expressing cyanobacterial or algae RuBisCO in any plant for now. One of the reasons is the differences and necessary of folding and assembling chaperones. RuBisCO has complex folding and assembling processes with chaperones involved (Hauser et al., 2015). In addition, RuBisCO activation and activity maintenance require an activase that belongs to the AAA+ family. Successful expression of cyanobacterial Synechococcus elongatus RuBisCO in tobacco was reported only recently (Lin et al., 2014). Synechococcus elongatus RuBisCO were assembled in tobacco chloroplasts both with and without the assembling chaperone RbcX, or the carboxysome protein CcmM35, and replaced the tobacco RuBisCO (Occhialini et al., 2016). Cyanobacterial RuBisCO functioned in tobacco, but the transformant line could grow autotrophically only under elevated CO2 concentration (3% v/v CO2/air) (Lin et al., 2014; Occhialini et al., 2016). Even though being an important step this transformant line is far away from an ideal high yield plant.

Fusion tag (protein tag) is widely used for protein purification (Kosobokova et al., 2016). Fusion tag can be a larger protein (like Glutathione S-Transferase, GST) or short peptides containing several amino acids (like FLAG tag, Streptavidin-binding tag). The short peptide tag may have positive effects on protein expression, protein solubility, protein efficiency or even folding (Kosobokova et al., 2016). There are reports that the presence of a His tag increased the expression of RuBisCO (FLAG-slride09-slride011-slride012, encoding large subunit N terminal tagged with FLAG tag (FLAG-slride09-slride011-slride012), encoding small subunit C terminal tagged with FLAG tag (slride09-slride011-slride012-FLAG), encoding large subunit C terminal tagged with FLAG tag (slride09-FLAG-slride011-FLAG), encoding large subunit N terminal tagged with strep-tag II (strept-slride09-slride011-slride012), and encoding small subunit C terminal tagged with strep-tag II (slride09-slride011-slride012-strep). To insert FLAG tag on C terminus of large subunit, overlap extension PCR was used. pPMQAK1 was digested with EcoRI and Psfl digesting sites. Then, five versions of RuBisCO genes were designed and constructed, encoding large subunit N terminus tagged with FLAG tag (FLAG-slride09-slride011-slride012), encoding small subunit C terminus tagged with FLAG tag (slride09-FLAG-slride011-FLAG), encoding large subunit C terminus tagged with FLAG tag (slride09-FLAG-slride011-slride012), encoding large subunit N terminus tagged with strep-tag II (strept-slride09-slride011-slride012), and encoding small subunit C terminus tagged with strep-tag II (slride09-slride011-slride012-strep). Conjugation was performed as described previously (Liang and Lindblad, 2016).

Synechocystis strain carrying pPMQAK1 (cutting off ccdB gene, WT +Km'-vector) is control to engineered strains introducing another rbc gene on pPMQAK1. PspsbA2 was flanked with EcoRI and XbaI digesting sites. RuBisCO gene was flanked with XbaI and Psfl digesting sites. In total, five versions of RuBisCO genes were designed and constructed, encoding large subunit N terminus tagged with FLAG tag (FLAG-slride09-slride011-slride012), encoding small subunit C terminus tagged with FLAG tag (slride09-FLAG-slride011-slride012), and encoding large subunit C terminus tagged with strep-tag II (strept-slride09-slride011-slride012). Fusion tag (protein tag) is widely used for protein purification (Kosobokova et al., 2016). Fusion tag can be a larger protein (like Glutathione S-Transferase, GST) or short peptides containing several amino acids (like FLAG tag, Streptavidin-binding tag). The short peptide tag may have positive effects on protein expression, protein solubility, protein efficiency or even folding (Kosobokova et al., 2016). There are reports that the presence of a His tag increased the expression of RuBisCO (FLAG-slride09-slride011-slride012, encoding large subunit N terminal tagged with FLAG tag (FLAG-slride09-slride011-slride012), encoding small subunit C terminal tagged with FLAG tag (slride09-FLAG-slride011-FLAG), encoding large subunit C terminal tagged with FLAG tag (slride09-FLAG-slride011-slride012), encoding large subunit N terminal tagged with strep-tag II (strept-slride09-slride011-slride012), and encoding small subunit C terminal tagged with strep-tag II (slride09-slride011-slride012-strep). To insert FLAG tag on C terminus of large subunit, overlap extension PCR was used. pPMQAK1 was digested with EcoRI and Psfl digesting sites. Three fragments were ligated using Quick ligase (New England Bio-labs). The resulting plasmids (pFL50, pFL52, pFL50C, pFL50strep, pFL52strep) were conjugated into wild type Synechocystis cells, resulting into engineered strains FL50, FL52, FL50C, FL50strep and FL52strep respectively. Conjugation was performed as described previously (Liang and Lindblad, 2016).

Synechocystis strain carrying a kanamycin cassette on the slride016 site (WT+Km'-genome) was used as a control to the engineered strains with genetic constructs on the chromosome. PspbsA2 together with the operon encoding RuBisCO (FLAG-slride09-slride011-slride012) were amplified from pFL50 and pFL52 and inserted onto pEERM3 Km (Englund et al., 2015) using EcoRI and Psfl sites. The resulting plasmids were conjugated into wild type Synechocystis, resulting in the engineered strains FL50G and FL52G. Complete segregation was confirmed with PCR. Kanamycin cassette and PspbsA2 (with ribosome binding site and an ATG-FLAG tag) were flanked with upstream sequences of slride09 and the initial part of slride09 (without start code) before introduced into Synechocystis wild type cells. Flanking sequence was about 800 bp. The resulting strain, named FL75 (Fig. 1A), was confirmed to be completely segregated using PCR (Fig. 1B). FLAG-TAA and kanamycin cassette flanked with downstream sequences of slride012 and the final part of slride012 (without stop codon) were introduced into Synechocystis wild type cells, resulting in strain FL76 (Fig. 1A). Flanking sequence was about 800 bp and full segregation was confirmed using PCR (Fig. 1C). All digestion enzymes were fast digestion enzymes from Thermo Fisher Scientific. Synechocystis strains used in this study are summarized in Table 1.

2.2. Plasmids and engineered strains

The RuBisCO gene operon (rbc, slride09-slride011-slride012) and PspbsA2 were amplified from wild type Synechocystis genome with corresponding primers (Supplementary Table 1) using Phusion polymerase (Thermo Fisher Science). rbc was expressed either on pPMQAK1 (Huang et al., 2010) or on the chromosome.

Synechocystis strain carrying pPMQAK1 (cutting off ccdB gene, WT +Km'-vector) is control to engineered strains introducing another rbc gene on pPMQAK1. PspbsA2 was flanked with EcoRI and XbaI digesting sites. RuBisCO gene was flanked with XbaI and Psfl digesting sites. In total, five versions of RuBisCO genes were designed and constructed, encoding large subunit N terminus tagged with FLAG tag (FLAG-slride09-slride011-slride012), encoding small subunit C terminus tagged with FLAG tag (slride09-FLAG-slride011-FLAG), encoding large subunit C terminus tagged with FLAG tag (slride09-FLAG-slride011-slride012), encoding large subunit N terminus tagged with strep-tag II (strept-slride09-slride011-slride012), and encoding small subunit C terminus tagged with strep-tag II (slride09-slride011-slride012-strep). To insert FLAG tag on C terminus of large subunit, overlap extension PCR was used. pPMQAK1 was digested with EcoRI and Psfl and larger fragment was recovered. These three fragments were ligated using Quick ligase (New England Bio-labs). The resulting plasmids (pFL50, pFL52, pFL50C, pFL50strep, pFL52strep) were conjugated into wild type Synechocystis cells, resulting into engineered strains FL50, FL52, FL50C, FL50strep and FL52strep respectively. Conjugation was performed as described previously (Liang and Lindblad, 2016).

Synechocystis strain carrying a kanamycin cassette on the slride016 site (WT+Km'-genome) was used as a control to the engineered strains with genetic constructs on the chromosome. PspbsA2 together with the operon encoding RuBisCO (FLAG-slride09-slride011-slride012) were amplified from pFL50 and pFL52 and inserted onto pEERM3 Km (Englund et al., 2015) using EcoRI and Psfl sites. The resulting plasmids were conjugated into wild type Synechocystis, resulting in the engineered strains FL50G and FL52G. Complete segregation was confirmed with PCR. Kanamycin cassette and PspbsA2 (with ribosome binding site and an ATG-FLAG tag) were flanked with upstream sequences of slride09 and the initial part of slride09 (without start code) before introduced into Synechocystis wild type cells. Flanking sequence was about 800 bp. The resulting strain, named FL75 (Fig. 1A), was confirmed to be completely segregated using PCR (Fig. 1B). FLAG-TAA and kanamycin cassette flanked with downstream sequences of slride012 and the final part of slride012 (without stop codon) were introduced into Synechocystis wild type cells, resulting in strain FL76 (Fig. 1A). Flanking sequence was about 800 bp and full segregation was confirmed using PCR (Fig. 1C). All digestion enzymes were fast digestion enzymes from Thermo Fisher Scientific. Synechocystis strains used in this study are summarized in Table 1.

2.3. Optical density, chlorophyll a content and oxygen evolution measurement

Synechocystis optical density was measured at 750 nm (OD750) using a spectrophotometer (Cary 50 UV–visible Spectrophotometer, Varian). Chlorophyll a was extracted with 90% methanol. Chlorophyll a content and oxygen evolution measurement as detailed previously (Liang and Lindblad, 2016). Three independent experiments were conducted, each experiment with biological replicates and technical
replicates. Obtained data were analyzed for statistical significance using one-way analysis of variance (one-way ANOVA).

2.4. Semi-quantitative reverse transcription polymerase chain reaction (RT-PCR)

Method and kit used for semi-quantitative RT-PCR were as detailed previously (Liang and Lindblad, 2016).

2.5. Crude protein extraction and relative RuBisCO content determination

*Synechocystis* cells were collected during log phase. Crude proteins were extracted with 50 mM Tris-HCl (pH 8.0) and concentration was determined using DC protein assay (BIO-RAD). For sodium dodecyl sulfate polyacrylamide gel (SDS-PAGE), 2.5 µg crude proteins were loaded. Western immunoblot was carried out following protocols detailed previously (Liang and Lindblad, 2016). Relative levels of RuBisCO content were determined using Quantity one.

2.6. *In vitro* RuBisCO activity measurement

*In vitro* RuBisCO activity measurement followed the protocol by Atsumi et al. (2009) with only minor modifications. Crude protein concentration was adjusted to 1 µg/µl. A 250 µl reaction mixture was used containing 125 µl 100 mM Tris-HCl (pH 8.0), 100 µl 50 mM MgCl₂, 5 µl 5 mM ethylenediaminetetraacetic acid (EDTA), 2.5 µl NaH¹⁴CO₃ (59 mCi/mmol), and 17.5 µl protein crude extraction. The reaction mixture was incubated for 5 min in 30 °C water bath to activate the enzyme. Then 1 µl 100 mM RuBP was added and the mixture incubated for another 5 min in 30 °C water bath. 100 µl 99% propionic acid was added to stop the reaction and unfixed 14C was evaporated by heating the mixture overnight at 65 °C. Acid-stable products were resuspended in 200 µl 2 mol/L HCl and 3 ml scintillation cocktail (Optiphase™ Hisafe III, PerkinElmer) added. Incorporation of ¹⁴C in the samples was analyzed using a scintillation counter (Tri-Carb® 2810 TR, PerkinElmer). RuBisCO activity in control strain was set as 1.0 and RuBisCO activities in the engineered strains were normalized to the activities in the control strain. Two independent experiments were conducted, each experiment with biological and technical replicates.

![Fig. 1. Genomic organization in the engineered *Synechocystis* strains FL75 and FL76 compared to in the wild type cells (A). Complete segregation of FL75 (B) and FL76 (C) were confirmed by PCRs. In FL75 and FL76, *rbc* on genome was tagged with a FLAG tag on the N terminus of the large subunit and on the C terminus of the small subunit, respectively separately. Numbers indicate the specific primers used in the PCRs to examine segregation. For further details, see Table 1.](image-url)

| Table 1 |
| --- |
| *Synechocystis* PCC 6803 strains used in this study. |
| | Strain | Expression construct | Expression site | Reference |
| | WT+Kmr-vector | pPMQAK1 without cdB |  |  |
| | rbc | Psdh2-RBS*-sr0009-slr0011-slr0012 |  |  |
| | FL50 | Psdh2-RBS*-FLAG-sr0009-slr0011-slr0012 |  |  |
| | FL50c | Psdh2-RBS*-sr0009-FLAG-slr0011-slr0012 |  |  |
| | FL52 | Psdh2-RBS*-sr0009-slr0011-slr0012-FLAG |  |  |
| | FL52strep | Psdh2-RBS*-strepB-sr0009-slr0011-slr0012 |  |  |
| | FL52strep | Psdh2-RBS*-strepB-sr0009-slr0011-slr0012 |  |  |
| | WT+Kmr-genome | Kanamycin cassette replaced sr0168 |  |  |
| | FL50G | Psdh2-RBS*-FLAG-sr0009-slr0011-slr0012 |  |  |
| | FL52G | Psdh2-RBS*-sr0009-slr0011-slr0012-FLAG |  |  |
| | FL75 | A kanamycin cassette with Psdh2 and FLAG tag inserted upstream of sr0009, after ATG of sr0009 |  |  |
| | FL76 | FLAG tag and a kanamycin cassette inserted downstream of sr0012, in front of TAA of sr0012 |  |  |

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Obtained data were analyzed for statistical significance using one-way analysis of variance (One-way ANOVA).

3. Results

3.1. Positive phenotypes of engineered Synechocystis strains with FLAG tag tagged RuBisCO

In a previous study, we demonstrated that increasing fructose-1,6/sedoheptulose-1,7-bisphosphatase, transketolase, and fructose-bisphosphate aldolase levels in Synechocystis PCC 6803 (Synechocystis) resulted in faster growth. Additionally, overexpressing the carboxysome protein CcmM resulted in higher RuBisCO level and faster growth under 100μmol m−2 s−1 light intensity (Liang and Lindblad, 2016). In this study, we used pPMQAK1 to introduce tagged Synechocystis rbc, encoding RuBisCO, either on the large subunit N terminus or the small subunit C terminus, into Synechocystis, resulting in engineered strains FL50 and FL52, respectively. The engineered strains were grown with air under 100μmol m−2 s−1 irradiance.

Strain FL50 grew faster than the control strain (WT+Kmr-vector), both when compared as OD750 and as chlorophyll a content (Fig. 2). However, strain FL52 showed higher chlorophyll a content than control strain but similar OD750 (Fig. 2).

Photosynthesis, examined by measuring maximal in vivo oxygen evolution rate standardized to chlorophyll a content, which increased by 54% and 42% respectively in strains FL50 and FL52 (Fig. 3A) and oxygen evolution rates standardized to OD750, which increased by 63% and 49%, respectively, in strains FL50 and FL52 (Fig. 3B). The Calvin cycle (CBB cycle) consumes ATP and NADPH from photosynthetic light reaction. In CBB cycle, RuBisCO is the initial carboxylation enzyme. But its dual function, carboxylation and oxygenation of RuBP, as well as low turnover rate makes it one of the bottlenecks for carbon fixation in photoautotrophic organisms (Whitney et al., 2011). It is possible that enhanced RuBisCO level resulted in increased flux within the CBB cycle and as consequence, increasing the photosynthesis rate.

3.2. FLAG tag increased both transcription and translation of RuBisCO in Synechocystis

Our earlier study showed that introducing an extra copy of rbc into Synechocystis wild type through pPMQAK1 (rbc strain) did not increase RuBisCO content even though the mRNA level of rbc increased. In addition, the engineered strain did not show any positive phenotypes (Liang and Lindblad, 2016). In the present study, we compared the mRNA and RuBisCO level of three engineered strains, rbc, FL50 and FL52, to the control strain (WT+Kmr-vector). The relative levels of the mRNA of rbc from the three engineered strains increased compared to the control strain while FL50 and FL52 showed even higher mRNA levels than rbc strain (Fig. 4). Even though the tags did not change the sequences of the 5′ untranslated region (5′-UTR) or 3′ untranslated region (3′-UTR), they changed the neighbor sequence of the two regions. Sequence changes on 5′ end of a coding gene may

![Fig. 2. Growth of engineered Synechocystis PCC 6803 strains control (WT+Kmr-vector), FL50, and FL52. A, Chlorophyll a content. B, Optical density at 750 nm (OD750). FL50 and FL52 had rbc gene, coding RuBisCO having FLAG tag fused to the large subunit N terminus or small subunit C terminus separately, overexpressed on pPMQAK1. Mean ± SD is from six biological replicates, six technical replicates. Asterisks indicate that the differences observed between the respective engineered strain and the control strain are significant (One-way ANOVA, P < 0.05). For strain information, see Table 1.](image)

![Fig. 3. Photosynthesis demonstrated by maximal oxygen evolution rate of engineered Synechocystis PCC 6803 strains control (WT+Kmr-vector), FL50, and FL52. A, Oxygen evolution rate per chlorophyll a content. B, Oxygen evolution rate per OD750. FL50 and FL52 had rbc gene, coding RuBisCO having FLAG tag fused to the large subunit N terminus or small subunit C terminus, respectively, overexpressed on pPMQAK1. Mean ± SD is from six biological replicates, six technical replicates. Asterisks indicate that the differences observed between the respective engineered strain and the control strain are significant (One-way ANOVA, P < 0.05). For strain information, see Table 1.](image)
have further effects on DNA polymerase binding and transcription efficiency (Mutalik et al., 2013). mRNA stability can also influence the relative level of mRNA. So it is possible that RuBisCO mRNA with FLAG tag sequence was more stable than that without FLAG tag sequence.

On SDS-PAGE and Western immunoblot images, it was clear that the RuBisCO protein level of rbc strain remained unchanged while FL50 and FL52 accumulated more RuBisCO protein (Fig. 5A). When the RuBisCO level of the control strain was normalized as 1.0, RuBisCO in FL50 and FL52 were 2.1 and 1.4 times of that in control strain respectively (Fig. 5B). Anti-FLAG IgG Western immunoblot showed that the tagged protein was successfully expressed (Supplementary Fig. 1A). rbc, FL50 and FL52 all had more RuBisCO mRNA. But only FL50 and FL52 accumulated more RuBisCO. It implied that the FLAG tag sequence may change the mRNA secondary structure and further affected ribosome binding and translation initiation (Mutalik et al., 2013). Since some RuBisCO large subunits from FL50 and small subunits from FL52 had extra 8 amino acids from FLAG tag, it is also possible that FLAG tag had some positive effects on protein stability and/or folding.

3.3. In vitro RuBisCO activity of engineered strain with FLAG tagged RuBisCO in Synechocystis

Theoretically, the small FLAG tag would not change the protein’s structure and catalytic ability under general conditions (Einhauer and Jungbauer, 2001). In order to examine the level of the RuBisCO activity per crude protein (total RuBisCO activity) and per RuBisCO protein (specific RuBisCO activity), we determined the in vitro total RuBisCO activity by measuring 14C carbon fixation efficiency. When the total RuBisCO activity of the control strain (WT+Km-vector) was normalized to 1.0, the total RuBisCO activity of FL50 and FL52 improved by 52% and 8.6% compared to the control strain, respectively (Fig. 6A). However, the specific RuBisCO activities were similar in the three strains: WT+Km-vector, FL50, and FL52 (Fig. 6B).

RuBisCO has complex activation and catalysis processes (Stec, 2012; Mueller-Cajar et al., 2014). It is easily inhibited by sugar phosphate as well as its own substrate ribulose-1,5-bisphosphate (RuBP). To reactive inhibited RuBisCO, an activase which belongs to the AAA+ (ATPases associated with various cellular activities) family is required (Mueller-Cajar et al., 2014). In this study, we did not engineer the activase. Total RuBisCO activity measurement using crude protein extracts without adding extra activase increased in FL50 and FL52 indicating that already available activase was not limiting the RuBisCO activity under our experimental conditions.

3.4. Strep-tag II did not have the same effect as FLAG tag on RuBisCO expression in Synechocystis

Flag tag and modified streptavidin binding tag (strept-tag II) (Ayala et al., 2013) both contain eight amino acids. Therefore, we explored if strept-tag II will have similar effects as the FLAG tag on RuBisCO engineering to see if the effects were sequence specific. In Synechocystis strains FL50strep and FL52strep, the introduced rbc was tagged with strept-tag II on the large subunit N terminus or small subunit C terminus separately (Table 1). RuBisCO mRNA levels in strains FL50strep and FL52strep were higher than in the control strain (WT+Km-vector), but similar to the rbc strain whose extra rbc was not tagged (Fig. 4). This meant that the presence of the strep-tag II did not have positive effects on transcription or mRNA
stability in comparison to the effect of the FLAG tag. Moreover, the RuBisCO level in the FL50strep and FL52strep strains remained unchanged, like the rbc strain (Fig. 5). This implies that the strep-tag II does not have any positive effects on translation, post-translation modification, or RuBisCO stability.

3.5. Expressing tagged rbc on the chromosome did not result in a changed phenotype in Synechocystis

Continuous antibiotic selection pressure is a widely used method to maintain genetically engineered microbial cells, specifically when introducing the non-native genetic construct using a self-replicating vector. However, it is neither economically and environmentally friendly nor practically feasible to use antibiotics in large-scale cultivation systems. Instead of using antibiotic free technologies (Liu et al., 2011; Tan et al., 2013), inserting the target gene together with antibiotic cassette into the chromosome and selecting for complete segregation is another option to avoid using antibiotics during cultivation. In the present study, the engineered *Synechocystis* strains FL50G, FL52G, FL75 and FL76 (Table 1) with insertion into the chromosome, were confirm by PCR to be still completely segregated after two weeks of growth in media without any antibiotics (data not shown).

*Synechocystis* strain FL50G expressed the 5′ end FLAG tag tagged rbc gene on chromosome while FL52G expressed the 3′ end FLAG tag tagged rbc gene. The genetic constructs were identical to that in strains FL50 and FL52 (Table 1). However, strains FL50G and FL52G did not show any increased level of RuBisCO (Fig. 7). It has been reported that heterologous gene expression on plasmids may be more efficient than the expression on the chromosome (Ng et al., 2015). Another explanation may be some unexpected effect caused by neighbor sequences in the chromosome.

Even though the total RuBisCO activity increased in strains FL50 and FL52, the specific RuBisCO activity was the same as that in the control strain (WT+Km′-vector) (Fig. 6). In fact, there are several versions of the RuBisCO holoenzyme in strains FL50 and FL52 based on numbers and positions of the tagged subunits. It is difficult to determine specific RuBisCO activity of each single version. Here, we measured the native version and the introduced tagged versions (all eight large subunits were tagged or all eight small subunits were tagged). The strains *Synechocystis* FL75 and FL76 (Table 1) were designed and constructed as only expressing tagged RuBisCO. Complete segregation was confirmed with PCR (Fig. 1B and 1C). All large subunits in strain FL75 were tagged with FLAG tag while all small subunits in strain FL76 were tagged. Interestingly, the specific RuBisCO activity was similar in control (WT+Km′-genome), FL75
and FL76 strains (Fig. 6B). This indicates that the FLAG tag did not result in enhanced specific Rubisco activity, and as a consequence, the increased total Rubisco activity observed in strains FL50 and FL52 was due to the increased level of Rubisco.

Except for the native rbc gene, strains rbc and FL52 both possessed an extra copy of rbc gene on shuttle vector. On mRNA level of rbc gene, FL52 had more mRNA compared to rbc strain. This meant rbc gene with FLAG on 3′ end resulted into more mRNA than the version without tag on shuttle vector. When comparing control strain (WT + Km-1-genome) and strain FL76, the rbc gene transcription and translation machineries were identical, meaning the promoter, ribosome binding sites and the neighbor sequences were identical. The only difference was that rbc gene in FL76 had FLAG on 3′ end. However, the mRNA level in FL76 was not higher than that in control strain (WT + Km-1-genome). This phenomenon was not the same as that on shuttle vector and indicated different expression pattern on shuttle vector and chromosome.

4. Discussion

Even though Rubisco is one of the most abundant proteins in photoautotrophic organisms, it may be considered as a bottleneck in photosynthesis and carbon fixation due to its low efficiency. Here we experimentally demonstrated and increased the level of Rubisco as well as Rubisco activity in Synechocystis with subsequently increased photosynthesis and growth by introducing a FLAG tag.

Here we demonstrated that the introduction of a FLAG tag increased transcription and maybe even translation or the stability after translation, when adding a FLAG tag on the N terminus of large subunit or the C terminus of small subunit of Rubisco. Interestingly, even though being of similar length, eight amino acids, strep-tag II in subunit or the C terminus of small subunit of RuBisCO. Interestingly, increased transcription and maybe even translation or the stability photosynthesis and growth by introducing a FLAG tag.

Here we demonstrated that the introduction of a FLAG tag increased transcription and maybe even translation or the stability after translation, when adding a FLAG tag on the N terminus of large subunit or the C terminus of small subunit of Rubisco. Interestingly, even though being of similar length, eight amino acids, strep-tag II in the same position did not have the same effect. Affinity tags may enhance recombinant protein yield as stated before (Walls and the same position did not have the same e-ffect). A FLAG tag on the N terminus of the large subunit or the C terminus of small subunit of RuBisCO allowed red-type form I RuBisCO chaperone–dependent assembly. This extension of 4 small subunits formed an 8–1278 complex core and stabilized it (Joshi et al., 2015). There is a possibility that the FLAG tag may transmit a similar function. However, the C terminus extension of the red-type Rubisco is 25 amino acids, which is much longer than the FLAG tag. In addition, a FLAG tag is a rigid tag with a low possibility to form a β-hairpin. It would be interesting to examine if there is any similarity on the amino acid interaction between the Rubisco protein and the FLAG tag or the β-barrel. In wheat leaves, the Rubisco large subunit was degraded after incubation in darkness. The degradation occurred on the N terminus of the large subunit (Kokubun et al., 2002). In the present study, an increased Rubisco level was found in Synechocystis strain FL50 where the FLAG tag was on N terminus of large subunit but not in strain FL50C where the FLAG tag was on C terminus of large subunit (Supplementary Fig. 2). These results indicated that the degradation of Rubisco large subunit in Synechocystis might also occur from the N terminus. Specific Rubisco activity measurement indicated that the FLAG tag did not increase fusion protein activity (Fig. 6B). Similar results have been reported for glutamate dehydrogenase (GDH). A FLAG tag on the N terminus of mouse GDH did not change the kinetic parameters while positioned on the C terminus, its sensitivity to ADP activation decreased (Pajjecka et al., 2014). Cyanobacterial Rubisco generally has a higher Ss/so (Rubisco specific factor) compared to plant Rubisco. However, correct folding, assembly and stability are issues to be solved when introducing cyanobacterial Rubisco into plants (Occhialini et al., 2016). The observations in the present study offer another approach to examine Rubisco assembly and stability.

Rubisco may be one of the key targets to increase efficiency of carbon fixation in photoautotrophic organisms. The present work offers an additional strategy to overexpress native Rubisco and other versions of Rubisco for further e-ffects. In this work we developed engineered strains of Synechocystis with increased level of Rubisco, which grew faster (Fig. 2) and had higher oxygen evolution rate (Fig. 3). This implies that increasing the performance of Rubisco is one strategy to increase photosynthesis and carbon fixation in cyanobacterial cells.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.metabol.2017.02.002.

References

Akbart, M.K., Turner, N.J., Jones, P.R., 2013. Carboxylic acid reductase is a versatile enzyme for the conversion of fatty acids into fuels and chemical commodities. Proc. Natl. Acad. Sci. USA 110, 87–92. doi:10.1073/PNAS.1216516110.
Atsumi, S., Higashide, W., Liao, J.C., 2009. Direct photosynthetic recycling of carbon dioxide to isobutyraldehyde. Nat. Biotechnol. 27, 1177–1180. http://dx.doi.org/10.1038/nbt.1586.
Ayal, J.C., Pimenta, E., Rodriguez, C., Anne, J., Vallin, C., Milanes, M.T., King-Batsios, E., Huang, K., Van Mellaert, L., 2013. Use of Strep-tag II for rapid detection and purification of Microbacterium tuberculosis recombinant antigens secreted from Streptomyces lividans. J. Microbiol. Methods. 94, 192–198. http://dx.doi.org/10.1016/j.mimet.2013.06.004.
Carmo-Silva, E., Scales, J.C., Madwick, P.J., Parry, M.A.J., 2015. Optimizing rubisco and its regulation for greater resource use efficiency. Plant Cell Environment. 38, 1817–1832. http://dx.doi.org/10.1111/pce.12425.
Durall, C., Lindblad, P., 2015. Mechanisms of carbon fixation and engineering for increased carbon fixation in cyanobacteria. Algal Res. 11, 261–270. http://dx.doi.org/10.1016/j.algal.2014.10.001.
Ehnerauer, A., Junghauner, A., 2001. The FLAG™ peptide, a versatile fusion tag for the purification of recombinant proteins. J. Biochem. Biophys. Methods. 49, 455–465. http://dx.doi.org/10.1016/S0165-222X(01)00215-5.
Egeland, E., Andersen-Ranberg, J., Miao, R., Hamberger, B., Lindberg, P., 2015. Metabolic engineering of Synechocystis sp. PCC 6803 for production of the plant diterpenoid manoyl oxide. ACS Synth. Biol. 4, 1270–1278. http://dx.doi.org/10.1021/acsacb.5b00070.
Hauser, T., Popilka, L., Hartl, F.U., Hayer-Hartl, M., 2015. Role of auxiliary proteins in Rubisco biogenesis and function. Nat. Plants 1, 150–165. http://dx.doi.org/10.1038/nplants.2015.65.
Huang, H.-H., Camsund, D., Lindblad, P., Heidorn, T., 2010. Design and characterization of molecular tools for a synthetic biology approach towards developing cyanobacterial biotechnology. Nucleic Acids Res. 38, 2577–2593, (doi: 0.1093/nar/gkq164).
Iwaki, T., Haranoh Kin Inoue, N., Kojima, K., Sato, R., Nishino, T., Wada, S., Ihara, H., Tanyumara, S., Kobayashi, H., Waddan, A., 2006. Expression of foreign type I ribulose-1,5-bisphosphate carboxylase/oxygenase (EC 4.1.1.39) stimulates photosynthesis in cyanobacterium Synechocystis PCC 7942 cells. Photosynth. Res. 88, 287–297. http://dx.doi.org/10.1007/s11120-006-9048-x.
Joshi, J., Mueller-Cajar, O., Tsai, Y.C., Hartl, F.U., Hayer-Hartl, M., 2015. Role of small subunit in mediating assembly of red-type Form I Rubisco. J. Biol. Chem. 290, 1066–1074. http://dx.doi.org/10.1074/jbc.M114.613091.
Kokubun, N., Ishida, H., Makino, A., Abe, T., 2002. The degradation of the large subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase into the 44-kDa fragment in the lysates of chloroplasts incubated in darkness. Plant Cell Physiol. 43, 1390–1395. http://dx.doi.org/10.1093/pcp/pcf159.
Korn dorfer, I.P., Skerra, A., 2001. Improved affinity of engineered streptavidin for the Strept-tag II peptide is due to a fixed open conformation of the lid-like loop at the binding site. Protein Sci. 11, 883–893. http://dx.doi.org/10.1002/pro.1159.
Kosobokova, E.N., Czernykin, K.A., Kosorukov, V.S., 2016. Overview of fusion tags for recombinant proteins. Biochem 81, 187–200. http://dx.doi.org/10.1016/j.thermal.2017.03.015.
Mueller-Cajar, O., Stotz, M., Bracher, A., 2014. Maintaining photosynthetic CO2 fixation via protein remodelling: the Rubisco activases. Photosynth. Res. 119, 191–201. http://dx.doi.org/10.1007/s11120-013-9819-0.

Ng, A.H., Berla, B.M., Pakrasi, H.B., 2015. Fine-Tuning of photoautotrophic protein assembly factors grow at near wild-type rates if provided with elevated CO2. Plant J. 85, 148–160. http://dx.doi.org/10.1111/tpj.13098.

Omata, T., Sugiura, M., Sugita, M., 2007. Complete nucleotide sequence of the Synechocystis sp. strain PCC 6803 chromosome: gene content and organization. Photosynth. Res. 93, 55–69. http://dx.doi.org/10.1007/s11120-006-9122-4.

Parry, M.A., Andralojc, P.J., Scales, J.C., Salvucci, M.E., Carmo-Silva, A.E., Alonso, H., Whitney, S.M., 2013. Rubisco activity and regulation as targets for crop improvement. J. Exp. Bot. 64, 717–730. http://dx.doi.org/10.1093/jxb/ers336.

Ruffing, A.M., 2014. Improved free fatty acid production in cyanobacteria with Synechococcus sp. PCC 7002 as host. Front Bioeng. Biotechnol. 2, 17. http://dx.doi.org/10.3389/fbioe.2014.00017.

Schmidt, T.G., Skerra, A., 2007. The Strep-tag system for one-step purification and high-affinity detection or capturing of proteins. Nat. Protoc. 2, 1528–1535. http://dx.doi.org/10.1038/nprot.2007.209.

Stec, B., 2012. Structural mechanism of Rubisco activation by carbamylation of the active site lysine. Proc. Natl. Acad. Sci. USA 109, 18785–18790. http://dx.doi.org/10.1073/pnas.1210754109.

Sugita, C., Ogata, K., Shikata, M., Jikuya, H., Takano, J., Furumichi, M., Kanchisa, M., Omata, T., Sugitaxa, M., Sugita, M., 2007. Complete nucleotide sequence of the freshwater unicellular cyanobacterium Synechococcus elongatus PCC 6301 chromosome: gene content and organization. Photosynth. Res. 91, 55–67. http://dx.doi.org/10.1007/s11120-006-9122-4.

Tabita, F.R., Satagopan, S., Hans, T.E., Kree1, N.E., Scott, S.S., 2008. Distinct form I, II, III, and IV Rubisco proteins from the three kingdoms of life provide clues about Rubisco evolution and structure/function relationships. J. Exp. Bot. 59, 1515–1524. http://dx.doi.org/10.1093/jxb/erm361.

Tan, X., Liang, F., Cai, K., Lu, X., 2013. Application of the FLP/FRT recombination system in cyanobacteria for construction of markerless mutants. Appl. Microbiol. Biotechnol. 97, 6373–6382. http://dx.doi.org/10.1007/s00253-013-4837-6.

Walls, D., Loughran, S.T., 2011. Tagging recombinant proteins to enhance solubility and aid purification. Methods Mol. Biol. 681, 151–175. http://dx.doi.org/10.1007/978-1-60327-475-3_11.

Waugh, D.S., 2005. Making the most of affinity tags. Trends Biotechnol. 23, 316–320. http://dx.doi.org/10.1016/j.tibtech.2005.03.012.

Whitney, S.M., Houtz, R.L., Alonso, H., 2011. Advancing our understanding and capacity to engineer nature's CO2-sequestering enzyme, Rubisco. Plant Physiol. 155, 27–35. http://dx.doi.org/10.1093/pltphysi100.155.27.

Wildman, S.G., Bonner, J., 1947. The proteins of green leaves. 1. Isolation, enzymatic properties and auxin content of spinach cytoplasmic proteins. Arch. Biochem 14, 381–413.