Research article

Title: Engineering cyanobacteria to improve photosynthetic production of alka(e)nes

Authors: Weihua Wang, Xufeng Liu, and Xuefeng Lu

Abstract

Background: Cyanobacteria can utilize solar energy and convert carbon dioxide into biofuel molecules in one single biological system. Synechocystis sp. PCC 6803 is a model cyanobacterium for basic and applied research. Alkanes are the major constituents of gasoline, diesel and jet fuels. A two-step alkane biosynthetic pathway was identified in cyanobacteria recently. It opens a door to achieve photosynthetic production of alka(e)nes with high efficiency by genetically engineering cyanobacteria.

Results: A series of Synechocystis sp. PCC6803 mutant strains have been constructed and confirmed. Overexpression of both acyl-acyl carrier protein reductase and aldehyde-deformylating oxygenase from several cyanobacteria strains led to a doubled alka(e)ne production. Redirecting the carbon flux to acyl-ACP can provide larger precursor pool for further conversion to alka(e)nes. In combination with the overexpression of alkane biosynthetic genes, alka(e)ne production was significantly improved in these engineered strains. Alka(e)ne content in a Synechocystis mutant harboring alkane biosynthetic genes over-expressed in both slr0168 and slr1556 gene loci (LX56) was 1.3% of cell dry weight, which was enhanced by 8.3 times compared with wildtype strain (0.14% of cell dry weight) cultivated in shake flasks. Both LX56 mutant and the wildtype strain were cultivated in column photo-bioreactors, and the alka(e)ne production in LX56 mutant was 26 mg/L (1.1% of cell dry weight), which was enhanced by 8 times compared with wildtype strain (0.13% of cell dry weight).

Conclusions: The extent of alka(e)ne production could correlate positively with the expression level of alkane biosynthetic genes. Redirecting the carbon flux to acyl-ACP and overexpressing alkane biosynthetic genes simultaneously can enhance alka(e)ne production in cyanobacteria effectively.

Keywords: Cyanobacteria, Synechocystis sp. PCC6803, Alka(e)ne, Fatty acid, Metabolic engineering

Background

Interest in engineering cyanobacteria for biofuel production has increased recently driven by using photosynthesis to directly convert carbon dioxide into a desirable fuel [1-6]. Additionally, cyanobacteria exhibit higher solar conversion efficiency and growth rate compared to plants and eukaryotic microalgae [7,8]. Synechocystis sp. PCC6803 was the first cyanobacterium for which the complete genome was sequenced in 1996 [9]. So far 126 genomic sequences of cyanobacteria strains are available [10]. Well established genetic manipulation techniques have been applied for cyanobacteria. The techniques make cyanobacteria highly tractable platforms to build efficient biosynthetic pathways for biofuel production by genetic engineering [11].

Alkanes with C4-C23 carbon chain length possess higher energy density, hydrophobic property and compatibility with existing liquid fuel infrastructure, which are the predominant constituents of gasoline, diesel, and jet fuels [12]. They can be produced by various organisms such as bacteria, yeasts, plants and insects [13]. In the late 1960s, production of alka(e)nes was reported in a diversity of cyanobacteria [14]. In 2010, a two-step alkane biosynthetic pathway in cyanobacteria was identified. Acyl-acyl carrier protein (ACP) can be reduced to aldehyde by an acyl-ACP reductase (AAR, EC 1.2.1.50), and then aldehyde can be oxidized to alkane or alkene by an aldehyde-deformylating oxygenase (ADO) [15].
Fatty acid substrates as acyl chains of membrane lipids are biosynthesized by fatty acid synthase (FAS). Acetyl-CoA is converted to malonyl-CoA by a multi-subunit acetyl-CoA carboxylase consisting of AccA, AccB, AccC and AccD, which is the rate-limiting step of fatty acid biosynthesis [16]. Acyl-ACPs synthesized by FAS can be incorporated into membrane lipids. Free fatty acids (FFAs) generated by lipolytic enzymes during degradation of membrane lipids can also be activated to acyl-ACPs by an acyl-ACP synthetase (AAS, EC 6.2.1.20) [17].

Kaczmarzyk and Fulda (2010) established that the only AAS gene in *Synechocystis* sp. PCC 6803 is *slr1609* [17]. The *slr1609*-knockout mutant was incapable of importing exogenous fatty acids and secreted fatty acids released from membrane lipids into the medium. This suggests a remarkable role for this cyanobacterial AAS in recycling released fatty acids [17]. Our previous study showed that the alka(e)ne production was significantly reduced in *slr1609* deletion mutant of *Synechocystis* sp. PCC6803, which indicates AAS plays an essential role in alka(e)ne production [18].

Alka(e)ne biosynthesis was reported in a diversity of cyanobacteria [19]. Heptadecane and heptadecene are the major constituents of alka(e)nes in *Synechocystis* sp. PCC6803, and the total alka(e)ne content was about 0.1% of the cell dry weight (DW) [5,20]. Alkanes biosynthetic genes from cyanobacteria were heterologously expressed in *Escherichia coli* and *Synechococcus* sp. PCC 7002, and alka(e)ne production ranged from 5–40 mg/L in *E. coli* and reached 5% of DW in *Synechococcus* sp. PCC 7002 [13,21]. Redirecting the glyceraldehyde 3-phosphate (3-PGA) originated from Calvin-Benson-Bassham cycle to acyl-ACP and enhancing the expression of alkane biosynthetic genes should improve efficiency and yield of alka(e)ne production in *Synechocystis* sp. PCC6803 (Figure 1).

In this study, metabolic engineering approaches were employed to construct a series of *Synechocystis* sp. PCC6803 mutant strains. Alka(e)ne production was enhanced by 8.3 times in one of these modified strains by overexpressing alkane biosynthetic genes in two different loci of the genome.

**Results and discussion**

Alka(e)ne production can be doubled in *Synechocystis* mutants overexpressing cyanobacteria alkane biosynthetic genes

*Synechocystis* mutants overexpressing either or both native alkane biosynthetic genes (*sll0208* and *sll0209*) were constructed. The production of alka(e)ne can be doubled in the mutant overexpressing both *sll0208* and *sll0209* (LX32, about 700 μg/L/OD) compared with parent strain (6803yu, about 300 μg/L/OD), and no significant changes was detected for the mutant strain only expressing either *sll0208* (LX31) or *sll0209* (LX33) as shown in Figure 2A.

Alkanes biosynthetic genes from *Synechococcus elongatus* PCC7942 (orf1593 and orf1594) and *Nostoc punctiforme* PCC73102 (*npun1710* and *npun 1711*) were also overexpressed in *Synechocystis* sp. PCC6803. Alka(e)ne production can also be doubled in these two mutants (LX34 and LX35, Figure 2B). These results indicate that enhanced activity of AAR and ADO resulting from overexpression can convert more acyl-ACP available to alka(e)ne.

Pentadecane can be produced in *Synechococcus elongatus* PCC7942, while no pentadecane was detected in the *Synechocystis* mutant overexpressing orf1593 and orf1594 (LX34). The *in vitro* enzyme activity assays performed by Eser et al. (2011) suggested that the *Nostoc punctiforme* PCC73102 ADO may possess higher activity than the *Synechocystis* sp. PCC6803 ADO [22]. The highest reported titers of alka(e)nes from this pathway involved *E. coli* overexpressing orf1594 (aar) and *Npun_R1711* (ado) among 16 different combinations of the two enzymes from a variety of cyanobacteria [13]. In the *Synechocystis* mutant overexpressing orf1594 and *Npun_R1711* (LX39), alka(e)ne production was not improved significantly compared with other three mutants
(LX32, LX34 and LX35), while about 60 μg/L/OD730 pentadecane can be produced in this mutant (Figure 2B).

Redirecting the carbon flux to acyl-ACP can enhance alka(e)ne production in cyanobacteria effectively

Since acyl-ACP is the immediate substrate for alka(e)ne biosynthesis, redirecting the carbon flux to acyl-ACP may enhance downstream alka(e)ne production in cyanobacteria [20]. AAS plays an essential role in recycling the released fatty acids to acyl-ACP [17]. Our previous work showed that native alka(e)ne production was not enhanced by overexpressing slr1609 alone. Maybe activities of AAR and ADO are too low to convert acyl-ACP to alka(e)ne efficiently [18]. A Synechocystis mutant overexpressing alkane biosynthetic genes and slr1609 (LX38) showed enhanced productivity of alka(e)ne by 130% and 60% when compared to Synechocystis sp. PCC6803 and LX32 mutant, respectively (Figure 3A). Overexpressing AAS, AAR and ADO gene simultaneously may improve acyl-ACP pool and downstream alka(e)ne production. Considering our previous observation of a 90% reduction of alka(e)ne content in slr1609 deletion mutant [18], acyl-ACPs from FFA activation by AAS may be predominant source of alka(e)ne in Synechocystis strains.

Synechocystis sp. PCC6803 can accumulate poly-β-hydroxybutyrate (PHB) as carbon and energy storage compound [23]. Acetyl-CoA and NADPH are required for PHB synthesis. The β-ketothiolase encoded by phaA (slr1993) condenses two molecules of acetyl-CoA to acetoacetyl-CoA, which is the first step of PHB biosynthesis. Alka(e)ne profiles of the Synechocystis mutant overexpressing slr0208 and slr0209 in acetyl-CoA carboxylase genes (LX57) and lipolytic enzyme gene (LX58) over-producing strain compared with wildtype strain (6803yu) and LX32 mutant. Error bars represent the standard deviation of three replicates.

Figure 2 Alka(e)ne production in Synechocystis mutants overexpressing cyanobacteria alkane biosynthetic genes. (A) Alka(e)ne production in Synechocystis mutants overexpressing slr0208 (LX31), slr0209 (LX33) and both genes (LX32) compared with Synechocystis sp. PCC6803 (6803yu). Error bars represent the standard deviation of three replicates. (B) Alka(e)ne production in Synechocystis mutants overexpressing orf1594 and Npun_R1711 (LX39), alkane biosynthetic genes from Synechococcus elongatus PCC7942 (LX34) and Nostoc punctiforme PCC73102 (LX35) compared with wildtype strain (6803yu) and LX32 mutant. Error bars represent the standard deviation of three replicates.

Figure 3 Alka(e)ne production can be enhanced effectively by redirecting the carbon flux to acyl-ACP. (A) Alka(e)ne production in Synechocystis mutants overexpressing slr0208 and slr0209 in slr1609 over-producing strain (LX38) and phaA gene deletion mutant (LX40) compared with wildtype strain (6803yu) and LX32 mutant. Error bars represent the standard deviation of three replicates. (B) Alka(e)ne production in Synechocystis mutants overexpressing slr0208 and slr0209 in acetyl-CoA carboxylase genes (LX57) and lipolytic enzyme gene (LX58) over-producing strain compared with wildtype strain (6803yu) and LX32 mutant. Error bars represent the standard deviation of three replicates.
Overexpression of AAR and ADO gene and deletion of PHB biosynthetic gene(s) simultaneously can divert acetyl-CoA and NADPH into production of fatty acid and enhance production of fatty acid-derived alka(e)nes. Acyl-ACP pool may also be improved by increasing activity of acetyl-CoA carboxylase (ACC), which is the bottleneck of fatty acid biosynthesis. In a Synechocystis mutant overexpressing accBCDA genes from our previous work, a 56% increase of native alka(e)ne production was obtained [5]. With this mutant, the sll0208 and sll0209 were overexpressed to yield LX57 strain. The alka(e)ne production of LX57 mutant was enhanced by 3.6 times when compared to Synechocystis sp. PCC6803 (Figure 3B).

The lipolytic enzymes are capable of hydrolyzing acyl chains from membrane lipids. FFAs released from membrane lipids can be activated to acyl-ACPs by an AAS. Cyanobacteria have plenty of membrane lipids and a dynamic lipid metabolism. Based on sequence identity analysis, sll1969 was annotated as a putative lipolytic enzyme gene (lipA). Liu and Curtiss (2012) showed that less FFAs were released in Synechocystis sll1969-deletion mutant than the wildtype strain [24]. Alka(e)ne productivity of the Synechocystis mutant overexpressing sll0208, sll0209 and sll1969 (LX58) was enhanced by 3 times in comparison with wildtype strain (Figure 3B). Increasing activities of LipA can degrade more membrane lipids into FFAs. FFA can be activated to acyl-ACP by AAS. Since acyl-ACP is the immediate substrate for alka(e)ne biosynthesis, overexpressing sll1969 would lead to an increased alka(e)ne biosynthesis.

Alka(e)ne production can be improved significantly in Synechocystis mutants overexpressing multiple copies of alkane biosynthetic genes

To investigate whether alka(e)ne productivity could be further enhanced by overexpressing multiple copies of alkane biosynthetic genes, Synechocystis mutants overexpressing two copies of sll0208 and sll0209 were constructed. Alka(e)ne productivity of a Synechocystis mutant overexpressing two copies of sll0208 and sll0209 in sllr0168 site (LX70) was 1.2 mg/L/OD730 (Figure 4A). Alka(e)ne productivity of LX56 strain overexpressing sll0208 and sll0209 in both sllr0168 and sllr1556 (2-hydroxyacid dehydrogenase gene, ddk) loci was 2.3 mg/L/OD730 (Figure 4A). The final DW of wildtype and LX56 strain in shake flasks was 0.44 and 0.5 g/L, respectively. Alka(e)ne production of wildtype strain in shake flasks was 0.14% of DW (0.64 mg/L). Alka(e)ne production of LX56 strain was enhanced by 8.3 times, up to 1.3% of DW (6.5 mg/L).

The transcriptional levels of sll0208 and sll0209 were steadily increased in wildtype, LX32, LX70 and LX56 mutant in semi-quantitative reverse transcription PCR analysis (Figure 5), which indicated alka(e)ne production could correlate positively with the expression of alkane biosynthetic genes to some extent. Transcription of two adjacent copies of sll0208 and sll0209 may interfere with each other, so transcription level of two copies of sll0208 and sll0209 in tandem in LX70 is lower than that of two copies of sll0208 and sll0209 in separate gene locus in
Overexpressing alkane biosynthetic genes in multiple gene loci can significantly improve the efficiency of alkane production in cyanobacteria. Similar effects can also be found when multiple-site overexpression was applied to cyanobacteria ethanol or ethylene production [6,25]. Small-scale photo-bioreactors for cultivation of cyanobacteria are required for precise characterization of wildtype and engineered strains to optimize the culture conditions and alkane production. The LX56 mutant and wildtype strain exhibited similar growth characteristics when cultivated in the bubble column photo-bioreactors. Both of the cultures in photo-bioreactors reached a much higher density (OD_{730}~13) than cultures in shake flasks (OD_{730} ~3 to 4) at stationary phase (Figure 4B), and the final DW of wildtype and LX56 strain in photo-bioreactors was 2.2 and 2.3 g/L, respectively. The alkane production of LX56 strain was 2.8 mg/L, which was 0.13% of DW. The alkane production of LX56 strain was enhanced by 8 times compared with the wildtype strain, up to 26 mg/L, which was 1.1% of DW (Figure 4C).

In the previous reports, alkane production ranged from 5 to 40 mg/L in E. coli [13] and reached 5% of DW in Synechococcus sp. PCC 7002 [21] by heterologously expressing cyanobacterial AAR and ADO genes. Integrating different strategies of genetic modifications used in these studies into a Synechocystis mutant strain will further enhance yield of alkane(s). Medium chain alkane is less toxic to cyanobacteria than alcohol, acid and other non-native products [26], which leaves huge room for engineering cyanobacteria to improve alkane production.

**Conclusions**
Overexpressing cyanobacteria alkane biosynthetic genes and redirecting the carbon flux to acyl-ACP can improve alkane production in cyanobacteria significantly. The results indicate that metabolic engineering strategies are powerful for engineering cyanobacteria to overproduce alkane(s). Low activities of AAR and ADO need to be enhanced by protein engineering for further improvement of alkane production. It is also important to understand the physiological roles and regulatory mechanism of native alkane(s) in cyanobacterial cell.

**Methods**

**Chemicals and reagents**
Eicosane was purchased from Sigma-Aldrich (USA). Other chemicals were from Merck (Germany) or Amresco (USA). Oligonucleotides and gene synthesis were carried out by Sangon (Shanghai, China). Taq DNA polymerase and all restriction endonucleases were obtained from Fermentas (Canada) or Takara (Japan). The DNA ladders were from Takara (Japan). The kits used for molecular cloning were from Omega (USA) or Takara (Japan).

**Plasmid construction**
Plasmids constructed and used in this study were listed in Table 1.

E. coli strain DH5α was used for molecular cloning. Synechocystis sp. PCC6803 and Synechococcus elongatus PCC7942 were generous gifts from Prof. Xudong Xu of Institute of Hydrobiology, Chinese Academy of Sciences. *Nostoc punctiforme* PCC73102 was a generous gift from Prof. John C. Meeks of UC Davis, USA.

Alkane biosynthetic genes *sll0208* along with *sll0209* were amplified from the genomic DNA of *Synechocystis* sp. PCC6803 with the primers 02089F1/02089R1. The primers 0208F1/0208R1 and 0209F1/0209R1 were used to amplify *sll0208* and *sll0209* gene respectively. Alkane biosynthetic genes *orf1593* along with *orf1594* were amplified from the genomic DNA of *Synechococcus elongatus* PCC7942 with the primers 9394F1/9394R1. The primers 1594F1/1594R1 were used to amplify *orf1594* gene. Alkane biosynthetic genes *npun1710* and *npun1711* were amplified from the genomic DNA of *Nostoc punctiforme* PCC73102 with the primers 1011F1/1011R1. The primers 1711F1/1711R1 were used to amplify *npun1711*. The *sll0208* gene, *sll0209* along with *orf1593* along with *orf1594*, *npun1710* along with *npun1711*, *orf1594* along with *npun1711* were subcloned into XbaI/SmaI site of the plasmid pFQ9R [5], resulting in pLX1, pLX3, pLX5, pLX9 plasmid, respectively. The *sll0209* gene was subcloned into *NdeI/XhoI* site of the plasmid pXT37b [5] to generate plasmid pLX4.

The *ddh* gene were amplified from the genomic DNA of *Synechocystis* sp PCC6803 by PCR using the primers ddh-F/ddh-R and inserted into the TA cloning site of pMD18-T-Simple, to generate the plasmids pXT119. The plasmid pLX3 was used as the template to amplify the 2.5 kb fragment of *Prbc* promoter, *sll0208* and *sll0209* and Rubisco terminator (*Trbc*) [5] using the primers rbcNF/rbcNR. The 2.5 kb fragment was inserted into the TA cloning site of pGEM-T Easy to generate the plasmids pLX27. The *ck2* cassette was excised with Sall and XbaI from pRLA46 [27] and

**Figure 5** Semi-quantitative reverse transcription PCR analysis of the transcriptional levels of *sll0208* and *sll0209* in wildtype, LX32, LX70 and LX56 mutant. The *rnpB* gene was used as the external standards. Lane 1, LX56; Lane 2, LX70; Lane 3, LX32; Lane 4: wildtype.
Table 1 Plasmids constructed and used in this study

| Plasmid | Relevant characteristics | Reference |
|---------|--------------------------|-----------|
| pFQ9R  | Ap<sup>+</sup> Spe<sup>-</sup>, pKW1188St derivative containing Omega, Tr<sub>rec</sub>, terminator, Pr<sub>rec</sub> promoter | [5] |
| pXT37b | Ap<sup>+</sup>, Spe<sup>-</sup>, pUC9<sup>+</sup> derivative containing upstream and downstream fragments of slr1609, Omega and Pr<sub>rec</sub> promoter. | [5] |
| pXT119 | Ap<sup>+</sup>, pMD18-T derivative containing upstream and downstream fragments of slr1556 | This study |
| pLX1   | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing sl0208 gene, P<sub>rec</sub> promoter | This study |
| pLX3   | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing sl0208 and sl0209 gene, P<sub>rec</sub> promoter | This study |
| pLX4   | Ap<sup>+</sup>, Spe<sup>-</sup>, pXT37b derivative containing sl0209 gene, P<sub>rec</sub> promoter | This study |
| pLX5   | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing orf1593 and orf1594 gene, P<sub>rec</sub> promoter | This study |
| pLX6   | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing npun1710 and npun1711 gene, P<sub>rec</sub> promoter | This study |
| pLX9   | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing orf1594 and npun1711 gene, P<sub>rec</sub> promoter | This study |
| pLX13  | Ap<sup>+</sup>, Kan<sup>+</sup>, pXT119 derivative containing sl0208 and sl0209 gene, CK2, and Pr<sub>rec</sub> promoter. | This study |
| pLX14  | Ap<sup>+</sup>, pGEM-T Easy derivative containing sl0208 and sl0209 gene, P<sub>rec</sub> promoter. | This study |
| pLX27  | Ap<sup>+</sup>, pGEM-T Easy derivative containing sl0208 and sl0209 gene, P<sub>rec</sub> promoter, T<sub>rec</sub>, terminator. | This study |
| pLX28  | Ap<sup>+</sup>, Kan<sup>+</sup>, pGEM-T Easy derivative containing sl0208 and sl0209 gene, P<sub>rec</sub> promoter, T<sub>rec</sub>, terminator, CK2. | This study |
| pLX59  | Ap<sup>+</sup>, Spe<sup>-</sup>, pFQ9R derivative containing two fragments of P<sub>rec</sub> promoter, sl0208 and sl0209 gene. | This study |

<sup>a</sup> Ap, Ampicillin. Spe, Spectinomycin. Kan, kanamycin.

<sup>b</sup> CK2 for kanamycin antibiotic resistance gene and Omega for spectinomycin antibiotic resistance gene.

inserted into the SalI/SpeI site of pLX27, to generate the plasmid pLX28. The 3.6kb fragment containing ck2, P<sub>rec</sub>, T<sub>rec</sub>, sl0208 and sl0209 was digested with NdeI and SpeI from pLX28 and cloned into BglII site of pXT119 with blunt ends, to generate the plasmid pLX13.

The P<sub>rec</sub> promoter and sl0208 along with sl0209 were amplified from the plasmid pLX3 by fusion PCR using the primers PrbcBX-F/PrbcK-R, 0809K-F/0809B-R. The above fragment with XbaI site and SpeI/SalI sites on the 5’ and 3’ ends respectively was inserted into the TA cloning site of pMD19-T-Simple to generate the plasmids pLX14. To utilize the isocaudarner pair XbaI and SpeI, the fragment containing two copies of P<sub>rec</sub>, sl0208 and sl0209 gene in tandem was digested with XbaI and SalI, and cloned into XbaI/SalI site of pFQ9R, to generate the plasmid pLX59. Plasmid maps were listed in Additional file 1: Figure S1.

Transformation and construction of *Synechocystis* sp. PCC6803 mutant strains

Strains constructed and used in this study were listed in Table 2.

All of the above plasmids were checked by enzyme digestion and then transformed to *Synechocystis* cells. The transformations of *Synechocystis* strains with plasmids were performed as described [28]. The plasmids pLX1, pLX3, pLX4, pLX5, pLX6, pLX9, pLX13 and pLX59 were transformed to *Synechocystis* sp. PCC6803 to generate the mutant strains LX31, LX32, LX33, LX34, LX35, LX39, LX55 and LX70, respectively. The plasmid pLX3 was transformed to *Synechocystis* mutant strain XT203 and GQ4 to generate LX40 and LX38 mutant strains, respectively. The plasmid pLX13 was transformed to *Synechocystis* mutant strain GQ10, Syn-20ACC and LX32 to generate LX58, LX57, and LX56 mutant strains, respectively. For the initial selection of transformants, the DNA/cell mixture was applied to BG11 agar plates. After 18 h the membrane filters were applied to fresh BG11 agar plates containing following antibiotics (10 μg mL<sup>-1</sup> spectinomycin, 10 μg mL<sup>-1</sup> erythromycin or 5 μg mL<sup>-1</sup> spectinomycin/kanamycin). Homogeneous mutants were obtained by successive streaking on BG11 plates with appropriate antibiotics. Homologous integration of the expressing cassette and complete segregation were confirmed by PCR using primers listed in Additional file 1: Table S1. Schematic diagrams for homologous recombination of different plasmids were listed in Additional file 1: Figure S2. PCR analysis of the genotype of *Synechocystis* mutant strains were displayed in Additional file 1: Figure S3.

Cultivation of *Synechocystis* strains

Normal liquid cultures of all *Synechocystis* strains in this study were grown at 30°C in 500 mL shake flasks containing 300 mL BG11 medium with aeration by sterile air under constant illumination at a photosynthetic photon flux density of approximately 30 μmol photons m<sup>-2</sup> s<sup>-1</sup> of white light. When necessary, the following antibiotics were added: kanamycin (20 μg mL<sup>-1</sup>) and spectinomycin (20 μg mL<sup>-1</sup>). Cell growth of each culture was monitored by measuring OD<sub>730</sub>.

*Synechocystis* sp. PCC6803 wild-type and the mutant strains exhibited similar growth rate and final cell density. The cultures cultivated in shake flasks were harvested at OD<sub>730</sub> of 3 to 4 after 14 days, when the stationary phase reached. All *Synechocystis* strains in this study were cultivated in shake flasks first to evaluate yields of alka(e)nes. The alka(e)ne
Table 2 Synechocystis strains constructed and used in this study

| Strain   | Genotype*                  | Reference                  |
|----------|----------------------------|----------------------------|
| 6803yu   | Synechocystis sp. PCC6803 Wild-type, Glucose-tolerance | Prof. Xudong Xu           |
| XT203    | slr1993::CK2 sacB           | This lab                   |
| GQ4      | psbA2::CK2 PpsbA2::slr1609 | This lab                   |
| GQ10     | slr0168:: Omega PpsbA2::slr1609 | This lab               |
| Syn-20ACC| slr1608:: Omega Prbc::accBCDA (PCC6803) Trbc (5) | [5]                        |
| LX31     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX32     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX33     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX34     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX35     | slr0168:: Omega npun1710&npun1711 | This study        |
| LX38     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX39     | slr0168:: Omega npun1710&npun1711 | This study        |
| LX40     | slr0168:: Omega PpsbA2::slr208 | This study               |
| LX55     | slr1556::CK2 PpsbA2::slr208 | This study               |
| LX56     | slr1556::CK2 PpsbA2::slr208 | This study               |
| LX57     | slr1556::CK2 PpsbA2::slr208 | This study               |
| LX58     | slr1556::CK2 PpsbA2::slr208 | This study               |
| LX70     | slr1556::CK2 PpsbA2::slr208 | This study               |

* PpsbA2. 0.4 kb DNA fragment containing the promoter of petE gene. PpsbA2. 0.3 kb DNA fragment containing the promoter of rbc operon. PpsbA2. 1.3 kb DNA fragment containing the promoter of psbA2 gene. PpsbA2. 0.2 kb downstream DNA fragment of rbc5 gene. PpsbA2. 1.5 kb DNA fragment containing the promoter of psbA2 gene. All promoters and terminators mentioned here are from Synechocystis sp. PCC 6803.

content of LX56 mutant with the highest alka(e)ne yield in this work and the wildtype control cultivated in shake flasks were calculated as a percentage of DW. Conversion between OD730 and DW of LX56 mutant and Synechocystis sp. PCC6803 cultivated in shake flasks was performed by regression analysis (Additional file 1: Figure S4).

The bubble column photo-bioreactor was a 580 mm×30 mm glass column with a silica gel plug. Synechocystis sp. PCC6803 and the LX56 mutant strain were grown in flasks to exponential phase and harvested by centrifugation. The harvested cells were re-suspended in 200 mL fresh BG11 media, and transferred to the column photobioreactors at 30°C under 50 μmol·m⁻²·s⁻¹ of white light with air bubbling for 24h, after which the light intensity was adjusted to 100 μmol·m⁻²·s⁻¹ and the aeration was switched to 5% (v/v) CO₂-enriched air. LX56 mutant was grown in the presence of 10 μg mL⁻¹ kanamycin and 10 μg mL⁻¹ spectinomycin. Cell growth of Synechocystis sp. PCC6803 and LX56 mutant was monitored by measuring OD730.

Extraction and GC-MS analysis of alka(e)nes

Alka(e)nes were extracted from Synechocystis cells. 200 mL normal culture or 50 mL culture of column photobioreactor at stationary phase was harvested by centrifugation. The cells were resuspended in 10 mL of TE buffer (pH8.0) and then lysed by sonication. 50 μL eicosane (1 mg/mL) was added to the cell lysate as the internal standard for alka(e)ne analysis. The lysate was extracted for 1h at room temperature with 10 mL chloroform–methanol (v/v, 2:1) [29]. A two-phase system (top: aqueous, bottom: organic) was generated after shaking for 1 h and centrifugation at 8000 rpm at room temperature for 15 min. The bottom organic phase was transferred to a new glass tube and evaporated to dryness under a stream of nitrogen at 55°C. The residue was dissolved in 1mL of n-hexane. Aliquots of this mixture were analyzed by GC-MS using an Agilent 7890A-5975C system equipped with a HP-INNOWax (30 m×250 μm×0.25 μm). Helium (constant flow 1 mL/min) was used as the carrier gas. The temperature of the injector was 250°C and the following temperature program was applied: 100°C for 1 min, increase of 5°C min⁻¹ to 150°C then increase of 10°C min⁻¹ to 250°C for 15 min. The internal standard was used to determine alka(e)ne yield, which was reported as the mean based on three independent experiments.

Semi-quantitative reverse transcription PCR

RNA was isolated from 50 mL cultures of Synechocystis cells in mid-exponential phase by using TRIZol Reagent (Life Technologies). The first-strand cDNA was synthesized from 1μg of total RNA using a RevertAid First
Strand cDNA synthesis Kit (Thermo SCIENTIFIC) according to the manufacturer’s protocol. PCR was performed using primers 0809RTF1/R1 (Additional file 1: Table S1) to amplify 350 bp of internal coding region of \textit{ssl0208} and \textit{ssl0209}. The RNase P subunit B (\textit{rnpB}) gene-specific primer pairs \textit{rnpB1/2} (Additional file 1: Table S1) were designed to amplify \textit{rnpB} as external standards. Thirty cycles were used for \textit{rnpB} cDNA, and 35 cycles were used for \textit{ssl0208} and \textit{ssl0209} cDNA.

**Additional file**

**Additional file 1: Table S1.** Primers used in this study. Figure S1. Plasmid maps. Figure S2. Schematic diagrams for homologous recombination of different plasmids. Figure S3. PCR analysis of the genotype of \textit{Synechocystis} mutant strains. Figure S4. Linear regression of cell dry weight (DW) versus OD730 for \textit{Synechocystis} sp. PCC6803(6803YU) and \textit{Lx56} strain cultivated in shake flasks.

**Abbreviations**

AACP: Acetyl carrier protein; AAC: AADC reductase; ADO: Aldehyde-deformylating oxygenase; ACS: Acyl-CoA synthetase; AAS: Acyl-ACP synthetase; ACC: Acetyl-CoA carboxylase; FAS: Fatty acid synthase; FFA: Free fatty acid; DW: Cell dry weight; PCR: Polymerase chain reaction; OD: Optical density.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

XL (Xuefeng Lu) conceived of the study. XL (Xuefeng Lu), WW and XL (Xufeng Liu) carried out the experiments including the construction and cultivation of \textit{Synechocystis} sp. PCC6803 mutant strains, extraction and analysis of alka(e)enes, and GC-MS analysis. XL (Xuefeng Lu), WW and XL (Xufeng Liu) wrote the manuscript. All authors read and approved the final manuscript.

**Acknowledgements**

This work was supported by grants from the National Basic Research Program of China (973: 2011CB709801, 2012CBA00907), the National Natural Science Foundation of China (90920308), and the National Key Scientific and Technological Special Project for the ‘11th Five-Year Plan’ of China (2006BAC03A04).

**Author details**

1Laboratory for Synthetic Biology, Key Laboratory of Advanced Bioengineering, Qingdao Institute of Bioenergy and Bioprocess Technology, Chinese Academy of Sciences, No. 189 Songling Road, Qingdao 266101, China. 2University of Chinese Academy of Sciences, Beijing 100049, China.

**Received:** 1 December 2012 **Accepted:** 29 April 2013 **Published:** 6 May 2013

**References**

1. Atsumi S, Higashide W, Liao JC: Photosynthetic conversion of CO2 to ethylene by recombinant cyanobacterium \textit{Synechocystis} 6803. *Energ Environ Sci* 2012, 5:8998–9006.
2. Lu XF: A perspective: Photosynthetic production of fatty acid-based biofuels in genetically engineered cyanobacteria. *BioTechnol Adv* 2010, 28:742–746.
3. Duca DC, Way JC, Silver PA: Engineering cyanobacteria to generate high-value products. *Trends Biotechnol* 2011, 29:95–103.
4. Kaneko T, Sato S, Tanaka A, Asamizu E, Nakamura Y, Miyajima N, Hiroswa M, Sugiyama M, Sasamoto S, et al: Sequence analysis of the genome of the unicellular cyanobacterium \textit{Synechocystis} sp. strain PCC6803. II. Sequence determination of the entire genome and assignment of potential protein-coding regions. *DNA Res* 1996, 3:109–136.
5. Shih PM, Wu DY, LaiF, Aksen SD, Fewer DP, Talla E, Calteau A, Cai F, de Marsac NT, Rippka R, et al: Improving the coverage of the cyanobacterial phyllum using diversity-driven genome sequencing. *Proc Natl Acad Sci USA* 2013, 110:1059–1058.
6. Huang HH, Carnrnud D, Lindblad P, Heidorn T: Design and characterization of molecular tools for a Synthetic Biology approach towards developing cyanobacterial biotechnology. *Nucleic Acids Res* 2010, 38:2577–2593.
7. Peralta-Tahya PP, Zhang FZ, del Cardayre SB, Keasling JD: Microbial engineering for the production of advanced biofuels. *Nature* 2012, 488:330–338.
8. Schirmer A, Rude MA, Li XZ, Popova E, del Cardayre SB: Microbial Biosynthesis of Alkanes. *Science* 2010, 329:559–562.
9. Han J, McCarthy ED, Heoewen WV, Calvin M, Bradley WH: Organic geochemical studies. I. A preliminary report on the distribution of aliphatic hydrocarbons in algae, in bacteria, and in a recent lake sediment. *Proc Natl Acad Sci U S A* 1968, 59:25–33.
10. Li N, Chang WC, Warui DM, Booker SJ, Krebs C, Bollinger JM: Evidence for Only Oxigenative Cleavage of Aldehydes to Alka(e)enes and Formate by Cyanobacterial Aldehyde Decarbonylases. *Biochemistry-US* 2012, 51:7908–7916.
11. Davis MS, Solfabti J, Cronan JE: Overproduction of acetyl-CoA carboxylase activity increases the rate of fatty acid biosynthesis in \textit{Escherichia coli}. *J Biol Chem* 2000, 275:28593–28598.
12. Kaczmazczyk D, Fulda M: Fatty acid activation in cyanobacteria mediated by acyl-acyl carrier protein synthetase enables fatty acid recycling. *Plant Physiol* 2010, 152:1598–1610.
13. Gao QQ, Wang WH, Zhao H, Lu XF: Effects of fatty acid activation on photosynthetic production of fatty acid-based biofuels in \textit{Synechocystis} sp PCC6803. *Biotechnology for Biofuels* 2012, 5.
14. Jia J, Zhang Z, Guo J, Li Y, Wang Y, et al: Biohydrogen production from cyanobacterial aldehydes at high temperatures. *ChemBioChem* 2010, 11:1461–1466.
15. Kaczmarek D, Fulda M: Fatty acid activation in cyanobacteria mediated by acyl-acyl carrier protein synthetase enables fatty acid recycling. *Plant Physiol* 2010, 152:1598–1610.
16. Gao QQ, Wang WH, Zhao H, Lu XF: Effects of fatty acid activation on photosynthetic production of fatty acid-based biofuels in \textit{Synechocystis} sp PCC6803. *Biotechnology for Biofuels* 2012, 5.
17. Ladygina N, Dedychkina EG, Vainshtein MB: A review on microbial synthesis of hydrocarbons. *Process Biochem* 2006, 41:1001–1014.
18. Hu P, Borglin S, Kamennaya NA, Chen L, Park H, Mahoney L, Kicaj A, Shan G, Chavaria KL, Zhang C, et al: Metabolic phenotyping of the cyanobacterium \textit{Synechocystis} 6803 engineered for production of alkanes and free fatty acids. *Appl Environ Microbiol* 2013, 102:850–859.
19. Reppas NB, Ridley CP, Reppas N, Ridley C, Rodley CP: Producing hydrocarbons comprises culturing engineered cyanobacterium in culture medium and exposing engineered cyanobacterium to light and carbon dioxide. *US: JOULE UNLIMITED INC; 2010. 7794969-B1.*
20. Eser BE, Das D, Han J, Jones PR, Marsh ENG: Oxygen-Independent Alkane Formation by Non-Heme Iron-Dependent Cyanobacterial Aldehyde Decarbonylase: Investigation of Kinetics and Requirement for an External Electron Donor. *Biochemistry-US* 2011, 50:10743–10750.
21. Hein S, Tran H, Steinbuchel A: \textit{Synechocystis} sp. PCC6803 possesses a two-component polyhydroxyalkanoic acid synthase similar to that of anoxygenic purple sulfur bacteria. *Arch Microbiol* 1998, 170:162–170.
22. Liu X, Curtis R: Thermo recovery of cyanobacterial fatty acids at elevated temperatures. *J Biotechnol* 2012, 161:445–449.
23. Gao ZX, Zhao H, Lu ZM, Tan XM, Lu XF: Photosynthetic production of ethanol from carbon dioxide in genetically engineered cyanobacteria. *Energ Environ Sci* 2012, 5:9857–9865.
24. Kamarainen J, Knoop H, Stanford NJ, Guerrero F, Akhtar MK, Aro EM, Steuer R, Jones PR: Physiological tolerance and stoichiometric potential of cyanobacteria for hydrocarbon fuel production. *J Biotechnol* 2012, 162:657–74.
27. Yin C, Li W, Du Y, Kong R, Xu X: Identification of a gene, ccr-1 (sll1242), required for chill-light tolerance and growth at 15°C in Synechocystis sp. PCC 6803. Microbiology 2007, 153:1261–1267.

28. Williams JGK: Construction of specific mutations in photosystem II photosynthetic reaction center by genetic engineering methods in Synechocystis-6803. Methods Enzymol 1988, 167:766–778.

29. Bligh EG, Dyer WJ: A rapid method of total lipid extraction and purification. Can J Biochem Physiol 1959, 37:911–917.