A novel gene expression system for *Ralstonia eutropha* based on the T7 promoter

Muzi Hu 1,2,3†, Bin Xiong 2,3†, Zhongkang Li 2,3, Li Liu 2,3, Siwei Li 2,3, Chunzhi Zhang 1†, Xueli Zhang 2,3 and Changhao Bi 2,3*

Abstract

**Background:** *Ralstonia eutropha* (syn. *Cupriavidus necator*) is a model microorganism for studying metabolism of polyhydroxyalkanoates (PHAs) and a potential chassis for protein expression due to various advantages. Although current plasmid systems of *R. eutropha* provide a basic platform for gene expression, the performance of the expression-inducing systems is still limited. In addition, the sizes of the cloned genes are limited due to the large sizes of the plasmid backbones.

**Results:** In this study, an *R. eutropha* T7 expression system was established by integrating a T7 RNA polymerase gene driven by the P_BAD promoter into the genome of *R. eutropha*, as well as adding a T7 promoter into a pBBR1-derived plasmid for gene expression. In addition, the essential DNA sequence necessary for pBBR1 plasmid replication was identified, and the redundant parts were deleted reducing the expression plasmid size to 3392 bp, which improved the electroporation efficiency about 4 times. As a result, the highest expression level of RFP was enhanced, and the L-arabinose concentration for expression induction was decreased 20 times.

**Conclusions:** The *R. eutropha* T7 expression system provides an efficient platform for protein production and synthetic biology applications.

**Keywords:** *Ralstonia eutropha*, *Cupriavidus necator*, T7 expression system, pBBR1 plasmid, Protein expression system

Background

*Ralstonia eutropha*, also known as *Cupriavidus necator*, is a facultative chemolithoautotrophic bacterium which is able to fix CO₂ via the Calvin-Benson-Bassham (CBB) cycle [1]. It is a model microorganism for studying the metabolism of polyhydroxyalkanoates (PHAs) [1, 2]. When phosphorus or nitrogen was limited, the weight of poly(3-hydroxybutyrate) (PHB) accumulated in the cells exceeded 80% of the cell dry weight [3], which makes *R. eutropha* a potential industrial strain for PHA production. Metabolic engineering and synthetic biology were used to engineer *R. eutropha* for the production of biofuels and other valued-added products, such as isobutanol [4], 3-methyl-1-butanol [4], methyl ketones [5], hydrocarbons [6], ethanol [7], isopropanol [8, 9], fatty acids [10] and α-humulene [11]. *R. eutropha* is also a potential chassis for production of industrial enzymes and proteins due to its ability to suppress the formation of inclusion bodies [12], lack of acidic byproducts, and high fermentation cell density [3]. When the enzyme organophosphohydrolase (OPH) was produced in engineered *R. eutropha*, its content reached more than 10 g/L, which

* Correspondence: bi_ch@tib.cas.cn
† Muzi Hu, Bin Xiong and Chunzhi Zhang contributed equally to this work.
2 Tianjin Institute of Industrial Biotechnology, Chinese Academy of Sciences, Tianjin 300308, People’s Republic of China
3 Key Laboratory of Systems Microbial Biotechnology, Chinese Academy of Sciences, Tianjin 300308, People’s Republic of China

Full list of author information is available at the end of the article

© The Author(s). 2020 Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.
was a more than 100-fold increase compared to OPH expression in *E. coli* [12]. This indicated that *R. eutropha* is a potential industrial protein production host.

A robust gene expression toolbox which includes suitable host strains, plasmid vectors and promoters is essential for metabolic engineering and synthetic biology. Plasmids derived from pBBR1, and from plasmids of *IncP* and *IncQ* can replicate in *R. eutropha* individually or compatibly and were used as its plasmid vectors [6]. Promoters used in *E. coli*, such as P lac and P tac [13], were found to be expressed constitutively in *R. eutropha*, while native promoters from *R. eutropha*, such as P phaC [14], P phb [14], P acoE [14], P acoD [14] and P pdhE [14], were also evaluated to be functional [13, 14]. In addition, promoters induced by L-arabinose [6], L-rhamnose [15], and anhydrotetracycline [16] were used in *R. eutropha*. However, most of the plasmid vectors and promoters mentioned above are not frequently used. Although broad-host-range plasmid vectors derived from pBBR1 with the L-arabinose induced promoter PBAD are currently the most popular ones, the vectors are very large, which limits the sizes of carried genes and decreases plasmid transformation efficiency.

In this study, a gene expression system based on the T7 promoter was constructed to improve the current gene expression technology for *R. eutropha* and make it a better protein expression system.

**Results**

**Construction of a T7 gene expression system for *R. eutropha***

The T7 RNA polymerase gene was cloned from *E. coli* BL21(DE3), modified to be driven by the L-arabinose induced promoter PBAD, and the *rfp* gene was cloned under the control of a T7 promoter (taattacgactcacttaggg). The resulting plasmid pj5_00019, based on pBBR1, was introduced into either *E. coli* S17–1 [17] or *R. eutropha* C5 (R. eutropha H16ΔH16_A0006ΔH16_A0008–9) [18]. However, we found that *rfp* was expressed in *E. coli* S17–1, but was not expressed in *R. eutropha* C5 with or without L-arabinose induction. We considered that the failure of expression in *R. eutropha* might be due to the multiple copies of the T7 RNA polymerase genes expressed from the multicopy plasmid.

To verify the hypothesis, *R. eutropha* strain C5T7 was constructed by integrating the RNA polymerase gene with the PBAD promoter and araC into the H16_A0666 locus on the *R. eutropha* C5 genome. H16_A0666 is annotated as lactate dehydrogenase of *R. eutropha*. Because *R. eutropha* strictly depends on respiratory chain for metabolism, the function of lactate dehydrogenase seems redundant. Therefore, we speculate that the removal of H16_A0666 may not affect its growth, which was verified by the experimental results. The expression plasmid pBBR1-P T7-rfp was constructed by cloning the T7-promoter-driven *rfp* into the pBBR1 backbone. After electroporation into *R. eutropha* C5T7, the strain C5T7(pBBR1-P T7-rfp) showed visible red color when induced (Fig. 1), which indicated a T7 gene expression system for *R. eutropha* was successfully constructed.

**Minimization of the expression plasmid vector pBBR1-P T7-rfp**

The pBBR1-P T7-rfp plasmid expressed in *R. eutropha* C5T7 had a size of 4751 bp, which limited the size of genes that can be cloned into the vector. Considering the origin of the plasmid, there might be redundant DNA regions that are non-essential for plasmid replication and could be deleted to minimize the plasmid. The DNA cassette containing acttaaaaatcaacatacttaggg is AT-

![Fig. 1 Schematic of the *R. eutropha* gene expression system. The system includes engineered *R. eutropha* C5T7 (a) and plasmid vector (b). a] Schematic illustration of the gene arrangement of changed genomic locus of *R. eutropha* C5T7. b] Profile of gene expression vector pBBR1-P T7-rfp. Bacterial colonies are visually red when *R. eutropha* C5T7(pBBR1-P T7-rfp) is induced by arabinose
rich with two direct repeats, which was assumed to be the replication origin of pBBR1-derived plasmids [19]. Thus, adjacent regions of the putative replication origin were gradually deleted via PCR followed by Golden Gate assembly to reassemble the plasmids. As a result, a series of circular DNA cassettes were constructed, among which only pJ5_00030 could be introduced into E. coli successfully, which confirmed that only this plasmid contained a functional minimized origin of replication (Fig. 2a). The putative replication origin with about 400 bp of its upstream sequence was conserved in pJ5_00030 and was essential for plasmid replication. The minimized plasmid vector pJ5_00030 was designated as pBBR1-P_{T7}-rfp(mini) (Fig. 2b, Additional file 2).

**Gene expression efficiency of the *R. eutropha* T7 expression system**

To determine the performance of the *R. eutropha* T7 expression system with the minimized expression plasmid, pBBR1-P_{T7}-rfp(mini) was transferred into *R. eutropha* C5T7 via electroporation for expression analysis. Rfp expression was induced by different concentrations of L-arabinose. The results showed that the expression level of Rfp increased with the increase of L-arabinose concentration in the range from 0 to 0.1 mg/mL L-arabinose. When 0.1 mg/mL L-arabinose was added for induction, the expression level of RFP was highest but decreased when L-arabinose exceeded that threshold (Fig. 3a). Compared with pBBR1-P_{BAD}-rfp in *R. eutropha* H16, the highest RFP expression level of *R. eutropha* C5T7(pBBR1-P_{T7}-rfp(mini)) was improved. Especially, due to the higher sensitivity of the T7 expression system, the L-arabinose concentration needed for the highest induction level was lowered 20X compared with the traditional P_{BAD} system (Fig. 3a), which can bring significant cost savings in industry.

**Electroporation efficiency of the plasmid**

To evaluate the electroporation efficiency of the pBBR1-derived plasmids, pBBR1-P_{BAD}-rfp, pBBR1-P_{T7}-rfp and...
pBBR1-P<sub>T7</sub>-rfp(mini) were extracted from E. coli S17–1, and then individually electroporated into R. eutropha C5T7 competent cells. The results showed that the electroporation efficiency of pBBR1-P<sub>T7</sub>-rfp(mini) was significantly higher than that of pBBR1-P<sub>BAD</sub>-rfp or pBBR1-P<sub>T7</sub>-rfp, probably due to the reduced plasmid size (Fig. 3b). Thus, the minimized vector could provide a better cloning efficiency.

### Discussion

**T7 expression system for R. eutropha** was constructed in this work. This system provides an alternative other than current systems, such as Escherichia coli and Bacillus subtilis, which also bring some improvements over current systems. The major advantages probably include R. eutropha’s ability to suppress the formation of inclusion bodies [12], lack of acidic byproducts, and high fermentation cell density [3]. In addition, since the presence of antibiotics resistance form the plasmid was stable during the all the culturing and experiment process, we considered the expression system was relatively stable.

### Conclusions

In this study, an optimized T7 expression system for *R. eutropha* was constructed for protein expression, synthetic biology manipulation and other applications. The system has a few advantages over previous ones. First, with the high sensitivity of the T7 system, the highest expression level was enhanced, and the L-arabinose concentration for induction was decreased 20 times, offering great cost saving. Secondly, due to the minimized size of the expression vector, the electroporation efficiency was improved 4 times ($p < 0.05$), which also makes the plasmid a better vector for accommodating large DNA cassettes.

### Methods

#### Strains and culture conditions

The strains used in this study are listed in Table S1. *E. coli* was cultured at 37°C in lysogeny broth medium (LB, 10 g/L peptone, 5 g/L yeast extract, 10 g/L NaCl). Streptomycin (100 μg/mL) or kanamycin (50 μg/mL) was added to the medium where appropriate. *R. eutropha* was cultured in LB at 30°C. Gentamycin (10 μg/mL) or kanamycin (200 μg/mL) was added where appropriate.

#### Primers and plasmids

Primers (Table S2) used for the construction of plasmids (Table S3) in this study were designed using j5 Device Editor [20]. The other primers were designed using Clonemanager software. High fidelity DNA polymerase was purchased from Takara (Dalian, China) or New England Biolabs (USA). BsaI restriction endonuclease and T4 ligase were purchased from New England Biolabs (USA) and Thermo-Fisher Scientific (USA), respectively. Plasmids were constructed using Golden Gate [21], Gibson [22] or CPEC [23] methods, and used to transform *E. coli* or *R. eutropha*. Plasmids were extracted using the AxyPrep Plasmid Miniprep Kit (AXYGEN, China) according to the manufacturer’s instructions, with minor modifications.

#### Genome integration of R. eutropha

Integration of araC and T7 RNA polymerase genes into *R. eutropha* C5 (*R. eutropha* H16ΔH16_A0006ΔH16_A0008–9) [18] was performed with the pk18mobsacb suicide plasmid with similar procedure as our previous work [18]. For construction of the integration plasmid pJ5_00015, the DNA cassettes of left homologous arm, AraC gene, PBAD promoter, T7 RNA polymerase gene and terminator were assembled into pk18mobsacb plasmid with the Golden gate method. Plasmid pJ5_00015 was then transferred into *R. eutropha* C5 strain by bacterial transconjugation. And the strain with kanamycin resistance was screened on agar plates containing 200 μg/ml kanamycin. The strain with genomic integrated pJ5_00015 was selected by colony PCR. To eliminate the DNA cassette containing antibiotic marker, the above selected strain was cultured in NaCl free LB liquid media containing 10% sucrose, and then plated agar plate containing the same content to screen for strain with no kanamycin. Integration of araC and T7 RNA polymerase genes into *R. eutropha* C5 (*R. eutropha* H16ΔH16_A0006ΔH16_A0008–9) [18] was performed with the pk18mobsacb suicide plasmid with similar procedure as our previous work [18]. For construction of the integration plasmid pJ5_00015, the DNA cassettes of left homologous arm, AraC gene, PBAD promoter, T7 RNA polymerase gene and terminator were assembled into pk18mobsacb plasmid with the Golden gate method. Plasmid pJ5_00015 was then transferred into *R. eutropha* C5 strain by bacterial transconjugation. And the strain with kanamycin resistance was screened on agar plates containing 200 μg/ml kanamycin. The strain with genomic integrated pJ5_00015 was selected by colony PCR. To eliminate the DNA cassette containing antibiotic marker, the above selected strain was cultured in NaCl free LB liquid media containing 10% sucrose, and then plated agar plate containing the same content to screen for strain with no kanamycin resistance. The obtained strain was confirmed again with colony PCR and sequencing of the PCR products, which was designated as *R. eutropha* C5T7 strain.

Integration of araC and T7 RNA polymerase genes into *R. eutropha* C5 (*R. eutropha* H16ΔH16_A0006ΔH16_A0008–9) [18] was performed with the pk18mobsacb suicide plasmid with similar procedure as
our previous work [18]. For construction of the integration plasmid pj5_00015, the DNA cassettes of left homologous arm, AraC gene, PBAD promoter, T7 RNA polymerase gene and terminator were assembled into pk18mobsacb plasmid with the Golden gate method. Plasmid pj5_00015 was then transferred into *R. eutropha* C5 strain by bacterial transconjugation. And the strain with kanamycin resistance was screened on agar plates containing 200 μg/ml kanamycin. The strain with genomic integrated pj5_00015 was selected by colony PCR. To eliminate the DNA cassette containing antibiotic marker, The above selected strain was cultured in NaCl free LB liquid media containing 10% sucrose, and then plated agar plate containing the same content to screen for strain with no kanamycin resistance. The obtained strain was confirmed again with colony PCR and sequencing of the PCR products, which was designated as *R. eutropha* C5T7 strain. 

Integration of araC and T7 RNA polymerase genes into *R. eutropha* C5 (*R. eutropha* H16ΔH16_A0006ΔH16_A0008–9) [18] was performed with the pk18mobsacb suicide plasmid with similar procedure as our previous work [18]. For construction of the integration plasmid pj5_00015, the DNA cassettes of left homologous arm, AraC gene, PBAD promoter, T7 RNA polymerase gene and terminator were assembled into pk18mobsacb plasmid with the Golden gate method. Plasmid pj5_00015 was then transferred into *R. eutropha* C5 strain by bacterial transconjugation. And the strain with kanamycin resistance was screened on agar plates containing 200 μg/ml kanamycin. The strain with genomic integrated pj5_00015 was selected by colony PCR. To eliminate the DNA cassette containing antibiotic marker, The above selected strain was cultured in NaCl free LB liquid media containing 10% sucrose, and then plated agar plate containing the same content to screen for strain with no kanamycin resistance. The obtained strain was confirmed again with colony PCR and sequencing of the PCR products, which was designated as *R. eutropha* C5T7 strain. 

Integration of araC and T7 RNA polymerase genes into *R. eutropha* C5 (*R. eutropha* H16ΔH16_A0006ΔH16_A0008–9) [18] was performed with the pk18mobsacb suicide plasmid with similar procedure as our previous work [18]. For construction of the integration plasmid pj5_00015, the DNA cassettes of left homologous arm, AraC gene, PBAD promoter, T7 RNA polymerase gene and terminator were assembled into pk18mobsacb plasmid with the Golden gate method. Plasmid pj5_00015 was then transferred into *R. eutropha* C5 strain by bacterial transconjugation. And the strain with kanamycin resistance was screened on agar plates containing 200 μg/ml kanamycin. The strain with genomic integrated pj5_00015 was selected by colony PCR. To eliminate the DNA cassette containing antibiotic marker, The above selected strain was cultured in NaCl free LB liquid media containing 10% sucrose, and then plated agar plate containing the same content to screen for strain with no kanamycin resistance. The obtained strain was confirmed again with colony PCR and sequencing of the PCR products, which was designated as *R. eutropha* C5T7 strain.
Assessment of RFP expression levels
Red fluorescence intensity was measured using a microplate reader (Infinite M200 PRO, TECAN) after the strains were induced with L-arabinose for 48 h. The excitation and emission wavelengths were 585 and 620 nm respectively [6]. Optical density at 600 nm (OD_{600}) was also measured using the microplate reader. The RFP expression levels were determined as the ratio of the red fluorescence intensity to the OD_{600}.

Identification of the region necessary for plasmid replication
Different primers for Golden Gate were designed using j5 DeviceEditor, and PCR was implemented using the plasmid pBBR1-P_{T7}-rfp as template. The products of the PCR were purified using the SanPrep DNA Purification Kit (Sangon Biotech, China), and then cyclized via Golden Gate assembly. As a result, some regions of the pBBR1-P_{T7}-rfp were deleted, and the minimized plasmids were used to transform E. coli. Transformed colonies would be obtained if the core region for plasmid replication was intact, while transformation would be unsuccessful if the core region were damaged.

Electroporation of R. eutropha and evaluation of transformation efficiency
Competent cells of R. eutropha were prepared according to the protocol described in our previous work [18]. Briefly, R. eutropha was cultured in 100 mL LB at 30 °C and 200 rpm, then chilled on ice for 5–10 min after the OD_{600} reached 0.8. The cells were collected by centrifugation at 3000×g for 5 min and washed with ice-cold sterile 10% (V/V) glycerol three times. Finally, the cells were suspended in about 0.5 mL 10% glycerol and 0.1 mL portions transferred to sterile 1.5-mL tubes. The competent R. eutropha cells were used immediately, or frozen in liquid nitrogen and preserved at −80 °C for a few weeks.

Electroporation of R. eutropha was implemented according to the protocol described in our previous work [18]. Briefly, appropriate plasmids were added to the competent cells and mixed gently. The mixture was transferred into a pre-chilled 2-mm electroporation cuvette (Bio-Rad, USA), and incubated on ice for 5 min, after which electroporation was performed at a voltage of 2.3 kV. Immediately afterwards, 1 mL of LB preincubated at 30 °C was added to the cells. All the cells were transferred to a sterile 1.5 mL tube and incubated at 30 °C for 2 h. Then, a portion of the cells was spread on LB agar plates with appropriate antibiotics, and cultured at 30 °C until colonies were visible. Electroporation efficiencies were calculated as the ratio of colony-forming units (CFU) to the amount of the used plasmid DNA.
6. Bi C, Su P, Müller J, Yeh Y-C, Chhabra SR, Beller HR, Singer SW, Hillson NJ. Development of a broad-host synthetic biology toolbox for Ralstonia eutropha and its application to engineering hydrocarbon biofuel production. Microb Cell Factories. 2013;12:107.

7. Lee HM, Jeon BY, Oh MK. Microbial production of ethanol from acetate by engineered Ralstonia eutropha. Biotechnol Bioprocess Eng. 2016;21:402–7.

8. Grousseau E, Lu J, Gorret N, Guillouet SE, Sinkey AJ. Isopropanol production with engineered Cupriavidus nercator as bioproduction platform. Appl Microbiol Biotechnol. 2014;98:6277–90.

9. Jillian M, Estelle G, Eric L, SA J, Nathalie G, Guillouet SE. Over expression of GroESL in Cupriavidus nercator for heterotrophic and autotrophic isopropanol production. Metab Eng. 2017;42:74–84.

10. Chen JS, Colon B, Dusel B, Ziesack M, C Way J, Torella JP. Production of fatty acids in Ralstonia eutropha H16 by engineering beta-oxidation and carbon storage. Peerl. 2015;3:e1468.

11. Li H, Liao JC. A synthetic anhydrotetracycline-controllable gene expression system in Ralstonia eutropha H16. ACS Synth Biol. 2015;4:101–6.

12. Simon R, Priefer U, Puher A. A broad host range mobilization system for in vivo genetic engineering transposon mutagenesis in gram negative bacteria. Nat Biotechnol. 1983;1:784–91.

13. Engler C, Gruetzner R, Kandzia R, Marillonnet S. Golden gate shuffling: a one-pot DNA shuffling method based on type IIs restriction enzymes. PLoS One. 2009;4:e5553.

14. Gibson DG, Young L, Chuang R-Y, Venter JC, Hutchinson CA, Smith HO. Enzymatic assembly of DNA molecules up to several hundred kilobases. Nat Methods. 2009;6:343–5.

15. Hillson NJ, Rosengarten RD, Keasing JD, JS DNA assembly design automation software. ACS Synth Biol. 2012;1:14–21.