Effect of Tween 40 and DtsR1 on L-arginine overproduction in Corynebacterium crenatum

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Abstract

Background: L-Glutamate is an important precursor in the L-arginine (L-Arg) biosynthetic pathway. Various methods, including polyoxyethylene sorbitan monopalmitate (Tween 40) addition and dtsR1 disruption, have been widely used to induce L-glutamate overproduction in Corynebacterium glutamicum. In this study, a novel strategy for L-Arg overproduction through Tween 40 trigger and ΔdtsR1 mutant were proposed in Corynebacterium crenatum.

Results: Corynebacterium crenatum mutant (CCM01) was selected as a host strain, whose argR was lethal via mutagenesis screening, the proB gene was knocked out, and argB was replaced by argB M4 (E19R, H26E, D311R, and D312R) to release L-Arg feedback resistance. After Tween 40 trigger in the logarithmic period, L-Arg production increased from 15.22 to 17.73 g/L in CCM01 strain. When NCgl1221 and dtsR1 disruption (CCM03), L-Arg production drastically increased to 27.45 g/L and then further to 29.97 g/L after Tween 40 trigger. Moreover, the specific activity of α-oxoglutarate dehydrogenase complex (ODHC) decreased, whereas the regeneration of NADP+/NADPH significantly increased after dtsR1 disruption and Tween 40 trigger. Results of real-time PCR showed that the transcriptional levels of odhA, sucB, and ipdA (encoding three subunits of the ODHC complex) were downregulated after Tween 40 trigger or dtsR1 disruption. By contrast, zwf transcription (encoding glucose-6-phosphate dehydrogenase) showed no significant difference among CCM01, CCM02 (ΔNCgl1221), and CCM03 (ΔNCgl1221ΔdtsR1) strains without Tween 40 trigger but evidently increased by 5.50 folds after Tween 40 trigger.

Conclusion: A novel strategy for L-Arg overproduction by dtsR1 disruption and Tween 40 trigger in C. crenatum was reported. Tween 40 addition exhibited a bifunctional mechanism for L-Arg overproduction, including reduced ODHC activity and enhanced NADPH pools accumulation by downregulated dtsR1 expression and upregulated zwf expression, respectively.

Keywords: Corynebacterium crenatum, L-Arginine, Tween 40, DtsR1, α-Oxoglutarate dehydrogenase complex, NADP+/NADPH

Background

L-Arginine (L-Arg) is a semi-essential amino acid that is widely used as an additive in food, cosmetic, and pharmaceutical industries because of its ability to promote secretion of growth hormones [1], insulin [2], and prolactin [3] and facilitate the synthesis of various immune active factors to prevent cancer cell growth [4]; L-Arg is a nitric oxide precursor for relaxing and dilating blood vessels [5]. Different microorganisms, such as Corynebacterium glutamicum [6, 7], Escherichia coli [8], Bacillus subtilis [9], and Saccharomyces cerevisiae [10], are used as model organisms for L-Arg overproduction.

In recent decades, various strategies based on genetic engineering technology have been designed to improve industrial levels of L-Arg. Ginesy et al. [8] reported...
an engineered *E. coli* strain for l-Arg overproduction by deleting the *speC, speF, argA, addA*, and *argR* genes, introducing feedback-resistant *argA214*, and overexpressing *argO* genes, whose l-Arg final production was achieved at 11.64 g/L in 1-L batch fermentation. Xu et al. [11] also performed site-directed mutagenesis of *N*-acetyl-l-glutamate kinase (including E19R, H26E, and H268D) to alleviate feedback inhibition by l-Arg; l-Arg production improved by about 41.7 % as compared with that of the initial strain. Moreover, Xu et al. overexpressed l-Arg biosynthetic genes, including the *argN-H* cluster [12] and the *lysE* gene [13] in *Corynebacterium crenatum* SYPA 5–5; l-Arg production was achieved at 45.3 and 35.9 g/L, respectively, by batch fermentation in a 5-L bioreactor. Lee et al. [14, 15] reported a metabolically engineered *C. glutamicum* for production of l-Arg at the industrial-scale based on systems metabolic engineering, including random mutagenesis to release feedback inhibition, knocking out *argR* and *farR* genes to remove repressors, increasing NADPH and carbamoyl phosphate pools, and deleting *NCgl1221* gene to avoid l-glutamate exporter. Subsequently, l-Arg production distinctly increased to 92.5 g/L by fed-batch fermentation in a 5-L bioreactor.

l-Glutamate is an important precursor in the l-Arg biosynthetic pathway. Previous research has demonstrated that reduced α-oxoglutarate dehydrogenase complex (ODHC) activity can significantly induce l-glutamate overproduction in *C. glutamicum* by increasing metabolic fluxes toward l-glutamate synthesis (as shown in Fig. 1). As a fermentation trigger, polyoxyethylene sorbitan monopalmitate (Tween 40) is widely used to enhance l-glutamate overproduction in *C. glutamicum* because it can decrease ODHC activity by downregulation of *dtsR1* expression [16]. DtsR1 protein is a homolog of the subunit of the biotin enzyme acetyl-CoA carboxylase complex, and disruption of the *dtsR1* gene can notably reduce ODHC activity, leading to an increase in l-glutamate production [17, 18]. However, to our knowledge, the strategy by adding Tween 40 and deleting *dtsR1* for l-Arg overproduction has not been published. *C. crenatum* is highly homologous to *C. glutamicum* and is frequently used for production of various amino acids, including l-Arg, because of its genetic tractability,

![Diagram](image.png)

**Fig. 1** Correlation biosynthetic pathway of l-Arg and possible signaling cascade involved in the regulation of ODHC specific activity under Tween 40 addition by *C. crenatum* (Brief). Red thick arrows indicate increased fluxes by overexpressing the corresponding genes. Dot-dashed lines represent decreased or cut off fluxes by reducing relevant enzyme activity and knocking out relevant genes, respectively. G-6-P glucose-6-phosphate, ACT-CoA acetyl-CoA, Zwf glucose-6-phosphate dehydrogenase, Gnd 6-phosphogluconate dehydrogenase, Pro8 gamma-glutamyl kinase, PutP l-proline exporter, *NCgl1221* glutamate exporter, *LysE* l-Arg and l-lysine exporter, *N-ACTGLU* N-acetylglutamate, *N-ACTORN* N-acetylornithine, *argBCDF-argGH*, the gene clusters of l-Arg biosynthetic pathway, DtsR1 acetyl-CoA carboxylase, AccBc biotin carboxylase and biotin carboxyl carrier protein, PknG serine/threonine protein kinase, Ppp protein phosphatase, OdhI signal transduction protein, FHA-domain-containing protein, OdhA α-oxoglutarate dehydrogenase, LpdA dihydrolipoamide dehydrogenase, SucB dihydrolipoamide acetyltransferase.
bio-safety, and robustness in fermentation [19, 20]. In this work, C. crenatum argB-M4 ΔproB, whose argR was lethal via mutagenesis screening, proB gene was deleted, and argB gene was replaced by C. crenatum argB M4 gene (E19R, H26E, D311R, and D312R) to release l-Arg feedback inhibition, was chosen as a host strain to study l-Arg production. Furthermore, NCgl1221 (encoded as l-glutamate exporter) and dtsR1 genes were knocked out using a marker-free system to increase l-Arg precursor. l-Arg production, ODHC specific activity, and NADPH pools accumulation by Tween 40 trigger and dtsR1 gene regulation were explored, and the related genes transcription in response to inducing l-Arg production was investigated to elaborate the relationship among dtsR1 gene deletion, Tween 40 addition, and l-Arg overproduction.

Results and discussion

Effect of Tween 40 and DtsR1 on l-Arg production in C. crenatum

Tween 40, as a fermentation trigger, can induce l-glutamate overproduction in C. glutamicum [21, 22]. To elaborate the effect of Tween 40 on l-Arg production in C. crenatum, we optimized Tween 40 addition and addition time in CCM01 strain. As shown in Fig. 2a, the biomass of C. crenatum gradually decreased with increasing concentration of Tween 40 from 0 to 8.0 mg/mL, and the l-Arg production reached a maximum value when Tween 40 concentration was 5 mg/mL in the fermentation medium. The addition time of Tween 40 on l-Arg production is shown in Fig. 2b, indicating that l-Arg overproduction could not be induced by Tween 40 addition before the 24 h fermentation time. During the logarithmic period (after 36 h fermentation time), Tween 40 addition significantly increased l-Arg production from 15.22 to 17.73 g/L (P < 0.05), while the yield of l-Arg per gram biomass was increased from 1.54 to 2.24 g/g. Moreover, the yield of l-Arg per gram glucose increased from 0.22 to 0.30 g/g, consequently increasing l-Arg productivity from 0.13 to 0.15 g/L/h in CCM01 strain (Table 1). The above results indicate that adding Tween 40 at the logarithmic period is conducive to l-Arg production. Meanwhile, the concentrations of other amino acids were analyzed using an amino acid analyzer. The results shown in Fig. 3 indicated that a large amount of l-glutamate was secreted into the fermentation supernatant (4.72 g/L). To avoid l-glutamate leak during l-Arg fermentation, NCgl1221 gene encoding l-glutamate transporter [23, 24] was removed from the genome of CCM01 strain (named CCM02) to promote l-Arg production. As expected, extracellular l-glutamate in CCM02 strain decreased to an undetectable level (Fig. 3), whereas l-Arg production increased to 19.56 g/L after Tween 40 trigger. Although Tween 40 addition increased l-Arg production in C. crenatum, the increased yield of l-Arg is very limited. A previous study has confirmed that disruption of dtsR1 can convert the metabolic fluxes from tricarboxylic acid cycle toward glutamate synthesis by repressing ODHC specific activity [18]. To further improve l-Arg production, we knocked out the dtsR1 gene in CCM02 strain (named CCM03). The results showed that extracellular l-Arg production in CCM03 strain drastically increased to 27.45 g/L, whereas l-lysine production evidently declined by 39.56 % to 1.06 g/L compared with 1.75 g/L in CCM01 + Tween 40 condition (Fig. 3). The yield of l-Arg per gram glucose increased to 0.34 g/g, improving by 54.55 % as compared with that in CCM01 strain (Table 1). Although l-Arg production after DtsR1 disruption is still low compared with the highest level.

Fig. 2 Effect of Tween 40 addition (0–8 mg/mL Tween 40, after 36 h incubation time) and addition time (0–48 h, 5 mg/mL Tween 40) on l-Arg production in CCM01 strain. l-Arg production and the DCW were monitored during shake flask cultivation for 120 h by Tween 40 addition. Results are the means ± standard deviations from three independent experiments. Compared with the control group, “ns” designates P > 0.05, *P < 0.05; **P < 0.01; ***P < 0.001.
(92.5 g/L) of that reported by Lee et al. [14, 15], it is notable that the yield of L-Arg per gram glucose is basically consistent with the previous report, suggesting the CCM03 strain exhibits great potential to improve the yield of L-Arg by fed-batch fermentation. In addition, L-Arg production could further improve to approximately 9.12 % (29.97 g/L) in CCM03 strain after Tween 40 addition. To further elaborate the function of Tween 40 and DtsR1 protein on L-Arg overproduction in C. crenatum, we investigated ODHC specific activity and NADPH pools accumulation.

**Table 1 Comparison of the performances of the different conditions for L-Arg production by fermentation**

| Conditions          | GlcC (g/L) | DCW (g/L) | Arginine (g/L) | Yield   | Vp (g/L/h) |
|---------------------|------------|-----------|----------------|---------|------------|
|                     |            |           |                | Y<sub>A/G</sub> (g arg/g glc) | Y<sub>A/D</sub> (g arg/g DCW) |
| CCM01               | 70.32 ± 3.35 | 9.92 ± 0.35 | 15.22 ± 0.27 | 0.22 ± 0.01 | 1.54 ± 0.01 | 0.13 ± 0.03 |
| CCM01 + Tween 40    | 60.33 ± 2.47 | 7.91 ± 0.42 | 17.73 ± 0.18 | 0.30 ± 0.02 | 2.24 ± 0.03 | 0.15 ± 0.04 |
| CCM02               | 60.96 ± 1.05 | 8.34 ± 1.07 | 17.58 ± 0.34 | 0.29 ± 0.04 | 2.11 ± 0.02 | 0.15 ± 0.04 |
| CCM02 + Tween 40    | 59.37 ± 2.91 | 7.36 ± 0.27 | 19.56 ± 0.19 | 0.33 ± 0.03 | 2.68 ± 0.02 | 0.16 ± 0.02 |
| CCM03               | 80.98 ± 1.05 | 8.17 ± 1.07 | 27.45 ± 0.58 | 0.34 ± 0.05 | 3.38 ± 0.13 | 0.23 ± 0.02 |
| CCM03 + Tween 40    | 89.34 ± 3.91 | 7.36 ± 0.27 | 29.97 ± 1.13 | 0.34 ± 0.09 | 4.10 ± 0.15 | 0.25 ± 0.05 |

Fermentations were performed in a 250 ml flask at 30 °C; the initial glucose concentration was 120 g/L. Results are the means ± standard deviations in three independent experiments.

GlcC: glucose consumed, Y<sub>A/G</sub> arginine yield vs. glucose, Y<sub>A/D</sub> arginine yield vs. DCW, Vp: volumetric productivity.

**Effect of Tween 40 and DtsR1 on ODHC specific activity and NADPH pools accumulation in C. crenatum**

ODHC is a branch-point enzyme complex between the tricarboxylic acid cycle and L-glutamate biosynthesis. Tween 40 addition can indirectly convert the metabolic fluxes into the L-glutamate biosynthetic pathway by downregulating dtsR1 gene expression; the disruption of dtsR1 gene can reduce the ODHC activity, thereby inducing L-glutamate overproduction in C. glutamicum. In addition, L-glutamate production induced by Tween 40 addition can completely be suppressed by dtsR1 gene.
that a large amount of NADPH pools for fatty acid biosynthesis (Table 2). The above results suggest that probably other mechanisms exist for l-Arg overproduction induced by Tween 40 trigger after dsr1 disruption in C. crenatum.

A recent study has confirmed that NADPH pools accumulation is also conducive to l-Arg production because the biosynthesis of 1 mol of l-Arg requires 3 mol of NADPH [14, 26]. DtsR1 protein is one of the subunits of acetyl-CoA carboxylase complex, which is involved in fatty acid biosynthesis. Fatty acid biosynthesis requires a large amount of NADPH pools for fatty acid elongation cycle [27]. After DtsR1 disruption, the intracellular level of NADPH notably increased to 0.59 mmol/L in CCM03 strain, which was approximately three times higher than that of CCM01 strain (Table 2). The above results suggest that a large amount of NADPH pools for fatty acid biosynthesis was converted for l-Arg biosynthesis. We also found that the NADPH level could further increase upon Tween 40 addition in CCM01, CCM02, and CCM03 strains. To date, relevant research about Tween 40 as a trigger for NADPH pools accumulation has not been reported.

Regulation mechanism of Tween 40 and DtsR1 on l-Arg overproduction

As shown in Fig. 4 and Table 2, l-Arg overproduction by Tween 40 trigger or DtsR1 disruption is attributed to reduced ODHC enzymatic activity and NADPH pools accumulation. ODHC complex consists of three subunits, including α-oxoglutarate dehydrogenase encoded by odhA gene [28], dihydrolipoamide S-succinyltransferase encoded by sucB gene, and dihydrolipoamide dehydrogenase encoded by lpdA gene [29]. A recent study has confirmed that ODHC activity regulation is not only related to the expression level of DtsR1 protein [17] but also with the phosphorylated and unphosphorylated forms of OdhI protein [30, 31]. The downregulated expression of DtsR1 protein is in favor of reducing ODHC activity, and unphosphorylated OdhI is also as an inhibitor of ODHC because the FDH domain of unphosphorylated OdhI can combine with the OdhA subunit inhibiting ODHC activity [32]. OdhI protein can be phosphorylated by serine/threonine protein kinases, including PknG and PknB, and dephosphorylated by phosphoserine/threonine protein phosphatase Ppp [33, 34]. In addition, intracellular NADPH is mainly generated from the pentose phosphate pathway, in which the gene cluster of tkt–tal–zwf–opcA–pgl is involved. Among them, glucose-6-phosphate dehydrogenase (encoded by zwf gene) is responsible for regeneration of NADP⁺ to NADPH [35]. To elucidate the regulation mechanism of Tween 40 and DtsR1 for l-Arg overproduction, we investigated the transcriptional levels of dsr1, ppp, pknG, odhA, sucB, lpdA, zwf, and argB, which is involved in the arginine biosynthetic pathway and is located in the gene cluster of argCJBDF-argGH, by real-time quantitative PCR method.

Table 2 Intracellular NADP⁺ and NADPH concentrations in different conditions

| Conditions         | NADPH (mmol/L) | NADP⁺ (mmol/L) | NADP⁺/NADPH |
|--------------------|----------------|----------------|-------------|
| CCM01              | 0.13 ± 0.01    | 0.25 ± 0.02    | 1.89        |
| CCM01 + Tween 40   | 0.30 ± 0.04    | 0.45 ± 0.02    | 1.49        |
| CCM02              | 0.15 ± 0.03    | 0.28 ± 0.04    | 1.90        |
| CCM02 + Tween 40   | 0.45 ± 0.02    | 0.68 ± 0.05    | 1.51        |
| CCM03              | 0.59 ± 0.03    | 0.78 ± 0.01    | 1.32        |
| CCM03 + Tween 40   | 0.75 ± 0.18    | 0.98 ± 0.05    | 1.31        |

Results are the means ± standard deviations in three independent experiments.
The results shown in Fig. 5 indicate that dtsR1 and pknG were downregulated by 3.12- and 3.14-fold, respectively, whereas ppp was upregulated by 2.80-fold in CCM01/02 strain after Tween 40 trigger, and the transcriptional levels of odhA, sucB, and lpdA were concomitantly downregulated by 3.20-, 2.47-, and 2.19-fold, respectively, which were consistent with the previous report by Kataoka et al. [36]. We also found that the expression level of argB was upregulated by 17.25-fold after removing the NCgl1221 gene in CCM02 strain. We suspected that the high expression of argB gene is associated with intracellular l-glutamate accumulation. After dtsR1 disruption, the transcriptional level of pknG notably downregulated by 6.00-fold, and ppp transcription was upregulated by 8.53-fold, which resulted in lesser transcriptional levels of odhA, sucB, and lpdA in CCM03 strain compared with those in CCM01 strain. Meanwhile, the expression levels of pknG, ppp, odhA, sucB, and lpdA remained unaltered in CCM03 strain after Tween 40 trigger. The above results demonstrated that reducing ODHC enzymatic activity induced by Tween 40 could be completely suppressed by dtsR1 disruption in C. crenatum. We also found that zwf transcription showed no significant difference among CCM01, CCM02, and CCM03 strains (P > 0.05). The above results indicated that NCgl1221 and dtsR1 expression were not associated with NADP+/NADPH regeneration, and the increase in intracellular NADPH level after DtsR1 deletion could come from fatty acid synthesis. However, after adding Tween 40, the transcriptional level of zwf gene increased by 5.50-fold in CCM01, CCM02, and CCM03 strains. This result suggests that Tween 40 could be a key trigger in inducing NADPH regeneration for l-Arg overproduction.

Conclusion

A novel strategy for overproducing l-Arg by reduction of ODHC enzymatic activity and promotion of NADPH accumulation in C. crenatum was reported. To avoid intracellular l-glutamate secretion and reduce ODHC specific activity, we successfully deleted NCgl1221 and dtsR1 by using a marker-free system in C. crenatum. l-Arg production significantly increased from 15.22 to 27.45 g/L, and l-Arg production further enhanced by 29.97 g/L after Tween 40 trigger. We also, for the first time, found that Tween 40 addition not only downregulated dtsR1 expression but also induced NADP+/NADPH regeneration by upregulating zwf expression. However, further experiments, such as transcriptomics,

![Fig. 5 Relative transcription level of genes possible involved in the regulation of ODHC specific activity and of genes involved in the biosynthetic pathway of l-Arg triggered by Tween 40 addition and dtsR1 disruption. Results are the means ± standard deviations from three independent experiments. Compared with the control group, "ns" designates P < 0.05.](image-url)
proteomics, and metabolic engineering, are necessary to elucidate the more detailed mechanism of L-Arg overproduction by Tween 40 trigger.

Methods
Strains, plasmids and primers
*Corynebacterium crenatum* argB-M4 ∆proB strain (named CCM01) used as a parent strain in this study was constructed in our previous work (data not shown), whose argR was lethal via mutagenesis screening, the proB gene was knocked out, and *argB* was replaced by *argB* M4 (E19R, H26E, D311R, and D312R) to release L-Arg feedback resistance. The strains and plasmids used in this study are listed in Table 3. The primers used for strain construction and RT-quantitative (q) PCR amplification are listed in Table 4.

Strain construction
*Escherichia coli* DH5α strain was used as a primary host for all gene cloning. *Corynebacterium crenatum* was grown in Luria–Bertani (LB) medium at 210 rpm and 30 °C, while *E. coli* was grown in LB medium at 180 rpm and 37 °C. Unless otherwise indicated, the concentration of Kanamycin was 25 μg/mL. CCM01 with deleted NCgl1221 gene (named CCM02) was obtained by knocking out NCgl1221 gene by using a marker-free system with *Bacillus subtilis* sacB gene via two rounds of recombination as described previously [37]. The fusion arms containing 5′-upstream and 3′-downstream fragments were obtained by overlapping PCR. These fusion arms were then used for construction of recombinant plasmid by inserting them into the pK18mobsacB vector. The resultant recombinant plasmid was transferred into *C. crenatum* by electroporation. The single crossover strains were selected on LBHIS (LB supplemented with brain heart infusion and sorbitol) agar plates containing 10 μg/mL kanamycin, whereas marker-free recombinants were screened on sucrose-resistant agar plates. Moreover, CCM02 with deleted *dtsR1* gene (named CCM03) underwent the same procedure as the previous strains.

L-Arg fermentation
The seed medium for *C. crenatum* cultivation (per liter) consisted of 30 g of glucose, 20 g of corn steep liquor, 20 g of (NH₄)₂SO₄, 1 g of KH₂PO₄, 0.5 g of MgSO₄·7H₂O, and 1.5 g of urea. For seed culture, cells on the LB medium plate were inoculated into a test tube containing 5 mL of the LB medium, and cultivated in a shaking incubator with 210 rpm at 30 °C for 24 h. One milliliter of the culture was transferred to a 250 mL flask containing 30 mL of the seed medium. After the optical density of seed culture (OD₅₆₂) was reached between 5.5 and 6.0, 2.0 mL seed culture was transferred into a 250 mL flask containing 25 mL the fermentation medium and cultured at 30 °C for 120 h. The fermentation medium (per liter) for L-Arg production was composed of 120 g of glucose, 25 g of corn steep liquor, 45 g of (NH₄)₂SO₄, 1 g of KH₂PO₄, 0.5 g of MgSO₄·7H₂O, and 30 g of CaCO₃ (pH 7.0). δ-Biotin was supplied from the corn steep liquor of the medium [20]. For the growth of CCM03 strain (∆proBDNSgl1221DΔtsR1), L-proline and oleic acid ester (Tween 80) were added to media with final concentrations of 2.5 and 1 mg/mL, respectively. Tween 40 was added to the medium to a final concentration of 5 mg/mL once cell growth reached the early exponential phase to further trigger L-Arg overproduction. Cell concentration was determined at 562 nm and measured by a pre-calibrated relationship (1 OD = 0.375 g/L). 3,5-Dinitrosalicylic acid colorimetry was used to measure glucose concentration in the culture [38]. Concentrations of L-Arg and other amino acids were determined using a Sykam S-433D amino acid analyzer (Sykam Co. Ltd., Germany). All the data are presented as means ± standard deviations in three independent experiments.

Table 3 Strains and plasmids used in this study

| Strains/plasmids | Function and relevant characteristics | Sources |
|------------------|--------------------------------------|---------|
| **Strains**      |                                      |         |
| DH5α             | General clone host strain            | Invitrogen |
| *C. crenatum* MT | Mutation strain with auxotrophic for biotin, and producing L-Arg | Lab stock |
| CCM01            | *C. crenatum* MT with proB gene deletion, which E-argB gene replaced by *C. crenatum* argB M4 gene (E19R, H26E, D311R, and D312R) | This work |
| CCM02            | Chromosomal deletion of the *Ncgl1221* gene of CCM01 | This work |
| CCM03            | Chromosomal deletion of the *dtsR1* gene of CCM02 | This work |
| **Plasmids**     |                                      |         |
| pK18mobsacB      | Mobilizable vector, allows for selection of double crossover in *C. crenatum*, Km®, sacB | Lab stock |
| pK18-Δ*Ncgl1221* | A derivative of pK18mobsacB, harboring Δ*Ncgl1221* fragment | This work |
| pK18-Δ*dtsR1*    | A derivative of pK18mobsacB, harboring Δ*dtsR1* fragment | This work |
Measurement of ODHC specific activity and concentrations of intracellular NADP⁺ and NADPH pools

Samples preparation and ODHC specific activity were conducted according to a previously described method [39, 40]. According to the previous work, the L-Arg production rate was kept highest during 36–60 h fermentation time (data not show), therefore, C. crenatum cells were harvested in the late exponential phase (after 48 h cultivation), and collected by centrifugation at 5,000 rpm at 4 °C for 10 min. The pellets were diluted with 0.2 M HCl to dissolve CaCO₃ and then washed twice with 0.2% KCl solution. The pure cells were suspended in 5 mL of 0.1 M N-tris(hydroxymethyl)methyl-2-aminoethanesulfonic acid (TES)·NaOH buffer (pH 7.7) containing 30% (v/v) glycerol and 10 mg/mL lysozyme. Following incubation at 37 °C for 3 h, the cells were disrupted by sonication and centrifuged to remove cell debris. The supernatant was collected, and the protein concentration was determined by Nanodrop 2000 (Thermo Scientific, Germany).

ODHC specific activity assay was performed by adding 100 µL of cell extract to 2.5 mL of reaction mixture. The reaction mixture contained 100 mM TES·NaOH buffer (pH 7.7), 5 mM MgCl₂, 3 mM cysteine, 0.3 mM thiamine pyrophosphate, 0.2 mM coenzyme A, and 1 mM 3-acetylpyridine adenine dinucleotide (APAD⁺). Upon adding 1 mM α-oxoglutarate to the reaction mixture, the initial increase in the absorbance of APADH at 365 nm was consecutively measured at 31.5 °C for 5 min with 30 s intervals. ODHC specific activity is defined as the amount of enzyme required to generate 1 µmol NADH per minute. The molar extinction coefficient of NAD⁺ was 9.1.

Intracellular NADP⁺ and NADPH concentrations were determined by enzymatic cycling reaction initiated with the EnzyChrom™ NADP⁺/NADPH Assay kit (BioAssay Systems, Hayward, CA 94545, USA). All the data are presented as the means ± standard deviations in three independent experiments.

Table 4 Sequences of oligonucleotide primers

| Primers   | Sequence (5'–3') | Purposes                      |
|-----------|------------------|-------------------------------|
| NCgl1221-up-F | CGCAAGCTTCAAGAAAGCCCTGTTCAAACACTG | Amplifying the left arm of NCgl1221 |
| NCgl1221-up-R | ATCACGCTCCTAAGGAATCAGAAAACGCCAAAGCCAGG | Amplifying the right arm of NCgl1221 |
| NCgl1221-down-F | GGCTTTTTTATCCCTAGGACGTGATACQCAGC | Amplifying the left arm of NCgl1221 |
| NCgl1221-down-R | GCCATCTAGAGAGGGGGATTGGTAGC | Amplifying the right arm of NCgl1221 |
| dtsR1-up-F | CGCAAGCTTCAAGAAAGCCCTGTTCAAACACTG | Amplifying the left arm of dtsR1 |
| dtsR1-up-R | GCCATCTAGAGAGGGGGATTGGTAGC | Amplifying the right arm of dtsR1 |
| dtsR1-down-F | CGCAAGCTTCAAGAAAGCCCTGTTCAAACACTG | Amplifying the left arm of dtsR1 |
| dtsR1-down-R | GCCATCTAGAGAGGGGGATTGGTAGC | Amplifying the right arm of dtsR1 |
| odhA-F | TATGAGCATATCG | RP of odhA |
| odhA-R | TCAGTGCTCAGAT | RP of odhA |
| lpdA-F | TGGGACTTAAACGGCCTGCT | RP of lpdA |
| lpdA-R | CACACGTTTAATCATCGGGAAC | RP of lpdA |
| sucB-F | TGTCAGAAGCGTGCCAGTTC | RP of sucB |
| sucB-R | GGCTCTCTGATTGTTGTCAGG | RP of sucB |
| pknG-F | GCCGCTGATGGTTGATTGATT | RP of pknG |
| pknG-R | GCCGCTGATGGTTGATTGATT | RP of pknG |
| ppp-F | CGATTGTTAGCG | RP of ppp |
| ppp-R | CCGGACCTAAAAGTGTGTT | RP of ppp |
| dtsR1-F | CCTCCATACCAAAATTTGGCT | RP of dtsR1 |
| dtsR1-R | TAAAGAAGGGCTCCGAGTAC | RP of dtsR1 |
| zwf-F | ACTGAGATCTGGTGGTTTGA | RP of zwf |
| zwf-R | ACGCGATAGCCACAACCTTC | RP of zwf |
| argB-F | GTCGGATTTGATGGTTGATTGAT | RP of argB |
| argB-R | GAGGCATGACATTAGGATTGAT | RP of argB |
| 16s rRNA-F | AAGAAGCACCAGCCTAACTAC | Reference Gene |
| 16s rRNA-R | CCGGCTTACGACAGCAGAC | Reference Gene |

Restriction sites were highlighted in bold; linker sequences for crossover PCR were shown in underline. RP RT-qPCR.
Real-time fluorescence reverse transcription quantitative PCR
The total RNA from CCM01, CCM01 + Tween 40, CCM02 + Tween 40, CCM03, and CCM03 + Tween 40 conditions after 48 h fermentation was extracted using a Trizol Plus RNA Purification Kit according to the manufacturer’s instructions (Invitrogen™, USA). cDNA was synthesized by the Primer Script™ RT Reagent Kit with gDNA Eraser (TaKaRa, Japan). The transcription levels of the argB, zwf, odhA, lpdA, sucB, dtsR1, pknG, and pgo genes were analyzed by the real-time fluorescence reverse transcription quantitative PCR (RT-qPCR) using SYBR® Premix Taq™ (Tli RNaseH Plus) Kit (TaKaRa, Japan) with the corresponding primers (listed in Table 4). 16S rRNA was adopted as a reference gene. The thermal cycling conditions were 95 °C for 30 s, followed by 40 cycles of 95 °C for 5 s and 60 °C for 30 s. A control reaction without template was implemented to evaluate primer dimmer formation. The relative transcription level for each gene was calculated through the 2−ΔΔCt method [41]. All the data are presented as the means ± standard deviations in three independent experiments.

Statistical analysis
The data were statistically compared using ANOVA, and significant differences were identified by Tukey- test (P < 0.05). These analyses were carried out in GraphPad Prism 5.0 software (GraphPad software Inc, California, USA).

Authors’ contributions
XYH and CXL conceived the study, participated in its design and coordination. CML designed and carried out the molecular genetic work, performed the analysis, and drafted the manuscript. WF carried out the fermentation experiments. ZB and CJC analyzed the data. All authors read and approved the final manuscript.

Acknowledgements
This work was supported by the Natural Science Foundation of China (No. 31360219 and No. 30960012).

Compliance with ethical guidelines
Competing interests
The authors declare that they have no competing interests.

Received: 22 May 2015 Accepted: 4 August 2015
Published online: 12 August 2015

References
1. Alba-Roth J, Müller OA, Schopohl J, Werder KV (1988) Arginine stimulates growth hormone secretion by suppressing endogenous somatostatin secretion. J Clin Endocrinol Metab 67:1186–1189
2. Thams P, Capito K (1999) l-Arginine stimulation of glucose-induced insulin secretion through membrane depolarization and independent of nitric oxide. Eur J Endocrinol 140:87–93
3. Davis S (1972) Plasma levels of prolactin, growth hormone, and insulin in sheep following the infusion of arginine, leucine and phenylalanine. Endocrinology 91:549–555
4. Bronte V, Zanovello P (2005) Regulation of immune responses by l-arginine metabolism. Nat Rev Immunol 5:641–654
5. Igaruo LJ, Cirim G, Casini A, Napoli C (1999) Nitric oxide as a signaling molecule in the vascular system: an overview. J Cardiovasc Pharmacol 34:879–886
6. Nakayama K, Yoshida H (1972) Fermentative production of l-arginine. Agric Biol Chem 36:1675–1684
7. Ramirez MM, Qin QZ, Dou W, Zhiming R, Xu Z (2013) Over-expression of NAD kinase in Corynebacterium crenatum and its impact on l-arginine biosynthesis. Trop J Pharm Res 11:909–916
8. Ginesy M, Belotserkovskiy J, Enman J, Isaksson L, Rova U (2015) Metabolic engineering of Escherichia coli for enhanced arginine biosynthesis. Microb Cell Fact 14:29
9. Kissumi M, Kato J, Sugimura M, Chibata I (1971) Production of l-arginine by arginine-hydroxamate-resistant mutants of Bacillus subtilis. Appl Microbiol 22:987–991
10. Yoshida H, Araki K, Nakayama K (1981) l-arginine production by arginine analog-resistant mutants of microorganisms. Agric Biol Chem 45:959–963
11. Xu M, Rao Z, Yang J, Jin J, Xu Z (2012) Site-directed mutagenesis and feedback-resistant N-acetyl-l-glutamate kinase (NAGK) increase Corynebacterium crenatum l-arginine production. Amino Acids 43:255–266
12. Xu M, Rao Z, Yang J, Xia H, Dou W, Jin J et al (2012) Heterologous and homologous expression of the arginine biosynthetic argC → H cluster from Corynebacterium crenatum for improvement of l-arginine production. J Ind Microbiol Biotechnol 39(3):495–502
13. Xu M, Rao Z, Yang J, Dou W, Xu Z (2013) The effect of a LYSE exporter overexpression on l-arginine production in Corynebacterium crenatum. Curr Microbiol 67(3):271–278
14. Park SH, Kim HU, Kim TY, Park JS, Kim S-S, Lee SY (2014) Metabolic engineering of Corynebacterium glutamicum for l-arginine production. Nat Commun 5:4618
15. Shin JH, Lee SY (2014) Metabolic engineering of microorganisms for the production of l-arginine and its derivatives. Microb Cell Fact 13:66
16. Kimura E (2002) Triggering mechanism of Glutamate overproduction in Corynebacterium glutamicum. J Microbiol Biotechnol 34:879–886
17. Kimura E, Usuda Y, Kawahara Y, Matsui K, Osumi T et al (2007) Microbial cell factory technology. Microb Cell Fact 14:29
18. Yao W, Deng X, Zhong H, Liu M, Zheng P, Sun Z et al (2009) Double deletion of dtsR1 and pgo induce efficient l-glutamate overproduction in Corynebacterium glutamicum. J Ind Microbiol Biotechnol 36(7):911–921
19. Xu H, Dou W, Xu H, Zhang X, Rao Z, Shi Z et al (2009) A two-stage oxygen supply strategy for enhanced l-arginine production by Corynebacterium glutamicum based on metabolic fluxes analysis. Biotechnol Bioeng 103(1):41–51
20. Dou W, Xu M, Cai D, Zhang X, Rao X, Xu Z (2011) Improvement of l-arginine production by overexpression of a bifunctional ornithine acetyltransferase in Corynebacterium glutamicum. Appl Biochem Biotechnol 165(34):845–855
21. Shimizu H, Tanaka H, Nakato A, Nagahisa K, Kimura E, Shioya S (2003) Effects of the changes in enzyme activities on metabolic flux redistribution around the 2-oxoglutarate branch in glutamate production by Corynebacterium glutamicum. Bioprocess Biosyst Eng 25(5):291–298
22. Asakura Y, Kimura E, Usuda Y, Kawahara Y, Matsui K, Osumi T et al (2007) Altered metabolic flux due to deletion of cdaA causes l-glutamate overproduction in Corynebacterium glutamicum. Appl Environ Microbiol 73(4):1308–1319
23. Nakamura J, Hirano S, Ito H, Wachi M (2007) Mutations of the Corynebacterium glutamicum NCgl1221 gene, encoding a mechanosensitive channel homolog, induce \(L\)-glutamic acid production. Appl Environ Microbiol 73:4491–4498

24. Hashimoto K, Nakamura K, Kuroda T, Yabe I, Nakamatsu T, Kawasaki H (2010) The protein encoded by NCgl1221 in Corynebacterium glutamicum functions as a mechanosensitive channel. Biosci Biotechnol Biochem 74(12):2546–2549

25. Kimura E, Abe C, Kawahara Y, Nakamatsu T, Tokuda H (1997) A dtsR gene-disrupted mutant of Brevibacterium lactofermentum requires fatty acids for growth and efficiently produces \(L\)-glutamate in the presence of an excess of biotin. Biochem Bioph Res Commun 234(1):157–161

26. Takeno S, Murata R, Kobayashi R, Mitsuhashi S, Ikeda M (2010) Engineering of Corynebacterium glutamicum with an NADPH-generating glycolytic pathway for \(L\)-lysine production. Appl Environ Microbiol 76:7154–7160

27. Lennen RM, Pfleger BF (2012) Engineering Escherichia coli to synthesize free fatty acids. Trends Biotechnol 30(12):659–667

28. Usuda Y, Tujimoto N, Abe C, Asakura Y, Kimura E, Kawahara Y et al (1996) Molecular cloning of the Corynebacterium glutamicum ('Brevibacterium lactofermentum') AJ12036) odhA gene encoding a novel type of 2-oxoglutarate dehydrogenase. Microbiology 142:3347–3354

29. Schwinde JW, Hertz PF, Sahm H, Eikmanns BJ, Guyonvarch A (2001) Lipoamide dehydrogenase from Corynebacterium glutamicum: molecular and physiological analysis of the lpd gene and characterization of the enzyme. Microbiology 147:2223–2231

30. Krawczyk S, Raasch K, Schultz C, Hoffelder M, Eggeling L, Bott M (2010) The FHA domain of OdhI interacts with the carboxyterminal 2-oxoglutarate dehydrogenase domain of OdhA in Corynebacterium glutamicum. FEBS Lett 584:1463–1468

31. Raasch K, Boccola M, Labahn J, Leitner A, Eggeling L, Bott M (2014) Interaction of 2-oxoglutarate dehydrogenase OdhA with its inhibitor OdhI in Corynebacterium glutamicum: mutants and a model. J Biotechnol 191:99–105

32. Niebisch A, Kabus A, Schultz C, Weil B, Bott M (2006) Corynebacterial protein kinase G controls 2-oxoglutarate dehydrogenase activity via the phosphorylation status of the OdhI protein. J Biol Chem 281(18):12300–12307

33. Schultz C, Niebsch A, Schwaiger A, Viets U, Metzger S, Bramkamp M et al (2009) Genetic and biochemical analysis of the serine/threonine protein kinases PknA, PknB, PknG and PknL of Corynebacterium glutamicum: evidence for non-essentiality and for phosphorylation of OdhI and FtsZ by multiple kinases. Mol Microbiol 74(3):724–741

34. Schultz C, Niebsch A, Gebel L, Bott M (2007) Glutamate production by Corynebacterium glutamicum: dependence on the oxoglutarate dehydrogenase inhibitor protein OdhI and protein kinase PknG. Appl Microbiol Biotechnol 76(3):691–700

35. Liu Z, Chen L, Hao N, Xu L, Li Y, Yan M et al (2015) Expression of glucose-6-phosphate dehydrogenase and 6-phosphogluconate dehydrogenase improve \(\alpha\)-citrulline biosynthesis in argG-deleted Corynebacterium glutamicum. In: Advances in applied biotechnology, vol 333. Springer, Berlin, pp 197–204

36. Kataoka M, Hashimoto KI, Yoshida M, Nakamatsu T, Horinouchi S, Kawasaki H (2006) Gene expression of Corynebacterium glutamicum in response to the conditions inducing glutamate overproduction. Lett Appl Microbiol 42:471–476

37. Xu J, Xia X, Zhang J, Guo Y, Qian H, Zhang W (2014) A method for gene amplification and simultaneous deletion in Corynebacterium glutamicum genome without any genetic markers. Plasmid 72:9–17

38. Miller GL (1959) Use of dinitrosalicylic acid reagent for determination of reducing sugar. Anal Chem 31:426–428

39. Shiio I, Ujigawa-Takeda K (1980) Presence and regulation of \(\alpha\)-ketoglutarate dehydrogenase complex in a glutamate-producing bacterium, Brevibacterium flavum. Agric Biol Chem 44(8):1897–1904

40. Kim J, Hirasa S, Tato Y, Nagahisa K, Furusawa C, Shimizu H (2009) Effect of odhA overexpression and odhA antisense RNA expression on Tween-40-triggered glutamate production by Corynebacterium glutamicum. Appl Microbiol Biotechnol 81(6):1097–1106

41. Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2(−\(\Delta\Delta C\)) method. Methods 25:402–408