GREACE-assisted adaptive laboratory evolution in endpoint fermentation broth enhances lysine production by *Escherichia coli*

Xiaowei Wang¹,²,³†, Qinggang Li²,³†, Cunmin Sun²,³, Zhen Cai⁴, Xiaomei Zheng²,³, Xuan Guo²,³, Xiaomeng Ni²,³, Wenjuan Zhou²,³, Yanmei Guo³, Ping Zheng²,³*, Ning Chen¹, Jibin Sun²,³*, Yin Li⁴ and Yanhe Ma³

**Abstract**

**Background:** Late-stage fermentation broth contains high concentrations of target chemicals. Additionally, it contains various cellular metabolites which have leaked from lysed cells, which would exert multifactorial stress to industrial hyperproducers and perturb both cellular metabolism and product formation. Although adaptive laboratory evolution (ALE) has been wildly used to improve stress tolerance of microbial cell factories, single-factor stress condition (i.e. target product or sodium chloride at a high concentration) is currently provided. In order to enhance bacterial stress tolerance to actual industrial production conditions, ALE in late-stage fermentation broth is desired. Genome replication engineering assisted continuous evolution (GREACE) employs mutants of the proofreading element of DNA polymerase complex (DnaQ) to facilitate mutagenesis. Application of GREACE coupled-with selection under stress conditions is expected to accelerate the ALE process.

**Results:** In this study, GREACE was first modified by expressing a DnaQ mutant KR5-2 using an arabinose inducible promoter on a temperature-sensitive plasmid, which resulted in timed mutagenesis introduction. Using this method, tolerance of a lysine hyperproducer *E. coli* MU-1 was improved by enriching mutants in a lysine endpoint fermentation broth. Afterwards, the KR5-2 expressing plasmid was cured to stabilize acquired genotypes. By subsequent fermentation evaluation, a mutant RS3 with significantly improved lysine production capacity was selected. The final titer, yield and total amount of lysine produced by RS3 in a 5-L batch fermentation reached 155.0 ± 1.4 g/L, 0.59 ± 0.02 g lysine/g glucose, and 605.6 ± 23.5 g, with improvements of 14.8%, 9.3%, and 16.7%, respectively. Further metabolomics and genomics analyses, coupled with molecular biology studies revealed that mutations SpeBA302V, AtpBS165N and SecYM145V mainly contributed both to improved cell integrity under stress conditions and enhanced metabolic flux into lysine synthesis.

**Conclusions:** Our present study indicates that improving a lysine hyperproducer by GREACE-assisted ALE in its stressful living environment is efficient and effective. Accordingly, this is a promising method for improving other valuable chemical hyperproducers.

**Keywords:** Lysine production, *Escherichia coli*, Adaptive laboratory evolution, GREACE, Fermentation broth
Background

Microbial strains are core element of industrial biotechnology. The rapid development of rational design methods and high-throughput screening assisted evolution systems have provided high efficiency strains for diverse industrial purposes [1–8]. Hyperproducers have been constructed to generate a large number of industrial chemicals including bulk amino acids, organic acids and other commodities, wherein many are present in over 100 g/L in final fermentation broths [9–13]. L-lysine, for example, is a microbial product predominantly used as food and animal feed additive, with approximately 2.4 million tons produced worldwide in 2015 [5]. Continuous efforts had been made to improve the production strains over the past several decades [1, 5, 14–18]. To date, the best reported lysine producers, *Escherichia coli* and *Corynebacterium glutamicum*, produced 136.5 g/L and 181.5 g/L lysine with different fermentation strategies, respectively [12, 17].

The environmental conditions at the late fermentation stage, such as in endpoint fermentation broth (EFB), is unfavourable for cell survival and metabolic activity. In addition to target chemicals produced at high concentrations, large amounts of supplements such as pH conditioners [9, 11, 12] and culture medium components such as (NH₄)₂SO₄ for lysine production [12, 17] are usually added to the broth. In a previous lysine fed-batch fermentation, 136.5 g/L lysine was produced by an *E. coli* strain MU-1, and about 28 g/L NH₃·H₂O and 57 g/L (NH₄)₂SO₄ were required to maintain the pH and provide ammonium for lysine biosynthesis [17]. High concentrations of chemicals result in high osmotic stress, which severely disturbs cell function and frequently leads to cell membrane damage [19–21]. In addition, a variety of extracellular molecules such as metal ions, organic acids, amino acids, sugars, nucleic acids, dipeptides and phospholipids are released into the broth due to cell lysis [22–24], which can perturb cellular metabolism. Therefore, host cells have to suffer complex stresses in late stage broths [19–26]. Increasing cell tolerance should be beneficial for improving production capacity.

Bacteria have naturally evolved multiple mechanisms to cope with environmental stress, such as increased synthesis of aquaporin to accelerate water export [27], regulated expression of potassium transporters to increase intracellular potassium concentration [28], expression of mechanosensitive channels [29], accumulation of osmo-protectants like glycine betaine, trehalose, ectoine and proline [30–32], and enhancement of cytoplasmic membrane stability [21]. However, these naturally evolved mechanisms are generally inadequate to prevent the strains from dramatically enhanced complex and combinatorial stress encountered in industrial conditions.

Adaptive laboratory evolution (ALE) has been demonstrated to be a powerful strategy to improve bacterial stress tolerance [33–35]. There have been some studies in many organisms to improve the stress responses and products improvement. For example, increased production of L-serine in *E. coli* was achieved through ALE in cultures with increasing amount of L-serine [36]. Elsewhere, hydrogen induced stress has been used to develop strains overproducing hydrogen in *Thermotoga maritima* [37–39]. Strain mutagenesis methods and environmental conditions provided for mutating cells and enriching mutants are two major factors affecting ALE efficiency and effectiveness. In a typical ALE process, spontaneous mutants which have survived after gradually increased stress are selected [36, 40]. In order to enhance ALE efficiency, different mutagenesis strategies have been applied, such as deletion of mutS or mutL to create a stress-induced mutagenesis system [41], utilization of a genetically modified proofreading element of the DNA polymerase complex (ε subunit encoded by dnaQ gene) to construct a “Genome Replication Engineering Assisted Continuous Evolution” (GREACE) system [42]. Using these methods, mutation occurs continuously at high rates during the ALE process, which, on one hand, enlarges the mutation diversity and may increase the probability of selecting expected mutants. However, on the other hand, the remaining mutagenesis system in the selected mutants can unfavourably reduce the genotype stability and phenotype homogeneity of the final strains.

The environmental conditions provided for mutants survival can largely impact the effectiveness of ALE process. Usually, during microbial ALE for improving stress tolerance, a microorganism is cultivated under clearly defined condition such as M9 minimal medium or Luria–Bertani (LB) medium containing a stress-providing chemical. However, due to complex unknown interactions of living organisms with their environment, the evolutionary trade-offs are ubiquitous [33]. It was shown for *E. coli* that increased tolerance to ethanol comes at the cost of decreased resistance to acidic conditions [43] and strains evolved under phosphate limitation conditions showed trade-offs in NaCl and oxidative stress survival [44]. Therefore, for biotechnological purposes, to avoid the trade-offs as far as possible, a real living stress condition such as the late-stage fermentation broth should be chosen as the ALE environment. To the best of our knowledge, ALE in such a stress condition has not been reported.

In this study, we used the EFB as the starting evolution condition to carry out a modified GREACE-assisted ALE in order to enhance the tolerance and lysine production of strain MU-1 [17]. The improved GREACE method includes expressing a DnaQ mutant
KR5-2 which can increase genome mutagenesis rate by 317-fold [42] with the control of an arabinose inducible promoter on a temperature-sensitive plasmid, so as to timely introduce mutations and stabilize acquired genotypes. A mutant RS3 was ultimately obtained, which produced 155.0 g/L lysine, improved by 14.8%. Further genomics and metabolomics analyses, coupled with molecular biology studies revealed that three genetic mutations of speB (agmatinase gene), atpB (gene of membrane subunit a of FoF1-type ATP synthase), and secY (gene of a subunit of preprotein translocase) may contribute to the improved stress tolerance and the enhanced metabolic flux into lysine. This study brings forward alternative targets to engineer lysine producer and provides an effective strategy involving ALE in EFB to improve production of other valuable hyperproducers.

**Methods**

**Chemicals and enzymes**

1-lysine and arabinose were supplied by Sinopharm Chemical Reagent Co., Ltd (Tianjin, China). Propidium iodide (PI) was purchased from Sigma-Aldrich (USA). Ampicillin and kanamycin were supplied by Solarbio (Beijing, China), and 3-morpholinopropanesulfonic acid (MOPS) was supplied by Amresco (USA). Other pure chemicals used in this study were of analytical grade or better. DNA polymerase was obtained from Transgene (Beijing, China). The ClonExpress II One Step Cloning kit was purchased from Vazyme (Nanjing, China). The Wizard Genomic DNA Purification Kit was obtained from Promega. Restriction endonucleases, T4 DNA kinase and T4 DNA ligase were purchased from New England Biolabs, Inc. (Beijing, China).

**Cultivation media**

The fermentation medium consists of (per liter) 40 g glucose, 5 g KH₂PO₄, 1 g MgSO₄·7H₂O, 10 g (NH₄)₂SO₄, 0.003 g FeSO₄·7H₂O, 0.003 g MnSO₄·4H₂O, 50 g corn syrup, and 0.7 g KCl [17]. The EFB of *E. coli* strain MU-1 was collected by centrifugation at 12,000 rpm for 10 min and filtration with a 0.22 μm syringe filter. The initial lysine concentration of the EFB was approximately 136.5 g/L [17]. The EFB was diluted with water or concentrated by vacuum distillation. Thereafter, 5 g/L yeast extract and 10 g/L peptone were supplied to prepare EFB media with lysine concentrations of 75 g/L, 100 g/L, 125 g/L, 140 g/L, and 150 g/L. Solid plates were prepared by adding 2% agar to the liquid medium. According to the resistance of cultivated strains, ampicillin and kanamycin were added at final concentrations of 100 mg/L and 25 mg/L, respectively.

**Plasmid and strain construction**

The strains, plasmids and primers used in this study are listed in Table 1. Other strains were constructed based on them. The plasmid pAG in MU-1 [17] was cured after serial transfers in LB medium without kanamycin, and the resulting strain was denoted as MU-11. The *dnaQ* mutant KR5-2 which is capable of inducing a 317-fold increased cell mutagenesis rate was amplified from the plasmid pQ-*dnaQ*-KR5-2 [42] using primers KR-F and KR-R. The resulting PCR product was digested with EcoRI and SmaI, and ligated to the plasmid pKD46 (GenBank accession no.: MF287367, kanamycin resistance, temperature sensitive) digested with the same restriction enzymes (with exo, bet and gam genes deleted, keeping the arabinose inducible promoter). The generated plasmid was named as pKAR, which was transformed into MU-11 to generate MU-11 (pKAR).

To construct the plasmid pSB and its derivatives, the pSB4K5 backbone was obtained by PCR amplification from the plasmid pSB4K5-152002 [45] with primers B4K5-F and B4K5-R. The speB and atpB genes (or their mutants) together with their promoter regions were amplified from MU-11 (or its mutant) using the respective primers speB-F and speB-R, atpB-F and atpB-R. The pSB4K5 backbone was treated with T4 DNA kinase, and self-ligated using T4 DNA ligase to obtain pSB. The speB gene and its mutant, atpB gene and its mutant were ligated separately with pSB4K5 backbone via homologous recombination using the ClonExpress II One Step Cloning kit to obtain the corresponding plasmids pSB-speB, pSB-speB<sup>C905T</sup>, pSB-atpB, pSB-atpB<sup>G494A</sup>. To construct plasmid pSB-sRNA<sup>secY</sup>-MicC, primers secY-F and secY-R were used to amplify the MicC scaffold and T1/TE terminator from plasmid pEC-sRNA<sup>secY</sup>-J23109 (http://parts.igem.org/Part:BBa_J23109). The primer sec-F contained the sequences of the corresponding 1–24th bases of *secY* gene and a constitutive promoter J23109 (http://parts.igem.org/Part:BBa_J23109). The obtained fragment was ligated with pSB4K5 backbone via homologous recombination to generate pSB-sRNA<sup>secY</sup>-MicC. The plasmids were transformed separately into LYS1 to generate LYS1 (pSB), LYS1 (pSB-speB), LYS1 (pSB-speB<sup>C905T</sup>), LYS1 (pSB-atpB), LYS1 (pSB-atpB<sup>G494A</sup>), and LYS1 (pSB-sRNA<sup>secY</sup>-MicC).

**Adaptive laboratory evolution process**

To improve the tolerance of MU-11 (pKAR) in its real living stress condition, diluted or concentrated EFB media were prepared for the GREACE-assisted ALE process. ALE was performed at 30 °C in 24-deep-well plates with 1 mL medium in each well, and 10 samples each in one well were subject to the ALE process. The process was initiated by cultivation of MU-11 (pKAR) overnight in
LB medium with 10 g/L arabinose to induce expression of the DnaQ mutant KR5-2. The culture was used as a seed to inoculate EFB media with 75 g/L lysine, followed by two rounds of transfers in EFB media with 100 g/L and 125 g/L lysine, respectively. In the first round in EFB medium with 125 g/L lysine, 10 g/L arabinose was supplied again to induce expression of KR5-2. Afterwards, cells were transferred into EFB medium with 140 g/L lysine for one round and then into EFB medium with 150 g/L lysine for another round. Three additional parallel samples without supplying arabinose during the whole process were used as the control groups. During the ALE, transfer was conducted when the OD600 (optical density at 600 nm) increase reached 0.2–0.3 compared to the control groups. The ALE process was stopped when the control culture almost could not grow in a certain EFB medium wherein the samples can obviously grow. The inoculation ratio was 1%. For measurement of cell growth, 150 μL culture from each well was sampled into Corning 96-Well Microplate, and OD600 was measured using a microplate reader (SpectraMax 190 from Molecular Devices, China). After the final transfer, samples were cultivated at 42 °C in EFB medium with 75 g/L lysine and without kanamycin to cure the plasmid pKAR. Thereafter, cells in each well were spread over agar plates of EFB media with 75 g/L lysine, and cultivated at 42 °C. Three colonies from each well were selected and further evaluated.

For testing growth of MU-11 and the ALE selected mutants in EFB media, strains were sequentially transferred at a ratio of 1% in the broths with lysine concentrations of 75 g/L, 100 g/L, 125 g/L and 150 g/L. Cultivation was carried out at 37 °C in 24-deep-well plates with 1 mL culture in each well. OD600 was measured using the microplate reader as above.

**Fermentation tests**

The fermentation medium was used for all the fermentation tests. Fermentation processes were carried out according to the methods described previously [17]. For fermentations in the 96-deep-well plate and shake flask, MOPS was supplemented at a final concentration of 0.4 mol/L to buffer the pH change. Lysine productions of MU-11 and its mutants in 96-deep-well plate with 300 μL culture in each well were tested at 16 h, while those of LYS1 derived strains were tested at 40 h.
tests of MU-11 and its mutants were also carried out in 500-mL flasks containing 20 mL fermentation medium shaking at 220 rpm. After cultivation for 16 h, 400 μL 25% NH₃·H₂O was supplied to each flask, and fermentation continued to 24 h. OD₆₀₀ of the samples was detected using a spectrophotometer (UV-1800, Shimadzu, Japan). Glucose and extracellular lysine were detected using SBA-40D (Biosensing Analyzer, Shandong, China) as described previously [17]. For fermentation in 5-L jar fermenters, two rounds of precultures were conducted to activate strains and prepare seed cultures. For the main fermentation process, the initial volume was 1.75 L. Continuous maintenance of pH at 7.0 with 25–28% ammonia, glucose at 5–10 g/L with 500 g/L glucose solution and ammonia–nitrogen at 0.05–0.1 g/L with 500 g/L ammonium sulfate solution were performed as described previously [17].

**Cell membrane damage analysis**

Cell membrane damage was analyzed by measuring the uptake of the fluorescent dye propidium iodide (PI). Cells were collected by centrifugation at 7000 rpm for 2 min. Pellets were resuspended in PBS (NaCl 8 g/L, KCl 0.2 g/L, Na₂HPO₄·12H₂O 3.58 g/L, KH₂PO₄ 0.24 g/L, pH 7.2) to an OD₆₀₀ of about 0.5. The suspensions were added with PI (prepared as 200 μg/mL stock solution in PBS) to the final concentration of 2 μg/mL, and incubated in the dark for 10 min at room temperature. Cells were subsequently washed once with PBS. The uptake of PI was analyzed using a MoFlo XDP flow cytometer (Beckman Coulter Inc., USA) according to a previous method [25] with minor modifications. The excitation wavelength and the emission wavelength were 488 nm and 620 nm, respectively. Data for at least 50,000 cells per single sample were collected. Percentage of positive cells for PI analysis was calculated using the statistics module of the Summit 5.2 software.

**Genome sequencing**

The processes for sample preparation and genome sequencing were carried out according to a previously reported method [47]. Genomic DNA was purified using the Wizard Genomic DNA Purification Kit. The average insert size of the library was 500 bp, and the average read length was 150 bp. Sequencing was performed using an Illumina Miseq 2000 system (Tianjin Institute of Industrial Biotechnology, Chinese Academy of Sciences, Tianjin, China). The raw sequence reads were sub-sampled and trimmed by removing low quality bases. The resulted reads were aligned onto the reference genome of MU-11 (unpublished data). The alignment result was visualized for assembly. SNP and Indel variants were analyzed, and relevant gene annotation was performed.

**Intracellular metabolomics analysis**

The intracellular metabolomics were analyzed as reported previously [48]. Cellular metabolism was quenched using fivefold volume of − 20 °C 40% methanol. Intracellular metabolites were extracted using ethanol/water (3:1, v/v) at 100 °C combined with acidic acetonitrile/water (1:1, v/v, with 0.1% formic acid) at − 20 °C. The LC–MS/MS analysis and data processing were all carried out accordingly.

**Results and discussion**

**Construction of a GREACE-assisted ALE method for improving MU-11 tolerance in the EFB media**

In a recent study, we obtained a high-level lysine producing *E. coli* strain MU-1. The EFB contained 136.5 g/L lysine, and about 28 g/L NH₃·H₂O and 57 g/L (NH₄)₂SO₄ were supplied, leading to high stress [17]. For many fermentation processes, the product generation rates obviously decrease in the late stages, especially for products with high concentrations [11, 12, 17]. One major reason for this reduction in product formation might be the cells encounter challenging stress in the late-stage fermentation environment. It is reasonable to assume that cells bearing enhanced stress tolerance might gain a survival advantage and maintain better metabolic activity to generate more products.

In this study, to accelerate the process of ALE, avoid excessive mutation, and timely stabilize the genotype, we reconstructed the GREACE system by expressing the DnaQ mutant KR5-2 [42] under the control of the arabinose inducible promoter on the temperature-sensitive plasmid pKD46. The constructed plasmid pKAR was transformed into MU-11 to generate MU-11 (pKAR). As shown in Fig. 1, during the transfer of control MU-11 (pKAR) in the EFB media with lysine concentration increased from 75 to 150 g/L, the cell growth level gradually decreased. In the medium with 150 g/L lysine, MU-11 (pKAR) almost could not grow. Comparatively, the final OD₆₀₀ of all 10 samples, named as MR1 to MR10, improved by around 10- to 18-folds in EFB medium with 150 g/L lysine (Fig. 1 and Additional file 1: Fig. S1). The whole process was performed within only 7 transfers, much fewer than previously reported ALE processes [33], indicating the high efficiency of improving bacterial stress tolerance in EFB medium using the established GREACE-assisted method in our study.

**Evaluation of ALE selected mutants by fermentation and growth test**

To test the lysine productivities of the evolved strains, three colonies from each of the 10 samples were randomly selected after cultivation in 42 °C to eliminate the plasmid pKAR, and the lysine productivities of the
selected strains were evaluated by fermentations in 96-deep-well plates. The best ones in each of the 10 samples were named as RS1 to RS10. Nine of the 10 samples contained mutants (except for RS9) with improved lysine productions (Fig. 2a). The 9 mutants were subjected to fermentation test in flasks. By comparing the lysine titer and yield with the control strain MU-11, the mutants RS2, RS3, RS4, RS5 and RS6 showed improved lysine production (Fig. 2b).

In order to test the stress tolerance of the selected mutants with improved lysine production, cell growth of RS2, RS3, RS4, RS5, and RS6 was measured in 24-deep-well plates at 37 °C. The cells were sequentially transferred in the EFB medium with lysine concentration from low to high (75, 100, 125 and 150 g/L). As shown in Fig. 3a, The OD600 of MU-11 at 24 h in EFB medium with 75 g/L lysine was higher than most of the tested mutants, indicating that the initial strain with high lysine producing capacity can also resist low stress and the mutants have no advantage under the minor stress conditions. In EFB media with 100 g/L lysine, even though the cell densities of MU-11 and the mutants differed slightly at 40 h, the PI uptake analysis, which has been frequently used to measure cell membrane integrity [20], revealed that MU-11 was the most damaged strain (Fig. 3b). When the lysine concentration in the EFB medium further increased to 125 g/L, the OD600 increase of all tested mutants were higher than that of MU-11. These data indicate that all the tested mutants showed improved tolerance in the EFB medium, suggesting that ALE under the cell living condition such as EFB medium could be an effective strategy. When even higher stress such as EFB medium with 150 g/L lysine was applied, all cells exhibited dramatically decreased growth. However, mutants RS3 and RS5 obviously grow better than other mutants and the MU-11 control (Fig. 3a).

Fed-batch fermentation tests and metabolomics analyses of the selected and control strains

After evaluation of fermentation performance and stress tolerance of selected mutants in deep-well plates and shake flasks, the best mutant RS3 was further examined in a 5 L jar fermenter with a mimetic industrial fed-batch fermentation process. As shown in Fig. 4a and Table 2, lysine production of RS3 was highly improved. The final titer, yield and total lysine amount in one batch reached 155.0 ± 1.4 g/L, 0.59 ± 0.02 g lysine/g glucose, and 605.6 ± 23.5 g, accounts for improvement of 14.8%, 9.3%, and 16.7%, respectively.

The PI uptake analysis was carried out to test the membrane integrity of RS3 and MU-11 during fermentation (Fig. 4b). At 9 h, a time point representing the early fermentation stage with less stress, the cell membrane damage degrees were slight for both strains. At 30 h, a time point representing mid to late fermentation stage with more than 120 g/L lysine produced, the
damage degrees of both strains increased, but MU-11 was obviously worse than RS3. The result implied that under a proper stress condition, RS3 was relieved from serious cell damage. At 42 h, the end of fermentation, an extremely stressful condition, the cell membranes of more than 90% of both strains were damaged, and the degree of RS3 was a little slighter than MU-11, which might also be the reason for better growth of RS3 after a long period of cultivation in the EFB media with 150 g/L lysine (Fig. 3).

The growth of the mutant RS3 in the 5 L fermenter delayed unexpectedly after 10 h in comparison to that of the control MU-11. This presumably was not caused by stress, since both strains did not face serious stress as shown by limited membrane damage at 9 h depicted in Fig. 4b. Another mutant RS5 even suddenly stopped growth when OD_{600} reached about 19 at 6 h in a 5 L fermenter (data not shown). Similar phenomena about the growth of mutants selected by ALE in one condition showing impediment in another condition has also been reported [33]. Therefore, to evaluate the potential of the obtained mutants for industrial purpose, it is better to test their performance in jar fermenters since more variants such as glucose and ammonia–nitrogen levels, and fermentation condition control were introduced. Nevertheless, the production capacity of RS3 was apparently enhanced. Comprehensive optimization of the fermentation medium in future work would further improve the production performance of RS3 as well as RS5.

To understand the metabolic mechanism for strain improvement, metabolomics analyses of RS3 and MU-11 cultures in the 5-L jar fermenters were carried out. In order to avoid the influence of cell damage to the analytical results, the 9 h fermentation cultures, wherein the damage degree was slight and almost the same for MU-11 and RS3 (Fig. 4b), were sampled and quenched immediately. Intracellular metabolites were extracted and then analyzed using LC–MS/MS. The LC–MS/MS data were normalized by cell density, and the concentrations of corresponding metabolites in MU-11 and RS3 were compared. As shown (Fig. 5 and Additional file 1: Table S1), compared with MU-11, the EMP, PPP, TCA left half, aspartate family amino acid synthesis, and redox equivalent and energy currency synthesis pathways of RS3 were all enhanced. The down regulation of the TCA right half pathway might be due to the cellular regulation to limit carbon flux into TCA cycle under enhanced carbon metabolism and energy generation condition [49]. The enhanced metabolic flux into lysine synthesis and increased redox equivalents especially NADPH synthesis are beneficial for lysine production [1, 14]. The improved ATP and NADPH supplies may provide more energy for cells to tolerate stressful conditions [50, 51].

The impact of genomic variations on stress tolerance and lysine production of RS3
Whole genome sequencing was performed on RS3 to discover genomic variations, and 3,655,204 high quality reads with an average read length of 150 bp were obtained. The sequence was compared with the genomic sequence of the parent strain MU-11. Three mutations occurred on genes including speB, atpB and secY, leading to corresponding amino acid sequence changes (Table 3). In addition, two transposases and three hypothetical proteins were mutated, each with a base change leading to an amino acid substitution (data not shown).
To evaluate the effects of A302V mutation in SpeB (corresponding to nucleotide change of C905T) (Table 3) on cellular stress tolerance and lysine production, the wild-type and mutant SpeB were over expressed respectively in strains LYS1 (pSB-speB) and LYS1 (pSB-speBC905T), which were subjected to fermentation test in a 96-deep-well plate incubated for 40 h. As shown in Fig. 6a, b, compared with the control strain LYS1 (pSB), LYS1 (pSB-speB) showed improved lysine production and cell membrane integrity. Since SpeB catalyzes the conversion of arginine to putrescine and spermidine, improved metabolic flux to polyamines biosynthesis is expected from SpeB overexpression. A previous study has demonstrated that the catabolism of these polyamines is a metabolic response to stress in bacteria such as E. coli [52]. Therefore, enhancing polyamine biosynthesis may be beneficial for lysine production and stress tolerance. Interestingly, overexpression of the SpeB mutant further increased lysine production and cell integrality, indicating improved performance of the SpeB mutant.

Numerous repair systems for stress induced cell damage are ATP-dependent. Improving ATP production is an important mechanism for strain tolerance [50, 51]. The enzyme AtpB is a subunit of the complex FoF1-type ATP

**Table 2** Lysine productions of MU-11 and RS3 in 5 L jar fermenters

| Strain | Lysine concentration (g/L) | Yield (g lysine/g glucose) | Total lysine in one batch (g) |
|--------|-----------------------------|-----------------------------|-----------------------------|
| MU-11  | 135.0 ± 2.8                 | 0.54 ± 0.01                 | 519.1 ± 27.9                |
| RS3    | 155.0 ± 1.4                 | 0.59 ± 0.02                 | 605.6 ± 23.5                |

**Fig. 3** Growth tests and PI uptake analyses of MU-11 and selected mutants in EFB media. *a* Growth of MU-11 and selected mutants. Cells were serially transferred at a ratio of 1% in EFB media with lysine concentrations from low to high. *b* PI uptake of MU-11 and selected mutant cells from the 40 h-samples cultivated in EFB media containing 100 g/L lysine. The growth curves of the control groups were linked to highlight the growth differences between controls and mutants. Data are the mean and standard deviation of independent triplicates.

**Fig. 4** Fermentation tests and PI uptake analyses of MU-11 and RS3 in 5 L jar fermenter. *a* Lysine productions and cell growth of MU-11 and RS3. Open circles and open squares represent the lysine productions of RS3 and MU-11, respectively. Filled circles and filled squares represent the growth of RS3 and MU-11, respectively. *b* PI uptake of MU-11 and RS3 cells from the samples at different fermentation time. Light grey bars and grey bars represent the PI positive rates of MU-11 and RS3, respectively. Data are the mean and standard deviation of independent triplicates.
synthase [53]. S165 of AtpB locates nearby an important site R169 whose mutation R169A can lead to the complete inactivation of the ATP synthase [54]. The effect of AtpB<sup>S165N</sup> (corresponding to nucleotide change of G494A) (Table 3) on lysine production and cell damage was examined. The control strain LYS1 (pSB) and the recombinant strain LYS1 (pSB-<i>atpB</i>) overexpressing the wild type AtpB showed similar lysine production and cell membrane integrity, suggesting that overexpression of a single component of the mature ATP synthase complex
might not affect the ATP synthase complex activity. However, the lysine production of LYS1 (pSB-atpB<sup>G494A</sup>), and cell integrity as well, improved (Fig. 6), indicating that the AtpBS<sub>165N</sub> could affect the activity of ATP synthase and should be beneficial for RS3. This is consistent with the increased level of ATP in RS3 compared to MU-11 revealed by metabolomics analysis (Fig. 5 and Additional file 1: Table S1).

SecY forms a channel of the important membrane protein secretory system complex SecY/E/G and SecA [55, 56]. SecY alteration that impaired membrane protein folding can up-regulate the Cpx/sigma(E) stress response pathways [57], which is important for acid stress and cell wall stability in <i>E. coli</i> [58]. To test the impact of decreased SecY expression on lysine production and cell membrane damage, strain LYS1 (pSB-sRNA<sub>secY</sub>-MicC) transcribing a small regulatory RNA (sRNA) to inhibit SecY synthesis was constructed. The sRNA contains a 24 bp sequence complementary to the 1–24th bases of secY gene, and contains a MicC scaffold for recruiting the Hfq protein [59]. Compared with the control strain LYS1 (pSB), LYS1 (pSB-sRNA<sub>secY</sub>-MicC) showed improved lysine production and decreased cell membrane damage (Fig. 6), which is consistent with the previous report. In addition, considering the location of the M145 site in the P2 region of SecY, which might be involved at the interface with SecG [60], the M145 V mutation of SecY is more likely prone to weaken the function of the protein complex SecY/E/G and SecA.

According to the results, it could be deduced that the SpeB<sub>A302V</sub>, AtpB<sub>S165N</sub> and SecY<sub>M145V</sub> mutants are likely to be the main contributors to the improvement of RS3 (Fig. 4a). However, it was difficult to deduce the mutants induced causality of the stress tolerance (Figs. 4b, 6b) and the overall metabolic enhancement of RS3 (Fig. 5).

Taken together, ALE is effective to improve stress tolerance of microbial cell factories but a time-consuming process that commonly requires tens of transfers and takes weeks up to months [33]. Strategies to increase mutagenesis rates are helpful for accelerating the ALE process [41, 42]. In this study, GREACE that facilitates mutagenesis coupled-with selection under stress conditions was modified and applied to accelerate the ALE process. Growth of all the 10 paralleled samples improved under stress conditions after only seven rounds of transfers (Fig. 1), indicating the high

---

### Table 3 Genomic variants of RS3 compared with MU-11

| Concerned gene | NCBI gene locus | Mutant site | Amino acid variation | Protein description |
|---------------|----------------|-------------|----------------------|---------------------|
| speB          | b2937          | C905T       | A302 V               | Arginase family enzyme |
| atpB          | b3738          | G494A       | S165 N               | FoF1-type ATP synthase, membrane subunit a |
| secY          | b3300          | A433G       | M145 V               | Preprotein translocase subunit |

---

![Fig. 6](image-url) Lysine productions and PI uptake analyses of LYS1 derived strains. **a** Lysine productions. **b** PI uptakes. Control represents LYS1 (pSB). speB represents LYS1 (pSB-speB). speB<sup>C905T</sup> represents LYS1 (pSB-speB<sup>C905T</sup>). atpB represents LYS1 (pSB-atpB). atpB<sup>G494A</sup> represents LYS1 (pSB-atpB<sup>G494A</sup>). secY-D represents LYS1 (pSB-sRNA<sub>secY</sub>-MicC), which can transcribe a small regulatory RNA (sRNA) to inhibit SecY synthesis. Cells were cultivated for 40 h in 96-deep-well plate with fermentation medium. Data are the mean and standard deviation of independent triplicates. The p value was calculated by Student’s two-tailed t-test. * represents a p value of < 0.05, and ** represents a p value of < 0.01, comparing with the data of control strain.
efficiency of the modified GREACE-based ALE method. Another improvement to traditional ALE process is the usage of the late-stage fermentation broth to mimic the actual stress conditions that industrial producers faces. By using this strategy, mutants with significant increase in lysine production and stress tolerance were obtained. The ALE strategy established here should also be helpful for improving hyperproducers of various valuable chemicals. Genome sequencing of the best mutant RS3 provided valuable information for understanding the mechanism of improved lysine production and stress tolerance. Mutations in genes involved in polyamine biosynthesis, energy supply, and protein secretory were proven to be beneficial, which highlighted the importance of these three cellular processes in enhancing bioproduction.

Conclusions
By reconstructing the GREACE system, we developed a mutagenesis controllable GREACE-assisted ALE process in EFB. The process was applied to improve the stress tolerance of the lysine hyperproducer *E. coli* MU-1 in lysine EFB medium. Tolerance enhanced mutants were obtained within only 7 transfers in EFB media, indicating the high efficiency of the established process. Because environmental stress is harmful to high production of chemicals by bacteria, we expect that this method established in the present study would be extended to improve other chemical hyperproducers.

Additional file

Additional file 1: Fig. S1. Growth comparison of the 10 ALE samples with the control samples. Table S1. Partial intracellular metabolites of RS3 from the 9-h fed-batch fermentation sample.

Abbreviations
ALE: adaptive laboratory evolution; GREACE: genome replication engineering assisted continuous evolution; EFB: endpoint fermentation broth; PI: propidium iodide; DnaQ: ε subunit of the DNA polymerase; SpeB: agmatinase; AtpB: membrane subunit a of the FoF1-type ATP synthase; SecY: one subunit of preprotein translocase; EMP: The Embden–Meyerhof–Parnas Pathway; PPP: the pentose phosphate pathway; TCA: the tricarboxylic acid cycle.

Acknowledgements
We thank Lixian Wang and Yu Lei (both from Technical Support Center, Tianjin Institute of Industrial Biotechnology, Chinese Academy of Sciences) for technical support in PI analysis and genome sequencing, respectively. We also thank Dr. Yu Wang and Dr. Timothy Cairns (both from Tianjin Institute of Industrial Biotechnology, Chinese Academy of Sciences) for their efforts to improve the quality of the manuscript.

Authors’ contributions
XW, QL, CS, XG,YG performed the experiments. QL, ZC, XZ and XN analyzed the data. QL, WZ, PZ, NC, JS, YL and YM designed the experiments and prepared the manuscript. All authors read and approved the final manuscript.

Funding
This research was supported by grants from the National Natural Science Foundation of China (31870081), the Special Program of Talents Development for Excellent Youth Scholars in Tianjin (TJYZH-QNBRC-2-10), the Youth Innovation Promotion Association of Chinese Academy of Sciences (2016164), and the Science and Technology Project of Tianjin (15RQCTYS00020 and 14ZCDZSY00157).

Availability of data and materials
All data generated or analyzed during this study are included in this published article and the Additional file 1. The authors are willing to provide any additional data and materials related to this research that may be requested for research purposes.

Ethics approval and consent to participate
Not applicable.

Consent for publication
All authors give consent to publish the research in Microbial Cell Factories.

Competing interests
The authors declare that they have no competing interests.

Author details
1 College of Biotechnology, Tianjin University of Science and Technology, Tianjin 300457, China. 2 Key Laboratory of Systems Microbial Biotechnology, Chinese Academy of Sciences, Tianjin 30008, China. 3 Tianjin Institute of Industrial Biotechnology, Chinese Academy of Sciences, Tianjin 300308, People’s Republic of China. 4 CAS Key Laboratory of Microbial Physiological and Metabolic Engineering, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China.

Received: 9 March 2019 Accepted: 1 June 2019 Published online: 11 June 2019

References
1. Contador CA, Rizk ML, Asenjo JA, Liao JC. Ensemble modeling for strain development of L-lysine-producing *Escherichia coli*. Metab Eng. 2009;11:221–33.
2. Esvelt KM, Wang HH. Genome-scale engineering for systems and synthetic biology. Mol Syst Biol. 2013;9:641.
3. Nisitani J, Kearing JD. Engineering cellular metabolism. Cell. 2016;164:1185–97.
4. Salis HM, Mirsky EA, Voigt CA. Automated design of synthetic ribosome binding sites to control protein expression. Nat Biotechnol. 2009;27:946–1112.
5. Lee JH, Wendisch VF. Production of amino acids—genetic and metabolic engineering approaches. Bioreasour Technol. 2017;245:1575–87.
6. Li Y, Wei H, Wang T, Yu Q, Zhang C, Fan X, Ma Q, Chen N, Xie X. Current status on metabolic engineering for the production of L-aspartate family amino acids and derivatives. Bioreasour Technol. 2017;245:1588–602.
7. Yu H, Wang N, Huo W, Zhang Y, Zhang W, Yang Y, Chen Z, Huo YX. Establishment of BmoR-based biosensor to screen isobutanol overproducer. Microb Cell Fact. 2019;18:30.
8. Liu Y, Li Q, Zheng P, Zhang Z, Liu Y, Sun C, Cao G, Zhou W, Wang X, Zhang D, et al. Developing a high-throughput screening method for threonine overproduction based on an artificial promoter. Microb Cell Fact. 2015;14:121.
9. Chung SC, Park JS, Yun J, Park JH. Improvement of succinate production by release of end-product inhibition in *Clostridium glutamicum*. Metab Eng. 2017;40:157–64.
10. Zheng P, Liu M, Liu XD, Du QY, Ni Y, Sun ZH. Genome shuffling improves thermo tolerance and glutamic acid production of *Corynebacterium glutamicum*. World J Microbiol Biotechnol. 2012;28:1035–43.
11. Xie X, Liang Y, Liu H, Liu Y, Xu Q, Zhang C, Chen N. Modification of glycolysis and its effect on the production of l-threonine in Escherichia coli. J Ind Microbiol Biotechnol. 2014;41:1007–15.

12. Xu JZ, Wu ZH, Gao SJ, Zhang W. Rational modification of tricarboxylic acid cycle for improving l-threonine production in Corynebacterium glutamicum. Microb Cell Fact. 2018;17:105.

13. Tsuge Y, Hasunuma T, Kondo A. Recent advances in the metabolic engineering of Corynebacterium glutamicum for the production of lactate and succinate from renewable resources. J Ind Microbiol Biotechnol. 2015;42:375–89.

14. Bommareddy RR, Chen Z, Rappert S, Zeng AP. A de novo NADPH generation pathway for improving lysine production of Corynebacterium glutamicum by rational design of the coenzyme specificity of glyceraldehyde-3-phosphate dehydrogenase. Metab Eng. 2014;25:30–7.

15. Becker J, Zelder O, Hafner S, Schroder H, Wittmann C. From zero to hero—design-based systems metabolic engineering of Corynebacterium glutamicum for l-lysine production. Metab Eng. 2011;13:519–68.

16. Geng F, Chen Z, Zheng P, Sun J, Zeng AP. Exploring the allosteric mechanism of dihydrodipicolinate synthase by reverse engineering of the allosteric inhibitor binding sites and its application for lysine production. Appl Microbiol Biotechnol. 2013;97:1963–71.

17. Wang Y, Li Q, Zheng P, Guo Y, Wang L, Zhang T, Sun J, Ma Y. Evolving the l-lysine high-producing strain of Escherichia coli using a newly developed high-throughput input screening method. J Ind Microbiol Biotechnol. 2016;43:1227–35.

18. Becker J, Kloppeogge C, Wittmann C. Metabolic responses to pyruvate kinase deletion in lysine producing Corynebacterium glutamicum. Microb Cell Fact. 2008;7.8.

19. Altdorfer K, Booth IR, Gralla J, Greie JC, Rosenthal AZ, Wood JMA. Osmotic stress. EcoSal Plus. 2009. https://doi.org/10.1128/ecosalplus.5.4.4.

20. Guo Y, Winkler J, Kao KC. Insights on osmotic tolerance mechanisms in Escherichia coli gained from an rpoC mutation. Bioengineering. 2017;4:61.

21. Sevin DC, Sauer U. Ubiquinone accumulation improves osmotic-stress tolerance in Escherichia coli. Nat Chem Biol. 2014;10:266–72.

22. Rice KC, Bayles KW. Molecular control of bacterial death and lysis. Microbiol Mol Biol Rev. 2008;72:109–109.

23. Liu X, Parales RE. Chemotaxis of Escherichia coli to pyrimidines: a new role for the signal transducer tap. J Bacteriol. 2008;190:972–9.

24. Soulský J. Receptor clustering and signal processing in E. coli chemotaxis. Trends Microbiol. 2004;12:569–76.

25. Li J, Ahn J, Liu D, Chen S, Ye X, Ding T. Evaluation of ultrasound-induced damage to Escherichia coli and Staphylococcus aureus by flow cytometry and transmission electron microscopy. Appl Environ Microbiol. 2016;82:1828–37.

26. Perez-Pardo MA, Ali S, Balasundaram B, Mannall GJ, Baganz F, Bracewell DG. Assessment of the manufacturability of Escherichia coli high cell density fermentations. Biotechnol Prog. 2011;27:1488–96.

27. Delamarche C, Thomas D, Rolland JP, Froger A, Gouranton J, Svelto M, Agre P, Calamita G. Visualization of AqpZ-mediated water permeability through the SigB-controlled betaine-choline-carnitine transporter and its effect on the production of l-threonine in Escherichia coli. J Bioenergy Biomembr. 2005;37:445–9.

28. Laimins LA, Rhoads DB, Epstein W. Osmotic control of cell envelope integrity. J Bacteriol. 1993;175:4699–708.

29. Clarke CF, Rowat AC, Gober JW. Osmotic stress: is CoQ a membrane stabilizer? Nat Chem Biol. 2012;8:242–3.

30. Chen X, An L, Fan X, Ju F, Zhang B, Sun H, Xiao J, Hu W, Qi T, Guan L, et al. A trehalose biosynthetic enzyme doubles as an osmotic stress sensor to regulate bacterial morphogenesis. PLoS Genet. 2017;13:e1007062.

31. Dragostis M, Mattanovich D. Adaptive laboratory evolution—principles and applications for biotechnology. Microb Cell Fact. 2013;12:64.

32. Gong Z. Nielsen J, Zhou YJ. Engineering robustness of microbial cell factories. Biotechnol J. 2017;12(10):1700014.

33. Long CP, Antoniewicz MR. How adaptive evolution reshapes metabolism to improve fitness: recent advances and future outlook. Curr Opin Chem Eng. 2018;22:209–15.

34. Mundhada H, Seoane JM, Schneider K, Koza A, Christensen HB, Klein T, Phanet PV, Herrgard M, Feist AM, Nielsen AT. Increased production of l-serine in Escherichia coli through adaptive laboratory evolution. Metab Eng. 2017;39:141–50.

35. Singh R, White D, Demirel Y, Kelly R, Noll K, Blum P. Uncoupling fermentative synthesis of molecular hydrogen from biomass formation in Thermotoga maritima. Appl Environ Microbiol. 2018;84:e00998-18.

36. Singh R, Tevatica R, White D, Demirel Y, Blum P. Comparative kinetic modeling of growth and molecular hydrogen overproduction by engineered strains of Thermotoga maritima. Int J Hydrogen Energy. 2019;44:7125–36.

37. Blum P, Rudrappa D, Singh R, McCarthy S, Pavlik B. Experimental microbial evolution of extremophiles. In: Rampelotto P, editor. Biotechnology of extremophiles: grand challenges in biology and biotechnology, vol. 1. Cham: Springer; 2016.

38. Sun XM, Ren LJ, Bi ZQ, Ji XJ, Zhao QY, Huang H. Adaptive evolution of microalgae Schizochytrium sp. under high salinity stress to alleviate oxidative damage and improve lipid biosynthesis. Bioreour Technol. 2018;267:438–44.

39. Zhu L, Cai Z, Zhang Y, Li Y. Engineering stress tolerance of Escherichia coli by stress-induced mutagenesis (SIM)-based adaptive evolution. Biotechnol J. 2014;9:120–7.

40. Luan G, Cai Z, Li Y, Ma Y. Genome replication engineering assisted continuous evolution (GREATE) to improve microbial tolerance for biofuels production. Biotechnol Biofuels. 2013;6:137.

41. Goodarzi H, Bennett BD, Amin S, Reaves ML, Hottes AK, Rabkowitz JD, Tavazoie S. Regulatory and metabolic rewiring during laboratory evolution of ethanol tolerance in E. coli. Mol Syst Biol. 2010;6:378.

42. Wang L, Spira B, Zhou Z, Feng L, Maharjan RP, Li X, Li F, McKenzie C, Reeves PR, Ferenci T. Divergence involving global regulatory gene mutations in an Escherichia coli population evolving under phosphate limitation. Genome Biol Evol. 2010;2:478–87.

43. Shetty RP, Endy D, Knight TF. Jr. Engineering BioBrick vectors from BioBrick parts. J Biol Eng. 2008;2:5.

44. Sun D, Chen J, Wang Y, Li M, Rao D, Guo Y, Chen N, Zheng P, Sun J, Ma Y. Metabolic engineering of Corynebacterium glutamicum with synthetic small regulatory RNAs. J Ind Microbiol Biotechnol. 2019;46:203–8.

45. Yang J, Yang S. Comparative analysis of Corynebacterium glutamicum genomes: a new perspective for the industrial production of amino acids. BMC Genomics. 2017;18:940.

46. Zhang Q, Zheng X, Wang Y, Yu J, Zhang Z, Dele-Oslanjo T, Zheng P, Sun J, Jia S, Ma Y. Comprehensive optimization of the metabolic methodology for metabolite profiling of Corynebacterium glutamicum. Appl Microbiol Biotechnol. 2018;102:713–21.

47. Lemuth K, Hardiman T, Winters S, Pfeiffer D, Keller MA, Lange S, Reuss M, Schmid RD, Siemann-Herzberg M. Global transcription and metabolic flux analysis of Escherichia coli in glucose-limited fed-batch cultivations. Appl Environ Microbiol. 2008;74:7002–15.

48. Adolfsen KJ, Brynildsen MP. futile cycling increases sensitivity toward oxidative stress in Escherichia coli. Metab Eng. 2015;29:26–35.

49. Cao H, Wei D, Yang Y, Shang Y, Li G, Zhou Y, Ma Q, Xu F. Systems-level understanding of ethanol-induced stresses and adaptation in E. coli. Sci Rep. 2017;7:44150.

50. Schneider BL, Hernandez VJ, Reitzer L. Putrescine catabolism is a metabolic response to several stresses in Escherichia coli. Mol Microbiol. 2013;88:537–50.

51. Vik SB, Ishmukhametov RR. Structure and function of subunit a of the ATP synthase of Escherichia coli. J Bioenergy Biomembr. 2005;37:445–9.

52. Kunuma Y, Suzuki T, Ono S, Yoshida M, Ueda T. Functional analysis of membrane-bound F-α of subunit F-Fo-ATP synthase by in vitro protein synthesis. Biochem J. 2012;442:631–8.

53. Park E, Menetret JF, Gumbart JC, Ludtke SJ, Li W, Whynot A, Rapoport TA, Ackey CW. Structure of the SecY channel during initiation of protein translocation. Nature. 2014;506:102–6.

54. Susano Y, Furukawa A, Nureki O, Tanaka Y, Tsuchazaki T, SecY-SecA fusion protein retains the ability to mediate protein transport. PLoS ONE. 2017;12:e0183434.
57. Shimohata N, Nagamori S, Akiyama Y, Kaback HR, Ito K. SecY alterations that impair membrane protein folding and generate a membrane stress. J Cell Biol. 2007;176:307–17.
58. Surmann K, Cudic E, Hammer E, Hunke S. Molecular and proteome analyses highlight the importance of the Cpx envelope stress system for acid stress and cell wall stability in *Escherichia coli*. Microbiologyopen. 2016;5:582–96.
59. Na D, Yoo SM, Chung H, Park H, Park JH, Lee SY. Metabolic engineering of *Escherichia coli* using synthetic small regulatory RNAs. Nat Biotechnol. 2013;31:170–4.
60. Mori H, Shimokawa N, Satoh Y, Ito K. Mutational analysis of transmembrane regions 3 and 4 of SecY, a central component of protein translocase. J Bacteriol. 2004;186:3960–9.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Ready to submit your research? Choose BMC and benefit from:
- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.
Learn more biomedcentral.com/submissions