Engineering a diaphorase via directed evolution for enzymatic biofuel cell application

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Abstract

Background: Diaphorase (DI) has received wide attention as the key anodic enzyme mediating the electron transfer and electric energy generation in enzymatic biofuel cells (EBFCs). Lowering the anodic pH may be a useful strategy for constructing high-performance in EBFCs. However, most DI suffered from the poor activity at low pHs. Therefore, it is necessary to modify the activity and its acidic tolerance to further improve the performance of the EBFC.

Results: This paper attempts to improve the enzyme activity of DI originated from Geobacillus stearothermophilus under acidic conditions through directed evolution. Three rounds of random mutagenesis by error-prone PCR of the Gs DI gene followed by high-throughput screening allowed the identification of the mutant 3–8 (H37Q, S73T, F105L, S68T, G61S, D74V) exhibiting a 4- or 7-fold increase in the catalytic activity at pH 5.4 or 4.5 compared to that of the wild type. And the pH stability of mutant 3–8 was significantly better than that of wild type and showed a 1.3 times higher in the stability at pH 5.4. The EBFC anode equipped with 0.5 mg of mutant 3–8 achieved a maximum current of 40 μA at pH 5.4, much higher than that with the same loading of the wild type enzyme.

Conclusion: The Gs DI has been improved in the specific activity and pH stability by directed evolution which leads to the improvement of the EBFC performance. Also, the enlarged catalytic channel of mutant and decreased B-factor may be beneficial for the activity and stability. These results suggest that this engineered DI will be a useful candidate for the construction of enhanced EBFCs.

Keywords: Diaphorase, Enzyme engineering, Directed evolution, Enzymatic biofuel cell, Bioelectrocatalysis

Introduction

Diaphorase (DI) is a type of flavin mononucleotide (FMN)-bound enzymes that catalyze the reduction of various dyes, such as methylene blue, resazurin, vitamin K₃ (VK₃), and azo dyes, which act as hydrogen acceptors from the reduced form of nicotinamide adenine dinucleotide (phosphate), i.e., NAD(P)H (Matsumoto et al. 2010; Rice et al. 1998; Shoji et al. 2016; Zhu et al. 2014). It can also be named as NAD(P)H dehydrogenase (EC 1.6.99.1). Because an oxidized FMN-DI has a strong absorbance at 452 nm and can be reduced to a FMNH₂-DI by accepting two electrons, it has been mainly applied to the coupled colorimetric determination of numerous dehydrogenase-mediated reactions (Collins et al. 2016). Since that the first DI was purified from mammalian heart muscle, many DIs have been obtained from various bacteria including Thermotoga maritima, Geobacillus stearothermophilus, Bacillus subtilis and Clostridium kluyveri (Bergsma et al. 1982; Kim and Kim 2018; Shoji et al. 2016; Zhu and Zhang 2017).

Recently, DI has also received wide attention as the key enzyme mediating the electron transfer and electric energy generation in enzymatic biofuel cells (EBFCs). EBFC is a device that generates electricity by oxidizing fuels at the anode and reducing oxygen at the cathode via enzymatic biocatalysis (Xiao et al. 2019). Due to its high safety, good biocompatibility, and the use of...
renewable biocatalysts and high-density fuels, EBFC is believed to hold the promise as a next generation milli- or micro-power source for wearable and implantable electronic devices in the future (Barton et al. 2004; Davis and Higson 2007; Mano et al. 2003). In EBFCs, the key step for electricity generation is the oxidation catalyzed by various oxidases or dehydrogenases at the anode (Kang et al. 2019; Kim et al. 2013; Liang et al. 2012; Scheiblbrandner and Ludwig 2019). Among such enzymes, NAD-dependent dehydrogenases account for a great portion. Many fuels such as glucose, xylose, formate, and glucose 6-phosphate (G6P), rely on their respective dehydrogenase to be oxidized, coupled with NADH regeneration catalyzed by DI (Gai et al. 2015; Wu et al. 2018; Zhu and Zhang 2017). For example, in our system, G6P as the fuel is oxidized by two enzymes, G6P dehydrogenase (G6PDH) and 6-phosphogluconate dehydrogenase (6PGDH). The NADH generated from NAD is further oxidized by DI, along with the electron transfer from NAD to the mediator, VK3, which is then oxidized at the anode and releases electrons. DI plays a vital role in this system in bridging the biological electron carrier, NAD, and the electrode, and mitigating the electron transfer potential barrier between them.

So far, there have been tremendous efforts focusing on improving the performance of EBFCs (e.g., power output and lifetime) and broadening their applications, such as constructing novel 3-dimensional nanomaterials for bioelectrodes, utilizing highly stable enzymes and biomimetic cofactors, as well as optimizing electrolyte composition and concentration (Campbell et al. 2012; Chen et al. 2016; Sakai et al. 2009). In our previous study, we demonstrated that lowering anodic pH may be a useful strategy for constructing high-performance in EBFCs, as the proton concentration at the lowered pH can be significantly increased at the anode and it therefore provided a high proton transport driving force (Ma et al. 2019). However, the DI used was derived from G. stearothermophilus with an optimum pH at 7.3 and still suffered from the poor activity at low pHs, severely limiting the long-term operation of the EBFC. Therefore, to further improve the performance of our EBFC, it is desirable to engineer this rate- and stability-limiting GSDI and increase its acidic tolerance.

Directed evolution is a powerful method for engineering of biological systems including proteins, pathways and genomes for various applications (Denard et al. 2015; Yang et al. 2019). In the EBFC field, the catalytic turnover and binding affinity of glucose oxidase have been tuned via directed evolution in order to achieve the fast oxidation of glucose (Mano 2019; Yu et al. 2011; Zhu et al. 2007). Alternatively, laccase has been directed evolved for improved catalytic activity, tailored substrate specificity, and enhanced tolerance to alkaline conditions (Camarero et al. 2012; Monza et al. 2015; Novoa et al. 2019). Recently, a DI has been engineered to increase its reactivity against a specific mediator, 2-amino-1,4-naphthoquinone, aiming to improve the power output of the EBFC (Sugiyama et al. 2010). A 6PGDH has been directed evolved toward increased acid-tolerance so that the operation conditions of the EBFC can be broadened (Ma et al. 2019).

In this study, the development of an acid-tolerant DI for the EBFC application is reported. The directed evolution method based on a colorimetric screening of DI variants has been developed. Three rounds of evolution bring us with the best mutant exhibiting a significantly increased activity and stability at the reduced pH. The improved electrochemical performance of the EBFC constructed is also observed.

**Materials and methods**

**Materials**

The primers were synthesized from GENEWIZ (Beijing, P.R. China). Taq DNA polymerase and PrimeSTAR Max DNA polymerase were purchased from TaKaRa (Dalian, P.R. China). Other enzymes for molecular biology experiments were purchased from New England Biolabs (Ipswich, MA, U.S.). *E. coli* Top10 was used for gene cloning and recombinant protein expression. Tetra-nitro blue tetrazolium (TNBT) was purchased from Solarbio (Beijing, P. R. China). Nicotinamide adenine dinucleotide (NAD\(^+\)) and glucose 6-phosphate (G6P) were purchased from Sigma-Aldrich (St. Louis, MO, U. S.).

**Random mutagenesis and library creation**

The plasmid pET28a-Ptac-di was used as template for variant library generation by error-prone PCR with a pair of primers DI_F/DI_R. The sequences were 5′-TTAAC TTTAA GAAGG AGAGG TACAT ATGAC GAAAG TTTAA GAAGG AGATA TACAT ATGAC GAAAG-3′ and 5′-TCAGT GGTGG TATTG TACAT CACCG-3′. The reaction solution with a total volume of 50 μL contained 0.2 mM dATP, 0.2 mM dCTP, 1 mM dGTP, 1 mM dTTP, 5 mM MgCl\(_2\), 0.01 mM MnCl\(_2\), 0.05 U/μL the Taq polymerase and 0.4 μM primer pairs. The PCR reaction was performed under the following conditions: initial denaturation at 94 °C for 2 min, followed by 18 consecutive cycles of denaturation at 94 °C for 30 s, anneal at 58 °C for 20 s, extension at 72 °C for 30 s, and a final extension at 72 °C for 5 min. The 50 μL reaction solution of the vector backbone was amplified from pET28a-Ptac-di with 25 μL PrimeSTAR Max and 0.4 μM primer pairs vector_F/vector_R. The sequences were 5′-CGAAA GACTT GGGGC ACACG TTTCT CGAGC ACCAC CACC CCACC ACTGA-3′; 5′-CGGTTG ATGTA CAATA CTTTC GTGAT ATGTA-3′.
The supernatant was loaded onto the column packed with adding 0.1 mM IPTG when the absorbance of the OD600 containing 50 μg/ml kanamycin at 37 °C and induced by wild type and its mutants were grown in LB medium Top10 for DNA sequencing and protein overexpression. Kit (Tiangen, Beijing, P. R. China) and transformed into of isolated colonies were then extracted by mini plasmid of the cell. When the plate temperature dropped to the incubation time.

Screening of the mutants with high activity at lowered pHs
The double-layer-based TNBT screening method was carried out as previously described with some modifications (Fig. 1a) (Ma et al. 2019). Transformed cells containing mutant plasmid libraries were firstly incubated on the LB kanamycin agar plate at 37 °C for 16 h. Then, the colonies were treated at 70 °C for 1 h to lyse the cell. When the plate temperature dropped to the room temperature, 5 mL of reagent solution constituted of 0.5% melted agarose solution, 0.1 mM NADH, 50 μM TNBT, and 100 mM acetate (pH 4.5) was poured on the treated colonies. After incubating at room temperature for 30 min, the mutants of GsDI were identified based on the formation of black color (Fig. 1b). The plasmids of isolated colonies were then extracted by mini plasmid kit (Tiangen, Beijing, P. R. China) and transformed into Top10 for DNA sequencing and protein overexpression.

Protein expression and purification
The Top10 strains harboring plasmids encoding the GsDI wild type and its mutants were grown in LB medium containing 50 μg/ml kanamycin at 37 °C and induced by adding 0.1 mM IPTG when the absorbance of the OD600 reached 0.8–1.0, and the culture was incubated at 18 °C for 16 h. Cells were harvested by centrifugation and then were resuspended in 20 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer (pH 7.5) containing 50 mM NaCl. After sonication and centrifugation, about 0.1 mM FMN was added into the supernatant for GsDI purification. And the His-tagged protein in the supernatant was loaded onto the column packed with HisPur Ni–NTA Resin (General Electric, NY, U. S.). Purified GsDI wild type and variants were analyzed by SDS-PAGE and used for enzymatic assay. The concentration of protein was determined by the Bradford assay using bovine serum albumin (BSA) as the standard.

Activity assays
The activity of DI was evaluated at 25 °C in 100 mM HEPES buffer (pH 7.3) or 100 mM sodium acetate buffer (pH 5.4 or 4.5) where 0.5 mM VK3 was reduced by 1 mM NADH. One unit of reduction activity was defined as the amount of enzyme required to consume 1 μmol NADH per minute. The decrease in the absorbance band of NADH at 340 nm was measured by a UV–Vis spectroscopy.

Optimum pH and acid-tolerance
The optimal pH was determined in 100 mM acetate buffers (pH 4.0–6.0), 100 mM HEPES buffers (pH 6.0–8.0), or 100 mM Tris–HCl buffers (pH 8.0–10.0). To evaluate the acid stability or thermostability, enzyme solutions (100 mg/L) were incubated at 25 °C in 100 mM sodium acetate buffer (pH 5.4) or 100 mM HEPES buffer (pH 7.3) at 60 °C for different hours, and at fixed time intervals, sample aliquots were tested for activity measurement at 25 °C in 100 mM HEPES buffer (pH 7.3). Three parallel experiments were performed to obtain the standard deviation.

Circular dichroism measurements
Far-UV circular dichroism (CD) measurements in the 190–260 nm region were performed on a Chirascan spectropolarimeter in a quartz cuvette cell with 0.1 cm path length at room temperature. The proteins were diluted in 5 mM acetate buffers (pH 5.0–6.0) or 2.5 mM HEPES buffers (pH 7.3), to a concentration of 0.2 mg/mL. The spectra presented are an average from triplicate measurements. Baseline correction was carried out with the appropriate blank (buffer) during the complete collection time.

Homology model analysis and reverse mutation
The three-dimensional homology model of GsDI wild type and mutant 3–8 were made by SWISS-MODEL based on the crystal structure of Bacillus sp. FMN-dependent NADH-azoreductase (PDB 3w7a, 57% sequence identity) (Yu et al. 2014). Structures were visualized using the PyMOL Molecular Graphics System (Schrödinger, NY, U. S.). The site-directed mutagenesis of mutant 3–8 was used to reverse 6 mutation sites to their original amino acid residues to determine whether mutated amino acids had the impact on activity and acidic stability. The reverse mutation was operated with...
a pair of primers that are 39 bp completely reversed complementary base sequence of original amino acid residues. All primers used in this study for the reverse mutation are listed in Table 1.

### Electrochemical measurement

Cyclic voltammetry was performed by a CHI 1000C potentiostat (CH Instruments Inc., Shanghai, P.R. China) at 25 °C at a scan rate 10 mV/s. A three-electrode system was used for CV with a glassy carbon working electrode, a Ag/AgCl reference electrode, and a Pt wire counter electrode. The same loading amount (0.1 mg or 0.5 mg) of GsDI wild type and mutant 3–8 was used at 25 °C in 100 mM sodium acetate (pH 5.4) in order to demonstrate the improved performance of the EBFC at the lowered pH.

### Results and discussion

#### Library construction and screening

In this study, directed evolution was used to improve the specific activity of the DI at lowered pHs. Error-prone PCR was employed in an attempt to generate the random mutant libraries of GsDI with an estimated average of one to three mutations per round. As an efficient high-throughput screening method a sensitive mediator tetra-nitro blue tetrazolium (TNBT) was applied to identify positive mutant enzymes in this work. The scheme of Petri-dish-based double-layer screening method was
shown as follows: the first layer strain was grown at 37 °C for 16 h, and then, the colonies were treated at 70 °C for 1 h to lyse the cell, degrading reduced coenzymes and deactivating the mesophilic redox enzymes, such as *E. coli* NADH dehydrogenase or thermal labile GsDI.

The second agarose layer containing NADH and TNBT was incubated at room temperature. In fact, TNBT was used in many directed evolution experiment as a sensitive and stable dye (Deimann et al. 1981; Kaewarsa et al. 2017; Shen et al. 2017). Only active stable mutants could reduce the oxidized colorless TNBT to the dark grey color TNBT-formazan (Fig. 1a). As the result, the color density of colonies was closely correlated with residual activity of mutants after heat treatment. Positive mutants with deeper black colors were identified easily for the following plasmid extraction and transformation (Fig. 1b). Three rounds of screening were performed in total, with a stepwise pH drop finally to 4.5. Approximate 20,000 mutants were screened per round with 10–20 mutants exhibiting deeper black colors identified. Finally, positive mutants 1–8 (H37Q, S73T, F105L), 2–12 (H37Q, S73T, F105L, S68T), 3–8 (H37Q, S73T, F105L, S68T, G61S, D74V) were selected for detailed characterization.

**Specific activity of the wild type and mutants**

The corresponding mutation sites, specific activities at different pHs of GsDI wild type and three selected mutants were summarized (Table 2). Remarkably, a new mutant 3–8 yielded a 4-fold increase in the activity at pH 5.4 (0.96 ± 0.12 U/mg) and a 7-fold enhanced activity at pH 4.5 (0.61 ± 0.1 U/mg) compared to wild type. To identify possible reasons for the increased performance of the mutant 3–8, reverse mutation was conducted for six mutation sites of 3–8.

It suggested that the sites had different effects on the activity, among which F105L had the greatest impact on the enzyme activity, as the activity of this reversed mutant dropped to 0.16 ± 0.02 U/mg while others were only slightly decreased (Table 3). The site-saturation mutagenesis of F105 was performed, while no better mutation was found other than F105L (data not shown).

**Optimum pH and stability of the wild type and mutants**

The relative pH profile of wild type and its mutants was obtained by using 100 mM different buffers for appropriate pHs. The optimal pHs of wild type, mutants 1–8, 2–12, 3–8 were found to be ~7.3, 7.0, 7.5 and 7.5, respectively (Fig. 2). Through three rounds of the random mutagenesis, the optimal pH seemed to be similar, probably because the mutant site that plays the main role does not involve charged amino acids. Further, the acid-tolerance of the mutants was evaluated. The pH stability of mutant 3–8 was significantly better than that of wild type and showed a 1.3 times higher in the stability at pH 5.4, allowing us to operate the EBFC with a long time. In fact, all three selected mutants exhibited an improved stability as compared to the wild type.

**Table 2** The specific activity (U/mg) of GsDI wild type and the mutants at different pHs

| Enzymes | New mutations based on wild type | pH 5.4 | pH 4.5 |
|---------|---------------------------------|--------|--------|
| Wild type | – | 0.25 ± 0.04 | 0.09 ± 0.01 |
| M 1–8 | H37Q, S73T, F105L | 0.44 ± 0.01 | 0.26 ± 0.04 |
| M 2–12 | H37Q, S73T, F105L, S68T | 0.80 ± 0.06 | 0.44 ± 0.06 |
| M 3–8 | H37Q, S73T, F105L, S68T, G61S, D74V | 0.96 ± 0.12 | 0.61 ± 0.10 |

**Table 3** The activity of reverse amino acid residue

| Site | Reverse amino acid residue | Sp.act (U mg⁻¹) |
|------|----------------------------|----------------|
| 37   |                            | 0.59 ± 0.05    |
| 61   |                            | 0.44 ± 0.05    |
| 68   |                            | 0.50 ± 0.06    |
| 73   |                            | 0.59 ± 0.09    |
| 74   |                            | 0.47 ± 0.02    |
| 105  |                            | 0.16 ± 0.02    |
| M 3–8|                            | 0.61 ± 0.1     |

Enzyme activity assayed at pH 4.5
at pH 5.4 (Fig. 3a). The mutants could retain greater than 90% activity at pH 5.4 for 20 h, however the wild type lost nearly 30% of its initial activity after 20 h. The residual activity of the wild type and mutants at 60 °C further demonstrated that the mutants maintained or had slightly less activity compared to the wild type at
high temperatures, suggesting that the acidic tolerance improvement of the mutants had nothing to do with their thermostability (Fig. 3b). To estimate the possible misfolding or alterations in the secondary structure of the mutants, circular dichroism (CD) measurements were performed. The far-UV CD spectra of wild type and mutants are shown in Fig. 4. The spectrum of the mutant 3–8 was different in the negative band intensity to that of the wild type at various pHs, being decreased by 1.5-fold. These results demonstrated that the mutants with a disruption of the protein structure might facilitate α helix formation and be benefit for the stability. Figure 5 is a B-factor view of wild type and mutant 3–8, where the closer the color is to red, the larger a B-factor value is. It can be observed that the corresponding B-factor values at the six positions are changed. The value increase indicates a theoretical benefit to the stability of the protein (Sun et al. 2019). Therefore, it can be speculated that the H37Q is likely to help the protein to maintain its activity at the acidic pH. Since histidine is an alkaline amino acid and glutamine is a neutral one, the substitution of glutamine for histidine might influence on the electrostatic force on the protein surface (Fukunishi et al. 2012; Xiang et al. 2019).

The homology model and molecular docking of mutant 3–8

The crystal structure of Bacillus sp. FMN-dependent NADH-azoreductase (AzrC) (PDB: 3w7a) which had 57% amino acid sequence identity was used for the three-dimensional homology model and molecular docking of
mutant 3–8. In this paper, the crystal structure of AzrC complexed with FMN and three different analogue of NAD(P)H were determined. A NAD(P)H analogue containing a triple ring which mimics the nicotinamide adenine, was sandwiched by Tyr127 (a conserved residue) and FMN, in the opposite direction to the isoalloxazine ring of FMN (Yu et al. 2014). The same result was found in our docking structure, as one mutant site, F105L, was located within 3.1 Å around the activity pocket, while other mutated residues were far away (Fig. 6). This finding also agreed with the reverse mutation result mentioned above, that F105L had the greatest impact on the enzyme activity. Obtained from the site-saturation mutagenesis of F105, the F105A exhibited similar kinetics data with F105L at pH 4.5 or pH 4.5 (Table 4). Their binding constants were much smaller than that of the wild type. Therefore, it can be speculated that the F105L mutation allowed for less steric hindrance of a smaller amino acid leucine and made the substrate easier to be combined (Yu et al. 2014, 2019). Based on Fig. 7, it clearly presents that the catalytic channel in mutant 3–8 became larger compared to that of wild type, which would be beneficial for the substrate access and metabolite release. These factors may contribute to the increase in the specific activity of mutant 3–8.

Table 4 Enzyme kinetics for GsDI wild type and mutants F105L and F105A at pH 5.4 or pH 4.5

| Enzyme      | pH 5.4 |                 |                 | pH 4.5 |                 |                 |
|-------------|--------|-----------------|-----------------|--------|-----------------|-----------------|
|             | $K_m$ (mM) | $k_{cat}$ (s$^{-1}$) | $k_{cat}/K_m$ (mM$^{-1}$ s$^{-1}$) | $K_m$ (mM) | $k_{cat}$ (s$^{-1}$) | $k_{cat}/K_m$ (mM$^{-1}$ s$^{-1}$) |
| Wild type   | 0.59 ± 0.12 | 0.18 ± 0.04 | 0.31 ± 0.05 | 1.36 ± 0.21 | 0.05 ± 0.01 | 0.04 ± 0.01 |
| F105L       | 0.31 ± 0.07 | 0.24 ± 0.03 | 0.77 ± 0.08 | 0.78 ± 0.17 | 0.13 ± 0.02 | 0.17 ± 0.02 |
| F105A       | 0.29 ± 0.05 | 0.26 ± 0.05 | 0.89 ± 0.12 | 0.66 ± 0.13 | 0.14 ± 0.02 | 0.21 ± 0.02 |

Fig. 7 Surface presentations of the 3D structure of GsDI wild type and mutant 3–8. Yellow: amino acid residues in WT; red: amino acid residues in mutant 3–8; cyan: the catalytic channel.
In comparison with F105L, G61S, S68T, D74V present slight influence on the stability, while H37Q and S73T are nonsense (Table 3). It is shown that residues G61S, S68T, D74V are closer to the activity pocket (Fig. 6a). Among them, residue G61S is located on alpha helix near the activity pocket and the change from glycine to more hydrophilic serine may benefit the water-soluble substrate reaction. Moreover, in addition to forming hydrogen bonds with F57 and S65, S61 also forms an additional hydrogen bond with F57 in mutant 3–8. The S68T and D74V located in a flexible motif adjacent to the a-helix mentioned above may contribute to more rationalized change of the activity pocket. However, the mechanism that how these residues affect the enzymatic activity and stability remains unclear and requires further efforts.

EBFC performance

In this work, the characterization of an anodic compartment containing DI and other components was evaluated. Since pH 4.5 was too harsh for the protein, the electrochemical test was conducted at pH 5.4. The CV results without or with the substrate G6P suggested that the oxidation potential of VK3 was shifted slightly to a more positive potential, which can be expected as the mediation of VK3 was a proton-coupled electron transfer process. For the wild type or mutant 3–8, with the increase of the amount of GsDI, the oxidation peak showed a significant improvement, indicating that this enzyme played an essential role in the redox process (Fig. 8). The higher oxidation peak height of mutant 3–8 from the CV plot indicated that the activity of the mutant 3–8 was better than the wild type. When 0.1 mg of protein was added, the current of wild type was increased by 4 μA compared to that without substrate G6P, and the mutant 3–8 was increased by 9 μA, 2.3 times higher compared to that of the wild type (Fig. 8a). When the 0.5 mg protein was supplemented under the same condition, the current values of wild type and mutant 3–8 were significantly improved, reaching about 40 μA (Fig. 8b). These results suggested that this engineered GsDI could be a potential candidate for a powerful EBFC.

Conclusion

In conclusion, directed evolution was applied to significantly increase the activity and stability of a GsDI in order to run the EBFC at lowered pHs. After three rounds of screening, the best GsDI mutant 3–8 exhibited a 4- or 7-fold increase in the catalytic activity at pH 5.4 or 4.5 compared to that of the wild type. The EBFC anode equipped with 0.5 mg of mutant 3–8 achieved a maximum current of 40 μA at pH 5.4, more than 1.3-fold that with the same loading of the wild type enzyme. This study demonstrates that directed evolution is a useful strategy for improving the activity and stability of an enzyme and can be used to construct high-performance EBFCs mediated with engineered enzyme mutants.

Abbreviations

DI: Diaphorase; EBFC: Enzymatic biofuel cells; TNBT: Tetra-nitro blue tetrazo- lium; VK3: Vitamin K3; IPTG: Isopropyl-β-d-thiogalactopyranoside; OD: Optical density; PCR: Polymerase chain reaction; SDS: Sodium dodecyl sulfate; PAGE: Polyacrylamide gel electrophoresis; POE-PCR: Prolonged overlap extension PCR; BSA: Bovine serum albumin; NAD(P)+: Nicotinamide adenine dinucleo- tide phosphate; G6P: Glucose 6-phosphate; G6PDH: G6P dehydrogenase; 6PGDH: 6-Phosphogluconate dehydrogenase; WT: Wild type.

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Authors’ contributions

CM and ML performed the experiments. CY and ZZ conceived the idea and supervised the research. CM and ZZ wrote and revised the manuscript. All authors read and approved the final manuscript.
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Ethical approval and consent to participate
Not applicable.

Consent for publication
All of the authors have read and approved to submit it to Bioresources and Bioprocessing.

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

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