Active Expression of Membrane-Bound L-Amino Acid Deaminase from *Proteus mirabilis* in Recombinant *Escherichia coli* by Fusion with Maltose-Binding Protein for Enhanced Catalytic Performance

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Abstract: L-amino acid deaminases (LAADs) are membrane flavoenzymes that catalyze the deamination of neutral and aromatic L-amino acids to α-keto acids and ammonia. LAADs can be used to develop many important biotechnological applications. However, the transmembrane α-helix of LAADs restricts its soluble active expression and purification from a heterologous host, such as *Escherichia coli*. Herein, through fusion with the maltose-binding protein (MBP) tag, the recombinant *E. coli* BL21 (DE3)/pET-21b-MBP-PmLAAD was constructed and the LAAD from *Proteus mirabilis* (PmLAAD) was actively expressed as a soluble protein. After purification, the purified MBP-PmLAAD was obtained. Then, the catalytic activity of the MBP-PmLAAD fusion protein was determined and compared with the non-fused PmLAAD. After fusion with the MBP-tag, the catalytic efficiency of the MBP-PmLAAD cell lysate was much higher than that of the membrane-bound PmLAAD whole cells. The soluble MBP-PmLAAD cell lysate catalyzed the conversion of 100 mM L-phenylalanine (L-Phe) to phenylpyruvic acid (PPA) with a 100% yield in 6 h. Therefore, the fusion of the MBP-tag not only improved the soluble expression of the PmLAAD membrane-bound protein, but also increased its catalytic performance.

Keywords: L-amino acid deaminases; membrane-bound protein; fusion protein; soluble expression; maltose-binding protein tag

1. Introduction

L-amino acid deaminases (LAADs; EC 1.4.3.2) belong to a family of amino acid dehydrogenases that catalyze the formation of α-keto acid from L-amino acid and release ammonium and H2O as well [1–4]. LAADs have been identified in several bacterial genera including *Proteus*, *Providencia*, and *Morganella* [2,5]. Some bacteria, such as *Proteus mirabilis*, express two types of LAAD which have similar sequences but have distinct substrate preferences: type I prefers aliphatic and aromatic amino acids, while type II shows significant activity with basic amino acids such as histidine and arginine [2,3,6–9]. Both types of LAAD contain a single membrane-spanning helix, which is anchored to the cytomembrane surface through the N-terminal transmembrane helix [2,9].
Recently, membrane-bound LAADs have been the focus of studies due to their potential biotechnological applications in industry [8–11]. A series of LAAD-based technologies has been developed to efficiently transform L-amino acids to α-keto acids such as ketoglutaric acid, methylthiobutyric acid, and phenylpyruvic acid (PPA) in an eco-friendly manner [3,7,12–16]. Among them, PPA is an important multi-functional organic acid which is widely used in the pharmaceutical, food, and chemical industries [6,17]. LAADs are also used to produce pure D-amino acids by removing L-amino acids from racemic mixtures [18–21]. Other potential applications, such as their use as biosensors of L-amino acid concentration in the medical and food industries, have also been proposed [22–24].

As the potential applications of LAADs have increased, many researchers have focused on the study of membrane-bound LAADs. Baek et al. cloned, expressed, and characterized a type II L-amino acid deaminase from *P. mirabilis* KCTC 2566 [2,14]. To facilitate the purification of a membrane-bound LAAD from *P. mirabilis*, Liu et al. removed the N-terminal transmembrane region (from 21 to 87 nucleotides) from the enzyme to block its binding to the membrane [3,6,10]. Similarly, to study the crystal structure of a membrane-bound LAAD from *P. vulgaris* (PvLAAD), Ju et al. also removed the N-terminal transmembrane region of the enzyme to obtain purified PvLAAD [9]. However, the LAAD was still expressed in the form of an inclusion body. Although it seems that the LAAD whole-cell biocatalysts are more suitable for catalysis, in order to better study the applicable feasibility of the enzyme, it is necessary to improve its soluble expression.

The maltose-binding protein (MBP) is the product of the *malE* gene in *Escherichia coli* K12. The protein consists of 396 amino acids [25–30] and has a molecular weight of 40 KDa [25,26]. The N-terminal of the MBP has a signal peptide sequence of 26 amino acids, which could guide the final expression of the MBP to locate in the periplasm [25,27]. According to the directional expression characteristics of the MBP protein, researchers constructed an MBP tag to promote the heterologous soluble expression of some proteins [27,30]. MBP has been used with a few membrane proteins and the MBP-tag has been employed for the membrane-binding protein to locate in the periplasm [25].

In this study, we fused the LAAD from *P. mirabilis* (PmLAAD) with the MBP-tag for soluble expression in recombinant *E. coli*. Then, the MBP-tag fused LAAD (MBP-PmLAAD) was expressed and purified as a soluble protein. After protease digestion and purification to remove the MBP-tag, the soluble purified PmLAAD was obtained. Then, the catalytic properties of the MBP-PmLAAD fusion, the soluble PmLAAD, and the MBP-PmLAAD cell lysate were studied. Finally, the catalytic efficiency of the MBP-fused LAAD and the membrane-binding LAAD were compared.

2. Results and Discussion

2.1. Construction of MBP-PmLAAD Recombinant Protein Expression Plasmids

The PCR products of PmLAAD were inserted into the vector pET-21b. The 1416 bp PmLAAD (Figure 1B) open reading frame was amplified from *P. mirabilis* KCTC 2566. The MBP-PmLAAD vector, designed to produce PmLAAD proteins in the cytoplasm, was constructed by fusing the *PmLAAD* gene to the MBP gene driven by a T7 promoter (Figure 1A). The MBP was generally used to improve the solubility of the fusion proteins in the *E. coli* system and expressed at the N-end of heterologous MBP-PmLAAD products. The vector encoding the MBP-PmLAAD products was used to transform *E. coli* BL21 (DE3) and produce fusion proteins.
2.2. Expression and Purification of MBP-PmLAAD

The MBP-PmLAAD and PmLAAD expressions were induced at an OD_{600} nm of 0.8 with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG), at 17 °C for 12 h. As shown in Figure 2A, when the PmLAAD was not fused to the MBP-tag, the molecular weight of the PmLAAD was 52 kDa. Only a small portion of the PmLAAD was present in the soluble fraction and most of it was found in the membrane fraction. These results are in agreement with Baek et al., who expressed a second L-amino acid deaminase type named Pm1 from *P. mirabilis* KCTC 2566 and reported that most of the Pm1 protein was present in the membrane fraction [2]. The molecular weight of the MBP was 41 kDa, and when the PmLAAD was fused to the MBP-tag, the molecular weight of the corresponding MBP-PmLAAD product was 93 kDa (Figure 2B). Lane 1 in Figure 2C also shows the protein band of purified MBP-PmLAAD, which is consistent with the MBP-PmLAAD protein band in Figure 2B. The SDS-PAGE showed that most of the MBP-PmLAAD protein was in the soluble fraction, thus the fusion to the MBP released the LAAD from the cell membranes and produced the soluble expression of the protein. Compared with previously reported methods of removing the N-terminal transmembrane region from the enzymes, such as the PmLAAD and PvLAAD which were still expressed in the form of an inclusion body [3,9], the fusion with the MBP-tag here could release the membrane binding state of the PmLAAD and promote its soluble expression.

The MBP-PmLAAD fusion protein was purified using affinity chromatography with a Ni-column; the results of the SDS-PAGE analyses are shown in Figure 2C. In order to better study the catalytic properties of the PmLAAD, an enzymatic digestion of the MBP-PmLAAD fusion with a rTEV protease followed by a purification of the reaction liquid with a Ni-column was used to remove the MBP-tag. The SDS-PAGE showed that once the MBP-tag was removed, the PmLAAD still remained a soluble protein. The molecular weight of the purified PmLAAD was 52 kDa.

To investigate the effect of the MBP fusion on the catalytic activity of PmLAAD, the activities of the PmLAAD and MBP-PmLAAD cell lysates, the MBP-PmLAAD fusion protein, and the purified PmLAAD were analyzed (Table 1). The purified MBP-PmLAAD fusion protein had a low activity of 0.025 μmol/mg PmLAAD·min using L-Phe as a substrate. The activity of the MBP-PmLAAD cell lysate was 0.17 μmol/mg PmLAAD·min, which was 2.2 times higher than the activity of the PmLAAD cell lysate. After removing the MBP-tag with the rTEV protease, the activity of the purified PmLAAD was not detected, which may be due to the instability of the PmLAAD without the fusion tag and/or

![Figure 1](image-url)  
**Figure 1.** Construction of the recombinant plasmid of pET-21b-MBP-PmLAAD. (A) A schematic representation of the pET-21b-MBP-PmLAAD plasmid. The *PmLAAD* gene was inserted into the pET-21b-MBP vectors. (B) PCR amplification of the *PmLAAD* gene.
membrane components. Since the fusion protein still has some activity, the catalytic properties of the fusion protein could be further investigated.

![Figure 2](image1.png)

**Figure 2.** The SDS-PAGE of the PmLAAD and MBP-PmLAAD cell lysates and the purified MBP-PmLAAD. (A) *E. coli* BL21 (DE3)/pET-21b cell lysate (pET-21b), total cell lysate (T) and soluble fraction (S) of *E. coli* BL21 (DE3)/pET-21b-PmLAAD. The samples were generated from 50 mg/mL *E. coli* BL21 (DE3)/pET-21b cells and 50 mg/mL *E. coli* BL21 (DE3)/pET-21b-PmLAAD cells. (B) Total cell lysate (T) and soluble fraction (S) of *E. coli* BL21 (DE3)/pET-21b-MBP-PmLAAD. Lane T, *E. coli* BL21 (DE3)/pET-21b-MBP-PmLAAD cell lysate before centrifugation; lane S, *E. coli* BL21 (DE3)/pET-21b-MBP-PmLAAD cell lysate after centrifugation. The samples were generated from 50 mg/mL *E. coli* BL21 (DE3)/pET-21b-MBP-PmLAAD cells. (C) Purified and rTEV protease-treated MBP-PmLAAD fusion protein. Lane 1, proteins purified with Ni-column affinity chromatography; lane 2, rTEV protease-treated MBP-PmLAAD fusion protein, i.e., the mixture of MBP-PmLAAD, PmLAAD, and MBP; lane 3, the sample after removal of PmLAAD by secondary purification with a Ni-NTA column, i.e., the mixture of MBP-PmLAAD and MBP; lane 4, purified PmLAAD.

| Table 1. Enzyme activities of the cell lysate, MBP-PmLAAD fusion protein, and PmLAAD. |
|-------------------------------------------------|
| **Specific activity (μmol/mg PmLAAD min)**       | **PmLAAD** | **MBP-PmLAAD** | **MBP-PmLAAD Purified Enzyme** | **PmLAAD Purified Enzyme** | **E. coli pET-21b Cell Lysate** |
| **Cell Lysate**                                   | **Cell Lysate** | **Purified** | **Enzyme** | **Purified** | **Enzyme** | **Purified** | **ND b** | **ND b** |
| 0.076                                            | 0.17        | 0.025        | ND b        | ND b        | ND b        |

*The reaction time was 30 min. b Not detected.*

2.3. Biochemical Characterization of the MBP-PmLAAD Fusion Enzyme

The effect of the pH and temperature on the MBP-PmLAAD fusion enzyme was investigated in order to study the catalytic properties of the enzyme. The range of temperatures at which the purified
MBP-PmLAAD showed activity was 30 to 50 °C. The enzyme activity increased as the temperature increased from 30 to 37 °C and then decreased with a temperature increase from 37 to 50 °C, reaching its maximum activity at 37 °C (Figure 3A). The purified MBP-PmLAAD was active at pH ranging from 7 to 10 with its optimal activity at pH 8.0 (Figure 3B). The activity of the MBP-PmLAAD decreased with a further increase in pH from 9 to 10. Subsequently, the stability of the MBP-PmLAAD was studied. As shown in Figure 3C, the MBP-PmLAAD fusion enzyme was relatively stable under the reaction conditions during the reaction for 10 h, even after the reaction reached equilibrium.

Figure 3. The effect of pH, temperature and storage time on the activity of the purified MBP-PmLAAD fusion enzyme. (A) The effect of temperature on fusion enzyme activity. (B) The effect of pH on fusion enzyme activity. (C) The stability of the purified MBP-PmLAAD.

The results show that the catalytic efficiency of the purified enzyme was much lower than the catalytic efficiency of the crude enzyme solution (Table 1). Therefore, the effect of different catalyst concentrations on the catalytic efficiency was explored. Figure 4B shows that for the same concentration, the amount of MBP-PmLAAD in the purified enzymes was much higher than in crude enzymes. However, the catalytic efficiency of the cell lysate with the same concentration was much higher than the catalytic efficiency of the purified enzyme. As shown in Figure 4A, the conversion of 30 mM L-Phe catalyzed by 4.25 mg/mL of the cell lysate (crude enzyme) was 100% in 2 h, while the conversion of 30 mM L-Phe catalyzed by 17 mg/mL of the pure enzyme was only 33% in 4 h. Therefore, the activity of the MBP-PmLAAD cell lysate was much higher than the one of the purified MBP-PmLAAD fusion enzyme, although the content of the target protein in the pure enzyme suspension was higher. At the same time, the mixture of 3 mg/mL purified MBP-PmLAAD and 50 mg/mL E. coli BL21 (DE3)/pET-21b cell lysate was also used to catalyze 30 mM L-Phe. As shown in Figure 4A, when the purified MBP-PmLAAD was mixed with the E. coli BL21 (DE3)/pET-21b cell lysate, 30 mM of L-Phe was completely transformed to PPA in 2 h. As the PmLAAD was the membrane-binding enzyme, the membrane components in the MBP-PmLAAD cell lysate could promote the catalytic reaction of MBP-PmLAAD. Similarly, it has been proposed that the membrane-bound LAADs are associated with the electron transport chain on the bacterial membrane and electrons are transferred to cytochrome oxidases to reduce O2 to H2O [9,12,31]. This may be the reason why the activity of the MBP-PmLAAD cell lysate was higher than the activity of the purified enzyme. Therefore, by comparing the catalytic efficiency between the MBP-PmLAAD purified enzyme and the cell lysate, the membrane components in the cell lysate were found to promote the catalytic efficiency of the MBP-PmLAAD. Although the purified MBP-PmLAAD had low enzyme activity, considering the feasibility of the LAAD-catalyzed conversion of L-Phe to PPA, the fusion of the MBP-tag to the PmLAAD promoted the soluble active expression of the PmLAAD in the E. coli system, and the obtained cell lysate system was more favorable for enhanced catalytic performance. At the same time, the fusion expression of the MBP-tag to the PmLAAD enabled us to obtain the cell lysate system which can be used for catalysis.
Figure 4. The effect of biocatalyst concentration and *E. coli* BL21 (DE3)/pET-21b cell lysate on the catalytic efficiency. (A) The catalytic conversion of L-Phe to PPA with different concentrations of the MBP-PmLAAD purified enzyme and cell lysate. The mixture of the MBP-PmLAAD and pET-21b cell lysate was composed of 3 mg/mL purified MBP-PmLAAD and 50 mg/mL *E. coli* BL21 (DE3)/pET-21b cell lysate. (B) The SDS-PAGE analysis of different concentrations of the MBP-PmLAAD purified enzyme and cell lysate. The reactions were all conducted in a 50 mM, pH 8.5 Tris-HCl buffer at 37 °C.

2.4. Conversion of L-Phe to PPA with MBP-PmLAAD Cell Lysate

In order to test the catalytic properties of the MBP-PmLAAD, the conversions of 50 mM and 100 mM L-Phe were catalyzed with the MBP-PmLAAD cell lysate and the PmLAAD whole-cell biocatalyst, respectively. As shown in Figure 5, when the catalyst was the MBP-PmLAAD cell lysate, 50 mM of L-Phe was fully converted to PPA after 2 h, while 100 mM of L-Phe was fully converted to PPA in 6 h. However, when 50 mM of L-Phe was catalyzed with the PmLAAD whole cell, the conversion to PPA was 66.3%. It is obvious that the catalytic efficiency of the PmLAAD whole cell was lower than the catalytic efficiency of the MBP-PmLAAD crude enzyme solution. Therefore, the MBP-tag allowed the soluble expression of the membrane-binding PmLAAD, and the soluble expression of the MBP-PmLAAD fusion protein would contribute to improving the catalytic activity of the membrane-binding PmLAAD.

Figure 5. The time course of the formation of PPA with different biocatalysts. A: the biocatalyst was the MBP-PmLAAD cell lysate at 50 mg/mL, with 100 mM L-Phe; B: the biocatalyst was the MBP-PmLAAD cell lysate at 50 mg/mL, with 50 mM L-Phe; C: the biocatalyst was the PmLAAD whole cell
at 100 mg/mL, with 50 mM L-Phe. The reactions were all conducted in a 50 mM, pH 8.5 Tris-HCl buffer at 37 °C.

3. Materials and Methods

3.1. Materials

The plasmid pET-21b-MBP and the PmLAAD gene from P. mirabilis (GenBank accession no. EU669819.1) were stored in our laboratory [2,4]. The enzymes, oligonucleotides, and other reagents for DNA cloning and amplification were obtained from the Takara-Bio Co., Kawasaki, Japan and the Novagen Co., Madison, WI, USA. The L-phenylalanine (L-Phe) and PPA were purchased from the Sinopharm Chemical Reagent Co., Ltd., Shanghai, China. The rTEV protease was purchased from Beijing Solarbio Science & Technology Co., Ltd., Beijing, China. The Bradford Protein Assay Kit was purchased from Beijing Solarbio Science & Technology Co., Ltd., Beijing, China.

3.2. Construction of the Expression Plasmid pET-21b-MBP-PmLAAD

The PmLAAD gene fragment flanked by the recognition sequences for Nde I and Sal I was inserted into the expression vector pET-21b-MBP, encoding an N-terminal maltose-binding protein, followed by a recognition site for the rTEV protease under the control of a T7 promoter. The primers used for plasmid construction were 5′-CIGTTTTTCAGACATATGATGGCAATAAGTAGAAGAAAATT TATT-3′ and 5′-ATCCCTCAGAAAGCTTTGCGTCGACTTAGAAACGATACAGACTAAATGTGT-3′. Then, the constructed plasmid pET-21b-MBP-PmLAAD was transformed into E. coli BL21 (DE3) cells. After plating, individual colonies were picked and the plasmids were sequenced.

3.3. Expression and Purification of MBP-PmLAAD Fusion Proteins

Recombinant E. coli BL21 (DE3) was grown in 1 L of Luria–Bertani medium, containing 100 mg/mL ampicillin at 37 °C. The culture was induced by the addition of isopropyl β-D-1-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM when the optical density (λ = 600 nm) was 0.6–0.8. Furthermore, the culture was incubated for an additional 16 h at 17 °C and 200 rpm. After centrifugation at 6000×g at 4 °C for 20 min, the recombinant cells were washed with a potassium phosphate buffer.

To purify the MBP-PmLAAD, the recombinant E. coli BL21 (DE3) cells were resuspended in a lysis buffer containing 20 mM Tris-HCl and 150 mM NaCl (pH 7.5). The cell lysates were produced with a high-pressure homogenizer (APV, Lubeck, Germany) and then centrifuged at 12000×g for 30 min at 4 °C to remove cell debris. The proteins were purified by Ni affinity chromatography with an Äkta Purifier 10 (GE Healthcare, Pittsburgh, PA, USA). The gradient elution buffer for the Ni-NTA column was composed of 20 mM Tris-HCl (pH 7.5), 150 mM NaCl, and 0–500 mM imidazole. The imidazole in the protein solution was removed using a PD-10 desalting column (GE Healthcare, USA). The MBP-tag was removed with the rTEV protease by adding 1000 μg of MBP-PmLAAD fusion protein and 10 μg of rTEV Protease to 1000 μL of buffer (50 mM NaH₂PO₄, 150 mM NaCl, 1 mM EDTA, 1 mM DTT, 10% glycerol, pH 8.0), then the reaction mixture was kept at 16 °C for 6 h. Subsequently, the MBP-tag and rTEV protease were removed with a Ni-NTA column. The protein concentrations were determined using a Bradford Protein Assay Kit (Solarbio Science & Technology, Beijing, China).

3.4. Activity Assay

The PmLAAD and MBP-PmLAAD cell lysates, the purified PmLAAD enzyme, and the MBP-PmLAAD purified enzyme were incubated in 50 mM Tris-HCl buffer (pH 8.0) with 30 mM L-Phe in a final volume of 5.0 mL at 37 °C. After reaction for 30 min, an equal volume of 10% (v/v) trichloroacetic acid solution was added to stop the reaction. The mixture was centrifuged at 12000×g for 5 min, and a 15 μL aliquot of the supernatant was added to 1 mL of ferric chloride solution and
incubated for 2 min at room temperature. The concentration of the PPA was determined by measuring the OD640 nm. One PmLAAD unit corresponds to the amount of enzyme that generates 1 μmol of PPA product per minute [6].

3.5. Effect of pH, Temperature and Storage Time on the Activity of MBP-PmLAAD

To investigate the effect of the reaction temperature on the MBP-PmLAAD catalyzing oxidative deamination of L-Phe, 30 mM L-Phe and 3.1 mg/mL MBP-PmLAAD were added to 2 mL Tris-HCl buffer (50 mM, pH 7.5), and the reaction was conducted at 220 rpm and different temperatures (30 °C, 37 °C, 40 °C, 45 °C, and 50 °C) for 30 min. The effect of the initial pH on the MBP-PmLAAD catalyzing oxidative deamination of L-Phe was studied using the same mixture of L-Phe and MBP-PmLAAD described above and added to 2 mL 50 mM Tris-HCl buffer with a pH varying from 7 to 10; the reaction was conducted at 220 rpm and 37 °C for 30 min. To investigate the stability of the MBP-PmLAAD, the MBP-PmLAAD was stored in a Tris-HCl buffer (50 mM, pH 8.5) at 37 °C for 1–12 h. Then, the activity of the MBP-PmLAAD was determined. The reaction buffer was a Tris-HCl buffer (50 mM, pH 8.5), the L-Phe was 30 mM, and the reaction was conducted at 220 rpm and 37 °C for 30 min. The PPA generated in the reaction process was measured using ferric chloride solution.

3.6. Conversion of L-Phe to PPA Using the MBP-PmLAAD Fusion Protein

The effect of the biocatalyst concentration on catalytic efficiency was assessed using different concentrations of the MBP-PmLAAD cell lysate (4.25–17 mg/mL) and the purified enzyme (3–17 mg/mL) that were added to the 2 mL, 50 mM, pH 8.5 Tris-HCl buffer; the L-Phe concentration was 30 mM. The effect of the E. coli BL21 (DE3)/pET-21b cell lysate on the catalytic efficiency was assessed using the mixture of 3 mg/mL purified MBP-PmLAAD and 50 mg/mL E. coli BL21 (DE3)/pET-21b cell lysate, that were added to the 2 mL, 50 mM, pH 8.5 Tris-HCl buffer; the L-Phe concentration was 30 mM. The reaction was conducted at 220 rpm and 37 °C and was stopped by adding an equal volume of 10% (w/v) trichloroacetic acid solution. The amount of PPA generated in the reaction was determined using ferric chloride solution. To compare the catalytic efficiency of the MBP-PmLAAD cell lysate and the PmLAAD whole-cell catalyst for catalyzing the transformation of L-Phe, 50 mg/mL of the MBP-PmLAAD whole-cell lysate or 100 mg/mL of the PmLAAD whole-cell biocatalyst was added to the 2 mL, 50 mM, pH 8.5 Tris-HCl buffer. The L-Phe concentrations were 50 and 100 mM, respectively. The reaction was conducted at 220 rpm and 37 °C and was stopped by adding an equal volume of 10% (w/v) trichloroacetic acid solution. The amount of PPA generated in the reaction was determined using ferric chloride solution.

4. Conclusions

In summary, the membrane-bound protein PmLAAD was successfully expressed as a soluble active protein and therefore lost its membrane-binding status. The key factor in this study was the usage of an MBP-fusion protein as a solubility tag. Then, the MBP-PmLAAD fusion protein was purified and the purified soluble PmLAAD was obtained after the rTEV protease digestion. The activities of the MBP-PmLAAD cell lysate, purified MBP-PmLAAD, and purified PmLAAD were determined. The purified MBP-PmLAAD fusion protein showed a low activity with L-Phe, while the purified PmLAAD lost its activity after the rTEV protease digestion. At the same time, the activity of the MBP-PmLAAD cell lysate was much higher than that of the purified MBP-PmLAAD. By comparing the catalytic process of the MBP-PmLAAD cell lysate and the PmLAAD whole cells, we found that fusion with MBP could improve the active expression of PmLAAD and thus enhance catalytic performance. L-Phe at 100 mM could be fully converted to PPA in 6 h by 50 mg/mL of MBP-PmLAAD-involved cell lysate. Therefore, the fusion expression of an MBP-tag to PmLAAD enables us to obtain a cell lysate system which can be used for highly efficient biocatalysis. In addition, this strategy provides a feasible way of improving the soluble expression of membrane-bound LAADs, which would be helpful in promoting the study of catalytic mechanisms of the enzyme.
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