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Keywords: Bioreduction, 3-oxoacyl-acyl-carrier-protein reductase, Short chain dehydrogenase, Duloxetine
Stereoselective bioreduction of ethyl 3-oxo-3-(2-thienyl) propanoate using the short-chain dehydrogenase/reductase ChKRED12

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Abstract

Ethyl (S)-3-hydroxy-3-(2-thienyl) propanoate ((S)-HEES) acts as a key chiral intermediate for the blockbuster antidepressant drug duloxetine, which can be achieved via the stereoselective bioreduction of ethyl 3-oxo-3-(2-thienyl) propanoate (KEES) that contains a 3-oxoacyl structure. The sequences of the short-chain dehydrogenase/reductases from Chryseobacterium sp. CA49 were analyzed, and the putative 3-oxoacyl-acyl-carrier-protein reductase, ChKRED12, was able to stereoselectively catalyze the NADPH-dependent reduction to produce (S)-HEES. The reductase activity of ChKRED12 towards other substrates with 3-oxoacyl structure were confirmed with excellent stereoselectivity (>99% ee) in most cases. When coupled with a cofactor recycling system using glucose dehydrogenase, the ChKRED12 was able to catalyze the complete conversion of 100 g/L KEES within 12 h, yielding the enantiopure product with >99% ee, showing a remarkable potential to produce (S)-HEES.

Keywords: Bioreduction; 3-oxoacyl-acyl-carrier-protein reductase; Short-chain dehydrogenase; Duloxetine
1. Introduction

Chiral compounds are very useful intermediates for the synthesis of numerous pharmaceuticals, agricultural chemicals, and specialty materials [1-5]. Among them, ethyl (S)-3-hydroxy-3-(2-thienyl) propanoate ((S)-HEES) is a useful chiral building block which is used in the synthesis of duloxetine [6-8] ((S)-(+) N-methyl-3-(1-naphthyloxy) -3-(2-thienyl) propylamine) which is a blockbuster antidepressant drug [9-11]. (S)-HEES is produced via the asymmetric reduction of ethyl 3-oxo-3-(2-thienyl) propanoate (KEES). In recent years, the enzymatic asymmetric reduction of prochiral ketones for the preparation of optically active alcohols has gained attention [5, 12-14]. The optically pure (S)-HEES can be prepared by biocatalytic reductions. However, the substrate concentration applied is limited with a typical loading of 10 g/L because at higher loadings often lead to the feedback inhibition [6, 8]. Therefore, it would be necessary to search for a biocatalyst with excellent enantioselectivity and high substrate tolerance for efficient synthesis of (S)-HEES.

3-oxoacyl-(acyl carrier protein) reductase (3-oxoacyl-ACP reductase, EC 1.1.1.100) is a short-chain dehydrogenase/reductase that catalyzes the pyridine nucleotide-dependent reversible reduction of a 3-oxoacyl form of ACP to its hydroxyl product (Fig.1) as the first reductive step in de novo fatty acid biosynthesis[15, 16]. The 3-oxoacyl-ACP reductase can also catalyze the reduction of the β-keto group of unsaturated acyl chain during the biosynthesis of unsaturated fatty acid in E. coli [17]. The stereoselectivity of dehydrogenases/reductases can be predicted by the Prelog rule [18]; this rule indicates that an enzyme has a large and a small pocket that form the active site in which the substrate binds and controls the stereoselectivity of the product based on the geometry of the substrate and coenzyme NAD(P)H. In 3-oxoacyl-ACP reductase, the hydride ion (H−) of NADPH is transferred from the si-face of ketones (3-oxoacyl-ACP) and yield anti-Prelog chiral alcohols ((R)-3-hydroxyacyl-ACP). This type of enzyme is required for the pharmaceutical industry [19, 20]. The target substrate (KEES) contains 3-oxoacyl structure (Fig. 1), it is expected to be reduced by 3-oxoacyl-ACP reductase. Using the Prelog rule to predict the stereoselectivity of 3-oxoacyl-ACP reductase toward some 3-oxoacyl ester substrates, it will be transformed to (S)-HEES that is anti-Prelog chiral alcohol (Fig. 1).

In a previous study, we discovered 27 microbial short-chain dehydrogenase/reductases from the genome of Chryseobacterium sp. CA49 and expressed them in E.
coli [21]. Because the target substrate KEES contains a 3-oxoacyl structure (Fig. 1), the sequences of these SDRs were then analyzed to seek the reductase that had a high sequence identity with 3-oxoacyl-ACP reductase and identified ChKRED12 to be an excellent catalyst to produce (S)-HEES with excellent enantioselectivity (>99% ee). A substrate concentration of up to 100 g/L was completely reduced with no product inhibition observed.

2. Materials and Methods

2.1 General

Substrates 2a-7a (Table 1) were purchased from Alfa-Aesar (Tianjin, China). Substrate 1a (Table 1) was synthesized from 2-acetylthiophene following established methods [22, 23]. The racemic alcohols were prepared by reducing the ketones with sodium borohydride [23, 24]. NADH, NADPH, NADP+, isopropyl β-D-1-thiogalactopyranoside (IPTG), and glucose dehydrogenase (GDH) were purchased from Sigma (St. Louis, USA). All other reagents were obtained from general commercial suppliers and used without further purification.

2.2 Expression and purification of ChKRED12 in Escherichia coli BL21 (DE3)

The plasmid, pET-28a-chKRED12, encoding ChKRED12 [21] was transformed into E. coli BL21(DE3) (Novagen, Madison, WI, USA) competent cells and selected on Luria Bertani (LB) agar plates containing 50 μg kanamycin/mL. Single colonies were grown overnight at 37 °C in LB medium containing 50 μg kanamycin/mL. Then, 2 mL of the culture was transferred into 200 mL TB medium containing 50 μg kanamycin/mL for protein expression. When OD600 of the culture reached 0.8, IPTG was added to a final concentration of 1.0 mM and the cultivation was continued at 30 °C for 5 h. After harvest by centrifugation, Cells were washed twice using 20 mM sodium phosphate buffer (pH 8.0), and resuspended in buffer A (20 mM sodium phosphate buffer, 500 mM NaCl and 10 mM imidazole, pH 8.0).

After disruption with a homogenizer (Nano, ATS-AH100B, ATS Engineering Inc., Canada), the cell debris was removed by centrifugation at 2×10⁴ g for 25 min at 4 °C. The resulting supernatant can be directly used as crude enzyme extracts or lyophilized to provide a dry powder of the crude enzyme. For purification, it was loaded onto Ni²⁺-nitrilotriacetic acid columns (Bio-Rad) pre-equilibrated with buffer A. Then, the
enzyme was eluted with buffer A containing a gradient of imidazole ranging from 10 to 500 mM at a flow rate of 1 mL/min. The fractions containing the target protein were collected and dialyzed against 20 mM potassium phosphate buffer (pH 7.0). Purified enzymes were analyzed by SDS-PAGE and used for enzymatic assays. The molecular mass of the reductase was determined by elution on a Sephacryl S-200 column (1.6×90 cm) calibrated with molecular mass standards (Sigma). Protein estimations were done with a commercial BCA Protein Assay kit with bovine serum albumin as a standard (Tiangen, Beijing, China). The purified enzyme solution and lyophilized powder were stored at −80°C.

2.3 Measurement of enzyme activity

All reactions were performed in triplicate. The reaction mixture comprised 100 mM potassium phosphate buffer (pH 7.0), 10 mM NADPH and 10 mM substrate in a total volume of 1.0 ml. The reaction was carried out at 30 °C, and terminated by extraction with 1.0 ml ethyl acetate after 10 min. The resulting organic phase was subjected to gas chromatographic (GC) analysis to determine the conversion rate and ee value. One unit of the enzyme activity was defined as the amount of enzyme that catalyzes the production of 1μmol (S)-3-Hydroxy-3-(2-thienyl) propanoate per minute.

To determine the pH optimum, standard assay method was applied except that different buffers were used for different pH ranges, which included sodium citrate (pH 5.0-6.0), potassium phosphate (pH 6.0-8.0), Tris-HCl (pH 7.0-9.0) and sodium carbonate (pH 9.0-10.0). The optimum temperature for ChKRED12 was determined at various temperatures ranging from 10 to 50 °C for 10 min. To investigate the thermostability of ChKRED12, the enzyme (1 mg/ml) was incubated at 35, 40, or 45 °C, withdrawn at intervals, cooled in ice, and the residual activity was assayed following the standard assay method.

The steady-state kinetic parameters toward the substrate 1a were investigated using standard assay method except varied substrate concentrations ranging from 0.25 to 50 mM for 5 min. The kinetic parameters toward cofactors were investigated in the presence of 10 mM 1a for 2 min at varied cofactor concentrations ranging from 0.1 to 10 mM for NAD⁺, and 1 to 200 μM for NADP⁺, respectively, and glucose dehydrogenase was employed for coenzyme regeneration cycle (glucose dehydrogenase 10 U/ml, glucose 100 mM). Data were fitted to the Michaelis-Menten
equation using Graph-Pad Prism v5.0 (GraphPad Software, San Diego, CA, USA) to generate estimates of $K_m$ and $k_{cat}$ values.

2.4 Preparative-scale bioreduction

The reaction was carried out at 30 °C in 50 ml potassium phosphate buffer (100 mM, pH 8.0) containing 0.2 mM NADP+, 12% (w/v) glucose, GDH (10 U/ml), crude enzyme extract or lyophilized powder of the crude enzyme (4 mg/ml, 8.4 U/mg) and the substrate concentration was 25-100 g/l. The pH of the reaction mixture was monitored and maintained at 7.0-8.0 by the addition of NaOH (1 M). To monitor the time-course of the biotransformation, samples were taken at intervals for analysis. The reaction was terminated by extraction with ethyl acetate, then the combined organic extracts were dried with anhydrous sodium sulfate, concentrated under reduced pressure and purified using silica gel column chromatography eluted with petroleum ether/ethyl acetate (10:1, v/v). The products were identified by 1H NMR analysis. The absolute configuration of the product was determined by comparing the optical rotation with the literature data.

2.5 Biotransformation of various ketones and product analysis

Standard reaction conditions were followed to convert 10 mM substrates in the presence of a coenzyme regeneration cycle (glucose dehydrogenase 10 U/mL, glucose 100 mM). The purified enzyme was applied at a concentration of 1 U/mL. After 1 h incubation at 30 °C, the reaction was terminated by extraction with ethyl acetate. The organic phase was dried with anhydrous sodium sulfate, concentrated under reduced pressure, and analyzed. The ee value and conversion of the product were determined by chiral GC analysis. Chiral GC analysis was performed on a Fuli 9790 II GC system connected to a flame ionization detector using a CP-Chirasil-DEX CB column (Varian, USA). The injector and detector were set at 260 °C and 280 °C, respectively. The column temperature was set at 160 °C ($1a$, $2a$), 120 °C ($3a$) and 90 °C ($5a$). The product of $4a$ was acetylated following literature method [25] and then analyzed with chiral GC.

3. Results and discussion

3.1 Sequence analysis of ChKRED12

The sequence comparisons have shown that the typical coenzyme-binding pattern for SDRs (TGXXXGXG, Gly motif, where ‘X’ denotes any residues) is located at the
N-terminus of the protein chain [26, 27] and Ser-Tyr-Lys triad motif involved in catalysis and substrate binding located further to the C terminus of the protein [27, 28] (Fig. 3). These results suggested that the ChKRED12 belonged to a short-chain dehydrogenase/reductase (SDR) family [21]. The sequence was compared with those in GenBank by using the BLASTp program and it was shown that the ChKRED12 shares maximal identities of around 90% with 3-oxoacyl-ACP reductase, such as FabG from Chryseobacterium daeguense (WP 027378905.1). The alignment of the ChKRED12 with 3-oxoacyl-ACP reductases, the known crystal structure of protein also showed high identity (around 40%) such as BKR from Brassica Napus (1EDO) [16]. Only 15-30% residue identity in pairwise comparisons, the three-dimension (3D) structures of SDR are quite similar [29]. These results indicated that the ChKRED12 belonged to 3-oxoacyl-ACP reductase in SDRs family [30]. 3-oxoacyl-ACP reductase catalyzes the reduction of a 3-oxoacyl form of ACP to (R)-3-hydroxyacyl-ACP [31]. Therefore, it is possible that the ChKRED12 has the activity to convert 3-oxoacyl substrate to (R)-3-hydroxyacyl isomer (Fig. 1).

3.2 Protein expression and purification

SDS-PAGE analysis indicated that the ChKRED12 was highly expressed in E. coli BL21 (DE3) cells and that most of the recombinant protein produced was in soluble form (Fig. 4). Expression in the pET28a(+) vector resulted in the protein with an N-terminal hexa-histidine tag that facilitated purification. Single-step affinity chromatography using Ni-NTA agarose yielded the homogeneous protein (Fig. 4). SDS-PAGE showed a single band around 30kDa. The molecular mass of the ChKRED12 was shown to be about 120 kDa by gel-permeation chromatography on Sephacryl S-200. Since SDS-PAGE showed a single band around 30kDa, native ChKRED12 appears to be a homotetramer.

3.3 Cofactor preference and catalytic activity of ChKRED12

The ChKRED12 belongs to the family of SDRs that require NADH or NADPH as a cofactor [32]. We investigated the kinetic parameters of the ChKRED12 towards two cofactors at varied concentrations in the presence of 10 mM substrate 1a. The results showed that NADPH was more efficient than NADH as a cofactor for ChKRED12. The $K_m$ values were two-folds higher and the $V_{max}$ values were five times lower when
NADH as cofactor than NADPH (Table 1). These results indicated that the ChKRED12 have a much higher binding affinity and more catalytic efficiency when NADPH as a cofactor [33], which were also consistent in cofactor preference of 3-oxoacyl-ACP reductase [27]. In biological systems, enzymes that catalyze oxidative exergonic reactions almost exclusively use NAD+/H as a cofactor and reductive endergonic reactive endergonic reactions generally utilize NADP+/H as a cofactor [34]. 3-oxoacyl-ACP reductase participates in fatty acid biosynthesis and polyunsaturated fatty acid biosynthesis, and most of them are NADPH-dependent [27, 35].

Steady-state kinetics towards substrate 1a was measured with NADPH as a coenzyme. The apparent $K_m$, $V_{max}$ and $k_{cat}$ values were 11.8±0.6 mM, 15.2±0.3 U/mg, and 7.8±0.1 s$^{-1}$, respectively.

3.4 Effect of pH and temperature

The pH dependence of activities of the ChKRED12 was measured using purified enzymes using 1a as the substrate at various pH values ranging from 5.0 to 9.5 (Fig. 5). In general, the enzyme displayed broad pH adaptability. Although the maximal catalytic activity was observed in phosphate buffer at pH 8.0, yet >80% of the maximal activity could be achieved over the range of pH 6.0-9.0.

The temperature dependence of the ChKRED12 was measured from 10 to 50 °C. The ChKRED12 displayed the maximal activity at 40°C and retained 28.6% of the maximal activity at 50 °C (Fig. 6). On the other hand, the thermostability of the enzyme should be a concern if a long-time reaction was applied. According to the thermal inactivation curve (Fig. 7), the half-lives of thermal inactivation ($t_{1/2}$) of the ChKRED12 were calculated to be 46, 22 and 1.8 h at 35, 40 and 45 °C, respectively.

The ChKRED12 showed a broad activity pH range, but activity was quickly lost at the temperature above 40 °C. The possible reason is that 3-oxoacyl-ACP reductase has a hydrophobic cave, which contains the active site, coenzyme binding site, and substrate binding site, and the enzymatic catalytic reaction occurs here [27, 36-38]. Because of this structure, the pH value affects the activity less before the conformation of the enzyme has been changed, but this structure cannot prevent the destruction of the conformation of the enzyme by heat when the temperature is above the optimal value.

3.5 Substrate specificity and catalytic properties
Enantiomerically pure alcohols are known as the important and valuable chiral synthons for the production of pharmaceutical and fine chemicals [39-41]. To investigate the substrate specificity of ChKRED12, some substrates (Table 2, 1a-7a, 10 mM) were tested. The biocatalyst reduced the substrates that contain a 3-oxoacyl structure (1a-4a) very effectively, the relative activity of over 80% were achieved for each substrate. Excellent enantioselectivity of ChKRED12 was observed for 1a, 2a, 3a, and 5a, resulting in chiral alcohols of over 99%. The presence of strongly electron-withdrawing chlorine (4a) enhanced the catalytic efficiency and decreases stereoselectivity. In addition, ChKRED12 has a higher catalytic activity for 2-oxoacyl structure substrate (5a) than 3-oxoacyl structure substrates (3a). For 2-Acetyl thiophene (6a) and acetophenone (7a) that only do not contain 3-oxoacyl structure compared with 1a and 2a, respectively, ChKRED12 shows very low relative activity, this result suggested that 3-oxoacyl structure has a role to promote the conversion efficiency.

3.6 Preparative-scale biotransformation

The preparative-scale biotransformation was performed with the substrate 1a in the presence of the GDH-catalyzed cofactor recycling system, using the crude enzyme that prepared from the cell extract of recombinant E. coli. Because the recycling system would result in the accumulation of stoichiometric gluconic acid, the pH of the reaction mixture was continuously monitored and maintained at 7.0-8.0 by the addition of 1 M NaOH. The substrate apparently did not result in inhibition at the tested substrate concentrations (125-500 mM) (Fig. 8). Complete conversion could be achieved within 2, 5, and 12 h at a concentration of substrate 25, 50, 100 g/L, respectively. Notably, there was no substrate inhibition observed. The reaction was terminated by extraction with ethyl acetate, and the combined organic extracts were dried with anhydrous sodium sulfate, concentrated under reduced pressure, and purified with column chromatography to yield (S)-Ethyl-3-hydroxy-3-(thiophen-2-yl) propanoate as a yellow oil (92% yield, >99% ee). 1H NMR (600 MHz, CDCl3): δ1.29 (t, 3H, J = 7.8 Hz, CH3), 2.61 (dd, 1H, J = 6.0 Hz, CH2), 2.86 (dd, 1H, J = 6.0 Hz, J = 4.8 Hz, CH2), 4.12 (q, 2H, J = 6.0 Hz, CH2), 5.14 (t, 1H, J = 6.0 Hz, CHOH), 6.95 (m, 2H, Ar-H), 7.23 (m, 1H, Ar-H). (lit. [42] D25= -17.2 (c = 1.0, CHCl3) for 99%ee, (S)).

The reaction afforded much higher substrate tolerance than previously reported bioreduction of duloxetine precursors [6-8, 43]. For the same substrate, the previous
study used KEES reductase, which yielded compromised stereoselectivity when substrate concentration exceeded than to g/L [8]. To the best of our knowledge, the \( ChKRED12 \) reported here is the best biocatalyst producing enantiopure 3-hydroxy-3-(2-thienyl) propanoate. The high-level production system for this enzyme established in this study might prove useful in the practical applications of this enzyme as a biocatalyst in the future.

4. Conclusion

This study aimed to search for a biocatalyst with excellent enantioselectivity and high substrate tolerance for KEES that contain a 3-oxoacyl structure. This type of substrate is expected to stereoselectively reduce by 3-oxoacyl-ACP reductase. The ketone reductase \( ChKRED12 \) had a high sequence identity with 3-oxoacyl-ACP reductase, indicating that the \( ChKRED12 \) belonged to 3-oxoacyl-ACP reductase in SDRs family and could convert 3-oxoacyl substrate to \((R)-3\)-hydroxyacyl isomer. Further study showed that the \( ChKRED12 \) had catalytic activities towards some 3-oxoacyl structure substrates with excellent anti-Prolog stereoselectivity. This reductase was successfully applied to catalyze the bioreduction of KEES to \((S)\)-HEES with >99% ee value and without product inhibition, the complete conversion could be achieved within 2, 5 and 12 h at a concentration of substrate 25, 50, 100 g/L, respectively. These results demonstrated that the \( ChKRED12 \) is a very promising biocatalyst to produce Ethyl \((S)\)-3-hydroxy-3-(2-thienyl) propanoate (HEES) and it is expected to screen the biocatalysts based on the comparison the specificities of the enzyme with the substrate structure.

Author contributions

Zhi-Qiang Ren, Yan Liu, Xiao-Qiong Pei designed and performed the experiments. Zhi-Qiang Ren and Zhong-Liu Wu analyzed the data and wrote the manuscript.

Conflict of interest

The authors declare that they have no conflicts of interest.

Ethical approval

This article does not contain any studies with human participants or animals performed by any of the authors.
Acknowledgments

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Figure Captions

**Fig. 1** The chemical reaction catalyzes by 3-oxoacyl-ACP reductase and using the Prelog rule to predict the stereoselectivity of 3-oxoacyl-ACP reductase toward some 3-oxoacyl ester substrates.

**Fig. 2** Transformation of KEES to HEES by *ChKRED12* coupled with a cofactor recycling system.

**Fig. 3** Multiple sequence alignment of *ChKRED12* with several members of the SDR family. The alignment was performed with the program DNAMAN. *ChKRED12*, short-chain dehydrogenase/reductase in this work (NCBI accession No. AHC30850.1); β-KACP-BN, β-keto acyl carrier protein reductase from *Brassica Napus* (PDB: 1EDO_A); CoA-AR-BP, Acetoacetyl-coA reductase from *Burkholderia Pseudomallei* 1710b (PDB: 3GK3_A); OR-MT, 3-oxoacyl-ACP reductase from *Mycobacterium Tuberculosis* H37Rv (PDB: 2NTN_A); ACPR-HM, β-ketoacyl-ACP reductase from *Haloarcula marismortui* (NCBI accession No. WP_004961317.1).

**Fig. 4** SDS-PAGE analysis for expression and purification of *ChKRED12*. SDS-PAGE was performed on a 12% gel under reduced conditions. Lane M, molecular weight markers; Lane 1, total proteins from *E.coli* cell containing pET28a (+) vector without induction; Lane 2, total proteins from *E.coli* cell containing pET28a (+) vector with addition 1mM IPTG; Lane 3, total proteins from *E.coli* cell containing pET28a (+)-*ChKRED12* construct without addition of IPTG; Lane 4, total proteins from *E.coli* cell containing pET28a (+)-*ChKRED12* construct with addition of 1mM IPTG; Lane 5, total soluble proteins from induced *E.coli* cell containing pET28a (+)-*ChKRED12* construct with addition of 1mM IPTG; Lane 6, purified recombinant *ChKRED12*.

**Fig. 5** Effect of reaction pH on the activity of the *ChKRED12*. Four different buffers, including sodium citrate (●), Potassium phosphate (○), Tris-HCl (□), and sodium carbonate (▼) were applied.

**Fig. 6** Effect of reaction temperature on the activity of the *ChKRED12*.

**Fig. 7** Thermal inactivation of the *ChKRED12*. The enzymes were incubated at 35°C (▲), 40°C (●) and 45°C (■) for varied times and cooled on ice before the assay.

**Fig. 8** Time courses of the bioreduction of ethyl 3-Oxo-3-(2-thienyl) propanoate (**1a**) at concentrations of 25 g/L (▼), 50 g/L (●), or 100 g/L (▲), using crude enzyme of the *ChKRED12* (4 mg/mL).
Fig. 1. The chemical reaction catalyzes by 3-oxoacyl-ACP reductase and using the Prelog rule to predict the stereoselectivity of 3-oxoacyl-ACP reductase toward some 3-oxoacyl ester substrates.
Fig. 2. Transformation of KEES to HEES by ChKRED12 coupled with a cofactor recycling system.
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