Structural and Functional Analysis of the Only Two Pyridoxal 5′-Phosphate-Dependent Fold Type IV Transaminases in Bacillus altitudinis W3

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Abstract: Aminotransferases are employed as industrial biocatalysts to produce chiral amines with high enantioselectivity and yield. BpTA-1 and BpTA-2 are the only two pyridoxal 5′-phosphate-dependent fold type IV transaminase enzymes in Bacillus altitudinis W3. Herein, we compared the structures and biochemical characteristics of BpTA-1 and BpTA-2 using bioinformatic analysis, circular dichroism spectroscopy, atomic force microscopy and other approaches. BpTA-1 and BpTA-2 are similar overall; both form homodimers and utilize a catalytic lysine. However, there are distinct differences in the substrate cofactor-binding pocket, molecular weight and the proportion of the secondary structure. Both enzymes have the same stereoselectivity but different enzymatic properties. BpTA-2 is more active under partial alkaline and ambient temperature conditions and BpTA-1 is more sensitive to pH and temperature. BpTA-2 as novel enzyme not only fills the building blocks of transaminase but also has broader industrial application potential for (R)-α-phenethylamines than BpTA-1. Structure-function relationships were explored to assess similarities and differences. The findings lay the foundation for modifying these enzymes via protein engineering to enhance their industrial application potential.

Keywords: aminotransferase; pyridoxal 5′-phosphate; secondary structure; conserved domain; AFM; catalytic activity; biochemical characteristics; (R)-α-phenethylamine

1. Introduction

Aminotransferases (ATAs), also known as transaminases (TAs), are a class of biological enzymes that can transfer amino groups from amino donors to amino acceptors in the presence of the coenzyme pyridoxal 5′-phosphate (PLP, a derivative of vitamin B₆) to produce chiral amines with high enantioselectivity and yield [1–3]. Broad substrate specificity and high activity under mild, environmentally friendly conditions make these enzymes attractive as industrial biocatalysts [4–9]. TAs have been employed for the full-scale production of chiral amines represented by phenylpropylamines, methylbenzylamines, aminotetralins and various polyfunctional and aromatic amines [10–13]. Savile and his colleagues reported the successful synthesis of Sitagliptin with a large substrate precursor using ATA-117, an (R)-selective variant of the ω-Tam from Arthrobacter sp. KNK 168 [14]. Hence, TAs are one of the most valuable enzymes in the field of the asymmetric synthesis of the chiral pharmaceutical intermediates. Obviously, the discovery and research of TAs [15–18], not only fill the amine transaminase building blocks but also have important implications for the industrial production of chiral amines.
TAms belong to PLP-dependent enzymes. According to the different structure, ATAs can be divided into two categories, namely PLP-dependent enzymes of fold type I and IV [19,20]. Although fold type I ATAs (aspartate ATA superfamily) can vary significantly in terms of substrate and reaction specificities, they are all (S)-selective enzymes [21,22]. Fold type IV ATAs (D-alanine ATA superfamily) are represented by three different subfamilies; (S)-selective branched-chain TAsms (BCATs), (R)-selective D-amino acid ATAs (DAATs) and R-TAsms [23–26]. Thus, fold type IV ATAs include both (S)-selective enzymes [27] and (R)-selective enzymes [28]. Fold type IV ATAs are more advantageous in terms of substrate specificity and stereoselectivity and have greater industrial application potential than fold type I ATAs. Furthermore, compared to (S)-selective TAsms [29,30], only a few (R)-selective counterparts have been studied [31–33]. Therefore, the mining and research of new (R)-selective TAsms still have important significance.

At present, various studies have been conducted on fold type IV ATAs but direct comparisons of related enzymes from the same organisms remain scarce. In the present work, we compared Bacillus altitudinis W3 (originally known as Bacillus pumilius W3) fold type IV ATAs BpTA-1 and BpTA-2. Amino acid sequence alignment and domain analysis were performed, genes were overexpressed and enzymes were purified to homogeneity. Structural and functional features of the enzymes were also explored.

2. Results and Discussion

2.1. Bioinformatic Analysis

B. altitudinis W3 aminotransferases BpTA-2 and BpTA-1 are the only two PLP-dependent fold type IV TAsms in this strain [34]. And ATA-117 is an (R)-selective TaM mutant from Arthrobacter sp. KNK 168. Phylogenetic tree analysis [35] and alignment of the amino acid sequences of aminotransferases ATA-117, BpTA-2 and BpTA-1 were performed and the results revealed high homology but low similarity (Figure 1). The sequence of BpTA-2 shares 18.4% sequence identity with that of BpTA-1. Moreover, the sequence identity between the sequences BpTA-2 and ATA-117 is even lower, only 14.6%. Although the amino acid sequence of BpTA-2 is longer than that of BpTA-1, the length of their conserved domains is similar; BpTA-2 spans residues 57–344 and BpTA-1 spans residues 6–292. As typical PLP-dependent fold type IV TAsms, both BpTA-2 and BpTA-1 utilize a catalytic lysine. Homology modelling of the above two enzymes showed that both form homodimers (Figure 2), with active sites shielded from solvent by lid residues in the liganded form. However, they also have differences. The substrate cofactor-binding pocket of BpTA-2 is composed of 10 amino acid residues (Y66, F71, R95, V151, K197, Y202, E233, I260, T261 and T303), while that of BpTA-1 is composed of 9 amino acid residues [35]. Compared with the amino acid residues that constitute the substrate cofactor-binding pocket in BpTA-1, BpTA-2 has an additional amino acid residue V151 in addition to the different positions of the residues. Therefore, this residue V151 may be the key reason for the difference in enzyme activity between BpTA-2 and BpTA-1.
Figure 1. Amino acid sequence alignment of the transaminase BpTA-2, BpTA-1 and ATA-117. The amino acid residues in the black box are the active site residues of these three enzymes.
2.2. SDS-PAGE and Gel Filtration Chromatography Analyses

BpTA-1 or BpTA-2, encoded by \( ota3 \) and \( ota8 \), were mainly expressed in soluble form and the recombinant proteins were both subsequently purified by immobilized metal affinity chromatography. The crude and pure enzymes are shown in Figure 3, respectively. Under the same experimental conditions, BpTA-1 was expressed at higher levels than BpTA-2. The molecular masses of BpTA-1 and BpTA-2 were \( \sim 33.4 \) kDa and \( \sim 40.0 \) kDa based on sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE), in accordance with the theoretical values and both yielded a single band following purification. To further confirm the molecular weight of BpTA-1 and BpTA-2, pure enzymes were loaded onto a size-exclusion chromatography column (Figure 4). Based on protein standards and the associated standard curve, the molecular weights of BpTA-1 and BpTA-2 were determined to be \( 65.84 \) kDa and \( 79.98 \) kDa, respectively. The results of SDS-PAGE and gel filtration chromatography indicated that BpTA-1 and BpTA-2 generally exist in the form of dimers. This result is consistent with the above bioinformatics analysis; BpTA-1 and BpTA-2 are typical PLP-dependent fold type IV TAms that are active in dimeric form.

Figure 2. Homology modelling structure of BpAT-2 showing conserved domains. The 10 residues in red (Y66, F71, R95, V151, K197, Y202, E233, I260, T261 and T303) constitute the catalytic site and the cofactor-binding pocket of BpAT-2.
Figure 3. Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) analysis of BpTA-1 and BpTA-2. Lane 1. Protein marker (14.3–97.2 kDa); Lane 2. Crude fractions of BpTA-1; Lane 3. Purification fractions of BpTA-1; Lane 4. Crude fractions of BpTA-2; Lane 5. Purification fractions of BpTA-2.

Figure 4. The size-exclusion chromatography elution curves of BpTA-1 and BpTA-2.

2.3. Secondary Structural Analysis

Secondary structure analysis of BpTA-1 and BpTA-2 was performed using circular dichroism (CD) spectroscopy and the results are shown in Figure 5. The CD spectrum has one positive peak near 198 nm and one negative trough near 218 nm, which are typical characteristics of α-helical and β-sheet structures, respectively. Based on experiments and theoretical calculations, BpTA-1 is composed of ~20% α-helix, 47% β-sheet and 33% random coil, whereas BpTA-2 is composed of ~30.5% α-helix, 14% β-sheet and 55.5% random coil. Although the two enzymes are similar overall, their secondary structural elements differ somewhat under the same experimental conditions. Compared with BpTA-1, BpTA-2 has a higher proportion of α-helical and random coil content and a smaller proportion of β-sheet content. From the perspective of secondary structure, α-helix structure shows the orderliness of protein, while its structure, such as β-sheet, β-turn and random coil, reflects the looseness of
protein. Therefore, the higher α-helical content of BpTA-2 indicates that it has lower sensitivity to the surrounding environment than BpTA-1.

Figure 5. Ultraviolet circular dichroism (UV-CD) spectrum of purified BpTA-1 and BpTA-2 at room temperature and pH 7.0.

2.4. Atomic Force Microscopy Imaging

Atomic force microscopy (AFM) can yield high-resolution images of enzyme molecules under physiological conditions that can reveal details of the morphology, particle diameter and height distribution. Figure 6 shows the 2D and 3D AFM imaging results for BpTA-1 and BpTA-2 at different dilutions. At a 20-fold dilution, BpTA-1 (Figure 6A) and BpTA-2 (Figure 6B) displayed aggregation. Although both enzyme conglomerates are spherical, BpTA-1 is flatter than that of BpTA-2. At a 200-fold dilution, BpTA-1 conglomerates (Figure 6C) and BpTA-2 conglomerates (Figure 6D) were dispersed. BpTA-1 conglomerates were observed as discrete particles with an average diameter of 125.3 nm and an average height of 5.45 nm. Similarly, BpTA-2 conglomerates formed discrete particles with an average diameter of 176.6 nm and an average height of 7.92 nm. Subsequently, even though the enzyme solution was further diluted, the average size of the protein aggregates did not change. The above results indicate that BpTA-2 and BpTA-1 may exist in nature as aggregations and perform the related transamination reactions in the same form.
BpTA-2 displayed high activity with the substrate (R)-α-phenethylamine but activity with (S)-α-phenethylamine was negligible, suggesting that BpTA-2 has (R)-selectivity and can accept specific chiral amines (Table 1). As the only two PLP-dependent fold type IV TAm s in B. altitudinis W3, the stereoselectivity of BpTA-2 was the same as that of BpTA-1. However, the activity of BpTA-2 with (R)-α-phenethylamine was higher than that of BpTA-1 [35], which may be due to differences in their structures. Thus, in terms of industrial applications using (R)-α-phenethylamine, BpTA-2 is more valuable than BpTA-1.

Table 1. BpTA-2 and other TAm s specificity with (R)-α-phenethylamine or (S)-α-phenethylamine as an amino donor.

| Protein | (R)-α-Phenethylamine Activity (U/mg) \( ^a \) | (S)-α-Phenethylamine Activity (U/mg) \( ^a \) |
|---------|---------------------------------|---------------------------------|
| Purified BpTA-2 | 2.1207 ± 0.20 | 0.0011 ± 0.19 |
| Purified BpTA-1 [35] | 1.1760 ± 0.19 | 0.0005 ± 0.15 |
| ATFo [36] | 1.3920 | 0.0004 |
| T. terrenum Tam [37] | 0.15 | n.d. |
| H. ochraceum Tam [37] | 0.10 | n.d. |
| HEWT [38] | <0.0705 | 1.4946 |
| C. violaceum ω-Tam [39] | <0.0564 | 0.97854 |
| V. fluvialis ω-Tam [40] | <0.0141 | 1.5792 |

\( ^a \) One unit of enzyme activity was defined as the amount of the enzyme that catalyzed the formation of 1 \( \mu \)mol of related ketones from amines in 1 min under the reaction conditions described above.

Also, as a newly discovered wild TAm, the (R)-selective TAm from Fusarium oxysporum [36] has higher activity on (R)-α-phenethylamine than BpTA-1 but lower than BpTA-2. The enzyme activities of the TAm s from Thermobaculum terrenum and Haliangium ochraceum to (R)-α-phenethylamine are 0.15 and 0.10 U/mg [37], respectively, which are lower than purified BpTA-2 and BpTA-1. Furthermore, compared with the purified BpTA-2 activity on (R)-α-phenethylamine, the transaminase HEWT from Halomonas elongate [38], Chromobacterium violaceum ω-TAm [39] and Vibrio fluvialis ω-TAm [40] as the (S)-selective TAm, have weaker enzyme activity on the substrate (S)-α-phenethylamine. The above...
results show that BpTA-2 has higher activity on (R)-α-phenethylamine and the industrial production potential of this substrate.

2.6. Effects of pH and Temperature on Enzyme Activity

The effect of pH on the activity of BpTA-2 was investigated with (R)-α-phenethylamine and sodium pyruvate as amino donors and acceptors, respectively, at 35 °C and pH ranging from 5.0 to 13.0 (Figure 7A). The results showed that BpTA-2 was maximally active at pH 9.0. In the range of pH 5.0 to 9.0, enzyme activity increased with increasing pH value. In the range of pH 9.0 to 13.0, enzyme activity decreased with increasing pH value. At pH values below 5.0 or above 13.0, BpTA-2 almost lost its activity.

![Figure 7. Effects of pH (A) and temperature (B) on BpTA-2 activity. Maximum activity in A (3.53 ± 0.11 U/mg) and B (3.50 ± 0.15 U/mg) was set as 100% and relative activity was calculated by comparison with the maximal activity.](image)

The effect of temperature on the activity of BpTA-2 was investigated at pH 7.0 across a temperature range from 20 °C to 75 °C (Figure 7B). The results revealed that the optimal temperature for BpTA-2 was 35 °C. Enzyme activity was increased with increasing temperature between 20 °C and 35 °C. When the temperature was above 35 °C, enzyme activity was decreased with the increase of temperature. Until the temperature was higher than 55 °C, enzyme activity was inactivated due to the high temperature. However, at temperatures below 20 °C, the enzyme remained weakly active, about 20% of the highest activity. To sum up, BpTA-2 exhibited the maximum activity under slightly alkaline and ambient temperature conditions, indicating that this condition would promote the enzymatic reaction.

By contrast, BpTA-1 was more active under neutral and high temperature conditions, which exhibits the best catalytic efficiency at pH 7.0 and 45 °C [35]. Both pH and temperature had a more pronounced effect on BpTA-1 than on BpTA-2. BpTA-2 retained approximately 90% and 80% activity at pH 8.0 and 10.0, respectively and 90% activity at 30 °C and 40 °C. By comparison, BpTA-1 retained only about 30% activity at pH 6.0 and 9.0 and activity was <30% at 40 °C and 50 °C. This demonstrates that BpTA-1 is more sensitive to pH and temperature than BpTA-2, so that the catalytic reactions involved in BpTA-1 have higher requirements on the surrounding environment. Moreover, BpTA-1 and BpTA-2 are the only two PLP-dependent fold type IV TAmS in B. altitudinis W3 and their optimal reaction conditions are so complementary, which may be important for the survival and proliferation of this strain.

2.7. Effects of pH and Temperature on Enzyme Stability

BpTA-2 was incubated in 50 mM phosphate buffer (pH 7.0) for 30 min at temperatures from 10 °C to 70 °C and its residual enzyme activity was measured (Figure 8A). Exposed to the temperature below 20 °C, BpTA-2 has always maintained the best catalytic activity. Nevertheless, at the temperature above 20 °C, the residual activity of the enzyme is decreased sharply as temperature increases until it is completely lost at 60 °C.
3. Materials and Methods

3.1. Bioinformatic Analysis of the ota8 Gene and the BpTA-2 Protein

The ota8 gene (GeneBank No: MN057780) was identified in the genome of Bacillus altitudinis W3 (GeneBank No: CP011150.1), which was isolated by our laboratory [41] and deposited in the China Center for Type Culture Collection (CCTCC No. M2015018). The deduced amino acid sequence of BpTA-2 (encoded by ota8) was submitted to the Swiss-Model online server (http://swissmodel.expasy.org/interactive) for homology modelling. Among the TAsms with a known 3D structure, an enzyme in Mycobacterium smegmatis (PDB ID: 3JZ6) shares 48.57% sequence identity with BpAT-2. The PDB file for this enzyme was obtained from the RCSB Protein Data Bank (http://www.rcsb.org/pdb/home/home.do) and assessed using a PDB Viewer, a powerful protein 3D-structure-viewing software. Meanwhile, the NCBI conserved domain database (CDD; http://www.ncbi.nlm.nih.gov/cdd/), sequence analysis software DNAMAN 5.0 and online software Weblogo3 (http://weblogo.threeplesone.com/) were employed to analyze the conserved domain sequence of BpTA-2 [42]. The amino acid sequence of BpTA-2 was compared with the sequences of aminotransferase ATA-117 (GeneBank No: AB638718.1) from Arthrobacter sp. KNK 168 and aminotransferase BpTA-1 (encoded by ota3; GeneBank No: MH196528) from B. altitudinis W3.

3.2. Materials

A MiniBEST Bacterial Genomic DNA Extraction Kit Version 3.0, PrimeSTAR HS DNA polymerase, 5× PrimeSTAR buffer (Mg²⁺ plus), protein markers and other molecular experimental products were purchased from TaKaRa (Otsu, Japan). PCR primers production and gene sequencing were performed by Hongxun (Suzhou, China). The pCold II vector was purchased from TaKaRa (Dalian, China). Ampicillin sodium salt was purchased from Molekula Ltd. (Gillingham, UK). All analytical grade chemical reagents, unless stated otherwise, were purchased from Sigma-Aldrich (Shanghai, China).
3.3. Gene Expression and Enzyme Purification

The ota8 gene was obtained by polymerase chain reaction (PCR) amplification from B. altitudinis W3 genomic DNA using PrimeSTAR HS DNA polymerase and oligonucleotide primers FWD-ota8 (5′-GGCG CTCGAGATGCCAAAACAACCCATCCCGGTGG-3′) andREW-ota8 (5′-GGCG TGCAGTTATGCGTCTGACGACCGAATCG-3′) incorporating XhoI and PstI restriction sites, respectively (underlined), according to the manufacturer’s instructions. The resulting recombinant plasmid (pCold II-ota8) was verified by sequencing, transformed into Escherichia coli BL21 (DE3) competent cells and transformants were culture in LB broth for protein expression [35]. The crude extract or lysate was obtained after induced and expressed.

Recombinant proteins were purified by chromatography using an Avant purifier (GE Healthcare, Little Chalfont, UK). The supernatant containing the target protein was passed through a 0.22 µm cellulose filter, loaded onto a 1 mL Ni-HisTrap TM column (GE Healthcare, Shanghai, China) and purified according to the manufacturer’s instructions. Enzyme-containing fractions were desalted using a HiTrap TM Desalting column (5 mL) and an ultra-filtration centrifuge tube. Purified enzymes were stored at 4 °C for short periods or at −20 °C for long periods. Both BpTA-2 and BpTA-1 were purified using these experimental procedures. Enzymatic activity was measured by high-performance liquid chromatography (HPLC) assay and crude and purified enzymes were analyzed by SDS-PAGE (see below).

3.4. SDS-PAGE and Gel Filtration Chromatography

The molecular mass of pure enzymes was determined by 12% SDS-PAGE [43] with staining by Coomassie Brilliant Blue R250 and broad range protein markers (14.3–97.2 kDa). The molecular mass of pure enzymes was further investigated by size-exclusion chromatography with a Superdex G-200 10/300GL column eluted using 50 mM phosphate buffer pH 7.0 [44]. Myoglobin (17 kDa), ovalbumin (44 kDa), bovine serum albumin (67 kDa) and γ-globulin (158 kDa) served as standards for calibrating molecular masses.

3.5. Circular Dichroism Spectroscopy

Circular dichroism (CD) spectroscopy measurements were performed on a Chirascan qCD from Applied Photophysics. Pure enzyme samples (0.11 mg/mL) were prepared in sodium phosphate buffer (2.5 mM, pH 7.0) and stored at 4 °C. A 1 mm path length cuvette was used for CD measurements. Each reported spectrum is the average of 10 measurements scanned between 190 and 250 nm. CD spectra of proteins was analyzed using CDtool software and the DichroWeb (http://dichroweb.cryst.bbk.ac.uk) online server [45–47].

3.6. Atomic Force Microscopy

Imaging experiments were carried out using a Nanoscope II atomic force microscopy equipped with a type-D head (Digital Instruments, Santa Barbara, CA, USA). Samples were spread on the mica surface and silicon nitride triangular cantilevers with wide legs (2 mm length; spring constant 0.12 N/m) were used for scanning the mica surfaces. Images were scanned from 80 to 14,000 nm at a 5.78 Hz scan speed and 0 V setpoint voltage and they included 400 sample points per line and 400 lines.

3.7. Enzyme Activity Assay

Measurement of enzyme activity was performed by incubating (R)-α-phenethylamine (20 mM) or (S)-α-phenethylamine (20 mM) with sodium pyruvate (20 mM) in purified (or crude) enzyme and coenzyme PLP at 40 °C and pH 7.0. The reaction was terminated after 15 min by adding an appropriate amount of ethyl acetate, centrifuged (12,000× g, 1min) and the organic phase was analyzed by HPLC using an Agilent C18 column (250 × 4.6 mm, Agilent, Santa Clara, CA, USA) at a UV absorbance of 254 nm with a mobile phase of acetonitrile/ultrapure water (50/50, v/v) and a flow rate of 0.6
mL/min. Enzyme activity was estimated based on the acetophenone yield and all the experiments were conducted with three replicates.

3.8. Effects of pH and Temperature on Enzyme Activity and Stability

The reaction between (R)-α-phenethylamine and sodium pyruvate was designedly carried out at temperatures from 20 to 75 °C at pH 9.0 aimed to determine the optimal temperature for pure enzymes. At the optimum temperatures, the specific buffers with pH values from 5.0 to 13.0 were prepared to determine the optimal pH of purified enzymes, which were respectively and accurately controlled with different pH values using sodium phosphate buffer (pH 5.0 to 8.0), glycine-NaOH buffer (pH 9.0 to 11.0) and KCL-NaOH buffer (pH 12.0 to 13.0). Aiming to evaluate the thermostability and pH stability of BpTA-2, the residual enzyme activities were measured after incubating the enzyme for 30 min with the optimal reaction conditions by separate single controlling the temperatures (varying from 10 to 70 °C) or pH values (varying from 5.0 to 13.0). Relative activities (%) were calculated using the maximal activity as 100%. All the experiments were conducted with three replicates.

4. Conclusions

Aminotransferases BpTA-2 and BpTA-1 are the only two PLP-dependent fold type IV TAMs in B. altitudinis W3. Through bioinformatics analysis, SDS-PAGE and gel filtration chromatography analyses, both enzymes are typical PLP-dependent fold type IV TAMs that form homodimers and utilize catalytic lysine. The catalytic characteristics of enzymes BpTA-2 and BpTA-1 were compared and they shared the same stereoselectivity but differed in terms of enzymatic properties. Combined with the comparison and analysis of the two enzyme substrate cofactor-binding pockets, we found that this residue V151 may be the key reason for the difference in enzyme activity between BpTA-2 and BpTA-1. Compared to BpTA-1, BpTA-2 was more active under slightly alkaline and ambient temperature conditions. And BpTA-1 was more sensitive to pH and temperature than BpTA-2, so that the catalytic reactions involved have higher requirements on the surrounding environment. Meanwhile, the results of CD spectroscopy measurements indicate that BpTA-2 has lower sensitivity to the surrounding environment than BpTA-1 due to its higher α-helical content. BpTA-2 as novel enzyme not only fills the building blocks of transaminase but also has more industrial application potential with (R)-α-phenethylamine than BpTA-1. The overall features of the two enzymes were characterized by AFM. And the 2D and 3D AFM imaging results indicate that BpTA-2 and BpTA-1 may exist in nature as aggregations and perform the related transamination reactions in the same form. The above results reveal that structural features and catalytic characteristics can differ between similar transaminases and differences can be investigated by probing structure-activity relationships. The findings will be of irreplaceable significance for future research on transaminases, especially protein engineering and industrial applications.

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