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Structural insights into the substrate stereospecificity of D-threo-3-hydroxyaspartate dehydratase from *Delftia* sp. HT23: a useful enzyme for the synthesis of optically pure L-threo- and D-erythro-3-hydroxyaspartate

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Abstract

D-threo-3-hydroxyaspartate dehydratase (D-THA DH) is a fold-type III pyridoxal 5’-phosphate-dependent enzyme, isolated from a soil bacterium of Delftia sp. HT23. It catalyzes the dehydration of D-threo-3-hydroxyaspartate (D-THA) and L-erythro-3-hydroxyaspartate (L-EHA). To elucidate the mechanism of substrate stereospecificity, crystal structures of D-THA DH were determined in complex with various ligands, such as an inhibitor [d-erythro-3-hydroxyaspartate (d-EHA)], a substrate (L-EHA), and the reaction intermediate (2-amino maleic acid). The Cβ-OH of L-EHA occupied a position close to the active site Mg$^{2+}$, clearly indicating a possibility of metal-assisted Cβ-OH elimination from the substrate. In contrast, the Cβ-OH of an inhibitor was bound far from the active site Mg$^{2+}$. This suggests that the substrate specificity of D-THA DH is determined by the orientation of the Cβ-OH at the active site, whose spatial arrangement is compatible with the 3R configuration of 3-hydroxyaspartate. We also report an optically pure synthesis of L-threo-3-hydroxyaspartate (L-THA) and D-EHA; promising intermediates for the synthesis of β-benzylxoyaspartate, by using a purified D-THA DH as a biocatalyst for the resolution of racemic DL-THA and DL-EHA. Considering 50% of the theoretical maximum, efficient yields of L-THA (38.9%) and D-EHA (48.9%) as isolated crystals were achieved with >99% enantiomeric excess (e.e.). The results of nuclear magnetic resonance signals verified the chemical purity of the products. We were directly able to isolate analytically pure compounds by the recrystallization of an acidified reaction mixtures (pH 2.0), and thus avoiding the use of environmentally harmful organic solvents for the chromatographic purification.
Introduction

3-Hydroxyaspartate and its derivatives have attracted attention because of their biological activity as competitive blockers against glutamate transporters [Excitatory Amino Acid Transporters 1-5 (EAAT1-5)] in the mammalian central nervous system (Arriza et al. 1994; Arriza et al. 1997; Shimamoto et al. 1998; Shimamoto et al. 2004; Shimamoto 2008). 3-Hydroxyaspartate has two chiral centers and thus it exhibits four stereoisomers, namely D-threo-3-hydroxyaspartate (2R,3R; D-THA), L-threo-3-hydroxyaspartate (2S,3S; L-THA), D-erythro-3-hydroxyaspartate (2R,3S; D-EHA) and L-erythro-3-hydroxyaspartate (2S,3R; L-EHA) (Fig. 1). Recently, we have identified and cloned a gene encoding D-THA dehydrating enzyme (D-THA DH) from a soil bacterium Delftia sp. HT23 (Maeda et al. 2010). D-THA DH displays a preference for the dehydration of D-THA and L-EHA, and is only slightly active against L-THA, while showing no activity on D-EHA. Although, there are several reported enzymes that act on 3-hydroxyaspartate (Wada et al. 1999; Liu et al. 2003; Wada et al. 2003; Murakami et al. 2009), D-THA DH is the only enzyme known to exhibit an efficient dehydratase activity towards D-THA. Thus, this enzyme was assigned a new Enzyme Commission (EC) number as 4.3.1.27.

We have also found that D-THA DH belongs to a fold-type III group of pyridoxal enzymes of which bacterial alanine racemases are the typical members (Schneider et al. 2000; Eliot and Kirsch 2004). Our previous work revealed that D-THA DH contains a pyridoxal 5′-phosphate (PLP) as its cofactor but its racemization activities on alanine, serine and aspartate were below the detection limit. To date, many crystal structures of alanine racemases from diverse bacteria have been deposited in the Protein Data Bank (PDB). Nevertheless, very little is known about the structure of dehydratases whose predictive structure closely resembles that of alanine racemases. Eukaryotic D-serine dehydratase has been recently identified as a common member of the fold-type III group of pyridoxal enzymes (Ito et al. 2008; Tanaka et al. 2008). D-THA DH shows amino acid sequence similarity (26-36%) with
D-serine dehydratases derived from *Saccharomyces cerevisiae* (scDSD; Ito et al. 2008) and chicken kidney (chDSD; Tanaka et al. 2008). The crystal structure of chDSD complexed with D-serine has been previously reported (Tanaka et al. 2011); however, no structural information is available regarding 3-hydroxyaspartate-recognition mechanism of this enzyme family.

Various effective methods for the preparation of 3-hydroxyaspartate as a racemate or an individual stereoisomer have been reported (Kaneko and Katsura 1963; Antolini et al. 1997; Cardillo et al. 1999; Khalaf and Datta 2008). It is of notable interest to develop a synthesis method for L-THA due to the obvious demand for its derivative L-threo-β-benzylxyaspartate (L-TBOA) in the field of neuroscience as a potent blocker of EAAT subtypes (Shigeri et al. 2004; Shimamoto 2008). L-THA acts as a high-affinity substrate for EAAT1-4 and as an inhibitor for EAAT5, thus providing a suitable lead compound as a blocker of EAATs (Arriza et al. 1994; Arriza et al. 1997; Shigeri et al. 2001).

Most of the L-THA synthetic routes relying on the chemical methods utilize a chiral template such as *trans*-methyl cinnamate as a starting material (Deng et al. 1995; Bionda et al. 2012). However, these routes require several reaction steps and are time-consuming. In contrast to the synthetic chemistry approach, few bacterial enzymes are known to efficiently produce L-THA. Strieker et al. (2008) have reported the bioconversion of L-aspartate to L-THA by using a single mutant of asparagine oxygenase (AsnOD241N) from *Streptomyces coelicolor*. However, the activity and the thermal stability of AsnOD241N were found to be lower than those of AsnOWT. Alternatively, Hara and Kino (2010) have reported a more efficient L-THA bioconversion by overexpression of native AsnO using *Escherichia coli*, coupled with *E. coli* asparaginase in vivo. Optically pure 3-hydroxyaspartate production by using a simple strategy is still a challenging process in both synthetic and biological chemistry. While many groups have attempted to increase the productivity, a dramatic improvement in the efficiency has not been realized, resulting in L-THA being an expensive product.

In this paper, we present crystal structures of D-THA DH complexed with its inhibitor...
D-EHA, a poorly active H351A mutant complexed with its substrates L-EHA, and H351A mutant complexed with the reaction intermediate 2-amino maleic acid. High-resolution structures provide a clear picture of how substrate/inhibitor binds to the enzyme, allowing us to propose a model for the catalytic reaction as well as the mechanism for the substrate stereospecificity of D-THA DH. We also report an enzymatic optical resolution by using purified D-THA DH, and show efficient yields of L-THA and D-EHA as isolated crystals with high enantiomeric excess (e.e.). This novel approach for the production of optically pure 3-hydroxyaspartate is simple, environment friendly and is applicable for the large-scale production.
Materials and methods

Materials

Oligonucleotides were obtained from Eurofins Genomics, Inc. (Tokyo, Japan). A (±)-trans-epoxysuccinic acid and DL-threo-3-hydroxyaspartate (DL-THA) were purchased from Tokyo Chemical Industry Co., Ltd. (Tokyo, Japan). L-EHA was purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). An isolated strain, identified as *Delftia* sp. HT23, has been deposited in the AHU Culture Collection of Hokkaido University under accession number AHU2003.

Site-directed mutagenesis

Plasmids expressing the H351A and C353A mutants were constructed using inverse PCR method with a KOD-Plus-Mutagenesis Kit (Toyobo Co., Ltd., Osaka, Japan) according to the manufacturer’s instructions. Previously described d-THA DH expression vector served as a template DNA (Matsumoto et al. 2013). Following synthetic oligonucleotide primers were used: H351A forward, 5′-GCCGCCTGCACGCCACGCGCAGTTCCCGGCC-3′, and reverse, 5′-GGTGAGGATGCGCCAGGGCGCAGGCCGTGCC-3′; C353A forward, 5′-GCCGCCACGGCGCCAGCCGTGCC-3′; C353A forward, 5′-GCCGCCACGGCGCCAGCCGTGCC-3′; and reverse, 5′-GCGCGCCACGGCGCCAGCCGTGCC-3′. The bold bases encode an Ala residue instead of His351 and Cys353, respectively. The inverse PCR conditions were as follows: initial 2-min denaturation at 94 °C, followed by 10 cycles of amplification at 98 °C for 10 s and at 68 °C for 5 min. The nucleotide substitutions were confirmed by DNA sequencing. The nucleotide sequence of the gene encoding d-THA DH from *Delftia* sp. HT23 has been deposited in the DDBJ/EMBL/GenBank database under the accession number AB433986. Each mutant was expressed in *Rhodococcus erythropolis* L88 cells according to a previously described method (Nakashima and Tamura 2004a,b).
Recombinant protein expression and purification

The recombinant enzymes used in this work contained additional Met-Gly-(His)$_6$-Ala-Met-Ser residues at the N-terminus. Recombinant d-THA DH and its mutants were overproduced using a *R. erythropolis* expression system and purified by Ni-affinity chromatography, as described previously (Matsumoto et al. 2013). The fractions containing recombinant proteins were collected, dialyzed against the buffer (10 mM Tris-HCl pH 8.0, 0.01 mM PLP, and 0.1 mM dithiothreitol), and subsequently concentrated to 15 mg/mL using a centrifugal filtration device (10,000 Da molecular-weight cutoff, Merck Millipore, Ltd., Darmstadt, Germany). Protein concentrations were determined by the Bradford method using the Bio-Rad protein assay (Bio-Rad Laboratories, Inc., Hercules, CA) with bovine serum albumin as a standard. Protein purity was checked using SDS-PAGE (Fig. S1).

Crystallization and X-ray diffraction studies

All crystallization experiments were performed at 20 °C by using either sitting drop or hanging drop vapor-diffusion method. The substrate-free wild-type enzyme was crystallized as previously described (Matsumoto et al. 2013). For the preparation of the substrate/inhibitor complex, the purified sample was mixed with molar excess of substrate/inhibitor (~50 mM) and incubated overnight before crystallization setup. All d-THA DH crystals used in this work were obtained in the solution consisting of 0.1 M Tris-HCl pH 8.5, 0.2 M MgCl$_2$, and 10-20% PEG 3350. Prior to X-ray diffraction studies, crystals were briefly soaked in a cryoprotectant solution containing a crystallization mother liquor and an additional 20% glycerol. For the structure analysis of a metal-free enzyme, the crystals were briefly soaked in the Mg$^{2+}$-free cryoprotectant solution consisting of 0.1 M HEPES-NaOH pH 8.0, 2.0 M ammonium formate, and 25% glycerol. Crystals were flash-cooled in a cold nitrogen gas stream or directly in the
liquid nitrogen. X-ray diffraction data were collected using a Charge Coupled Device (CCD) detector (ADSC) at the Photon Factory (PF; Tsukuba, Japan). Raw diffraction images were processed with the program iMosflm/SCALA (Battye et al. 2011; Winn et al. 2011) or HKL2000 suite (Otwinowski and Minor 1997).

Structure solution and model refinement

The structure of D-THA DH was solved by a single-wavelength anomalous diffraction (SAD) method with bromide-soaked crystals as described (Matsumoto et al. 2013). Bromide sites were determined using the program SHELXC/D (Sheldrick 2008). Density modification and model building were performed using the program SOLVE/RESOLVE (Terwilliger and Berendzen 1999; Terwilliger 2000). Model refinement was performed using the program REFMAC 5 (Murshudov et al. 2011), and the manual model fitting was done with the program Coot (Emsley and Cowtan 2004). Ligand models not found in the Chemical Component Dictionary were created using the program SKETCHER and the geometrical restraints files were generated using the program LIBCHECK provided in the CCP4 program package (Winn et al. 2011). Molecular drawings were prepared using the program PyMOL (DeLano 2002). Atomic coordinates and the structure factor amplitudes of all the structures reported in this paper were deposited in the RCSB Protein Data Bank under accession codes 3WQC, 3WQD, 3WQF, 3WQG, 4PB3, 4PB4, and 4PB5. Data collection and refinement statistics are summarized in Table 1.

Enzyme assay

3-Hydroxyaspartate dehydratase activity was assayed spectrophotometrically by measuring a change in the absorbance of NADH at 340 nm using a DU800 spectrophotometer (Beckman Coulter, Inc., Brea, CA). Assay was performed using a coupling system with NADH-dependent malate dehydrogenase (MDH), as described previously (Wada et al. 1999).
The standard assay mixture consisted of 100 mM Tris-HCl buffer (pH 8.0), 0.01 mM PLP, 0.1 mM MnCl₂, 10 mM D-THA or L-EHA as a substrate, 0.32 mM NADH, 10 units of MDH, and an appropriate amount of the enzyme in a total volume of 0.5 mL. The reactions were carried out at 30 °C with addition of the substrate. Reaction mixtures without substrate served as controls. One unit of the enzyme is defined as 1 μmol of NADH utilized per minute at 30 °C on the basis of an absorption coefficient of 6.22 mM⁻¹ cm⁻¹ at 340 nm.

Chemical synthesis of DL-erythro-3-hydroxyaspartate

DL-erythro-3-Hydroxyaspartate (DL-EHA) was synthesized by ammonolysis of a (±)-trans-epoxysuccinic acid at 60 °C. Reaction product was evaporated to dryness, and the resulting residue was dissolved in an acidified water (pH 2.0 adjusted with HCl). Water solution was allowed to stand for 16 h at 4 °C. The appeared crystals of 3-hydroxyaspartate were collected by filtration, washed with a small amount of the acidified water, and subsequently dried in vacuum for 24 h. Resulting product was confirmed by thin layer chromatography (TLC) and high performance liquid chromatography (HPLC) as described in the analytical methods.

Enzymatic optical resolution of DL-racemic 3-hydroxyaspartate

Purified D-THA DH was used to prepare L-THA or D-EHA from the corresponding racemates by an enzymatic optical resolution. The reaction mixture consisted of 100 mM Tris-HCl buffer (pH 8.0), 0.01 mM PLP, 0.1 mM MnCl₂, 200 mM DL-THA or DL-EHA, and purified D-THA DH in a total volume of 100 mL. Reactions were performed at 30 °C with reciprocal shaking at a speed of 90 rpm and started by addition of 0.66 mg of the enzyme to DL-THA solution and 0.87 mg to DL-EHA solution. Incubation was continued for 9 h or 12 h, respectively. During incubation, D-form and L-form of 3-hydroxyaspartate in the reaction mixture were monitored using HPLC and an amino-acid analyzer as described below.
Samples taken for the analyses were heated at 70 °C for 30 min to inactivate the enzyme.

After incubation, L-THA or D-EHA in the reaction product were isolated by crystallization as described below.

**Isolation of L-THA or D-EHA**

The reaction product was acidified to pH 2.0 using HCl. Resulting solution was evaporated to about 10 mL and subsequently allowed to stand for 16 h at 4 °C. Appeared crystals of 3-hydroxyaspartate were collected with the same procedure as for the preparation of DL-EHA. TLC, HPLC and nuclear magnetic resonance (NMR) spectroscopy as described in the analytical methods, confirmed the resulting product.

**Analytical methods**

**TLC**: Crystallized DL-EHA, L-THA and D-EHA were confirmed by Silica Gel TLC using 7:1:2 (v/v/v) ethanol/28% aqueous ammonia/water as a mobile phase. After developing the TLC plates, compounds were visualized by using a Ninhydrin Spray (Wako Pure Chemical Industries), followed by heating.

**HPLC**: For chiral analysis of amino acids, 2,3,4,6-tetra-O-acetyl-β-D-glucopyranosyl isothiocyanate (GITC) was used for the amino acid derivatization. In a GITC derivatization procedure, 0.1 mL of 0.2% (w/v) GITC (dissolved in acetonitrile) and 0.02 mL of 2% (v/v) triethylamine (dissolved in water) were added to 0.01 mL of reaction samples that were subsequently diluted with acetonitrile to make a total volume of 0.2 mL. The resulting solution was allowed to stand for 30 min at 30 °C. The amino acid derivatives were analyzed using HPLC at a flow rate of 1.0 mL min⁻¹ with a CAPCELL PAK C18 MGIII column (4.6 × 250 mm; Shiseido Company, Ltd., Tokyo, Japan) at 40 °C. Thirty percent (v/v) methanol or 35% (v/v) methanol in water (pH 2.5 adjusted with H₃PO₄) were used as the mobile phases for the analysis of DL-THA or DL-EHA, respectively. Detection was done with a UV detector.
tuned at 250 nm. The enantiomeric excess (e.e.) was calculated from the peak areas of the stereoisomers.

**Amino acid analyzer:** Total concentration of DL-THA or DL-EHA in the reaction mixture was determined with a JLC-500S amino acid analyzer (JEOL, Ltd., Tokyo, Japan).

**NMR spectroscopy:** All NMR spectra were recorded on a Bruker AMX-500 spectrometer (Bruker Corporation, Billerica, MA, $^1$H NMR: 500 MHz; $^{13}$C NMR: 125 MHz). Samples taken for the analysis were dissolved in deuterium oxide (D$_2$O). Chemical shifts were determined relative to the reference signal of an external 3-(trimethylsilyl)propionic-2,2,3,3-d$_4$ acid (TSP; $\delta_H$, 0 ppm; $\delta_C$, 0 ppm).
Results

Overall structure of d-THA DH

The crystal structure of d-THA DH was determined in the substrate-free form at a resolution of 1.5 Å. The asymmetric unit comprises a dimer formed by a head-to-tail association of two monomers (Fig. 2a). Each monomer is divided into N- and C-terminal domains (Fig. 2b). The N-terminal domain (residues 17-257) consists of eight α-helices (α1-α8) and eight β-strands (β2-β9) which constitute a well-known “TIM-barrel” fold, providing a binding site for PLP. The first β-strand of the monomer, β1, does not form the TIM barrel and is a part of the C-terminal domain since β1 is located at the extreme N-terminus. The C-terminal domain, comprising of residues 1-16 and 258-380, is mainly composed of β-strands. All the nine β-strands (β1 and β10-β17) of the domain form a β-sheet in which the strands are arranged in an antiparallel orientation except for β1 and β16. This β-sheet creates a six-stranded β-barrel structure in which the strands (β10-β11 and β14-β15) are arranged in a Greek-key topology. The PLP-binding site is located at the center of the TIM barrel, between two domains. The overall folding topology is typical of a bacterial alanine racemase. Thus, the current structure provides a conclusive evidence that d-THA DH belongs to the fold-type III group of pyridoxal enzymes.

Cofactor binding site

The active-site pockets harboring PLP are located at the dimer interface, between the TIM barrel of one monomer and the C-terminal domain of the homodimeric partner. Accordingly, residues bound to PLP (Lys43) yield two active sites per dimer. A highly defined electron density was observed for the entire PLP molecule in each active site. The PLP cofactor binds covalently via an imine bond to the ε-amino group of Lys43, forming an internal aldime (Schiff base) (Fig. 2c; Fig. S2a). The Lys residue is also highly conserved in
scDSD and chDSD (Ito et al. 2008; Tanaka et al. 2008). The internal aldimine and the guanidine group of Arg141 are within hydrogen-bonding distance of the phenolic O atom of PLP. The phosphate group of PLP is involved in forming hydrogen bonds to the side-chain atoms of the residues Tyr177 and Thr218, and the main-chain atoms of the residues Thr218, Gly236 and Val237. The pyridine ring of PLP makes a hydrogen bond between its nitrogen atom and the guanidine group of Arg234.

We observed a strong electron density peak between side-chains of His351 and Cys353 in each subunit. The electron density was assigned as fully occupied Mg$^{2+}$, and the model was refined with the reasonable B-factor values comparable to those of nearby residues. Mg$^{2+}$ was probably bound due to the high concentration of MgCl$_2$ (200 mM) in the crystallization solution. The metal-binding site is a part of the C-terminal domain and is formed by His351 and Cys353, which coordinate the metal ion through the N-$\varepsilon$ atom of the imidazole ring and the thiol group, respectively (Fig. 2c; Fig. S2a). In addition to the two residues, four water molecules complete the octahedral coordination of the Mg$^{2+}$. The PLP is located near Mg$^{2+}$ and its phosphate group lies at a distance of about ~5 Å from the metal center. The sequence comparison of D-THA DH with scDSD and chDSD demonstrates that the His and Cys residues coordinating the metal ion are well conserved in these proteins (Ito et al. 2008; Tanaka et al. 2008). All the related proteins show a strong dependency on Zn$^{2+}$, while D-THA DH works with a various divalent cations including Mg$^{2+}$ as discussed later (Fig. S3).

**D-EHA (inhibitor) binding mode of D-THA DH**

Our previous work has revealed that the specific activity of D-THA DH towards D-EHA is below the limits of detection. D-EHA inhibits D-THA DH with the $K_i$ value of 0.114 mM, determined by replotting the slopes of the Lineweaver-Burk plots. Considering both the results, it can be concluded that D-THA DH has a significant binding affinity towards D-EHA.
but does not recognize it as a substrate. To elucidate how D-EHA binds in the active site of D-THA DH, its structure was determined in the presence of D-EHA. The structure was determined to a resolution of 1.5 Å and a clear electron density map was observed for the whole D-EHA molecule bound to the PLP cofactor. The active site of each subunit includes a D-EHA molecule and two Mg\(^{2+}\) (Fig. 3). One Mg\(^{2+}\) is located between His351 and Cys353 as described in the ligand-free structure. The other Mg\(^{2+}\) is chelated by the β-carboxyl O atom and the β-hydroxyl O atom of D-EHA, forming a D-EHA-Mg\(^{2+}\) complex. Thus, the latter Mg\(^{2+}\) is most likely accompanied by the presence of a ligand containing a β-carboxyl and β-hydroxyl group. Electron density allowed the modeling of a covalent linkage between the C4A atom of PLP and the α-amino group of D-EHA, generating an external aldimine bond. The internal aldimine linkage between the cofactor and Lys43 appeared to be broken in the electron density (Fig. S2b). Although significant structural differences were not observed, an approximately 16° rotation of the cofactor pyridine ring was seen accompanied by the Schiff-base interchange. The α-carboxyl group of D-EHA forms hydrogen bonds to the guanidine group of Arg141 from the PLP-binding subunit and the main-chain N atom of Gln319 from the neighbouring subunit. Note that the β-hydroxyl O atom of D-EHA is located at a distance of 5.5 Å from the Mg\(^{2+}\) positioned between His351 and Cys353.

Crystallographic studies of chDSD have demonstrated that the Zn\(^{2+}\) sandwiched by His347 and Cys349 probably comes in contact with the β-hydroxyl group of D-serine and may have a catalytic role in leaving the hydroxyl as a water molecule (Tanaka et al. 2011). Therefore, the spatial arrangement of the hydroxyl group that lies far from the enzyme-coordinated Mg\(^{2+}\) presumably results in loss of the dehydration activity against D-EHA.

**Insights into the substrate binding**

To elucidate the structural mechanism of the substrate recognition of D-THA DH, we have attempted to determine the structure of inactive mutant in complex with its substrate. We
generated the H351A and C353A mutants of D-THA DH and performed an enzyme assay using D-THA and L-EHA as substrates. The results demonstrated that both H351A and C353A mutations significantly decrease the activity towards D-THA and L-EHA (Table 2), similar to the results obtained with scDSD (Ito et al. 2012). In particular, the specific activity of C353A mutant towards D-THA and L-EHA was below the detection limit, indicating that Cys353 is a key determinant of the enzymatic activity of D-THA DH. We have first determined the crystal structures of H351A and C353A in the substrate-free form at resolutions of 1.7 and 1.55 Å, respectively. Crystals of each mutant were obtained under the same crystallization condition as for the wild-type D-THA DH. In the structure of the H351A mutant, Mg$^{2+}$ occupies a position near the side chain of Cys353. It is coordinated in an octahedral geometry by the thiol S atom of Cys353 and five water molecules, positioned from the metal center at distances of about 3.6 and 2.0-2.1 Å, respectively. These observations indicate that the H351A mutant utilizes only one cysteine residue for the metal binding. In contrast, no electron density for a metal bound to the active-site pocket was observed in the structure of C353A mutant. It is thus suggested that the Mg$^{2+}$ is held by the side chain of Cys353 rather than by that of His351. Even in the absence of the metal, the structure of C353A mutant displays a conformation similar to the wild-type enzyme. It should also be noted that the short soak of wild-type D-THA DH crystal into the metal-free solution led to the elimination of Mg$^{2+}$ from the active site (Table 1; PDB code, 3WQF). These suggest a low binding affinity of Mg$^{2+}$, and that the Mg$^{2+}$ is not required for the structural stabilization of the enzyme.

Structures of the H351A co-crystallized with D-THA and L-EHA were also determined at resolutions of 1.8 and 1.9 Å, respectively. The covalent linkages appear to be formed between the PLP cofactor and the $\alpha$-amino groups of substrates from the electron density map. Attempts to solve the crystal structures of the C353A in complex with D-THA and L-EHA were not successful. Mg$^{2+}$ could not bind to the active site of C353A mutant, leading to a significant decrease in its substrate binding affinity. The electron density map of the structure
of H351A co-crystallized with D-THA unexpectedly showed that the N-Cα-Cβ-Cγ of a bound
substrate is roughly coplanar with the lacking β-hydroxyl group. This prompted us to build a
model of 2-amino maleic acid (Fig. 4, a and b). 2-Amino maleic acid is a likely product of the
dehydration of D-THA, and thus the current complex structure represents its intermediate state
in the catalytic cycle of D-THA DH (Fig. 5). The water molecule (Wat709A/Wat712B) located
near the intermediate Cβ atom at a distance of 2.0 Å was probably generated by the
dehydratase reaction catalyzed by the H351A mutant. Water also interacts with the liberated
ε-amino group of Lys43 (2.9 Å), the phosphate group of PLP (2.5 Å) and the
cysteine-coordinated Mg2+ (2.3 Å). These observations indicate that the Cβ-OH orientation of
D-THA would be an important factor contributing to the substrate recognition of D-THA DH.
In addition, Mg2+ might directly interact with the β-hydroxyl group of D-THA to promote the
dehydration reaction. Not only D-THA but also L-EHA is a good substrate for D-THA DH
(Maeda et al. 2010). In the structure of H351A co-crystallized with L-EHA, the electron
density corresponding to L-EHA was clearly visible (Fig. 4, c and d). The hydroxyl O atom of
L-EHA occupies roughly the same position as Wat709A/Wat712B liberated from D-THA in
the reaction intermediate complex, and is also in direct contact with the Mg2+. Thus, the
substrate stereospecificity of D-THA DH is determined by the Cβ-OH orientation of the
substrate at the active site.

In summary, we have determined three complex structures that shed light on the
3-hydroxyaspartate-recognition mechanism of D-THA DH: the wild-type enzyme in complex
with D-EHA, the H351A mutant in complex with 2-amino maleic acid originating from
D-THA, and the H351A mutant in complex with L-EHA. Obtaining the complex structure
with L-THA was tried with both the wild-type and the H351A mutant enzymes, however,
there was no visible electron density for L-THA in the active-site pockets. This was explained
by the high $K_m$ value of D-THA DH for L-THA ($K_m = 6.16 \text{ mM}$; Maeda et al. 2010).
Interestingly, the current structural studies revealed that chirality at the Cβ position of
3-hydroxyaspartate (3R) can be well recognized by D-THA DH as substrates. The Cβ configuration of D-THA superimposed well onto that of L-EHA in the present models, which is associated with the high levels of activities against D-THA and L-EHA. Thus, the active-site conformation of D-THA DH would be compatible with the 3R configuration of 3-hydroxyaspartate. This finding provides an explanation for the efficient production of optically pure 3-hydroxyaspartate having the 3S configuration as described below.

Biocatalytic preparation of L-THA from DL-THA using D-THA DH

We have developed a novel method for producing optically pure L-THA, which is (2S,3S) form of 3-hydroxyaspartate and a promising intermediate for the synthesis of L-TBOA, an important compound in understanding the regulatory functions of glutamate transporters in the human brain (Shimamoto 2008). Enzyme kinetics and the structural analysis have indicated that D-THA DH is useful in the dehydratase-catalyzed optical resolution of DL-THA. Therefore, we evaluated the biocatalytic preparation of optically pure L-THA by using purified D-THA DH. After 6 h, the enantiomeric excess (e.e.) of L-THA in the reaction mixture reached >99% as calculated from the peak areas in the HPLC analysis (Fig. 6a). In addition, the total concentration of DL-THA determined with an amino-acid analyzer decreased to about half the initial value at the same time (Fig. S4). These results indicate that D-THA in the racemate solution was enantioselectively converted to oxaloacetate and ammonia by the enzymatic reaction. Remaining L-THA in the reaction product was crystallized under the acidified condition (pH 2.0). The resulting residue was an enantiopure L-THA as determined by HPLC analysis (Fig. S5). The isolation yield as dry weight was 38.9% against the amount of DL-THA initially added to the reaction mixture. Moreover, chemical purity of the final product was verified by NMR analysis and it showed the following signals (Fig. S6); 1H NMR (500 MHz, TSP = 0 ppm): 3.98 (1H, d, J = 1.8 Hz), 4.52 (1H, d, J = 1.8 Hz) ppm; 13C NMR (125 MHz, TSP = 0 ppm): 60.02, 74.08, 176.47, 179.77 ppm.
ppm. These signals indicate the *threo* isomer of 3-hydroxyaspartate.

**Biocatalytic preparation of D-EHA from DL-EHA by D-THA DH**

Similarly, biocatalytic preparation of optically pure D-EHA, which is (2*R*,3*S*)-3-hydroxyaspartate, from DL-EHA was achieved by using purified D-THA DH. The derivative of D-EHA, D-*erythro*-β-benzylxoyaspartate (D-EBOA) was also found to act as a blocker for EAATs although the effect was relatively weak (Shimamoto et al., 2000). After 10 h of bioconversion, the reaction mixture contained only the D-enantiomer of *erythro*-3-hydroxyaspartate with >99% *e.e.* and 48.9% of the isolated yield (Fig. 6b; Fig. S7). The result of NMR signals verified the chemical purity of the final product as *erythro* isomer of 3-hydroxyaspartate (Fig. S8); $^1$H NMR (500 MHz, TSP = 0 ppm): 4.00 (1H, d, $J$ = 3.6 Hz), 4.33 (1H, d, $J$ = 3.6 Hz) ppm; $^{13}$C NMR (125 MHz, TSP = 0 ppm): 60.86, 74.47, 175.12, 179.16 ppm.
Discussion

Structure comparison with D-serine dehydratase from chicken kidney

Pairwise structure comparison using the *DALI* server (Holm and Park 2000) showed that D-THA DH is structurally homologous to D-serine dehydratase from chicken kidney (chDSD; Tanaka et al. 2011; PDB code, 3ANU; Z score, 45.2; rmsd, 1.9 Å for 358 C$^{\alpha}$ atoms; sequence identity for fit regions, 37%). Although the overall fold of D-THA DH and chDSD are identical, there are apparent differences that need to be discussed. The first difference is the oligomerization state in the solution. Tanaka et al. based on the gel-filtration chromatography results have reported that chDSD forms a dimer in the solution. Even though crystallographic studies of chDSD demonstrate that the asymmetric unit comprises of a single monomer, the biological assembly is assumed head-to-tail dimer. It is created by a crystallographic 2-fold axis involving large and hydrophilic interface between the subunits. In contrast to chDSD, judging from its apparent molecular mass analyzed by the gel filtration, D-THA DH is a monomeric enzyme as previously reported (Maeda et al. 2010). However, as shown in the present study, all the crystal forms of D-THA DH have the head-to-tail dimer in the asymmetric unit, similar to the biological assembly of chDSD. The dimeric state of D-THA DH revealed that the amino-acid residues Asn318-His321, which form a loop in the C-terminal domain, contact neighboring PLP-binding pockets and include a Gln319 that is expected to be utilized for D-EHA, D-THA and L-EHA binding. Each domain is also responsible for the dimerization through hydrogen bonds. Therefore, the current symmetric dimer is likely to be one of the possible conformational states of D-THA DH and the crystal lattice forces may stabilize this conformation. The second difference is found in the arrangement of residues that directly interact with the pyridine nitrogen, designated as N1. The structure of chDSD revealed that the side chain of Tyr174 is within hydrogen-bonding distance of N1. This environment may preclude the protonation of N1 that possibly functions
as a driving force for the catalysis of transamination. Therefore, the N1-unprotonated state of chDSD is needed for the catalysis of β-elimination and for avoiding other PLP-catalyzed reactions. Likewise, although the Tyr residue is substituted with Leu170 in d-THA DH based on the primary structure alignment, the alternate Arg residue at position 234 is hydrogen bonded to N1, thereby preventing proton transfer to N1 due to the high pKₐ value of the arginine side-chain (~12.5 in water).

In this structural study, Mg²⁺ was observed between the side chains of His351 and Cys353, probably due to artificial crystallization solution containing 200 mM MgCl₂. Although it is uncommon that Mg²⁺ is coordinated by imidazole and thiol ligands, the atomic coordinates and the B-factor values were reasonably refined without extra Fₒ-Fc density map, indicating that the bound metal is conclusively Mg²⁺. The Mg²⁺-free structure (Table 1; PDB code, 3WQF) also suggests that d-THA DH has a low affinity for Mg²⁺. To examine which metals occupy the active site prior to crystallization, an inductively coupled plasma mass spectrometry (ICP-MS) analysis was conducted on purified d-THA DH. The results showed that a large amount of Zn²⁺ was detected in the purified enzyme (Fig. S9). A relatively high content of Ni²⁺ and Co²⁺ was also detected after Ni²⁺- and Co²⁺-affinity purification, respectively. Therefore, it can be concluded that d-THA DH naturally contains Zn²⁺, but under the given experimental conditions, the substitution of Zn²⁺ for other metals takes place easily. Tanaka and co-workers have reported that chDSD naturally possesses Zn²⁺ based on its X-ray structure, and only Mn²⁺ can partially substitute Zn²⁺ for the catalytic activity against d-serine. In contrast, we have previously reported that d-THA DH works with a various divalent cations such as Co²⁺, Mn²⁺, Ni²⁺, Ca²⁺, Zn²⁺, and Fe²⁺ (Maeda et al. 2010). Furthermore, addition of MgCl₂ to the enzyme solution after treatment with ethylenediaminetetraacetic acid (EDTA) gave a low but significant recovery of the specific activity of d-THA DH from 0.1 to 1.2 μmol/min/mg protein (Fig. S3). This clearly shows that the Mg²⁺ is functional and is involved in the enzymatic activity.
Insights into the catalytic mechanism and the substrate stereoselectivity

Note that the substitution of alanine for Cys353 precludes the binding of Mg$^{2+}$ to D-THA DH (Table 1; PDB code, 3WQC). Structure of C353A mutant also showed no binding affinity towards D-EHA, D-THA and L-EHA (data not shown). Considering these results, it can be concluded that the substrate/inhibitor binding affinities of D-THA DH are greatly decreased by the absence of the metal. In contrast, crystallographic analysis have revealed that the pretreatment of EDTA allows chDSD to bind with D-serine even in the absence of Zn$^{2+}$. This suggests that chDSD does not require metal for the initial occurrence of the Schiff-base interchange from lysine-PLP to PLP-substrate.

On the basis of the X-ray structure of H351A cocrystallized with D-THA/L-EHA, a possible mechanism for the catalytic reaction of D-THA DH has been proposed (Fig. 5). Initially, the substrate covalently binds to the PLP molecule in each active-site pocket of the dimeric state of D-THA DH. Metal ion is also in direct contact with the $\beta$-carboxyl and $\beta$-hydroxyl O atoms, presumably playing a role in the stabilization of the bound substrate during the catalytic reaction. Upon abstracting a proton from the $\alpha$-carbon of D-THA by Lys43 to generate a quinonoid resonance structure, the $\beta$-elimination reaction occurs by a proton donation to the substrate hydroxyl group. In previous structural studies on chDSD, the reported distance between the NZ atom of the Lys45 and the $\beta$-hydroxyl of D-serine was an optimal distance (3.2 Å) for the proton donation, suggesting that the abstracted $\alpha$-proton could be transferred to the substrate hydroxyl via the lysine residue as an acid/base catalyst (Tanaka et al. 2011). In contrast, the current complex structures of the H351A mutant indicate that another candidate, a phosphate group of PLP, is located in the proper position contributing to the $\beta$-elimination reaction. We can assume that the protonated phosphate O atom acts as a general acid to donate a proton to the leaving hydroxyl group of the substrate. Indeed, the side chain of Tyr177 is in contact with the phosphate O atom, thus maintaining the protonation
state in transferring proton throughout the reaction. Also, the distances from the hydroxyl
group of substrate L-EHA to the ε-amino group of Lys43 (2.8 Å) and the phosphate group of
PLP (2.6 Å) are shorter than the corresponding distances from the hydroxyl group of inhibitor
d-EHA to the ε-amino group (5.8 Å) and the phosphate (5.7 Å). Considering that the ε-amino
group of Lys43 exhibits conformational dynamics in response to the catalytic cycle, the
association between the protonated phosphate group and the substrate hydroxyl is more
responsible for leaving the hydroxyl group as a water molecule. We observed the
PLP-2-amino maleic acid complex as the subsequent intermediate. An imine/enamine
released after the internal Schiff-base regeneration is thought to be hydrolyzed
non-enzymatically. Thus, tautomeric change of free 2-amino maleic acid into 2-imino succinic
acid and subsequent hydrolysis eliminate ammonia from the imine to give oxaloacetate.

\[ \text{C}^{\alpha}-\text{H deprotonation is commonly carried out by the } \varepsilon\text{-amino group of a lysine that is} \]
liberated from the internal aldimine with PLP (Toney 2005). In this study, the ε-amino group
of Lys43 could access the α-proton of d-THA, while it could not interact with that of L-EHA.
Alternatively, side-chains of Arg141 and His172 are situated close to the α-proton of L-EHA
and would play a role in the proton abstraction. Our previous kinetic analysis based on the
\[ k_{\text{cat}}/K_m \] values indicates that d-THA DH has 2.6-fold higher binding affinity for L-EHA than
for d-THA, thereby acting more effectively on L-EHA (Maeda et al. 2010). Since the only
difference between d-THA and L-EHA is the chirality at the C\( ^\alpha \) position (Fig. 1), it is
suggested that the α-proton abstraction from L-EHA by Arg141 and/or His172 might increase
the dehydratase activity. This raises the possibility that d-THA DH possesses the epimerase
activity between d-THA and L-EHA. However, the HPLC chromatograms can preclude this
possibility.

**Synthesis of optically pure 3-hydroxyaspartate using d-THA DH**

In this study, we have developed a novel synthetic approach leading to L-THA and
D-EHA via the dehydratase-catalyzed optical resolution. The separation of L-THA from the corresponding racemate, DL-THA, was achieved by converting D-THA into oxaloacetate using recombinant D-THA DH. Additionally, L-EHA also serves as a good substrate; therefore, D-THA DH was also used for the synthesis of D-EHA starting from DL-EHA. To date, there have been no published studies on an enzymatic optical resolution of racemic 3-hydroxyaspartates, although convenient synthetic routes for DL-THA and DL-EHA starting from (±)-trans-epoxysuccinic acid were established many years ago (Kaneko and Katsura 1963). Thus, our data confirm for the first time the utility of D-THA DH as a biocatalyst for the resolution of DL-threo/erythro-3-hydroxyaspartate.

The racemic mixture contains both enantiomers in equal amounts, and thus the maximum yield in an optical resolution is 50%. Considering this theoretical maximum, the synthetic approaches are quite efficient as judged from the isolated yields of L-THA and D-EHA. However, D-THA DH was found to possess a slight activity against L-THA (Maeda et al. 2010), resulting in a lower yield of L-THA (38.9%) compared to D-EHA (48.9%). There is a need for further studies to create the mutant enzyme showing no activity on L-THA. In addition, the reaction time of L-EHA degradation was longer than expected. This is likely due to inhibition of the dehydratase by binding of D-EHA, which is the remaining product. Thus, a short reaction time requires use of the mutant enzyme capable of overcoming the inhibition, which could be accomplished by the structure-based engineering of D-THA DH whose structure in complex with D-EHA has been reported here.

We confirmed the final products as pure L-THA and D-EHA by analyzing them with TLC, HPLC, and NMR spectroscopy. We were able to isolate pure compounds directly by recrystallization of the reaction mixtures; utilizing the lower solubility of 3-hydroxyaspartate in acidic water (pH 2.0) compared to oxaloacetate and ammonia. Therefore, this synthetic approach has an advantage for obtaining pure products without any chromatographic purification. It is efficient and friendly to the environment because of minimal or no use of
organic solvents.

Strieker et al. (2008) reported that a single mutant of asparagine oxygenase (AsnO_D241N) from *Streptomyces coelicolor* catalyzes one-step hydroxylation of L-aspartate to produce L-THA. The bioconversion represents one of the simplest strategies for preparing L-THA from a low-cost material. However, it is important to note that 1.16 g of the isolated yield of L-THA per 100 mL of the reaction mixture was achieved in this study, which is a considerable amount compared to the production data of Strieker et al. (14.8 mg of the isolated yield per 10 mL of the reaction mixture). Strieker and co-workers have also reported that the conversion reaction was carried out under optimized conditions including a low temperature of 16 °C most probably due to the reduced enzyme stability of AsnO_D241N. In contrast, D-THA DH was found to be thermally stable up to 45 °C (Fig. S10), which would be more acceptable for industrial-scale reactions demanding moderate to high-temperature conditions.
Acknowledgments

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References


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**Figure Captions**

**Figure 1**
Structures of the four 3-hydroxyaspartate stereoisomers. L-THA, L-threo-3-hydroxyaspartate; D-THA, d-threo-3-hydroxyaspartate; L-EHA, L-erythro-3-hydroxyaspartate; D-EHA, d-erythro-3-hydroxyaspartate.

**Figure 2**
Structure of D-THA DH from *Delftia* sp. HT23 (a) Ribbon diagram of the dimeric structure where one of the monomers is colored in green and the other in yellow. The PLP cofactor is shown as a stick model colored in magenta, and the Mg$^{2+}$ between side-chains of His351 and Cys353 is shown as a sphere in blue. (b) Ribbon diagram of the monomeric structure. The $\alpha$-helices and $\beta$-strands, and loops are colored in red, yellow, and green, respectively. The $\alpha$-helices and $\beta$-strands are labeled numerically from the N- to C-terminus ($\alpha_1$-$\alpha_8$ and $\beta_1$-$\beta_{17}$). (c) Stereo-view showing the cofactor-binding site of D-THA DH in the substrate-free form. Side-chains interacting with PLP and Mg$^{2+}$ are shown as sticks and labeled. The surrounding polypeptide chain is represented as a semi-transparent ribbon diagram colored in green.

**Figure 3**
Stereo-view of the active-site structure of D-THA DH with the bound inhibitor (D-EHA). The functionally important residues (in green), PLP (in magenta), and bound D-EHA (in yellow) are shown as sticks and labeled. Mg$^{2+}$ is shown as blue spheres.

**Figure 4**
2-Amino maleic acid and L-EHA binding to the H351A mutant of D-THA DH. (a) An
Figure 5

Proposed catalytic cycle of D-THA DH based on the structure of the H351A mutant in complex with 2-amino maleic acid. Initial addition of the substrate converts lysine-PLP (internal aldimine) to PLP-D-THA (external aldimine). Next, proton abstraction from the α-carbon of bound D-THA occurs by Lys43 (Arg141 and/or His172 in the case of L-EHA) to generate a quinonoid intermediate. The PLP-2-amino maleic acid complex is subsequently generated by the β-elimination reaction promoted by the interaction between the hydroxyl O atom and the active-site metal ion (represented as M²⁺). The protonated phosphate of PLP acts as an acid to donate its proton to the hydroxyl, which is then released as water. Finally, regeneration of the internal aldimine linkage and non-enzymatic hydrolysis of the resulting free imine/enamine release oxaloacetate and ammonia.

Figure 6

Optically pure production of L-THA and D-EHA. (a) Biocatalytic preparation of L-THA from DL-THA by purified D-THA DH. The reaction mixture consisted of 100 mM
Tris-HCl buffer (pH 8.0), 0.01 mM PLP, 0.1 mM MnCl$_2$, 200 mM DL-THA, and 0.66 mg of the enzyme in a total volume of 100 mL. During incubation, the D-form and L-form of 3-hydroxyaspartate in the reaction mixture were monitored using HPLC. The enantiomeric excess ($e.e.$) was calculated from the peak areas of the stereoisomers. (b) Biocatalytic preparation of D-EHA from DL-EHA by purified D-THA DH. The experiment and calculation were performed with the same procedure as for the production of L-THA except for an enzyme amount of 0.87 mg.
Figure S1 SDS-PAGE of the wild-type and mutants of d-THA DH. Lanes 1-3 correspond to the wild-type, H351A, and C353A mutants, respectively. Lane M contains marker proteins, including (from top to bottom), phosphorylase ($M_r$, 97,400), bovine serum albumin (66,300), aldolase (42,400), carbonic anhydrase (30,000), trypsin inhibitor (20,100), and lysozyme (14,400). Gel was stained for protein with Coomassie Brilliant Blue R-250 (Wako Pure Chemical Industries).
Figure S2 Final 2mFo-DFc map of the wild-type d-THA DH around active site in (a) substrate-free form (PDB code, 3WQC) and in (b) complex with d-EHA (PDB code, 3WQD), clearly showing the Schiff-base interchange from lysine-PLP to PLP-d-EHA. Each map was shows contours at 1.4σ level.
Figure S3 Effect of Mg$^{2+}$ on the specific activity of d-THA DH. EDTA-treated enzyme was prepared using the method described earlier (Maeda et al. 2010). It was incubated for 10 min with the reaction mixture in the absence of any metal, or in the presence of 0.1 or 200 mM MgCl$_2$. The dehydratase activity was then measured. Addition of MgCl$_2$ increases the specific activity of the EDTA-treated d-THA DH.
Figure S4 Time-dependent changes in the total concentration of (a) DL-THA and (b) DL-EHA in the reaction mixture. The concentration in the sample of 0 h incubation was taken as 200 mM, and the relative values for following samples were obtained. Total concentration of DL-THA and DL-EHA decreases to about half the initial value when the enantiomeric excess (e.e.) of L-THA and D-EHA in the reaction mixture reaches >99%, respectively.
Figure S5 HPLC chromatograms recorded at 250 nm of (a) DL-THA purchased from Tokyo Chemical Industry and (b) L-THA obtained by enzymatic optical resolution. Samples were derivatized with GITC and the derivatives were analyzed using HPLC at a flow rate of 1.0 mL min⁻¹ with 30% methanol in water as the mobile phase. Retention times of the D/L-enantiomer of threo-3-hydroxyaspartate were 19.25/20.84 min.
Figure S6 (a) $^1$H NMR spectra recorded at 500 MHz and (b) $^{13}$C NMR spectra recorded at 125 MHz of L-THA obtained by enzymatic optical resolution. Note that the $^1$H signal at 4.77 ppm is attributed to deuterium oxide (D$_2$O).
Figure S7 HPLC chromatograms recorded at 250 nm of (a) DL-EHA synthesized by the protocol described in the Materials and methods and (b) d-EHA obtained by enzymatic optical resolution. Samples were derivatized with GITC and the derivatives were analyzed using HPLC at a flow rate of 1.0 mL min\(^{-1}\) with 35% methanol in water as the mobile phase. Retention times of the D/L-enantiomer of erythro-3-hydroxyaspartate were 17.75/18.56 min.
Figure S8 (a) $^1$H NMR spectra recorded at 500 MHz and (b) $^{13}$C NMR spectra recorded at 125 MHz of d-EHA obtained by enzymatic optical resolution. Note that the $^1$H signal at 4.79 ppm is attributed to deuterium oxide (D$_2$O).
Figure S9 The metal concentration in the solution of recombinant d-THA DH purified by (a) HisTrap Ni^{2+}-affinity column (GE Healthcare, Little Chalfont, UK) and (b) HisTALON Co^{2+}-affinity column (Clontech Laboratories, Inc., Mountain View, CA). These were determined by inductively coupled plasma mass spectrometry (ICP-MS; ELAN DRC-e; Perkin Elmer, Waltham, MA). Enzyme concentrations were taken as 100 μmol/L. Shown are the five top-ranked metals for each analysis (Only four metals were detected in the sample of a). Large amount of Zn^{2+} was detected along with a relatively high content of Ni^{2+} and Co^{2+} after purification by Ni^{2+}- and Co^{2+}-affinity chromatography, respectively. There was a slight or no concentration of Mg^{2+}.
**Figure S10** Thermal stability of d-THA DH. Purified d-THA DH solution was incubated for 15 min at different temperatures (0-65 °C). After incubation, it was placed on ice for 5 min and then the residual dehydratase activity was assayed by the protocol described in the Materials and methods. The activity of the enzyme incubated at 0 °C was taken as 100%. Over 80% of the residual activity was present in the range of temperature between 0 and 45 °C.
Fig. 1 Matsumoto et al.
Fig. 2 Matsumoto et al.
Fig. 3 Matsumoto et al.
Fig. 4 Matsumoto et al.
Fig. 5 Matsumoto et al.
Fig. 6 Matsumoto et al.
## Table 1 Crystallographic parameters and model refinement statistics.

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1. Values in parentheses refer to data in the highest resolution shell.
2. $R_{merge} = \frac{\sum_{i} \sum_{h} I_{h,i} - \langle I_{h} \rangle \sum_{i} \sum_{h} I_{h,i}}{\sum_{i} \sum_{h} I_{h,i}}$, where $\langle I_{h} \rangle$ is the mean intensity of a set of equivalent reflections.
3. $R_{work} = \frac{\sum |F_{obs} - F_{calc}|}{\sum F_{obs}}$ for 95% of the reflection data used in the refinement. $F_{obs}$ and $F_{calc}$ are observed and calculated structure factor amplitudes, respectively.
4. $R_{free}$ is the equivalent of $R_{work}$, except that it was calculated for a randomly chosen 5% test set excluded from refinement.
5. The residues found in the disallowed regions of the Ramachandran plot are Arg141, Phe241 and Ser284; all of these residues have well-defined electron density and their main-chain conformations are reliable.
<table>
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<th>Variant</th>
<th>3-Hydroxyaspartate dehydratase activity (µmol/min/mg protein)</th>
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<td>D-THA</td>
<td>L-EHA</td>
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<td>Wild-type</td>
<td>21.1 ± 0.79 (100)</td>
<td>16.1 ± 0.18 (100)</td>
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<tr>
<td>H351A</td>
<td>0.12 ± 0.001 (0.57)</td>
<td>0.46 ± 0.006 (2.86)</td>
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<td>C353A</td>
<td>N.D.</td>
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Values in parentheses refer to the relative activity (%)

N.D.: not detected (less than 0.01 µmol/min/mg protein)
Supplementary Material for *Applied Microbiology and Biotechnology*

Structural insights into the substrate stereospecificity of *d*-threo-3-hydroxyaspartate dehydratase from *Delftia* sp. HT23: a useful enzyme for the synthesis of optically pure *l*-threo- and *d*-erythro-3-hydroxyaspartate

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**Figure S1** SDS-PAGE of the wild-type and mutants of d-THA DH. *Lanes 1-3* correspond to the wild-type, H351A, and C353A mutants, respectively. *Lane M* contains marker proteins, including (from top to bottom), phosphorylase (*M*ₙ, 97,400), bovine serum albumin (66,300), aldolase (42,400), carbonic anhydrase (30,000), trypsin inhibitor (20,100), and lysozyme (14,400). Gel was stained for protein with Coomassie Brilliant Blue R-250 (Wako Pure Chemical Industries).
Figure S2 Final 2mFo-DFc map of the wild-type d-THA DH around active site in (a) substrate-free form (PDB code, 3WQC) and in (b) complex with d-EHA (PDB code, 3WQD), clearly showing the Schiff-base interchange from lysine-PLP to PLP-d-EHA. Each map was shows contours at 1.4σ level.
Figure S3 Effect of Mg$^{2+}$ on the specific activity of d-THA DH. EDTA-treated enzyme was prepared using the method described earlier (Maeda et al. 2010). It was incubated for 10 min with the reaction mixture in the absence of any metal, or in the presence of 0.1 or 200 mM MgCl$_2$. The dehydratase activity was then measured. Addition of MgCl$_2$ increases the specific activity of the EDTA-treated d-THA DH.
**Figure S4** Time-dependent changes in the total concentration of (a) DL-THA and (b) DL-EHA in the reaction mixture. The concentration in the sample of 0 h incubation was taken as 200 mM, and the relative values for following samples were obtained. Total concentration of DL-THA and DL-EHA decreases to about half the initial value when the enantiomeric excess (e.e.) of L-THA and D-EHA in the reaction mixture reaches >99%, respectively.
**Figure S5** HPLC chromatograms recorded at 250 nm of (a) DL-THA purchased from Tokyo Chemical Industry and (b) L-THA obtained by enzymatic optical resolution. Samples were derivatized with GITC and the derivatives were analyzed using HPLC at a flow rate of 1.0 mL min$^{-1}$ with 30% methanol in water as the mobile phase. Retention times of the D/L-enantiomer of *threo*-3-hydroxyaspartate were 19.25/20.84 min.
Figure S6 (a) $^1$H NMR spectra recorded at 500 MHz and (b) $^{13}$C NMR spectra recorded at 125 MHz of L-THA obtained by enzymatic optical resolution. Note that the $^1$H signal at 4.77 ppm is attributed to deuterium oxide (D$_2$O).
Figure S7 HPLC chromatograms recorded at 250 nm of (a) D,L-EHA synthesized by the protocol described in the Materials and methods and (b) D-EHA obtained by enzymatic optical resolution. Samples were derivatized with GITC and the derivatives were analyzed using HPLC at a flow rate of 1.0 mL min\(^{-1}\) with 35% methanol in water as the mobile phase. Retention times of the D,L-enantiomer of erythro-3-hydroxyaspartate were 17.75/18.56 min.
**Figure S8** (a) $^1$H NMR spectra recorded at 500 MHz and (b) $^{13}$C NMR spectra recorded at 125 MHz of d-EHA obtained by enzymatic optical resolution. Note that the $^1$H signal at 4.79 ppm is attributed to deuterium oxide (D$_2$O).
Figure S9 The metal concentration in the solution of recombinant d-THA DH purified by (a) HisTrap Ni$^{2+}$-affinity column (GE Healthcare, Little Chalfont, UK) and (b) HisTALON Co$^{2+}$-affinity column (Clontech Laboratories, Inc., Mountain View, CA). These were determined by inductively coupled plasma mass spectrometry (ICP-MS; ELAN DRC-e; Perkin Elmer, Waltham, MA). Enzyme concentrations were taken as 100 μmol/L. Shown are the five top-ranked metals for each analysis (Only four metals were detected in the sample of a). Large amount of Zn$^{2+}$ was detected along with a relatively high content of Ni$^{2+}$ and Co$^{2+}$ after purification by Ni$^{2+}$- and Co$^{2+}$-affinity chromatography, respectively. There was a slight or no concentration of Mg$^{2+}$. 
Figure S10 Thermal stability of D-THA DH. Purified D-THA DH solution was incubated for 15 min at different temperatures (0-65 °C). After incubation, it was placed on ice for 5 min and then the residual dehydratase activity was assayed by the protocol described in the Materials and methods. The activity of the enzyme incubated at 0 °C was taken as 100%. Over 80% of the residual activity was present in the range of temperature between 0 and 45 °C.