Three-Dimensional Structure of 4-Amino-4-Deoxychorismate Lyase from Escherichia coli

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4-Amino-4-deoxychorismate lyase (ADCL) is a member of the fold-type ‡W of PLP dependent enzymes that converts 4-amino-4-deoxychorismate (ADC) to p-aminobenzoate and pyruvate. The crystal structure of ADCL from Escherichia coli has been solved using MIR phases in combination with density modification. The structure has been refined to an R-factor of 20.6% at 2.2 Å resolution. The enzyme is a homo dimer with a crystallographic twofold axis, and the polypeptide chain is folded into small and large domains with an interdomain loop. The coenzyme, pyridoxal 5-phosphate, resides at the domain interface, its re-face facing toward the protein. Although the main chain folding of the active site is homologous to those of D-amino acid and L-branched-chain amino acid aminotransferases, no residues in the active site are conserved among them except for Arg59, Lys159, and Glu193, which directly interact with the coenzyme and play critical roles in the catalytic functions. ADC was modeled into the active site of the unliganded enzyme on the basis of the X-ray structures of the unliganded and liganded forms in the D-amino acid and L-branched-chain amino acid aminotransferases. According to this model, the carboxylates of ADC are recognized by Asn256, ArgO7, and Lys97, and the cyclohexadiene moiety makes van der Waals contact with the side chain of Leu258. ADC forms a Schiff base with PLP to release the catalytic residue Lys159, which forms a hydrogen bond with Thr38. The neutral amino group of Lys159 eliminates the ß-proton of ADC to give a quinonoid intermediate to release a pyruvate in accord with the proton transfer from Thr38 to the olefin moiety of ADC.

Key words: aminodeoxychorismate lyase, pyridoxal enzyme, reaction mechanism, tertiary structure, X-ray structure.

In bacteria, chorismate serves as the branch point precursor for metabolites essential for the biosynthesis of many important aromatic products, such as anthranilate (precursor of tryptophan), prephenate (precursor of tyrosine or phenylalanine), p-aminobenzoate (precursor of folic acid), and p-hydroxybenzoate (precursor of ubiquinone) (7). Chorismate is converted into 4-amino-4-deoxychorismate (ADC) by p-aminobenzoate synthase encoded by pabA and pabB. Aminodeoxychorismate lyase (ADCL), which is a product of pabC and contains a tightly bound pyridoxal 5-phosphate (PLP) as a cofactor, then converts ADC to p-aminobenzoate and pyruvate (Scheme 1) (1-5). Thus, ADCL is a key enzyme along with p-aminobenzoate synthase in the biosynthesis of p-aminobenzoate.

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Based on the similarities in sequence, secondary structure, and hydropathy profiles, many PLP-dependent enzymes are placed in fold type I, while ADCL is placed in fold type ‡W together with D-amino acid aminotransferase (DAAT) and branched-chain amino acid aminotransferase (BCAT) (6). The sequence identities of Escherichia coli ADCL (eADCL) with Bacillus species YM-1 DAAT (bs-DAAT) and E. coli BCAT (eBCAT) are 23 and 22%, respectively (7).

In most PLP-dependent enzymes, the catalytic residue lysine shuttles protons on the si-face of the planar ß-system of the substrate-cofactor complex. However, enzymes belonging to fold type IV transfer protons on the re-face of the cofactor (8). The re-face specificity of proton transfer in bsDAAT and eBCAT have been further confirmed by X-ray crystallographic studies of these enzymes (9-11). Because the configuration between the cofactor and catalytic base at the active site is considered to reflect the stereospecificity of
against 400\(^{\circ}\)C of reservoir solution at 293 K. After a few propanol, 100mM citrate buffer, pH 6.5) and equilibrated reservoir solution (17% polyethylene glycol 6000, 17% 2-phosphate buffer, pH 6.5) was mixed with an equal volume of mg/ml protein, 0.1mM PLP, and 20mM potassium phosphate. Thus, this is the first structure elucidation of a PLP-dependent lyase belonging to fold type \(\alpha\) and having \(\beta\)-face specificity Here we report the results of an \(\alpha\)-ray crystallographic study of eADCL in the unliganded form at 2.2 A and propose a reaction mechanism based on the crystallographic study of eADCL in the unliganded form.

the hydrogen transfer, ADCL, DAAT, and BCAT are expected to have a common structure different from those of the other fold types of PLP-dependent enzymes. In contrast to bsDAAT (12) and eBCAT (13, 14) that catalyze transamination, ADCL catalyzes a unique reaction, the elimination of pyruvate from ADC with aromatization of the ring. Structure determination of ADCL should help in understanding the structure-function relationship of not only ADCL but also the PLP-dependent enzymes. X-ray crystallographic studies of PLP-dependent lyases have been carried out on tyrosine phenol-lyase (15), tryptophan indole-lyase (16) and cystathionine \(\beta\)-lyase (17), all of which belong to fold type I and have \(\sigma\)-face specificity of proton transfer. Thus, we report the results of an X-ray crystallographic study of eADCL in the unliganded form at 2.2 A and propose a reaction mechanism based on the model structure for the ADC-PLP external aldimine intermediate.

MATERIALS AND METHODS

Protein Expression, Purification, and Crystallization—
The structural gene of eADCL, was inserted between the EcoRl and HindIII restriction sites of plasmid pKK223-3. E. coli JM109 was transformed with the resultant plasmid, pADCL. The enzyme was purified by ammonium sulfate fractionation and three-step column chromatography; first on a DEAE-Toyopearl column (TOSOH) with a linear gradient from 0 to 0.3 M KCl, followed by a Butyl Toyopearl column (TOSOH) using a linear gradient of saturated ammonium sulfate (30 to 0%), and finally by a MonoQ column (Pharmacia) with a linear gradient from 0.1 to 0.5 M KCl. Selenomethionyl ADCL was prepared by overexpressing pADCL in DL41metA-1 cells grown in the presence of selenomethionine. The selenomethionyl protein was purified and crystallized similar to the wild type one.

Preliminary crystallization conditions for the PLP-type eADCL in the unliganded form were determined using the sparse-matrix method (18) along with the hanging-drop vapor-diffusion method (19). Several crystal forms were obtained and one of the most promising crystallization conditions was optimized. A droplet of 5 \(\mu\)l protein solution (5 mg/ml protein, 0.1 mM PLP, and 20 mM potassium phosphate buffer, pH 6.5) was mixed with an equal volume of reservoir solution (17% polyethylene glycol 6000, 17% 2-propanol, 100 mM citrate buffer, pH 6.5) and equilibrated against 400 \(\mu\)l of reservoir solution at 293 K. After a few days, small yellow prisms started to grow and were used as seeds for growing large crystals under the same hanging drop conditions. Within 2 weeks, crystals had grown to dimensions of about 0.2 \(\times\) 0.2 \(\times\) 0.8 mm.

Data Collection—An X-ray diffraction data set for a native crystal was collected to 2.2 A resolution at 287 K on the BL6A station at the Photon Factory, KEK (Tsukuba), using an X-ray beam of wavelength 1.00 A and Fuji imaging plates with a screenless Weissenberg camera (20). The space group is P2\(_1\)2\(_1\)2\(_1\) with cell dimensions of \(a = 40.5\), \(b = 73.9\), and \(c = 83.7\) A. There is one subunit in the asymmetric unit, and approximately 42% of the crystal volume is occupied by solvent. The data sets for crystals soaked in 1 mM methylmercury chloride and 1 mM p-chloromercuribenzoate, and a selenomethionyl protein crystal were collected to 2.5, 4.0 and 2.5 A resolution, respectively, at 293 K with a Rigaku R-Axis IIC imaging plate detector using monochromated Cu K\(\alpha\) radiation (40 kV, 100 mA). All data were processed and scaled using the programs DENZO and SCALEPACK (21) (Table I).

Structure Determination—The primary sequence of eADCL is 22 and 23% identical to those of eBCAT and bsDAAT, respectively (7). The molecular replacement method was used to solve the structure of ADCL using the known structures of bsDAAT and eBCAT as the search models. One marked solution was obtained for both models, and the electron density maps calculated from either model gave the assembly of \(\beta\)-strands in the protein core region. However, the assembly could not be extended to the overall folding of the protein. Therefore, the structure of ADCL was solved by the MIR method using three isomorphous data sets. Data scaling and map calculations were performed with the CCP4 program suite (22). The difference Patterson map calculations for methylmercury chloride using the data from 15.0-3.5 A resolution and p-chloromercuribenzoate using the data from 15.0-4.0 A resolution allowed a clear interpretation of the two and one mercury sites, respectively. A common origin for these derivatives was determined by calculating the difference Fourier summation for p-chloromercuribenzoate based on the phases calculated by the mercury sites of methylmercury chloride. Two other minor mercury sites of methymercury chloride derivative were determined from the difference Fourier maps based on the phase calculated using the major mercury sites. The positions of six selenium sites out of seven in the selenomethionyl ADCL were determined from the difference Fourier maps based on mercury phasing. Refined of the heavy atom parameters and calculation of the initial phases were performed with the program MLPHARE (22). The resulting MIR map has a mean figure-of-merit of 0.46 at a resolution of 15 to 3.0 A. The map was significantly improved by the process of solvent flattening (23) and histogram matching (24) using the program DM (22). The mean figure-of-merit reached 0.78 with the same resolution range.

Although, at this stage, the quality of the electron density map was not good enough to build an entire molecule, the \(\beta\)-sheets in the core region were located at the same position as that of the assembly of \(\beta\)-strands observed in the map calculated on the basis of molecular replacement. Moreover, the selenium sites determined by the difference Fourier map coincided with the positions with methionines expected for the models determined by molecular replacement. Thus, a partial model consisting of 127 out of the 289

\[\begin{align*}
\text{O} & \quad \text{O} \\
\text{O} & \quad \text{O} \\
\text{NH}_2 & \quad \text{p-aminoacetate}
\end{align*}\]

Scheme 1 Reaction catalyzed by ADCL.
residues and PLP was built using the electron density of the core region with the program O (25). The phases calculated from this model were combined with the MIR phases by the program SIGMAA (26). The mean figure-of-merit reached 0.78 at a resolution of 15–30 Å. Two cycles of this procedure allowed us to build a model consisting of 207 out of the 269 residues. The structure was refined by simulated annealing using the program X-PLOR (27, 28) using X-ray data from 8.0–3.0 Å resolution. The structure including the PLP molecule was scrutinized by successively omitting 10 residue segments of the model from the phasing calculation and inspecting the map to give an additional 47 residues. Refinement by simulated annealing and rebuilding was alternated until no further improvements in structure and statistics were apparent with an R_free of 27.2% and R_ref of 38.7%. The resolution was progressively increased to 2.2 Å, and after several rounds of refinement and manual rebuilding, the R_free and R_ref values were reduced to 24.6 and 32.3%, respectively. Water molecules were picked up on the basis of peak heights and distance criteria from the difference map. Water molecules whose thermal factors were above 49 Å² (maximum thermal factor of the main chain) after refinement were removed from the list. Further model building and refinement cycles resulted in an (R_fo - R_calc) map. However, this model lacks the following 15 residues due to poor electron density: Tyr126 to Gin137, the residues connecting two domains (Pro 116 and Pro260) were found in the cis-conformation. Two proline residues (Pro116 and Pro260) were found in the cis-conformation. Two proline residues (Pro116 and Pro260) were found in the cis-conformation.

**Quality of the Structure**—The refined model of eADCL contains 1964 non-hydrogen protein atoms and a covalently bound molecule of PLP. Additionally, 126 water molecules were included (Table I). Almost all amino acid residues were found to have good electron densities in the final 2Fo – Fcalc electron density map. However, this model lacks the following 15 residues due to poor electron density: Tyr126 to Gin137, the residues connecting two domains (Pro 116 and Pro260) were found in the cis-conformation.

**Table 1. Data collection, MIR, and refinement statistics.**

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<td>13.5</td>
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<tr>
<td>Water atoms (Å^2)</td>
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\( R_{	ext{free}} = \frac{\sum_{i} \left| I_{\text{obs}} - I_{\text{calc}} \right|}{\sum_{i} I_{\text{obs}}} \), where \( I \) = observed intensity and \( \langle I \rangle \) = average intensity for multiple measurements. \( R_{\text{free}} = \frac{\sum_{i} F_{\text{obs}} - F_{\text{calc}}}{\sum_{i} F_{\text{obs}}} \), where \( F_{\text{obs}} \) and \( F_{\text{calc}} \) are the derivative and calculated structure factor amplitudes, respectively. Phasing power is the ratio of the root mean square r.m.s.) of the heavy atom scattering amplitude and the lack of closure error. The values in parentheses are for the highest resolution shell (2.3–2.2 Å). p-chloromercuribenzoate.
Fig. 1. Alignments of bsDAAT, eBCAT, and four ADCLs with secondary structures. α-Helices are denoted as α1-α17 and β-strands as β1-β17. bsDAAT and eBCAT are from Bacillus sp. YM-1 and E. coli, respectively. eADCL, vHAADCL, bsADCL, and aaADCL are from E. coli, Vibrio harveyi, Bacillus subtilis, and Aquifex aeolicus, respectively. In the consensus sequence, capital letters show conserved residues, and small letters denote homologous residues in ADCLs. The numbering of the primary sequence of bsDAAT is from Ref. 9.

RESULTS AND DISCUSSION

Primary Structure—The primary sequence of eADCL was aligned versus those of bsDAAT and eBCAT based on the three-dimensional structures. The details of the procedure was previously described (33). Sequence alignment of eADCL with other ADCLs was performed with the program CLUSTAL W (34). As a result of these alignments (Fig. 1), the sequence identities of eADCL with respect to bsDAAT, eBCAT, Vibrio harveyi ADCL (35), Bacillus subtilis ADCL, and Aquifex aeolicus ADCL (pir:C64593) are 21, 20, 35, 26, and 20%, respectively. Helicobacter pylori ADCL (pir:C64593) is omitted from the alignments because the sequence identity with that of eADCL is too low (<10%). The 269 amino acid residues of eADCL are numbered from Met7 to Asn292 according to the sequence of eBCAT (13).

Overall Structure—The eADCL is folded into a dimeric form with a crystallographic two-fold axis, and each subunit is composed of 269 amino acid residues with a subunit molecular weight of 29,715 and PLP. The overall structure of eADCL is shown in Fig. 2. The Cα atoms of eADCL were superimposed against those of bsDAAT and eBCAT by a least-squares fitting with r.m.s. deviation of 1.93 Å for 92.9% structurally equivalent Cα atoms and 2.10 Å (94.4%), respectively.

In the consensus sequence, capital letters show conserved residues, and small letters denote homologous residues in ADCLs. The numbering of the primary sequence of bsDAAT is from Ref. 9.

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The arrangement of the α-helices and β-sheets of eADCL as a whole is similar to those of bsDAAT and eBCAT of fold type IV (Fig. 1), indicating that among these molecules the secondary and overall structures of the enzymes are conserved.

The subunit of eADCL is divided into one small domain comprising the N-terminus to Ala125 and one large domain comprising Ile139 to the C-terminus. The interdomain loop connecting the small and large domains (Tyr126–Gly137) was disordered in the unliganded form of eADCL, and could not be modeled. Also, in eBCAT, the corresponding interdomain loop in the unliganded form is disordered.

The N-terminal small domain is an α/β structure with an open β-sheet structure. The six antiparallel β-strands designated as β2, β1, β6, β5, β3, and β4 form a twisted β-sheet structure as a central core surrounded by two α-helices (α2 and α3) from the surface side of the protein and a short α-helix (α1) around the twofold axis from the interior side (Figs. 1 and 3). The C-terminal large domain is also an α/β structure with a pseudo β-barrel structure. The β-sheet comprising four strands designated as β9, β8, β7, and β15 (all parallel except for β9) and that comprising five strands designated as β12, β11, β10, β13, and β14 (all antiparallel except for β12) form a pseudo β-barrel surrounded by three
α-helices (α5, α6, and α7) from the surface side of the protein and an α-helix (α4) from the interior side.

The β-strand (β0) and that from the other subunit related by the twofold axis (β0' shown in Fig. 3) form an inter-subunit anti-parallel β-sheet. The α-helix (α1) and the loop between the β-strands (β5 and β6) are also located around the molecular twofold axis participating in the formation of the subunit interface.

Fig. 4. Stereoview of the 2Fo–Fc electron density map calculated using data between 8.0–2.2 Å resolution for PLP and residues close to PLP (a, top). A close-up view of the active site of the unliganded eADCL (b, middle) and the eADCL-ADC complex model (c, bottom) from nearly the same direction as that in Fig. 3. PLP (pale yellow), ADC (pale red) and selected residues of the active site are shown for clarity. Two loops (pink) of the other subunit participate in the formation of the active site.
The Active Site Structure of the Unliganded eADCL—The active site cleft of the enzyme is located at the domain and subunit interface. The active site comprises the residues from both domains of one subunit and the small domain (two loops shown in Fig. 3) of the other subunit of the dimer unit. The 2Fo–Fc electron density map for PLP and the residues around it is shown in Fig. 4a. The structure and hydrogen bonding scheme of the active site are shown in Figs. 4b and 5a, respectively.

PLP binds to the pocket by forming an internal aldimine bond (Schiff base linkage) with the catalytic residue Lys159 and by extensive noncovalent interactions with the residues of the active site. The imine nitrogen of the internal aldimine bond is displaced by 0.65 Å from the pyridine ring of PLP with the dihedral angle of C3-C4-C4'-N = -34°, and makes a hydrogen bond with O3' of PLP. PLP is sandwiched between the main chain of Ala196-Ala197 and the side chain of Val217 from above and below. Glu193 forms an ion pair with the protonated nitrogen atom of the pyridine ring of PLP, and is considered to strengthen the electron withdrawing effect of the pyridine ring of PLP as an electron sink, as has been observed for enzymes of fold types I and IV (38-40). Tyr109 is hydrogen bonded to the O3' of PLP to regulate the electronic state of PLP (41, 42). Residue 109 is Val in eBCAT (11) and His in bsDAAT (9, 10), and Tyr164 and a water molecule take the place of Tyr109, respectively. Some ADCLs in Fig. 1 do not have Tyr109, but have a Tyr164 as in eBCAT. The phosphate group of PLP is involved in seven hydrogen bonds and acts as an anchor to fix the cofactor to the active site. The negative charge of the phosphate group is well balanced with the positive charge of Arg59 and the dipole of the α-helix (α5), whose N-terminus is close to the phosphate group.

The PLP-dependent enzymes have ω-face specificity of proton transfer with respect to the C4' of the cofactor except for enzymes of fold type IV. The re-face specificity of the proton shuttle was first clarified for bsDAAT and eBCAT by hydrogen transfer experiments (8), and further confirmed by X-ray crystallographic studies of these enzymes. eADCL was then proposed to have ω-face specificity (43). An X-ray study of eADCL showed that PLP has the same orientation toward the protein as that found in bsDAAT or eBCAT (9, 11), and rotates by 180° compared with other PLP-dependent enzymes (38). The re-face of the PLP-ring faces the protein side, and the catalytic residue Lys159 transfers protons on the re-face of PLP. Thus, the ω-face specificity of eADCL was unambiguously determined.

The bottom of the active site pocket consists of Phe36, Arg59, Lys159, Arg164, Glu193, Val217, Ile220, and Met221. The side or wall of the active site is made up of Thr37, Thr38, Arg40, Ala219, Lys97, Ala196, Asn256,

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Fig. 5. Schematic diagram showing hydrogen bond and salt bridge interactions of the active site residues in the unliganded eADCL (a, left) and the eADCL-ADC complex model (b, right). Putative interactions are shown by dotted lines if the acceptor and donor are less than 3.5 Å apart.
Ala257, Leu258, Phe31, Arg107, Gly108, and Tyr109. Three water molecules (W1, W2, and W3) are located in the vicinity of the PLP ring, and are involved in the formation of the hydrogen bonding network as shown in Fig. 5a. When the active site residues of eADCL are compared with those of eBCAT (11) and bsDAAT (9), the only residues conserved are Arg59, Lys159, and Glu193, all in the bottom of the active site, and conserved in all known ADCLs. Arg59 is the key residue fixing the phosphate group of PLP, and Lys159 and Glu193 are catalytically-critical residues as described above. None of the residues from the wall of the active site are conserved. In spite of the poor conservation of the active site residues, the main chain atoms of the active site residues of eADCL fit those in eBCAT and bsDAAT with r.m.s. deviations of 0.89 and 0.78 Å, except for the residues 258 and 107, respectively. The eADCL, eBCAT, and bsDAAT enzymes belonging to fold type IV are divergently evolved from the same ancestor to have different substrate and reaction specificities by replacing most of the active site residues except for the critical Arg59, Lys159, and Glu193 residues for enzyme functions, but by maintaining the main chain folding of the active site. This result indicates that in principle PLP-dependent enzymes with quite different substrate specificities and/or different reaction specificities can be designed by protein engineering starting from fold type IV enzymes, but in practice, the engineering is quite difficult since too many residues must be rationally replaced.

The primary sequences of ADCLs depicted in Fig. 1 show that in addition to Arg59, Lys159, Glu193, and Val217, the roles of which have already been described, Phe36, Thr38, Lys or Arg97, Asn256, Leu or Val258, and Arg or Lys107 are conserved in almost all ADCLs as the active site residues. These residues are possibly concerned with substrate recognition or the catalytic action of the enzyme.

Interestingly, upon binding of a substrate analogue, the interdomain loop of eBCAT shows its ordered structure to close the active site and shield the analogue from the solvent region, and plays important roles in substrate recognition (30). Similarly, the disordered interdomain loop of eADCL might be ordered to close the active site upon binding of a substrate, but the exact behavior of the loop can not be seen until the structure of eADCL in complex with a substrate or inhibitor is determined.

The Active Site Structure of the eADCL-ADC Complex Model—A model of eADCL in complex with the substrate

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Fig. 6. The proposed reaction mechanism of eADCL with ADC to produce p-aminobenzoate and pyruvate.
ADC (the external aldimine model) was constructed and the energy minimized with X-PLOR (31) as described in "MATERIALS AND METHODS". The active site structure and hydrogen bonding scheme of the final model are displayed in Figs. 4c and 5b, respectively. On binding of ADC, two water molecules, W2 and W3, are expelled from the active site of the unliganded eADCL, and PLP forms a new Schiff base with ADC to release Lys159 with a neutral amino group, which is hydrogen bonded to the hydroxy group of Thr38. All the residues of the active site in the complex model have essentially the same positions as those in the unliganded eADCL except for Arg107 and Lys159, whose side chains move by more than 1 Å. Almost all the hydrogen bonds found in the active site of the unliganded eADCL are thus maintained in the complex model. Glu193 makes a salt bridge with the protonated nitrogen atom of the pyridine ring of PLP. Tyr109 is hydrogen bonded to the O3' of PLP, and the phosphate group of PLP is fixed to the active site by hydrogen bonds.

The amino group of Asn256 is hydrogen bonded to the carbonyl group of the cyclohexadiene moiety of the bound ADC. Arg107 changes its side chain direction toward the active site to form salt bridges with two carboxylates of ADC. The directional change in Arg107 is reminiscent of the behavior of Arg292 in aspartate aminotransferases upon the binding of a substrate (44-47). The protonated amino group of Lys97 interacts with the carbonyl of the olefin moiety. The side chain of Leu258 makes van der Waals contact with the cyclohexadiene moiety of ADC.

Mechanistic Implications—A possible reaction mechanism of eADCL with ADC to give p-aminobenzozoate and pyruvate is shown in Fig. 6 based on the stereochemical consideration of the external aldimine model described above and the proposed mechanism (7). The eADCL binds ADC to give a Michaelis complex (a in Fig. 6). ADC is reasonably postulated to occupy the same position as that of the ADC moiety in the external aldimine form, since the X-ray structures of the aspartate aminotransferases and eBCAT show that substrate analogues in Michaelis complexes do not change their locations during the formation of the external aldimines (30, 44-46). The amino group of ADC is directed toward the PLP-Lys159 Schiff base linkage, and the C4' of PLP undergoes nucleophilic attack by the amino group of ADC. Through the tetrahedral intermediate, a PLP-ADCL Schiff base (external aldimine) is formed between PLP and ADC to release Lys159 (b). The neutral amino group of Lys159 makes a hydrogen bond with Thr38, which will decrease the free energy level of the external aldimine form of the enzyme. The OH group of Thr38 makes van der Waals contact with the methylene group of ADC. The α-carbon of the substrate is activated by the protonated Schiff base and the protonated pyridine ring. The α-proton of the substrate is nearly perpendicular to the plane defined by the Schiff base, pyridine ring and α-carbon, and is directed toward Lys159, which is the only base able to abstract the α-proton. Thus, good conditions for α-proton elimination are achieved to give a quinonoid intermediate (b and c). The quinonoid intermediate is stabilized by the delocalization of the cofactor-substrate π-system and the hydrogen bond between N1-H of the cofactor ring and Glu193. The electron would migrate toward the substrate from the cofactor in accord with the proton transfer from Thr38 to the olefin moiety of the substrate, resulting in the release of a pyruvate with aromatization of the six-membered ring of the substrate (c and d). Then p-aminobenzozoate will be liberated to recover the starting Schiff base by the reverse of the reaction process from step b.

The final elucidation of the substrate recognition and reaction mechanism of ADCL awaits the X-ray crystallographic determinations of eADCL in complex with substrate analogues, which are now in progress. The results reported here form the basis for future investigations on the structure-function relationships of eADCL and other related members of fold type IV PLP-dependent enzymes, including studies of the catalytic action by inhibitor binding and site-directed mutagenesis.

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REFERENCES

philic Bacillus species and its correlation with L-amino acid aminotransferases. J. Biol. Chem. 264, 2450-2454