Structural Basis of Substrate Recognition by Aldehyde Dehydrogenase 7A1

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ABSTRACT: Aldehyde dehydrogenase 7A1 (ALDH7A1) is part of lysine catabolism and catalyzes the NAD⁺-dependent oxidation of α-aminoadipate semialdehyde to α-aminoadipate. Herein, we describe a structural study of human ALDH7A1 focused on substrate recognition. Five crystal structures and small-angle X-ray scattering data are reported, including the first crystal structure of any ALDH7 family member complexed with α-aminoadipate. The product binds with the ε-carboxylate in the oxyanion hole, the aliphatic chain packed into an aromatic box, and the distal end of the product anchored by electrostatic interactions with five conserved residues. This binding mode resembles that of glutamate bound to the proline catabolic enzyme ALDH4A1. Analysis of ALDH7A1 and ALDH4A1 structures suggests key interactions that underlie substrate discrimination. Structures of apo ALDH7A1 reveal dramatic conformational differences from the product complex. Product binding is associated with a 16 Å movement of the C-terminus into the active site, which stabilizes the active conformation of the aldehyde substrate anchor loop. The fact that the C-terminus is part of the active site was hitherto unknown. Interestingly, the C-terminus and aldehyde anchor loop are disordered in a new tetragonal crystal form of the apoenzyme, implying that these parts of the enzyme are highly flexible. Our results suggest that the active site of ALDH7A1 is disassembled when the aldehyde site is vacant, and the C-terminus is a mobile element that forms quaternary structural interactions that aid aldehyde binding. These results are relevant to the c.1S2delG genetic deletion associated with pyridoxine-dependent epilepsy, which alters the C-terminus of ALDH7A1.

Aldehyde dehydrogenase 7A1 (ALDH7A1, also called antiquitin) plays a role in lysine catabolism by catalyzing the NAD⁺-dependent oxidation of α-aminoadipate semialdehyde (AASA) to α-aminoadipate (AA) (Figure 1A). ALDH7A1 and other ALDHs are important biomarkers of cancer stem cells and play a functional role in cancer stem cell-mediated metastasis and cancer drug resistance. Over-expression of ALDH1A1, ALDH1A3, ALDH2, ALDH4A1, and ALDH7A1 isoforms has been found in several types of cancers. A high level of expression of ALDH7A1 occurs in prostate cancer cells lines, prostate cancer tissue, and matched bone metastasis samples, suggesting that ALDH7A1 plays a functional role in prostate cancer bone metastasis. In addition, patients with ALDH7A1-expressing non-small cell lung carcinoma tumors have a significantly increased incidence of lung cancer recurrence, and the level of ALDH7A1 expression is increased in ovarian tumors.

Mutations in ALDH7A1 cause the inherited seizure disorder pyridoxine-dependent epilepsy (PDE). PDE is an autosomal recessive epileptic encephalopathy caused by mutations in the gene encoding ALDH7A1. PDE is characterized by intractable seizures that are not controlled with antiepileptic drugs but that respond to large daily supplements of pyridoxine, a form of vitamin B6. Disruption of ALDH7A1 activity in PDE patients causes a buildup of AASA, which forms a spontaneous equilibrium with Δ¹-piperidine-6-carboxylic acid (Figure 1B). The accumulating Δ¹-piperidine-6-carboxylic acid inactivates pyridoxal S'-phosphate via Knoevenagel condensation (Figure 1B), leading to a pyridoxal S'-phosphate deficiency. Because pyridoxal S'-phosphate is a cofactor in many enzymes, this chemically induced pyridoxal S'-phosphate deficiency has major metabolic consequences, including changes in brain chemistry that lead to seizures. The mutational spectrum of PDE is very large. More than 60 different mutations within the 18 exons of the ALDH7A1 gene have been reported, including splice site mutations, insertions, deletions, nonsense mutations, and more than two dozen missense mutations.

Herein, we describe a structural and biochemical study of ALDH7A1 focusing on substrate recognition and conformational change. We report the first structures of the full-length human enzyme, including the first structure of any ALDH7 family member complexed with the product AA. The structures show that the C-terminus is a mobile active site element that facilitates substrate recognition.

EXPERIMENTAL PROCEDURES

Crystalization of Apo ALDH7A1 in Space Group C2. Full-length recombinant human cytosolic ALDH7A1 isoform 2 (511 residues, NCBI RefSeq NP_001188306.1) was expressed and purified as described previously.

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Figure 1. Reactions related to ALDH7A1 and PDE. (A) Reaction catalyzed by ALDH7A1. (B) Inactivation of pyridoxal 5′-phosphate by Δ1-piperidine-6-carboxylic acid.

Centered monoclinic crystals of apo ALDH7A1 were grown in sitting drops (CrystalEX 96-well plates from Hampton Research) at 22 °C using drops formed by mixing 3 μL of each of the protein and reservoir solutions. The protein stock solution contained 3 mg/mL ALDH7A1 in a buffer containing 50 mM Tris (pH 7.8), 0.5 mM tris(3-hydroxypropyl)phosphine (THP), 50 mM NaCl, and 5% glycerol (precrystallization buffer). The reservoir contained 0.2 M ammonium sulfate, 20% (w/v) polyethylene glycol (PEG) 3350, and 0.1 mM Bis-Tris (pH 6.5), which was obtained by optimizing a hit from crystal screening trials using commercially available kits (Hampton Research). The crystals were cryoprotected in the reservoir solution supplemented with 16% (v/v) PEG 200, picked up with Hampton nylon loops, and plunged into liquid nitrogen. The space group is C2 with the following unit cell dimensions: a = 156 Å, b = 162 Å, c = 159 Å, and β = 95°.

Cocrystallization of ALDH7A1 with L-2-Aminoadipate in Space Group C2. L-2-Aminoadipate (AA) was purchased from Sigma-Aldrich (catalog no. A7275). A stock solution of 300 mM AA in 1 M HCl was prepared, and then AA was added to precryostallization buffer to a final concentration of 100 mM with the pH adjusted to 7.5. A volume of the 100 mM AA solution was mixed with an equal volume of a 6 mg/mL ALDH7A1 stock solution in precryostallization buffer supplemented with 1.5% DMSO. The pH of the resulting solution was adjusted to 7.5, and the solution was stored overnight at 4 °C. Although the enzyme/AA solution turned cloudy, it was nevertheless used in matrix microseeding crystal screening trials using commercially available kits (Hampton Research). The crystal screens were performed in sitting drops at 22 °C in CrystalEX 96 trays. The sitting drops were formed by mixing 3 μL of each of the reservoir and protein solutions. The microseed stock for these trials was prepared by crushing crystals of C2 apo ALDH7A1 in a buffer consisting of 0.2 M ammonium sulfate, 20% (w/v) PEG 3350, and 0.1 mM Bis-Tris (pH 6.5). After the protein and reservoir solutions were mixed, the microseeds were added to the drops by streaking with a horsehair. Following optimization of hits from the crystal screening trials, the C2 crystal form described above was obtained using a reservoir consisting of 0.1 M MgCl2, 0.1 M sodium acetate trihydrate, 0.1 M Tris-HCl (pH 8.0), and 20% (w/v) PEG 4000. The crystals were cryoprotected with 0.1 M MgCl2, 0.1 M sodium acetate trihydrate, 0.1 M Tris-HCl (pH 8.0), 20% (w/v) PEG 4000, and 15% (v/v) PEG 200, picked up with Hampton nylon loops, and plunged into liquid nitrogen. The space group is C2 with the following unit cell dimensions: a = 154 Å, b = 163 Å, c = 159 Å, and β = 95°.

Cocrystallization of Apo ALDH7A1 in Space Group P42,2 and Soaking with NAD+. Tetragonal crystals of apo ALDH7A1 were grown using the sitting drop method of vapor diffusion (CrystalEX 96-well plates from Hampton Research) at 22 °C with drops formed by mixing 3 μL of each of the protein and reservoir solutions. The protein stock solution consisted of 3 mg/mL ALDH7A1 in a buffer containing 50 mM Tris (pH 7.8), 0.5 mM THP, 50 mM NaCl, and 5% glycerol. The best crystals were grown in 0.1 M Hepes (pH 7.5), 0.1 M MgCl2, and 20% (w/v) PEG 3350. The crystals were cryoprotected using the reservoir supplemented with 18% (v/v) PEG 200, picked up with Hampton nylon loops, and plunged into liquid nitrogen. The space group is P42,2 with the following unit cell dimensions: a = b = 159 Å, and c = 80 Å. The asymmetric unit includes two protein molecules, which implies 46% solvent content and a V_m of 2.28 Å³/Da. The ALDH7A1–NAD⁺ complex was obtained by soaking apo tetragonal ALDH7A1 crystals with the cryobuffer containing 1 mM NAD⁺.

X-ray Diffraction Data Collection, Phasing, and Refinement. X-ray diffraction data for C2 apo ALDH7A1 crystals were collected on Northeastern Collaborative Access Team beamline 24-ID-E at the Advanced Photon Source using a Q315r detector. The 2.4 Å resolution data set used for refinement consisted of 180 frames with an oscillation width of 1° per image, a detector distance of 310 mm, and an exposure time of 1 s per image.
X-ray diffraction data from monoclinic and tetragonal crystals of the ALDH7A1—NAD⁺ complex were collected on beamline 4.2.2 at the Advanced Light Source using a Taurus-1 CMOS detector in shutterless mode. Each data set consisted of 1800 frames covering a total rotation range of 180° with the detector distance set to 240 mm. The total exposure times were 540 s for the C2 data set and 180 s for the tetragonal data set.

X-ray diffraction data for crystals of the ALDH7A1−AA complex and tetragonal apo ALDH7A1 were collected on Structural Biology Center beamline 19-ID-C at the Advanced Photon Source using a Q315r detector. The 1.76 Å resolution data set used for refinement of the ALDH7A1−AA complex consisted of 720 frames with an oscillation width of 0.25° per image, a detector distance of 230 mm, and an exposure time of 1 s per image. The 1.90 Å resolution data set used for refinement of tetragonal apo ALDH7A1 consisted of 360 frames with an oscillation width of 0.5° per image, a detector distance of 260 mm, and an exposure time of 1 s per image.

The data sets were processed with XDS15 and AIMLESS16 via CCP4i.17 PHENIX18 and COOT19 were used for refinement and model building, respectively. Crystallographic structure refinements against the C2 data sets were initiated from the coordinates of truncated ALDH7A1 [Protein Data Bank (PDB) entry 2J6L].14 Initial phases for the tetragonal data sets were determined using molecular replacement as implemented in PHASER20 with a search model derived from a dimer of human ALDH7A1 (PDB entry 2J6L). MolProbity was used for structure validation.21 Data collection and refinement statistics are listed in Tables 1 and 2.

Small-Angle X-ray Scattering (SAXS). SAXS experiments were performed at beamline 12.3.1 of the Advanced Light Source via the mail-in program.22,23 Prior to data collection, an ALDH7A1 sample was passed through a Superdex 200 size exclusion column. The column buffer consisted of 50 mM Tris, 5% glycerol, 0.5 mM TTH, and 50 mM NaCl (pH 7.8). Scattering intensities were measured at three nominal protein concentrations using exposure times of 0.5, 1.0, 3.0, and 6.0 s.
Scattering curves collected from the protein samples were corrected for background scattering using intensity data collected from the Superdex 200 column effluent. Composite scattering curves for each protein concentration were generated with PRIMUS by scaling and merging the background-corrected high-q region from the 3 s exposure with the low-q region from the 0.5 or 1.0 s exposure. PRIMUS was also used for Guinier analysis. Shape reconstruction calculations were performed with the sastbx.shapeup module of the Small Angle Scattering ToolBox server. The shape reconstruction calculations employed the PISA database of shapes. The FoXS server was used to calculate SAXS profiles from atomic coordinates.

## RESULTS

**ALDH7A1 Adopts the Standard ALDH Superfamily Fold.** Crystal structures of ALDH7A1 were determined in space groups C2 (Table 1) and P42,2 (Table 2). ALDH7A1 exhibits the ALDH superfamily fold consisting of NAD⁺-binding, catalytic, and oligomerization domains (Figure 2A).

### Table 2. X-ray Diffraction Data Collection and Refinement of Tetragonal Structures

|                   | apo                  | NAD⁺                 |
|-------------------|----------------------|----------------------|
| beamline          | APS 19-ID-C          | ALS 4.2.2            |
| space group       | P42,2                | P42,2                |
| unit cell parameters | a = 159.3 Å         | a = 158.9 Å         |
| wavelength        | 0.97 Å               | 1.00Å                |
| resolution (Å)    | 46.0–1.90 (1.93–1.90) | 53.0–1.90 (1.94–1.90) |
| no. of observations | 1175993              | 1175225              |
| no. of unique reflections | 81138              | 80249                |
| R<sub>merge</sub>(I) | 0.097 (1.209)      | 0.102 (1.445)        |
| R<sub>meas</sub>(I) | 0.100 (1.256)      | 0.105 (1.504)        |
| mean I/σ          | 20.9 (2.4)          | 23.7 (2.5)           |
| mean CC1/2        | 0.999 (0.743)       | 0.999 (0.697)        |
| completeness (%)   | 99.8 (96.5)         | 99.7 (95.0)          |
| multiplicity      | 14.5 (13.5)         | 14.6 (13.3)          |
| no. of protein chains | 2                   | 2                    |
| no. of protein residues | 948                | 946                  |
| no. of protein atoms | 7076                | 6994                 |
| no. of NAD⁺ atoms | –                   | 23                   |
| no. of water molecules | 312                | 385                  |
| R<sub>crys</sub> | 0.169 (0.281)       | 0.182 (0.269)        |
| R<sub>free</sub> | 0.195 (0.329)       | 0.213 (0.305)        |
| rmsd for bond lengths (Å) | 0.007              | 0.007                |
| rmsd for bond angles (deg) | 0.973              | 0.967                |
| Ramachandran plot (%) | favored | 98.30 | 98.51 |
|                     | outliers | 0.11 | 0.00 |
|                     | all-atom | 1.3  | 2.2  |
|                     | Clashcore © | 34.8 | 35.2 |
|                     | average B factor (Å²) | 35.0 | 32.8 |
|                     | protein | 0.19 | 0.20 |
|                     | NAD⁺ | – | 35.1 |
|                     | water | 35.0 | 32.8 |
| coordinate error (Å) | PDB entry | 4ZVX | 4ZVY |
|                     | PDB entry | 4ZVX | 4ZVY |

“Values for the outer resolution shell of data are given in parentheses. The 5% test set. *Generated with MolProbity via PHENIX refine. The maximum likelihood-based coordinate error estimate reported by PHENIX refine.”

The NAD⁺-binding domain adopts the Rossmann fold. As in other ALDHs, the core of this domain is a parallel five-stranded β-sheet having a 32145 strand order. This topology differs from the classic Rossmann fold sheet, which has six strands arranged in a 321456 strand order. The catalytic domain has an α/β structure and contains the nucleophilic Cys302 residue that attacks the carbonyl of the aldehyde substrate. The oligomerization domain is a β-substructure that mediates domain-swapped dimerization. The active site is located in the crevice between the NAD⁺-binding and catalytic domains.

Two sections of the polypeptide chain link the NAD⁺-binding and catalytic domains. These sections are notable because they form part of the active site. The short linker consisting of residues 269–271 connects strand S of the Rossmann sheet to the beginning of the catalytic domain. This linker interacts with the carboxamide of the cofactor.
other linker is longer and consists of residues 461–482 (yellow in Figure 2A). As described below, this linker interacts with the product AA and adopts multiple conformations.

**ALDH7A1 Forms the Classic ALDH Dimer-of-Dimers Tetramer in Solution.** Small-angle X-ray scattering (SAXS) was used to determine the oligomeric state and quaternary structure of ALDH7A1 in solution. Guinier analysis of three ALDH7A1 samples having different protein concentrations yields a radius of gyration ($R_g$) of 37.7 ± 0.3 Å (Figure 3A).

ALDHs are known to form domain-swapped dimers, and in some cases, the dimers assemble into tetramers or hexamers. The classic ALDH tetramer is present in both the $C_2$ and $P4_2_1_2$ lattices (Figure 2B). The tetramer is a dimer of dimers having 222 symmetry, which can be described by three mutually perpendicular 2-fold axes (Figure 2B). This tetramer is also present in the $F_{222}$ and $I_{422}$ crystal forms of covalently inactivated ALDH7A1 that we recently reported. The $R_g$ calculated from the ALDH7A1 tetramer is 36 Å, which is close to the experimental SAXS value of 37.7 Å. In contrast, the dimer has an $R_g$ of only 30 Å, while hexameric ALDHs have an $R_g$ of 43 Å. Thus, analysis of $R_g$ and crystal packing suggests that ALDH7A1 is tetrameric in solution.

Additional analysis of the SAXS data also indicates a tetramer in solution. The SAXS curve calculated from the tetramer has good agreement with the experimental curve ($\chi = 1.5$), while that calculated from a domain-swapped dimer shows large deviations ($\chi = 12.4$) (Figure 3A). Multiple Ensemble Search analysis marginally improves the fit ($\chi = 1.2$) and indicates 96% tetramer and 4% dimer. Also, the tetramer is consistent with the SAXS shape reconstruction (Figure 3B). In summary, SAXS suggests that ALDH7A1 forms the classic ALDH dimer-of-dimers tetramer in solution, which is also observed for ALDH1 and ALDH2.

**The Structure of ALDH7A1 Complexed with AA Provides Insight into Substrate Recognition.** The structure of ALDH7A1 complexed with the product AA was determined at 1.76 Å resolution (Table 1). The active site is located in the crevice between the NAD$^+$-binding and catalytic domains, below the long interdomain linker peptide (Figure 2A). Catalytic Cys302 is torsioned away from AA and not in the nucleophilic attack rotamer, which is consistent with the presence of the product AA and the absence of NAD$^+$ (Figure 4).

AA forms several interactions with the enzyme (Figure 4). One of the oxygen atoms of the $\epsilon$-carboxylate of AA occupies the oxyanion hole, forming a hydrogen bond with the backbone of catalytic Cys302. This atom represents the carbonyl O of the substrate AASA. Notably, this O atom apparently does not hydrogen bond with the conserved oxyanion hole residue Asn167, as the potential hydrogen bond distance is 3.9 Å. At the $\alpha$-carbon end of the product, the carboxylate of AA is anchored by hydrogen bonds to the main chain of Gly461-Ala462 of the interdomain linker, another hydrogen bond with Thr303, and an ion pair with Arg301. The amino group of AA forms an ion pair with Glu121. The aliphatic chain of AA packs into an aromatic box formed by Phe168, Trp175, and Phe468. We note that aromatic boxes are often found in ALDH active sites.

The product is essentially inaccessible to solvent. Calculations of the solvent accessibility with CNS show that the product exposes only 2% of its total surface area when bound to the enzyme. This result indicates that AA is buried inside the protein and implies that protein conformational changes must occur to release the product. As described below, this conformational change likely involves the C-terminus.

**The C-Terminus Contributes to the Active Site through Quaternary Structural Interactions.** The AA complex structure shows that the C-terminus plays a role in stabilizing the active site. The 13 C-terminal residues form an...
L-shaped appendage that is connected to the final $\beta$-strand of the oligomerization domain (Figure 2A). In the domain-swapped dimers that make up the tetramer, this appendage forms quaternary structural interactions that stabilize the aldehyde-binding site in the AA complex. One protomer of the dimer is colored white with bound AA colored pink. The C-terminus of the other protomer is colored green. (C) Superposition of the dimers of the ALDH7A1–AA complex (gray with a cyan C-terminus), apo ALDH7A1 in space group $C_2$ (gray with a red C-terminus), and the ALDH7A1–NAD$^+$ complex in space group $C_2$ (also gray with a red C-terminus). The arrow indicates the 16 Å movement of the C-terminus from the open conformation (AA-free) to the closed conformation (AA-bound). (D) Close-up of a superposition of the AA complex (cyan) and the apoenzyme in space group $C_2$ (gold). The arrows indicate movement of the active site from the open conformation (AA-free) to the closed conformation (AA-bound). AA is colored pink.

The Active Site Conformation Is Variable in the Absence of AA. The structures of ALDH7A1 without bound AA were determined to gain insight into the conformational flexibility of the active site. Structures of the apoenzyme and the NAD$^+$ complex were determined in the same $C_2$ lattice as the AA complex and also in space group $P4_2_2$ (Tables 1 and 2).

NAD$^+$ binds in the expected location at the C-terminus of the Rossmann fold domain. Electron density is strong for the AMP portion of the cofactor but weak and variable for the nicotinamide half of the cofactor. Accordingly, the nicotinamide riboside could not be modeled in any of the protomers of the two NAD$^+$ complex structures. We note that density for the nicotinamide of NAD$^+$ bound to maize ALDH7 is also weak ($^\text{33}$ (PDB entry 4PXN), but the entire cofactor is ordered in a structure of truncated human ALDH7A1 complexed with NADH (PDB entry 2J6L). Disorder of the nicotinamide half of the cofactor has been observed in other ALDHs.$^\text{34,35}$

The conformation of the C-terminus appears to be highly variable in the absence of AA. In the $C_2$ apo enzyme and C2 NAD$^+$ complex, the C-terminus is rotated out of the active site (Figure 5C). Compared to the AA complex, the crook of the C-terminus shifts by 16 Å, which leaves the active site open and solvent-exposed. Intermolecular interactions appear to stabilize the open conformation of the C-terminus. Ile508 and Phe510 pack into a hydrophobic pocket located between two helices of the Rossmann fold domain, while the backbone of Lys509 forms two hydrogen bonds with the side chain of Gln328 (Figure 5D).

Comparison of the structures reveals additional differences between the closed (AA-bound) and open (AA-free) active sites (Figure 5D). In the open state, Phe292, Arg301, and the aldehyde anchor loop are displaced from the active site. In the closed state, Phe292 has rotated into the space occupied by Arg301 of the open state, while Arg301 and the anchor loop converge to engage the carboxylate of AA. Hydrogen bonds to the backbone of the C-terminus also differ in the two active site
conformations. As noted above, the open conformation is stabilized by intermolecular hydrogen bonding between Lys509 and Gln328. Upon closure, these interactions are broken, and a new intramolecular hydrogen bond between the backbone of Gln506 and the side chain of Glu144 is formed (Figure 5D).

The tetragonal structures, which were determined in the absence of bound AA, suggest a more severe conformational variation of the active site. In these structures, electron density for the 12 C-terminal residues (500–511) is absent in both chains in the asymmetric unit. Similarly, electron density for the long interdomain linker (residues 459–481) is missing. The absence of these parts of the structure leaves the active site disassembled and highly solvent-exposed. The disordered sections are notable because they contain residues that bind AA, including the anchor loop, the crook of the C-terminus, and aromatic box residue Phe468.

![Image](biochemistry/p422.png)

**Figure 6.** Crystal packing of ALDH7A1 in P42_2. The lattice is viewed down the a-axis, with the b-axis horizontal and c-axis vertical. The two chains of the asymmetric unit are colored gold and blue. Neighboring chains are colored pale cyan. The C-terminus from the C2 structure (open conformation) has been grafted onto the P42_2 structure and is colored red. Note that the modeled C-terminus occupies a solvent channel.

![Image](biochemistry/p423.png)

**Figure 7.** Comparison of substrate recognition in ALDH7A1 and ALDH4A1. (A) Superposition of ALDH7A1 complexed with AA (white) and ALDH4A1 complexed with Glu (green). Black dashes indicate interactions that are unique to ALDH7A1. Magenta dashes denote those unique to ALDH4A1. (B) Substrate interaction diagrams for ALDH7A1 and ALDH4A1 inferred from the enzyme–product complex structures. Dashes denote hydrogen bonds and ion pairs. Squares denote van der Waals interactions between aromatic box residues and the aliphatic chain of the substrate. Interactions unique to either enzyme are colored red. W in a circle represents water-mediated interactions.
The two ALDH7A1 crystal lattices were analyzed to understand the influence of crystal packing on the conformation and disorder of the C-terminus. In particular, we wondered whether crystal packing in space group C2 artificially stabilizes the C-terminus. This analysis showed that the C-terminus is free of crystal contacts (5.0 Å cutoff). This result is consistent with the fact that the C2 lattice accommodates both the open and closed conformations, which differ by as much as 16 Å (Figure 5C). We also entertained the possibility that steric hindrance in space group P42_2 prevents the C-terminus from adopting the conformations observed in the C2 structure. This idea was tested by grafting the C-terminus from the C2 structure (open conformation) onto the P42_2 structure and inspecting the crystal packing interactions in the hypothetical structure. The grafted C-terminus extends into a solvent channel between two tetramers and does not clash with neighboring molecules in the lattice (Figure 6). Thus, the P42_2 lattice seems to be compatible with the open (and closed) conformation of the C-terminus. The absence of electron density implies that the C-terminus adopts multiple conformations in the P42_2 crystal. Apparently, the solvent channel is large enough to accommodate multiple, diverse conformations of the C-terminus.

**Comparison to ALDH4A1.** ALDH4A1 and ALDH7A1 recognize structurally similar substrates. ALDH4A1 is the last enzyme in the proline catabolic pathway and catalyzes the oxidation of glutamate-γ-semialdehyde (GSA) to glutamate. GSA and AASA are nearly isostructural, differing by just one extra methylene group in the aliphatic chain of AASA. Because we had previously determined the structure of ALDH4A1 complexed with the product glutamate (PDB entry 3V9K),34 we decided to compare that structure to that of the ALDH7A1–AA complex to understand how these enzymes discriminate between closely related substrates.

The active sites of ALDH7A1 and ALDH4A1 are very similar in terms of amino acid composition and structure, despite sharing only 24% global sequence identity (Figure 7). Several active site residues are identically conserved: the Asn residue of the oxynion hole, two Phe residues of the aromatic box, a Gly in the anchor loop, and a Glu near the substrate amino group. Two conservative substitutions are also evident: Arg301 and Thr303 of ALDH7A1 correspond to Lys347 and Ser349 of ALDH4A1. The active sites also have two notable sequence differences that may facilitate discrimination between AASA and GSA (Figure 7). First, Ala463 in the anchor loop of ALDH7A1 is replaced with Ser513 in ALDH4A1. Ser513 forms hydrogen bonds with the amino group and the α-carboxylate of the product in ALDH4A1. Second, the aromatic box residue Trp175 of ALDH7A1 corresponds to Phe169 of ALDH4A1. The bulkier Trp175 contacts the aliphatic chain of AA, whereas Phe169 is too small for such interaction with glutamate in ALDH4A1. Furthermore, the indole of Trp175 forms a water-mediated interaction with the aldehyde anchor loop residue Gly461, while Glu506 hydrogen bonds directly to aldehyde anchor loop residue Ala462 and indirectly to AA (Figure 5B). We note that Gly461, Ala462, Ala505, and Glu506 are identically conserved in family 7 of the ALDH superfamily, which includes enzymes from diverse organisms that are related by as little as 50% global sequence identity. The observed interactions among these residues and their strong conservation imply a functionally important role, which we suggest is aldehyde substrate recognition.

Our structures and proposed role for the C-terminus in AASA recognition are consistent with the generally accepted mechanism of ALDHs36 in which NAD⁺ binds before the aldehyde substrate. The C-terminus of the NAD⁺ complex adopts the open state conformation (C2 form) or is disordered (P42_2). Thus, the aldehyde-binding site is accessible when NAD⁺ is bound, which allows formation of the ternary enzyme–NAD⁺–AASA complex. In summary, our structures are consistent with the C-terminus being a dynamic active site element that facilitates the binding of AASA.

The discovery that the C-terminus of ALDH7A1 is part of the active site aids our understanding of the c.1512delG genetic deletion, which is implicated in PDE.14 Deletion of G1512 from exon 18 mutates six of the last seven amino acids of the wild-type polypeptide and extends the C-terminus by 10 residues (Figure 8). It is possible that the mutated C-terminus is incapable of forming the closed state that we observed in the AA complex structure. If so, the mutation may cause a defect in AASA recognition, which would impact catalytic function. In particular, c.1512delG changes the crook residues Ala505 and Glu506 to Pro and Lys, respectively. Mutation of Ala505 to Pro may alter the shape and reduce the flexibility of the C-terminus, and the longer lysine side chain at position 506 may be unable to interact favorably with AASA or the anchor loop. Also, because the C-terminus forms quaternary structural interactions, it is possible that c.1512delG affects oligomerization. The longer C-terminus could also affect protein solubility.
Site-directed mutagenesis of recombinant ALDH7A1 could be used to improve our understanding of the structural and biochemical consequences of c.1512delG.

The AA complex structure permitted an examination of how ALDH7A1 and ALDH4A1 discriminate between the structurally similar substrates AASA and GSA. These two substrates differ only in the number of methylene groups. Our analysis suggests that ALDH7A1 and ALDH4A1 accommodate the different aliphatic chain lengths by modulating the position of the distal end of the substrate, with the amino group shifted 3 Å closer to the anchor loop in ALDH4A1 (Figure 7A). Two residues appear to underlie this subtle difference in substrate pose: Ala/Ser in the aldehyde anchor loop and Trp/Phe in the aromatic box. One may be able to alter the substrate specificities of ALDH7A1 and ALDH4A1 by mutating these residues.

Finally, the tetragonal structure of ALDH7A1 provides another example of an ALDH crystal structure with a disordered active site. Electron density is lacking for the interdomain linker and C-terminus in the two tetragonal ALDH7A1 structures. The interdomain linker is also disordered in structures of human and yeast ALDH4A1 (PDB entries 4OE6 and 4OES) and ALDH1A2 (PDB entry 1B9). This disorder is significant because the interdomain linker contains the aldehyde anchor loop and one of the aromatic box residues (Figure 4). The interpretation of these disordered structures is not obvious. Perhaps they indicate that the interdomain linker is highly flexible and samples multiple conformations when the aldehyde site is vacant. This interpretation is consistent with a conformational selection mechanism of substrate recognition. With the list of ALDHs exhibiting disordered active sites growing, it may be worthwhile to consider whether this phenomenon is a global aspect of the superfamily.

ASSOCIATED CONTENT
Accession Codes
Atomic coordinates and structure factor amplitudes have been deposited in the Protein Data Bank as entries 4ZUL (AA complex), 4ZUK (C2 NAD+ complex), 4ZVW (C2 apo), 4ZVX (P42,2 apo), and 4ZVY (P42,2 NAD+ complex).

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ABBREVIATIONS
ALDH, aldehyde dehydrogenase; AASA, α-aminoadipate semialdehyde; AA, α-aminoadipate; PDE, pyridoxine-dependent epilepsy; THP, tris(3-hydroxypropyl)phosphine; SAXS, small-angle X-ray scattering; GSA, glutamate-γ-semialdehyde.

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