Crystalllographic and spectroscopic snapshots reveal a dehydrogenase in action

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Aldehydes are ubiquitous intermediates in metabolic pathways and their innate reactivity can often make them quite unstable. There are several aldehydic intermediates in the metabolic pathway for tryptophan degradation that can decay into neuroactive compounds that have been associated with numerous neurological diseases. An enzyme of this pathway, 2-aminomuconate-6-semialdehyde dehydrogenase, is responsible for ‘disarming’ the final aldehydic intermediate. Here we show the crystal structures of a bacterial analogue enzyme in five catalytically relevant forms: resting state, one binary and two ternary complexes, and a covalent, thioacyl intermediate. We also report the crystal structures of a tetrahedral, thiohemiacetal intermediate, a thioacyl intermediate and an NAD⁺-bound complex from an active site mutant. These covalent intermediates are characterized by single-crystal and solution-state electronic absorption spectroscopy. The crystal structures reveal that the substrate undergoes an E/Z isomerization at the enzyme active site before an sp³-to-sp² transition during enzyme-mediated oxidation.
The dominant route of tryptophan catabolism, the kynurenine pathway, has recently garnered increased attention given its apparent association with numerous inflammatory and neurological conditions, for example, gastrointestinal disorders, depression, Parkinson's disease, Alzheimer's disease, Huntington's disease and AIDS dementia complex. Though the precise mechanism by which the kynurenine pathway influences these diseases has not yet been fully elucidated, it has been determined that several metabolites of this pathway are neuroactive. Notably, the concentration of quinolinic acid, a non-enzymatically derived decay product of an intermediate of the kynurenine pathway used for NAD⁺ biosynthesis, is elevated over 20-fold in patients' cerebrospinal fluid with AIDS dementia complex, aseptic meningitis, opportunistic infections or neoplasms, and more than 300-fold in the brain of human immunodeficiency virus-infected patients. This NAD⁺ precursor has also been shown to be an agonist of N-methyl-D-aspartate receptors, and an increase of its concentration may lead to over-excitation and death of neuronal cells.

The apparent medical potential of the kynurenine pathway warrants detailed study and characterization of its component enzymes and their regulation. One enzyme in particular, 2-aminomuconate-6-semialdehyde dehydrogenase (AMSDH), is responsible for oxidizing the unstable metabolic intermediate 2-aminomuconate-6-semialdehyde (2-AMS) to 2-aminomuconate (2-AM) (Fig. 1a). On the basis of sequence alignment, AMSDH is a member of the hydroxymuconic-semialdehyde dehydrogenase (HMSDH) family under the aldehyde dehydrogenase (ALDH) superfamily. ALDHs are prevalent in both prokaryotic and eukaryotic organisms and are responsible for oxidizing aldehydes to their corresponding carboxylic acids. They use NAD⁺ as a hydride acceptor to harvest energy from their primary substrate and generate NADH, which provides the major reducing power to maintain cellular redox balance. In addition to being commonly occurring metabolic intermediates, aldehydes are reactive electrophiles, making many of them toxic. Enzymes of the ALDH superfamily are typically promiscuous with regards to their substrates; however, in recent years, this superfamily has had several new members identified with greater substrate fidelity, especially when the substrate is identified as a semialdehyde.

The putative native substrate of AMSDH, 2-AMS, is a proposed metabolic intermediate in both the 2-nitrobenzoic acid degradation pathway of Pseudomonas fluorescens KL-7 (ref. 15) and the kynurenine pathway for l-tryptophan catabolism in mammal. The presence of NAD⁺ and AMSDH, 2-AMS is oxidized to 2-AM (Fig. 1a); however, it can also spontaneously decay to picolinic acid and water with a half-life of 35 s at neutral pH. Due to its instability, 2-AMS has not yet been isolated, leaving its identity as the substrate of AMSDH an inference based on decay products and further metabolic reactions. There are several reasons for the poor understanding of this pathway: it is complex with many branches, some of the intermediates are unstable and difficult to characterize, and several enzymes of the pathway, including AMSDH, are not well understood. Hence, the structure of AMSDH will help to address questions such as what contributes to substrate specificity for the semialdehyde dehydrogenase and how 2-AMS is bound and activated during catalysis.

In the present study, we have cloned AMSDH from Pseudomonas fluorescens, generated an E. coli overexpression system and purified the target protein for molecular study. We also constructed several mutant expression systems to characterize the role of specific active site residues. Enzymatic assays were performed for all forms of the enzyme, and crystal structures were solved for the wild type and one mutant. We were able to capture several catalytic intermediates in crystallo by soaking protein crystals in mother liquor containing either the primary organic substrate or a substrate analogue and discovered that in addition to dehydrogenation, the substrate undergoes isomerization at the active site.

### Results

#### Catalytic activity of wild-type AMSDH.

Due to the unstable nature of its substrate, 2-AMS, the activity of AMSDH was detected using a coupled-enzyme assay that employed its upstream partner, α-amino β-carboxymuconate ε-semialdehyde decarboxylase (ACMSD), to generate 2-AM in situ. ACMSD transforms α-amino β-carboxymuconate ε-semialdehyde (ACMS) (λₘₓ at 360 nm) to 2-AMS (λₘₓ at 380 nm). As seen in Fig. 1b, in an assay that uses only ACMSD, the absorbance peak of its substrate, ACMS, red-shifts to 380 nm as 2-AMS is formed. The absorbance at 380 nm then quickly decreases as 2-AMS decays to picolinic acid, a compound with no absorbance features above 200 nm. In a coupled-enzyme assay, ACMSD, AMSDH and NAD⁺ are included in the reaction system. As shown in Fig. 1c, ACMS is still consumed; however, there is no red shift observed because 2-AMS is enzymatically converted to 2-AM (λₘₓ at 325 nm) rather than accumulating and decaying to picolinic acid. The production of 2-AM requires that an equimolar amount of NAD⁺ be reduced to NADH (λₘₓ at 339 nm). A stable alternative substrate, 2-hydroxymuconate-6-semialdehyde (2-HMS), was used to pursue kinetic parameters (Fig. 1d), when using saturating NAD⁺ concentrations (≥1 mM), the kₐcat and Kₘ of AMSDH for 2-HMS were 1.30 ± 0.01 s⁻¹ and 10.4 ± 0.2 μM, respectively (Fig. 1e).

#### Structural snapshots of the dehydrogenase catalytic cycle.

We solved five crystal structures of wild-type AMSDH, including the ligand-free (2.20 Å resolution), NAD⁺-bound binary complex (2.00 Å), ternary complex with NAD⁺ and substrate 2-AMS (2.00 Å) or 2-HMS (2.20 Å) and a thioacyl intermediate (1.95 Å). All five structures belong to space group P2₁2₁2₁. Data collection and refinement statistics are listed in Supplementary Table 1. The complete AMSDH model includes four polypeptides per asymmetric unit describing one homotetramer (Supplementary Fig. 1a). Each monomer of AMSDH contains three domains: a subunit interaction domain, a catalytic domain and an NAD⁺ binding domain (Supplementary Fig. 1b). For details of the secondary structure, see Supplementary Discussion.

In the structure of the co-crystallized binary complex, an NAD⁺ molecule is present in an extended, anti-conformation in the amino-terminal, co-substrate-binding domain of each monomer (Fig. 2a). The electron density map of NAD⁺ is well defined, and the interactions between the protein and NAD⁺ are equivalent in all four subunits as shown in Fig. 2e. The NAD⁺-bound AMSDH structure is similar to the ligand-free structure with an aligned r.m.s.d. of 0.239 Å. Residues that belong to the NAD⁺-binding pocket are also well aligned with the exception of Cys302, Arg108 and Leu116 (Supplementary Fig. 2). On binding NAD⁺, the thiol moiety of Cys302 rotates so that the sulfur is 2.3 Å closer to the substrate-binding pocket and away from the nicotinamide head of NAD⁺.

#### Crystal structures of enzyme–substrate ternary complexes.

Structures of AMSDH in ternary complex with co-substrate NAD⁺ and its primary substrates were obtained by soaking co-crystallized AMSDH-NAD⁺ crystals with 2-AMS and 2-HMS, respectively. Extra density that fits with the corresponding substrate molecule was observed in the active site of each subunit. The co-substrate NAD⁺ in the ternary complex structures is...
Arg464 or by a failure of the enzyme to properly bind and stabilize the imine form of the substrate.

**Catalytic intermediates trapped after ternary complex formation.**

Enzyme–NAD$_{+}$ binary complex crystals were soaked in mother liquor containing 2-HMS for a range of time points from 5 min to more than 3 h before flash cooling in liquid nitrogen. In a crystal that was soaked for 40 min, an intermediate was trapped and refined to a resolution of 1.95 Å (Fig. 2d). Crystals soaked for longer time points gave a similar intermediate with poorer resolution. In this structure, 2-HMS is observed in the 2-Z,4-E isomer rather than the 2-E,4-E isomer as seen in the substrate-bound ternary structure. Also, the substrate interacts with Arg120 and Arg464 with both of its carboxyl oxygens rather than one carboxyl oxygen and the 2-hydroxy oxygen as shown in the 2-HMS ternary complex structure. Additionally, the distance between its sixth carbon and Cys302’s sulfur is now at 1.8 Å, which is within covalent bond distance for a carbon–sulfur bond. Also, the continuous electron density extends, and the distance between its sixth carbon and Cys302’s sulfur is now at 1.8 Å, which is within covalent bond distance for a carbon–sulfur bond. Also, the continuous electron density

**Figure 1 | Activity of AMSDH.** (a) Reaction scheme showing the enzymatic generation of 2-AMS, the reaction catalysed by AMSDH, and the competing non-enzymatic decay of 2-AMS to picolinic acid. (b) Representative assay showing the ACMSD (1 μM)-catalysed conversion of ACM (λ$_{\text{max}}$ 360 nm) to 2-AMS (λ$_{\text{max}}$ 380 nm), which decays to picolinic acid (transparent). (c) Coupled-enzyme assay in which AMSDH (200 nM) oxidizes 2-AMS, produced in situ as shown in b in 50 s, to 2-AM (λ$_{\text{max}}$ 325 nm). (d) Reaction scheme showing 2-HMS oxidation by AMSDH. (e) Representative assay showing the activity of AMSDH (200 nM) on 2-HMS (λ$_{\text{max}}$ 375 nm) in 50 s. The inset is a Michaelis–Menten plot.
The thioacyl intermediate in AMSDH consists of the reaction intermediate trapped by soaking crystals with 2-HMS for 10 min before flash cooling. (b) the ternary complex of AMSDH-NAD⁺ crystals soaked with 2-AMS for 5 min before flash cooling, (c) the ternary complex of AMSDH-NAD⁺ soaked with 2-HMS for 10 min before flash cooling, (d) the trapped thioacyl, NADH-bound intermediate obtained by soaking AMSDH-NAD⁺ crystals with 2-HMS for 40 min before flash cooling, (e) Two-dimensional interaction diagram for NAD⁺ binding. (f) Close-up of the thioacyl intermediate in d. (g) Single-crystal electronic absorption spectrum of d. Protein backbone and residues are shown as light blue cartoons and sticks, respectively. The substrates and intermediate are shown as yellow sticks, and NAD⁺ and NADH are shown as green sticks. The omit map for ligands is contoured to 2.0σ and shown as a grey mesh.

Figure 2 | Crystal structures of wild-type AMSDH and single-crystal electronic absorption spectrum of a catalytic intermediate. AMSDH was co-crystallized with NAD⁺ to give AMSDH-NAD⁺ binary complex crystals that were used for soaking experiments. (a) Active site structure of the binary AMSDH-NAD⁺ complex, (b) the ternary complex of AMSDH-NAD⁺ crystals soaked with 2-AMS for 5 min before flash cooling, (c) the ternary complex of AMSDH-NAD⁺ soaked with 2-HMS for 10 min before flash cooling, (d) the trapped thioacyl, NADH-bound intermediate obtained by soaking AMSDH-NAD⁺ crystals with 2-HMS for 40 min before flash cooling, (e) Two-dimensional interaction diagram for NAD⁺ binding. (f) Close-up of the thioacyl intermediate in d. (g) Single-crystal electronic absorption spectrum of d. Protein backbone and residues are shown as light blue cartoons and sticks, respectively. The substrates and intermediate are shown as yellow sticks, and NAD⁺ and NADH are shown as green sticks. The omit map for ligands is contoured to 2.0σ and shown as a grey mesh.

between Cys302-SG and 2-HMS-C6 indicates the presence of a covalent bond (Fig. 2f). Another feature of this intermediate is that the nicotinamide ring of NAD⁺ has moved 4.6 Å away from the active site and adopted a bent conformation (Fig. 2d) compared with the position in the binary or ternary complex structures (Fig. 2a–c). The structural changes of NAD⁺ associated with reduction has been observed and well documented. In the oxidized state, NAD(P)⁺ lies in the Rossman fold in an extended conformation, allowing for hydride transfer from the substrate to its nicotinamide carbon during the first half of the reaction. Reduced NAD(P)H then adopts a bent conformation in which the nicotinamide head moves back towards the protein surface. This movement provides more space in the active site for the second half of the reaction, acyl-enzyme adduct hydrolysis, to take place. Thus, the coenzyme in this intermediate structure is likely to have been reduced to NADH and, as such, the structure is assigned as a thioacyl-enzyme–substrate adduct. The single-crystal electronic absorption spectrum of the sample has an absorbance maximum at 394 nm (Fig. 2g). The same absorbance band was observed in crystals soaked with 2-HMS from 30 min to 2 h (Supplementary Fig. 5a). However, this long-lived intermediate in the crystal was not observed in solution with millisecond-to-second time resolution in stopped-flow experiments (Supplementary Fig. 6a). Thus, it is either present in an earlier time domain (sub-milliseconds), or alternatively, it may not accumulate in solution because NADH can readily dissociate in solution, whereas it may be trapped in the active site when in the crystalline state.

Another notable change in the intermediate structure is the movement of the side chain of Glu268, which rotates 73° towards the active site (Fig. 2c,d). To probe the function of Glu268, we constructed an alanine mutant and found that it exhibited no detectable activity in steady-state kinetic assays. Interestingly, E268A exhibits completely different pre-steady state activity than the wild-type enzyme. As shown in Supplementary Fig. 6b, an absorbance band at 422 nm was formed concomitant with the decay of the 2-HMS peak within 1 s of the reaction. This new species is generated stoichiometrically on titration of 2-HMS with E268A (Fig. 3d). The moiety that gives rise to this new absorbance band is stable for minutes at room temperature and can be separated from the protein by membrane filtration-based methods, suggesting that it is covalently bound to the protein. The formation of an enzyme–substrate adduct in the E268A mutant was investigated by mass spectrometry. For the isolated E268A, the resultant multiply charged states (Supplementary Fig. 7) were deconvoluted to obtain a molecular weight (MW) of 56,252 Da (Fig. 4a). This value is in good agreement with the predicted MW of E268A AMSDH plus an amino-terminal His-tag and linking residues, 56,251 Da. The
second largest peak in the deconvoluted spectrum has a MW 177 Da greater than that of the most abundant signal. This is likely due to post-translational modification of the His-tag: α-N-Gluconoylation of His-tags has been observed in E. coli-expressed proteins, which cause 178 Da extra mass. The mutant protein light blue cartoons and sticks, respectively. The substrate and intermediate are shown as yellow sticks, and NAD (top panel) and single-crystal electronic absorption spectrum of the intermediate in (bottom panel). Protein backbone and residues are shown as light blue cartoons and sticks, respectively. The substrate and intermediate are shown as yellow sticks, and NAD$^+$ and NADH are shown as green sticks. The omit map for ligands is contoured to 2.0 $\sigma$ and shown as a grey mesh.

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**Figure 3 | Crystal structures of the E268A mutant and its solution and single-crystal electronic absorption spectra.**

(a) Structure of the active site of the co-crystallized E268A-NAD$^+$ binary complex, (b) a thiohemiacetal intermediate obtained by soaking the E268A-NAD$^+$ crystals with 2-HMS for 30 min before flash cooling and (c) a thioacyl intermediate obtained by soaking the E268A-NAD$^+$ crystals with 2-HMS for 180 min before flash cooling. (d) Solution electronic absorption spectra of a titration of 2-HMS with E268A. (e) Single-crystal electronic absorption spectrum of the intermediate in (top panel) and single-crystal electronic absorption spectrum of the intermediate in (bottom panel). Protein backbone and residues are shown as light blue cartoons and sticks, respectively. The substrate and intermediate are shown as yellow sticks, and NAD$^+$ and NADH are shown as green sticks. The omit map for ligands is contoured to 2.0 $\sigma$ and shown as a grey mesh.

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**Figure 4 | Deconvoluted positive-mode electrospray ionization mass spectra of as-isolated E268A (a) and 2-HMS treated-E268A (b).** The two major components are labelled with their respective molecular weights.

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overall structure aligns very well with the wild-type binary complex structure with an r.m.s.d. of 0.139 Å. The active site of E268A also resembles the native AMSDH structure (Supplementary Fig. 8). The nature of the absorbing species at 422 nm was further investigated by soaking co-crystallized E268A-NAD$^+$ crystals in mother liquor containing 2-HMS. By doing so, two temporally, structurally and spectroscopically distinct intermediates were identified.

When E268A-NAD$^+$ crystals are soaked with 2-HMS for 40 min or less, their single-crystal electronic absorption spectra show an absorbance maximum at 422 nm (Supplementary Fig. 5b), as was observed in the solution-state titration and the stopped-flow assays. An individual electronic absorption spectrum for an E268A-NAD$^+$ crystal soaked with 2-HMS for 15 min can be found in Fig. 3e (top). The structure of E268A-NAD$^+$ soaked with 2-HMS for 30 min before flash cooling was solved and refined to 2.15 Å resolution (Fig. 3b). In this structure, a continuous electron density between Cys302-SG and 2-HMS-C6 is observed, similar to the thioacyl intermediate observed in the wild-type enzyme. However, in contrast to the thioacyl intermediate, the density around C6 is less flat, indicating an sp$^3$-rather than sp$^2$-hybridized carbon (Fig. 5a). The angle between the plane of the carbon backbone of the substrate and the newly aldehydic oxygen is 55 ± 9°, compared with the angle of the wild-type thioacyl intermediate at 26 ± 4° (Supplementary Table 2). More importantly, the C6 of 2-HMS and the C4N of NAD$^+$ are very close (2.4–2.8 Å), making it unlikely that the hydride has been transferred from the substrate. Taken together, these data allow us to assign this intermediate to a thiohemiacetal enzyme adduct (Fig. 3b). A similar intermediate has only been trapped once previously
in a crystal that contains no co-substrate. Hence, this is the first time for this intermediate to be trapped in the presence of NAD$^+$. If the E268A-NAD$^+$ crystals are soaked with 2-HMS for longer than 1 h, their single-crystal electronic absorption spectra begin to resemble that of the wild-type, thioacyl intermediate with a corresponding absorbance maximum at 394 nm (Supplementary Fig. 5b), as seen in wild-type, thioacyl intermediate crystals. An individual electronic absorption spectrum for an E268A-NAD$^+$ crystal soaked with 2-HMS for 120 min can be found in Fig. 3e (bottom). The structure of an E268A-NAD$^+$ crystal soaked with 2-HMS for 180 min was solved and refined to 2.20 Å (Fig. 3c). The structure of this intermediate is also similar to the wild-type, thioacyl-enzyme adduct with NADH, rather than NAD$^+$ found at the active site. The distance between the C4N of NADH and C6 of 2-HMS is longer than 6.1 Å (Fig. 3c). The electron density around C6 is flatter (Fig. 5b) compared with the thiohemiacetal intermediate and similar to the thioacyl intermediate trapped in the wild-type AMSDH structure (Fig. 2f), and the angle between the plane of the carbon backbone of the substrate and the carbonyl oxygen is 20 ± 5°/C176, which is statistically indistinguishable from that of the wild-type, thioacyl intermediate, 26 ± 4° (Supplementary Table 2). On the basis of the similarities in their absorbance and structures, we conclude that this latter intermediate is equivalent to the wild-type, thioacyl intermediate. It is also worth noting that the strictly conserved asparagine 169 (Fig. 5) is seen to stabilize both the thiohemiacetal and thioacyl intermediates through hydrogen-bonding interactions.

Investigation of isomerization by computational modelling. The isomerization of 2-AMS from the $2E$ to $2Z$ isomer implied by the solved crystal structures was probed with density functional theory calculations. The free energy profiles obtained were used to help illuminate the nature of 2-AMS and gain insight into how the active site of AMSDH may facilitate the isomerization. The total energies of different isomers and rotamers of 2-AMS in its enamine/alddehyde and imine/enol tautomers and the rotational barriers about their respective 2–3 bond were compared. For the imine/enol tautomer, additional computations were performed with the side groups from Arg120 and Arg464 to investigate what effect, if any, they will have on the free energy profile for rotation about the 2–3 bond of 2-AMS.

First, 2-AMS was constructed and optimized in its $2E$ isomer with a negatively charged 2-carboxylate group (Fig. 6a). To estimate the energy barrier for an uncatalysed
isomerization from the 2E to the 2Z isomer, the 2–3 double bond was then restrained at 10° intervals from 180 to 0°, and the structure was optimized at each point. On the basis of the free energy profile (Fig. 6a), the uncatalysed isomerization barrier is 31.9 kcal mol$^{-1}$. The profile also shows that the 2Z isomer, as might be expected, is lower in energy than the 2E isomer by 4.2 kcal mol$^{-1}$. Next, the rotational barrier about the 2–3 bond of 2-AMS when in its 2-imine, 6-enol tautomer, as is suggested by the ternary complex structure, was calculated in the same manner. The barrier was found to be 9.2 kcal mol$^{-1}$, and opposite to the enamine tautomer, the ‘2Z-like’ rotamer is higher in energy than the ‘2E-like’ rotamer by 1.7 kcal mol$^{-1}$ (Fig. 6b). Unsurprisingly, the rotational barrier about the 2–3 bond is much lower in the imine tautomer; however, the ‘2Z-like’ rotamer of the imine tautomer is 21.8 kcal mol$^{-1}$ higher in energy than the 2Z isomer of the enamine tautomer.

Possible influences of the two active site arginines on the free energy profile for rotation were also considered. To mimic the conditions of the enzyme active site, similar calculations as those above were performed, which included the guanidinium heads of Arg120 and Arg464. The starting model was built using the active site geometry of the ternary complex crystal structure (PDB entry: 4I25), and on inspection, it is immediately apparent that with two arginines in such close proximity to the substrate, there is insufficient space for two hydrogen atoms on the nitrogen at the 2-position of 2-AMS, and attempts to optimize an enamine tautomer with the hydrogen-bonding pattern of the ternary complex produced structures within which the entire 2-AMS molecule rotates so that only the carboxylate group interacts with the guanidinium moieties. The absolute positions of the guanidinium groups were fixed and the structure of 2-AMS in the imine tautomer was optimized. The dihedral angle of the 2–3 bond of 2-AMS was then increased in 45° increments and the structure optimized while restraining the position of the guanidinium groups and the 2–3 bond to build a rough free energy profile to estimate the rotational barrier. In the presence of the active site arginines, the barrier about the 2–3 bond of 2-AMS is further reduced to 8.5 kcal mol$^{-1}$ (Supplementary Table 3). Another interesting finding is that in the presence of the guanidinium groups, the ‘2E-like’ and ‘2Z-like’ rotamers of 2-AMS are nearly isoenergetic, with a free energy difference of 0.2 kcal mol$^{-1}$ (Supplementary Table 3).

Discussion
The substrate of AMSDH, 2-AMS, contains an unstable aldehyde in conjugation with an enamine and can decay to picolinic acid and water, presumably through an electrocyclization reaction similar to its metabolic precursor, ACMS$^{23}$. To assay the enzymatic activity, the upstream enzyme was utilized in the reaction mixture to generate substrate, and it was shown that AMSDH is catalytically active. Unfortunately, no kinetic parameters can be reliably determined because the concentration of 2-AMS is not well defined in the coupled-enzyme assay. To circumvent this issue, a previously-identified, stable alternative substrate, 2-HMS$^{24,25}$, in which a hydroxyl group replaces the amino group in 2-AMS to prevent cyclization, was used to characterize the activity of AMSDH and to examine the activity of the mutants.

Substrate-bound, ternary complex structures were obtained by soaking co-crystallized protein and NAD$^+$ with 2-AMS or 2-HMS. 2-AMS is an unstable compound which decays with a $t_{1/2}$ of about 9s at pH 7.5 and 37°C or 35s at pH 7.0 and 20°C. Notably, this is its first reported structure. It appears to be stabilized in the enzyme active site in its imine tautomer by forming two sets of bifurcated hydrogen bonds with Arg120 and Arg464 so that the electrocyclization reaction cannot occur. Both arginine residues are close to the protein surface and in good positions to serve as gatekeepers, bringing the substrate into the active site. As a residue residing on a loop, Arg464 should be relatively flexible. The electron density for the side chain of Arg120 is partially missing in the binary complex structure but very well resolved in both ternary complex structures. This observation indicates that the presence of substrate can stabilize what may be a flexible residue. It becomes evident from the coordinates that Arg120 and Arg464 play an important role in substrate recognition, stabilization and possibly product release.

Two arginine residues are rarely observed in such close proximity, stabilizing one end of the same molecule with multiple hydrogen bonds. With the exception of the hydrogen bonds provided by Arg120 and 464, the substrate-binding pocket is mostly composed of hydrophobic residues. On the basis of sequence alignment (Supplementary Fig. 9), these two arginine residues are strictly conserved throughout the HMSDH family but are not found in other members of the ALDH superfamily. We propose that these dual arginines combined with the size restrictions provided by the hydrophobic pocket endow this enzyme with its specificity towards small $\alpha$-substituted carboxylic acids with an aldehyde moiety, such as 2-AMS and 2-HMS. Furthermore, our computational work suggests that these arginines are crucial for stabilizing the imine tautomer of 2-AMS to allow for rotation about its 2–3 bond.

Two strictly conserved catalytic residues, Cys302 and Glu268, are present at the interior of the substrate-binding pocket. General features regarding these residues in the ALDH superfamily are (1) that the cysteine serves as a catalytic nucleophile, which is anticipated to form a covalent-adduct intermediate with the substrate by a nucleophilic addition during catalysis$^{26–28}$ and (2) that the glutamate serves as a base to activate water for hydrolysis of the thioacyl-enzyme adduct$^{29–32}$. Previous studies indicate that the catalytic cysteine can adopt two conformations, resting and attacking$^{19}$. In the ligand-free structure, Cys302 is far from where the carbonyl carbon of the substrate should be and is in the resting state. In the ternary complex structures, Cys302 is located at an ideal position to initiate catalysis, which is the attacking state. It is proposed to attack the aldehydic carbon (C6) of the substrate. In the two ternary complex structures, the distance between the sulfur of Cys302 and the C6 of the substrate is $\sim3.3\,\AA$. Cys302 and the aldehydic carbon form a covalent bond in both thioacyl and thiohemiacetal intermediates. Mutation of Cys302 to serine led to enzyme with no detectable dehydrogenase activity, further confirming its catalytic significance.

Examining the wild-type AMSDH structures shows that in the NAD$^+$-bound binary complex, Glu268 adopts a ‘passive’ conformation, pointing away from the substrate-binding pocket, and forms hydrogen bonds with both NE of Trp177 (3.2 Å distance) and the backbone oxygen of Phe470 (3.6 Å) to leave space for the reduction of NAD$^+$. Its electron density is very well resolved and the side chain B-factor is close to average: 28.2 Å$^2$/28.5 Å$^2$. The thiol moiety of Cys302 is 7.14 Å from Glu268 and is unlikely to form interactions. Interestingly, in both substrate-bound structures, Glu268 becomes more flexible and exhibits much weaker electron density and increased side chain B-factors compared with average protein B-factors: 37.8 Å$^2$/28.5 Å$^2$ and 66.37 Å$^2$/39.7 Å$^2$. In the thioacyl intermediate structure, the electron density of Glu268 becomes very well defined again, but its side chain rotates 73° towards the bound substrate and seems to be in an ‘active’ position to abstract a proton from a deacylating water (Fig. 2d). At this point in the reaction cycle, the NADH molecule needs to leave the active site to make room for the catalytic water molecule. Movement of the nicotinamide ring of
NAD\(^{+}\) coupled with the rotation of an active site glutamate has previously been observed in other ALDHs during catalysis\(^{30-32}\).

Mutation of Glu268 to alanine led to the accumulation of the thiohemiacetal intermediate in both solution and crystalline states. The strictly conserved glutamate residue in the active site of ALDH enzymes has been proposed to play up to three possible roles during catalysis. It is strictly required to activate the deacylating water that allows for product release, it is in a ‘passive’ conformation during NAD(P)\(^{+}\) reduction, and in some cases, it may serve to activate cysteine for nucleophilic attack\(^{33}\). On the basis of these roles, mutation to alanine would be expected to decrease the rate of hydrolysis of the thioacyl adduct, have no effect on the rate of reduction of NADH and possibly decrease the rate of nucleophilic attack by cysteine. With this understanding, deletion of the active site glutamate should cause an accumulation of the thioacyl intermediate. However, in this work, the E268A mutant is shown to accumulate the preceding thiohemiacetal intermediate both in crystal and in solution. This finding suggests an additional catalytic role for this residue: rotation of Glu268 towards the active site facilitates the hydride transfer from the tetrahedral thiohemiacetal adduct to NAD\(^{+}\). The rapid formation of the intermediate in solution indicates that Glu268 of AMSDH does not play a role in activating cysteine. However, it does appear necessary to complete hydride transfer from the substrate to NAD\(^{+}\), and its removal turns the native, primary substrate into a suicide inhibitor.

On the basis of previous studies of the ALDH mechanism, the eight high-resolution crystal structures solved (Supplementary Table 1) as well as our biochemical and computational studies, we propose a catalytic mechanism for AMSDH. As shown in Fig. 7, NAD\(^{+}\) binds to the enzyme, 1, to form an NAD\(^{+}\)-bound AMSDH complex, 2. The substrate, 2-AMS, is then recognized by Arg120 and Arg464 through multiple hydrogen-bonding interactions, and its imine tautomer is stabilized in the active site, 3. At this point, the order of the rotation, tautomerization and nucleophilic attack by C302 on the aldehydic carbon to produce the tetrahedral, thiohemiacetal intermediate, 4, is not yet clear. The isomerization and nucleophilic attack drive a translation of the substrate away from Arg120 and Arg464 so that they are only able to interact with the carboxylate group of the substrate. Next, NAD\(^{+}\) is reduced to NADH by abstraction of a hydride from 4, forming a thioacyl intermediate, 5, a process which involves an sp\(^{3}\)-to-sp\(^{2}\) transition during oxidation of the organic substrate by NAD\(^{+}\). On reduction, the nicotinamide portion of NADH moves away from the substrate as Glu268 rotates into position to activate a water molecule to perform a nucleophilic attack on the same carbon that was previously attacked by Cys302, forming a second tetrahedral intermediate, 6. Finally, the second tetrahedral intermediate collapses, breaking the C–S bond and releasing the final products, 2-AM and NADH. Species 1–5 are spectroscopically and structurally characterized, while intermediate 6 was not seen to accumulate.

In this work, five catalytically relevant structures of the wild-type AMSDH and three mutant structures yield a comprehensive understanding of the protein’s overall structure, co-substrate-binding mode and elucidate the primary residues responsible for substrate specificity among the HMSDH family of the ALDH superfamily. The structural and spectroscopic snapshots capture the crystal structure of an unstable kynurenine metabolite, 2-AMS, and two catalytic intermediates, including stabilizing a tetrahedral intermediate in a mutant protein, which was further verified by mass spectrometry. Capture of a thiohemiacetal intermediate upon deletion of E268 also points to a new role for this well-established active site base in hydride transfer from the substrate to NAD\(^{+}\). Another interesting finding revealed through solving the ternary complex and intermediate crystal structures and supported by computational studies is that an E to Z isomerization of the substrate occurs in this dehydrogenase before hydride transfer. To the best of our knowledge, this is the first piece of structural evidence illustrating an ALDH that proceeds via an E/Z isomerization of its substrate during catalysis.

**Figure 7 | Proposed catalytic mechanism for the oxidation of 2-AMS by AMSDH.** The primary substrate (2E, 4E)-2-aminomuconate-semialdehyde binds to the enzyme in its imine tautomer to form the ternary complex (3). An isomerization and attack by cysteine on the aldehydic carbon form the (2Z, 4E)-2-aminomuconate-thiohemiacetal adduct (4). AMSDH-mediated oxidation of 4 concomitant with reduction of NAD\(^{+}\) to NADH follows, generating a thioacyl-enzyme intermediate (5). Both 4 and 5 are the catalytic intermediates covalently attached to the enzyme. Hydrolysis of 5 then allows the release of the products 2-AM and NADH, restoring the ligand-free enzyme for the next catalytic cycle.
Methods

General methods. The cloning, expression, purification and site-directed mutagenesis of AMSDH are described in the Supplementary Methods.

Preparation of ACMs and 2-HAMS. ACMs was generated by catalysing the insertion of molecular oxygen to 3-hydroxyanthranilic acid by purified, Fe²⁺-reconstituted 3-hydroxyanthranilate 3,4-dioxigenase as described previously. 2-HAMS is generated non-enzymatically from ACMs following a previously established method. The pH of solutions containing ACMs was adjusted to ~2 by the addition of dehydrogenase acid. 2-HAMS formation was monitored at an Agilent 8453 diode-array spectrophotometer at 315 nm. The solutions were then neutralized with sodium hydroxide once the absorbance at 315 nm stopped increasing. 2-HAMS at neutral pH has a maximum absorbance at 375 nm (ref. 24).

Enzyme activity assay using 2-HMS as substrate. Steady-state kinetics analyses were carried out at room temperature on an Agilent 8453 diode-array spectrophotometer. Reaction buffer contains 25 mM HEPES and 1 mM NAD⁺, pH 7.5. Consumption of 2-HMS by 20 mM AMSDH was detected by monitoring the decrease of its absorbance at 375 nm with a molar extinction coefficient of 43,000 M⁻¹ cm⁻¹ (ref. 24) for 15 s with a 0.3 s integration time. For mutants, 700 nM protein and a wavelength of 420 nm, ε₂₅₀, 11,180 M⁻¹ cm⁻¹, was used. Absorbance at 375 nm decreased and blue shifted to 295 nm, the maximum ultraviolet absorbance for the product, 2-hydroxyxanadonic acid. This is consistent with previous reports in which the ending compound was purified and verified as the correct product. The pre-steady state spectra were obtained with an Applied Photophysics Stopped-Flow Spectrometer SX20 (UK) with the mixing unit hosted inside an anaerobic chamber made by Coy Laboratory Products (MI, USA). Pre-steady state activity used the same reaction buffer but with 23 mM AMSDH or E268A and 25 μM 2-HMS and were carried out at 10°C. The change in absorbance was monitored for 1 s.

X-ray crystallographic data collection and refinement. Purified AMSDH samples at a final concentration of 10 mg ml⁻¹ containing no NAD⁺ or 10 equiv. of NAD⁺ were used to set up sitting-drop vapour diffusion crystal screening trays in Art Robbins 96-well Inteli-Plates using an ARI Gryphon crystalization robot. The initial crystallization conditions were obtained from PEG-1/2 (Hampton Research) screening kits at room temperature. The screened conditions were optimized by increasing protein concentration to 40 mg ml⁻¹ and lowering crystalization temperature to 18°C. NAD⁺-bound AMSDH crystals were obtained from drops assembled with 1.5 μl of protein (preincubated for 10 min with 10 equiv. of NAD⁺) and 1.5 μl of reservoir solution containing 20% polyethylene glycol 3350 and 0.2 M sodium phosphate dibasic monohydrate, pH 9.1, by hanging drop diffusion in VDX plates (Hampton Research). Pyramid shaped crystals that diffract up to ~1 Å appeared overnight. The reservoir solution for crystallizing the cofactor-free AMSDH crystals contains 12% polyethylene glycol 3350, 0.1 M sodium formate, pH 7.0. Crystals belonging to the same space group formed within 2–3 days as complete when the last droplet solution containing 25 μM 2-HMS and incubated for 10–180 min before crystallization. Crystals were transferred to mother liquor solution containing 1 mM 2-HMS and incubated for 10–180 min before flash cooling in liquid nitrogen. Soaking 2-AMS as a substrate is more complicated because of its instability. Crystalization solution containing ~1.5 mM AMSDH was used for soaking. After transferring several crystals to the soaking solution (8 μl), 2 μl of 1 mM purified ACMS was added to calculate the conversion of ACMS to 2-AMS. Crystals were flash frozen after a 5 min-incubation. Crystallization solution containing 20% glycerol or ethylene glycol was used as cryoprotectant. X-ray diffraction data were collected on a MAR345 image plate or on a Rigaku R-AXIS image plate and processed with the HKL crystallographic suite of programs. The chemical structures were optimized using the ternary complex crystal structure (PDB entry: 4GDG) as a starting model. To calculate the isomerization barrier, the dihedral angle about the 2-3 bond was restrained and the rest of the molecule was optimized. For the calculations that included the guanidinium heads of Arg120 and Arg464, the geometry was obtained from the crystal structure, and their positions were fixed while the substrate was optimized.

Mass spectrometry. To prepare samples for ESI mass spectrometry, as-isolated E268A AMSDH was buffer-exchanged to 10 mM Tris (pH 8.0) by running through a desalting column (GE Healthcare). Intermediate bound E268A was obtained by mixing E268A with 3 equiv. of 2-HMS. Excess 2-HMS was removed by desalting chromatography using the same buffer. Desalted proteins were concentrated to a final concentration of 20 μM. Freshly prepared samples were rinsed with acetonitrile and 0.1% formic acid (1:1 ratio) before injection. Mass spectrometry experiments were conducted using a Waters (Milford, MA) Micromass Q-TOF micro (ESI-Q-TOF) instrument operating in positive mode. The capillary voltage was set to 3,500 V, the sample cone voltage to 35 V and the extraction cone voltage to 2 V. The source block temperature and the desolvation temperature were set to 100 and 120°C, respectively. The samples were introduced into the ion source by direct injection at a flow rate of 5 μl min⁻¹. The raw data containing multiple positively charged protein peaks were deconvoluted and smoothed using Masslynx 4.1.

Computational studies. All ground-state density functional theory calculations were performed with Gaussian 03 Revision-E01 at the B3LYP/6-31G* level of theory. The chemical structures were optimized using the ternary complex crystal structure (PDB entry: 4GDG) as a starting model. To calculate the isomerization barrier, the dihedral angle about the 2-3 bond was restrained and the rest of the molecule was optimized. For the calculations that included the guanidinium heads of Arg120 and Arg464, the geometry was obtained from the crystal structure, and their positions were fixed while the substrate was optimized.

References
1. Kezhetly, D., Troost, F. J. & Mascele, A. A. Understanding the role of tryptophan and serotonin metabolism in gastrointestinal function. *Neurogastroenterol. Motil.* 21, 1239–1249 (2009).
2. Myint, A. M. et al. Kynurenine pathway in major depression: evidence of impaired neuromodulation. *J. Affect. Disord.* 98, 143–151 (2007).
3. Ogawa, T. et al. Kynurenine pathway abnormalities in Parkinson’s disease. *Neurology* 42, 1702–1706 (1992).
4. Guillenin, G. I. et al. Quinolinic acid in the pathogenesis of Alzheimer’s disease. *Adv. Exp. Med. Biol.* 527, 167–176 (2003).
5. Guidetti, P. & Schwarz, R. 3-Hydroxykynurenine and quinolinate: pathogenic synergism in early grade Huntington’s disease? *Adv. Exp. Med. Biol.* 527, 137–145 (2003).
6. Kerr, S. I., Armati, P. J., Guillenin, G. I. & Brew, B. J. Chronic exposure of human neurons to quinolinic acid results in neuronal changes consistent with AIDS dementia complex. *AIDS* 12, 355–363 (1998).
7. Heyes, M. P. et al. Quinolinic acid in cerebrospinal fluid and serum in HIV-1 infection: relationship to clinical and neurological status. *Ann. Neurol.* 29, 202–209 (1991).
8. Heyes, M. P. et al. Sources of the neurotoxic quinolinic acid in the brain of HIV-1-infected patients and retrovirus-infected macaques. *FASEB J.* 12, 881–896 (1998).
9. Schwarz, R. Kynurenine pathway of tryptophan degradation as a drug target. *Curr. Opin. Pharmacol.* 4, 12–17 (2004).
10. Stone, T. W. & Darlington, L. G. Endogenous kynurenines as targets for drug discovery and development. *Nat. Rev. Drug Discov.* 1, 609–620 (2002).
11. Perozik, J., Nicholas, H., Wang, B. C., Lindahl, R. & Hempel, J. Relationships within the aldehyde dehydrogenase extended family. *Protein Sci.* 8, 137–146 (1999).
12. Under, G. & Bengaerts, J. Alternative regulatory pathways of *Escherichia coli*: energetics and transcriptional regulation in response to electron acceptors. *Biochim. Biophys. Acta* 1320, 217–234 (1997).
13. Nicholls, D. G. Mitochondrial function and dysfunction in the cell: its relevance to ageing and age-related disease. *Int. J. Biochem. Cell Biol.* 34, 1372–1381 (2002).
14. Hempel, J., Nicholas, H. & Lindahl, R. Aldehyde dehydrogenases: widespread structural and functional diversity within a shared framework. *Protein Sci.* 2, 1890–1900 (1993).

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15. Hasegawa, Y. et al. A novel degradative pathway of 2-nitrobenzoate via 3-hydroxyanthranilate in *Pseudomonas fluorescens* strain KU-7. *FEMS Microbiol. Lett.* **190**, 185–190 (2000).

16. Li, T., Walker, A. L., Iwaki, H., Hasegawa, Y. & Liu, A. Kinetic and spectroscopic characterization of ACMSD from *Pseudomonas fluorescens* reveals a pentacoordinate mononuclear metallocofactor. *J. Am. Chem. Soc.* **127**, 12282–12290 (2005).

17. Li, T., Ma, J., Hosler, J. P., Davidson, V. L. & Liu, A. Detection of transient intermediates in the mammalian non-oxidative decarboxylation catalyzed by *α*-aminobutyrylcarbamoyl-ε-semialdehyde decarbamylase. *J. Am. Chem. Soc.* **129**, 9278–9279 (2007).

18. Perez-Miller, S. J. & Hurley, T. D. Coenzyme isomerization is integral to catalysis in aldehyde dehydrogenase. *Biochemistry* **42**, 7100–7109 (2003).

19. Muñoz-Clare, R. A., González-Segura, L. & Díaz-Sánchez, A. G. Crystallographic evidence for active-site dynamics in the catalytic aldehyde dehydrogenases. Implications for the deacylation step of the catalyzed reaction. *Chem. Biol. Interact.* **191**, 137–146 (2011).

20. Huo, L., Davis, I., Chen, L. & Liu, A. The power of two: arginine 51 and arginine 239* from a neighboring subunit are essential for catalysis in *α*-aminobutyrylcarbamoyl-ε-semialdehyde decarbamylase. *J. Biol. Chem.* **288**, 30862–30871 (2013).

21. Geoghegan, K. F. et al. Spontaneous *α*-N-6-phosphogluconolysis of a ‘His tag’ in *Escherichia coli*: the cause of extra mass of 258 or 178 Da in fusion proteins. *Anal. Biochem.* **267**, 169–184 (1999).

22. Blanco, J., Moore, R. A. & Viola, R. E. Capture of an intermediate in the catalytic cycle of L-sparagine-*β*-semialdehyde dehydrogenase. *Proc. Natl Acad. Sci. USA* **100**, 12613–12617 (2003).

23. Colabroy, K. L. & Begley, T. P. The pyridine ring of NAD is formed by a nonenzymic pericyclic reaction. *J. Am. Chem. Soc.* **127**, 840–841 (2005).

24. Ichiyama, A. et al. Studies on the metabolism of the benzene ring of tryptophan in mammalian tissues. II. Enzymic formation of *α*-aminomuconic acid 3-hydroxyanthranilic acid. *J. Biol. Chem.* **240**, 740–749 (1965).

25. He, Z., Davis, J. K. & Spain, J. C. Purification, characterization, and sequence analysis of *α*-aminomuconic 6-semialdehyde dehydrogenase from *Pseudomonas pseudocaldogenes* JS45. *J. Bacteriol.* **180**, 4591–4595 (1998).

26. Abriola, D. P., Fields, R., Stein, S., MacKerril, J. A. D. & Pietruszko, R. Active site of human liver aldehyde dehydrogenase. *Biochemistry* **26**, 5679–5684 (1987).

27. Kitzson, T. M., Hill, J. P. & Midwinter, G. Identification of a catalytically essential nucleophilic residue in sheep liver cytosplasmic aldehyde dehydrogenase. *Biochem. J.* **275**, 207–210 (1991).

28. Farres, J., Wang, T. T., Cunningham, S. J. & Weiner, H. Investigation of the active site cysteine residue of rat liver mitochondrial aldehyde dehydrogenase by site-directed mutagenesis. *Biochemistry* **34**, 2592–2598 (1995).

29. Steinmetz, C. G., Xie, P., Weiner, H. & Hurley, T. D. Structure of mitochondrial aldehyde dehydrogenase: the genetic component of ethanol aversion. *Structure* **5**, 701–711 (1997).

30. Moore, S. A. et al. Sheep liver cytosolic aldehyde dehydrogenase: the structure reveals the basis for the retinal specificity of class 1 aldehyde dehydrogenases. *Structure* **6**, 1541–1551 (1998).

31. D‘Ambrosio, K. et al. The first crystal structure of a thiocyclenyl enzyme intermediate in the ALDH family: new coenzyme conformation and relevance to catalysis. *Biochemistry* **45**, 2978–2986 (2006).

32. Park, J. & Bhee, S. Structural basis for a cofactor-dependent oxidation protection and catalysis of cytochromeb3cemic semialdehyde dehydrogenase. *J. Biol. Chem.* **288**, 15760–15770 (2013).

33. Wang, X. & Weiner, H. Involvement of glutamate 268 in the active site of human mitochondrial (class 2) aldehyde dehydrogenase as probed by site-directed mutagenesis. *Biochemistry* **34**, 237–243 (1995).

34. Adams, P. D. et al. PHENIX: a comprehensive Python-based system for macromolecular structure solution. *Acta Crystallogr. D Biol. Crystallogr.* **66**, 213–221 (2010).

35. Emsley, P. & Cowtan, K. Coot: model-building tools for molecular graphics. *Acta Crystallogr. D Biol. Crystallogr.* **60**, 2126–2132 (2004).

36. Orville, A. M. et al. Correlated single-crystal electronic absorption spectroscopy and X-ray crystallography at NSLS beamline X26-C. *J. Synchrotron Radiat.* **18**, 358–366 (2011).

37. Pothineni, S. R., Strutz, T. & Lamzin, V. S. Automated detection and centring of cryo-cooled protein crystals. *Acta Crystallogr. D Biol. Crystallogr.* **62**, 1358–1368 (2006).

38. Frisch, M. J. et al. Gaussian 03, Revision E.01 (Gaussian, Inc., 2004).

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Author contributions
A.L. conceived of and led the study. H.I. and Y.H constructed the initial expression system. F.L. optimized the expression and established protein isolation and activation procedures. I.D. solved the apo-AMSDH structure. L.H. solved all complex and intermediate structures. L.H. and I.D. performed the kinetic assays. L.H. performed the mass spectrometry experiment. S.E. constructed the mutant expression systems. B.A. and A.M.O. collected the single-crystal electronic absorption spectra. I.D. performed the quantum chemical calculations. The manuscript was written by L.H. I.D. and A.L. All authors approved the final submitted manuscript.

Additional information
Accession codes: Coordinates and structure factors for apo-AMSDH, NAD+ bound AMSDH, NAD+ and 2-AMS-bound AMSDH, NAD+ and 2-HMS-bound AMSDH, thioyf intermediate AMSDH, E268A AMSDH, E268A thiochymoic intermediate, and E268A thioacyl intermediate have been deposited in the RCSB Protein Data Bank (Gaussian, Inc., 2004).

Supplementary Information
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