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Characterization of Cd36_03230p, a putative vanillin dehydrogenase from *Candida dubliniensis*†

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A coding sequence (CD36-03230) from the yeast *Candida dubliniensis* had been previously annotated as a vanillin dehydrogenase (VDH). The corresponding protein (CD36-03230p) was recombinantly expressed in *Escherichia coli* and analysed. The protein is most likely a tetramer in solution as judged by crosslinking and gel filtration experiments. CD36-03230p is an active aldehyde dehydrogenase favouring cyclic and aromatic substrates. Positive cooperativity and substrate inhibition were observed with some substrates. The redox cofactor NADP⁺ and substrates affected the thermal stability of the protein. Interestingly, the enzyme had no detectable activity with vanillin suggesting that the annotation is incorrect. It has been previously hypothesized that a methionine residue at a key position in the active site of yeast aldehyde dehydrogenases sterically hinders cyclic substrates and restricts specificity to aliphatic aldehydes. Molecular modeling of CD36-03230p demonstrates that it has an isoleucine residue (Ile-156) at this position, further strengthening this hypothesis.

Introduction

Most mammalian aldehyde dehydrogenases (ALDHs) have a broad specificity for aliphatic aldehydes, as well as some aromatic and polycyclic aldehydes,^{1–3} thus rendering an important protective enzymatic function against these xenobiotics. Eubacterial ALDHs, on the other hand, exhibit relatively narrow substrate specificity depending on their natural habitat and exposure to endogenous and exogenous aldehyde reactive elements. Hence, characterization of ALDHs capable of catalyzing the oxidation of aromatic aldehydes has been well documented in bacteria.^{4–13}

Vanillin dehydrogenase (VDH), a sub-class of benzaldehyde dehydrogenases,³ is a critical enzyme for the degradation of lignin

derived phenylpropanoids (such as vanillin, vanillate, caffeate, *p*-coumarate, cinnamate and benzaldehyde). These aromatic aldehydes, especially vanillin, are abundant as flavour and aromas in food and cosmetic industries. It is important to elucidate the structural and functional characteristics of these enzymes given their potential role in food chemistry and biotechnology. This group of enzymes has been characterized in a number of eubacterial species, including *Pseudomonas fluorescens*,⁶ *Pseudomonas putida* KT2440,¹³ *Corynebacterium glutamicum*,⁷ *Rhodococcus jostii* RHA1,⁴ *Amycolatopsis* sp. strain ATCC 39116,⁸ *Sphingomonas paucimobilis* SYK,⁹ and *Micrococcus* sp.¹⁰ Mammalian epithelial ALDH1A and salivary ALDH3A1 typically show activity towards a wide range of aromatic aldehyde substrates (including vanillin, benzaldehyde, cinnamaldehyde, 4-hydroxynonenal).¹⁴ Plant ALDH2 family members have also been observed to show a broad aromatic substrate specificity range, but with no report of activity with vanillin.¹⁵ Two ALDHs from white rot fungus *Phanerochaete chrysosporium* which are translationally up-regulated with exogenous addition of vanillin are active as VDHs.¹⁶

There are reports of recombinant vanillin production in metabolically engineered baker's yeast harbouring heterologous genes,¹⁷ but no studies are available focusing on the characterization of endogenous vanillin dehydrogenase enzymes in yeast species. To date, a single protein sequence CD36-03230p (accession number: XP_002416995) from *Candida dubliniensis* genome has been provisionally annotated as a putative vanillin dehydrogenase (Cd36_03230p). The purpose of this study was to characterise the substrate specificity and oligomeric structure of recombinant Cd36_03230p, in order to validate (or otherwise) its putative role. The results were interpreted, in part, based on results from our previous study which described two ALDHs from *Saccharomyces cerevisiae* var. *boulardii*.¹⁸

Methods

Recombinant expression and purification of Cd36_03230p

The coding sequence for *Cd36_03230* (based on accession number: XM_002416950) was synthesised following optimization

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of the sequence for expression in *Escherichia coli* (GenScript NJ, USA). The coding sequence was PCR-amplified and amplicons were inserted into the *E. coli* expression vector pET46 Ek/LIC (Merck-Millipore, Nottingham, UK) according to the manufacturer's instructions (note that this vector introduces bases coding for the amino acid sequence MAHHHHHHVDDDDK at the 5' end of the coding sequence). Correct insertion into the vector was verified by PCR and by DNA sequencing (GATC, London, UK) of the insert.

The expression vector was used to transform competent *E. coli* Rosetta™ (DE3) cells (Merck-Millipore) and colonies resulting from this transformation were used to inoculate cultures (5 ml of Luria Bertani medium (LB) supplemented with 100 µg ml⁻¹ ampicillin and 34 µg ml⁻¹ chloramphenicol) which were grown at 37 °C overnight (17–18 h) with orbital shaking. Each culture was then diluted into 1 l of LB (supplemented with 100 µg ml⁻¹ ampicillin and 34 µg ml⁻¹ chloramphenicol), grown (with orbital shaking) until A_{600} reached 0.6 to 1.0 (typically 5–6 h) at 30 °C, followed by a slow induction by adding 1.3 mM IPTG overnight (12–16 h) at 16 °C. These induction conditions were based on our previous experience of working with a wide variety of recombinant proteins. Cells were harvested by centrifugation (4200g for 15 min), resuspended in cell resuspension buffer (50 mM HEPES-OH, pH 7.5, 150 mM NaCl, 10% (v/v) glycerol) and stored frozen at –80 °C until the purification step.

For purification, cell suspensions were thawed, disrupted by sonication on ice (three pulses at 100 W for 30 s with 30 s gaps for cooling) and clarified by centrifugation (20 000g, 20 min, 4 °C). The supernatant was applied to a cobalt agarose column (1 ml, His-Select, Sigma, Poole, UK) which had been pre-equilibrated in buffer A (cell resuspension buffer, except 500 mM NaCl) and allowed to pass through by gravity. The column was washed with 40 ml of buffer A and the protein eluted with three 2 ml aliquots of buffer C (buffer A plus 250 mM imidazole). Protein containing fractions were identified by SDS-PAGE and dialysed overnight at 4 °C against cell resuspension buffer supplemented with 1 mM DTT. The concentration of Cd36_03230p was determined by the method of Bradford¹⁹ using BSA as a standard. The purified fractions were frozen at –80 °C in 20 µl aliquots.

Bioinformatics and modeling

Multiple sequence (structure-based) alignments were carried out for Cd36_03230p with known structures of aromatic aldehyde dehydrogenases (class 3) such as benzaldehyde dehydrogenase from *Pseudomonas putida* (PDB ID 3LV1), *Corynebacterium glutamicum* (PDB ID 3R64) and a salicylaldehyde dehydrogenase from *Pseudomonas putida* G7 (PDB ID 4JZ6) using T-Coffee algorithm in Expresso template mode available at <http://www.tcoffee.org>.^{20–22} Human retinal dehydrogenase 1 (PDB ID 4WB9)²³ and human liver mitochondrial dehydrogenase (PDB ID 1CW3)²⁴ were also incorporated into the alignment as members of class 1 and class 2 ALDHs respectively to have an insight into their relatedness with Cd36_03230p if any. The sequence homology was evaluated using ESPript 3.0 available at <http://esprict.ibcp.fr>.²⁵ The phylogenetic tree was constructed using

ClustalW Phylogeny (version 2.1), a web-based service available at <http://www.ebi.ac.uk/Tools/msa/clustalw2/>²⁶ by neighbour-joining method.²⁷

An initial molecular model of the protein was generated using Phyre2 (ref. 28) and energy minimized using YASARA.²⁹ A model of dimeric Cd36_03230p was generated by aligning two copies of the model to the ALDH domains of *Geobacter sulfurreducens* PutA (PDB 4NMB³⁰) and saving the two monomers into a single protein structure (pdb) file. This initial, dimeric model was then subjected to a second round of energy minimization using YASARA. A tetrameric model was generated in the same way using the tetrameric structure of sheep liver class 1 aldehyde dehydrogenase (1BXS³¹) as the template. These models are available as ESI† to this paper.

Cross-linking

Crosslinking with bis(sulfosuccinimidyl)suberate (BS³; 50–800 µM) was carried out with 18 µM protein (diluted as required in 100 mM sodium phosphate buffer pH 7.4) in a total volume of 10 µl. Reaction mixtures were incubated at 30 °C for 30 min before addition of the crosslinker and then incubated at the same temperature for a further 35 min. Reactions were stopped by addition of an equal volume of SDS-loading buffer (120 mM Tris-HCl, pH 6.8, 4% (w/v) SDS, 20% (v/v) glycerol, 5% (w/v) bromophenol blue, 1% (w/v) DTT) and analysed by 10% SDS-PAGE.

Analytical gel filtration

Cd36_03230p (200 µl of a 60 µM purified protein aliquot) was chromatographed on a Sephacryl S-300 (Pharmacia) column (total volume, $V_t = 65.2$ ml; void volume, $V_0 = 15.1$ ml) at a flow rate of 1 ml min⁻¹. The column was equilibrated and developed in buffer G (50 mM Tris-HCl, 17 mM Tris base, 150 mM sodium chloride, pH 7.4).^{32–34} Fractions (1 ml) were collected and analysed for protein content by measuring the absorbance at 280 nm. Standard proteins (thyroglobulin, 669 kDa; albumin, 67 kDa and chymotrypsinogen, 25 kDa) were used to calibrate the column. Their elution volumes (V_e) were used to calculate K_{av} according to the equation:

$$K_{av} = (V_e - V_0)/(V_t - V_0)$$

The Stoke's radius (R_s) was estimated from the inverse correlation of this parameter with K_{av} and the sedimentation coefficient ($S_{20,w}$) was estimated from the molecular models using WinHydroPRO 1.00.³⁵ The sub-unit stoichiometry (n) was then estimated using the equation:

$$nM = S_{20,w}N_A(6\pi\eta R_s)/(1 - v_2\rho)$$

where M is the molecular mass of a monomer (52 100 Da), N_A is Avogadro's number (6.023×10^{23} mol⁻¹), η is the viscosity of the solvent (0.01 g cm⁻¹ s⁻¹), v_2 is the partial specific volume (0.73 g cm³ g⁻¹)³⁶ and ρ is the density of the solvent (1.0 g cm⁻³). Values obtained with both the dimeric and tetrameric models of Cd36_03230p were compared in order to see which fit better to the experimental data.



ALDH activity measurements and enzyme kinetic analysis

The enzyme assay was performed as described previously.^{7,37–39} Aldehyde dehydrogenase enzyme activity was monitored at 30 °C using a ThermoScientific Multiskan™ Microplate spectrophotometer. The reactions contained 100 mM potassium phosphate buffer (pH 7.3), 0.5 mM NADP⁺, varied concentrations of substrates (10–1200 μM) and 0.6 μM enzyme. The long-chain (C₈–C₁₃) and phenolic aldehydes were dissolved in DMSO (1.7% (v/v), final concentration) as a solvent carrier.

Steady-state kinetic data was obtained in triplicates in 96-well plates with readings taken every 5 s. The initial, linear portion of the progress curve was identified by visual inspection and fitted to linear regression to give the initial rates (v) of change in absorbance at 340 nm. These rates were converted to molar units using the extinction coefficient of NADPH (6.22 mM⁻¹ cm⁻¹)⁴⁰ to give rates of reactions in micromolar concentration of NADH formed per second.

The kinetic parameters (k_{cat} , $K_{0.5}$ and Hill coefficient, h) were obtained by plotting the rates of reaction against substrate concentration and fitting the data to the equation below using non-linear regression as implemented in GraphPad Prism 6.0 (GraphPad Software Inc, CA). All points were weighted equally.

$$v = k_{\text{cat}}[E][S]^h / (K_{0.5}^h + [S]^h)$$

where k_{cat} is the turnover number, $[E]$ is the enzyme concentration, $[S]$ is the concentration of substrate, $K_{0.5}$ is the concentration of substrate that produces a half-maximal enzyme velocity (analogous to the Michaelis constant, K_m , in non-cooperative enzymes) and h is the Hill coefficient.^{41,42}

Differential scanning fluorimetry (DSF)

Enzyme aliquots were diluted in 50 mM HEPES, pH 7.3, to a concentration of 5–7 μM to a final volume of 20 μL. Sypro Orange (10×; manufacturer's concentration definition) was as previously described.^{43,44} Cofactor (NADP⁺) and substrates were added as appropriate. Where required, substrates were initially dissolved in 100% DMSO and diluted in buffer R (50 mM HEPES–OH, pH 7.5, 150 mM NaCl, 10% v/v glycerol) as required. The concentration of DMSO never exceeded 1% (v/v).

Results and discussion

Cd36_03230p is an unusual aldehyde dehydrogenase-like protein

Structure-based multiple sequence alignment showed conserved residues at both NAD(P)⁺ binding and catalytic domains. However, the Cd36_03230p sequence has gaps and substitutions at otherwise conserved residues when aligned to class 1, class 2 and class 3 ALDHs which suggests structural and functional disparity (ESI Fig. S1a†). It also forms an independent cluster on the phylogenetic tree and exhibits clear evolutionary distance with the already known structures of class 3 benzaldehyde and salicylaldehyde dehydrogenases with verified VDH activities,^{7,13} besides the class 1 and 2 ALDHs (non-VDH) (ESI Fig. S1b†). Therefore, it is evident that Cd36_03230p sequence is not

sufficiently similar to the salicylaldehyde dehydrogenases or benzaldehyde dehydrogenases to be classified as the member of the class 3 vanillin dehydrogenase family.⁴⁵

However, molecular modelling of Cd36_03230p predicted that it has a similar fold to other aldehyde dehydrogenases (Fig. 1a). The highest ranked template used in Phyre2 was PutA from *G. sulfurreducens* PutA (PDB 4NMB³⁰) with a root mean squared deviation (rmsd) of 1.0 Å over 1922 equivalent atoms. The fold is also predicted to be similar to mammalian aldehyde dehydrogenases (for example, the rmsd when compared to sheep liver aldehyde dehydrogenase is 0.7 Å over 1957 equivalent atoms).

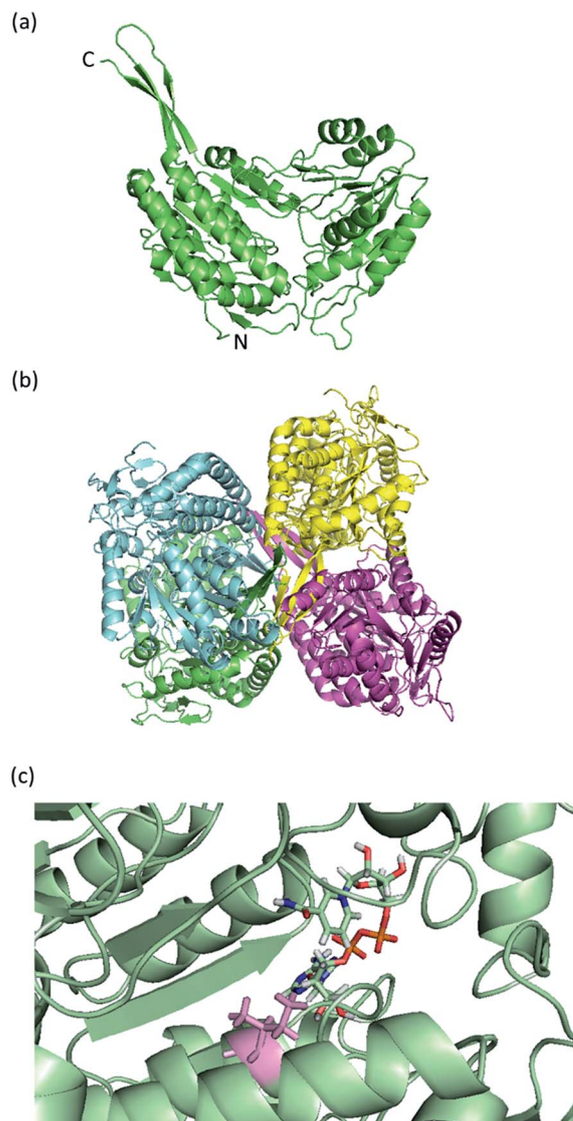


Fig. 1 Predicted structure of the putative vanillin dehydrogenase from *Candida dubliniensis*. (a) A view of the monomer structure with an NAD⁺ cofactor bound shown in stick format at the active site. (b) The predicted tetrameric structure with each subunit in a different colour. (c) The predicted active site showing the bound cofactor and residue Ile-156 (pink) which is structurally equivalent to the Met-177 in *S. cerevisiae* var. *boulardii* Ald6p. This residue is believed to be partly responsible for controlling access by cyclic aldehydes to the active site.



The predicted structure is largely α -helical with a protruding β -sheet region at the C-terminus which is, by comparison to oligomeric aldehyde dehydrogenase structures, likely to be involved in homo-oligomer assembly. Since there are both dimeric and tetrameric aldehyde dehydrogenases known, we built both oligomeric versions in order to assist with the interpretation of gel filtration experiments (see below). Comparison with the structure of sheep liver aldehyde dehydrogenase (which was solved with NAD⁺ bound) enabled the prediction of the cofactor binding site. This lies near the surface of each subunit, adjacent to a cleft which is the likely aldehyde substrate binding site (Fig. 1b). Previously, we have suggested that a bulky amino acid residue (Met-177 in *S. cerevisiae* var. *boulardii* Ald6p) is partly responsible for restricting access to bulkier, cyclic aldehydes.¹⁸ In Cd36_03230p, the structurally equivalent residue is Ile-156. Therefore, we hypothesized that this smaller residue may enable Cd36_03230p to accommodate cyclic aldehydes.

Expression, purification and oligomeric structure of Cd36_03230p

Cd36_03230p could be expressed in, and purified from, *E. coli* Rosetta™ (DE3) cells. Typical yields were approximately 1.5 mg l⁻¹ of bacterial cell culture (Fig. 2a). Unlike other yeast ALDHs,¹⁸ this protein did not show multiple bands resulting from

oligomerisation on 10% SDS-PAGE suggesting that any oligomeric form(s) are less resistant to heat and SDS denaturation.

The enzyme was able to form dimers and tetramers as demonstrated by chemical crosslinking with BS³. Resolution of the crosslinked products by 10% SDS-PAGE revealed bands corresponding primarily to a homotetramer (~210 kDa), with some higher order oligomers (Fig. 2b). The intensity of these bands was greater following treatment with increasing concentrations of BS³. Dimeric ALDHs have previously been reported in, for example, human ALDH3 due to an extended C-terminal tail which prevents tetramerisation.⁴⁶ However, our predicted structure of Cd36_03230p suggests that there is no such tail in this protein. Gel filtration chromatography was used to estimate the native molecular mass in solution and, thus the subunit stoichiometry. Sedimentation coefficients of models of the dimeric and tetrameric models of Cd36_03230p were computationally estimated as 6.3 and 10.0 S respectively in order to allow for effects due to the shape of the protein. The Stoke's radius was estimated from the gel filtration data as 5.0 nm. This yielded an estimated subunit composition of using the dimeric model of 2.5 and 4.0 using the tetrameric model. The tetrameric model is clearly a better fit to the data, suggesting the Cd36_03230p exists predominantly as a tetramer in solution. However, given that higher molecular mass species were detected by crosslinking, higher oligomeric forms may also be present.

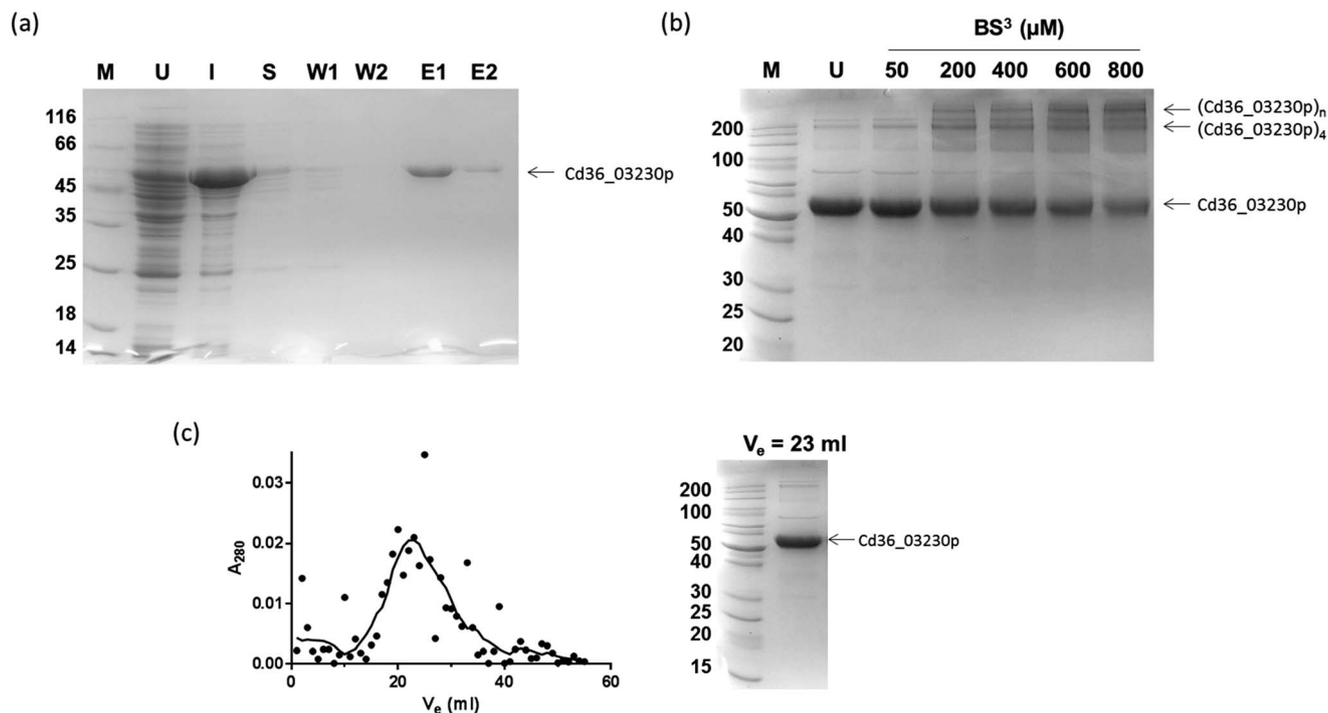


Fig. 2 Expression purification and oligomeric state of Cd36_03230p. (a) SDS-PAGE showing the stages in expression and purification of the protein. M, molecular mass markers (sizes shown to the left of the gel in kDa); U, cell extract from bacteria prior to induction with IPTG; I, cell extract from bacteria after induction and immediately prior to harvesting by centrifugation; S, extract of soluble proteins released on sonication; W1 and W2, the first and second washes of the cobalt affinity column; E1 and E2, the first and second elutions from the column. For details of buffers etc., see Methods. (b) Cross-linking of Cd36_03230p (18 μ M) with BS³. M, molecular mass markers (sizes shown to the left of the gel in kDa); U, untreated protein; the remaining lanes had increasing concentrations of BS³ as indicated above the gel. (c) Analytical gel filtration of Cd36_03230p. On the left, the elution profile of the protein. On the right, SDS-PAGE of the fraction with the highest absorbance demonstrating the presence of Cd36_03230p.



Cd36_03230p exhibits unusual kinetic patterns with cyclic and aromatic substrates

Cd36_03230p demonstrated activity with cyclohexanecarboxyaldehyde (a cyclic aliphatic aldehyde), benzaldehyde and 4-hydroxybenzaldehyde (an aromatic aldehyde), showing highest activity towards 4-hydroxy benzaldehyde judged by its k_{cat} to $K_{0.5}$ ratio (Table 1 and Fig. 3a). Interestingly, Cd36_03230p kinetics also showed substantial substrate inhibition by 4-hydroxy benzaldehyde at a substrate concentration higher than $\sim 10 \mu\text{M}$. Despite attempts to fit these data to various kinetic models, it was not possible to obtain a good fit with corresponding estimates of kinetic constants for the data with this substrate. Previously, substrate inhibition of benzaldehyde dehydrogenases from *Acinetobacter calcoaceticus* by benzaldehyde and betaine aldehyde dehydrogenase from *Staphylococcus aureus* by betaine aldehyde has been reported.^{47,48}

Apart from this, Cd36_03230p failed to show any activity towards aliphatic (short-chain and long-chain) and most aromatic aldehydes used in this study. These included acetaldehyde, propionaldehyde, butyraldehyde, isobutyraldehyde, valeraldehyde, hexanaldehyde, heptanaldehyde, octanaldehyde, nonanaldehyde, decyl aldehyde, undecyl aldehyde, dodecyl aldehyde, tridecyl aldehyde, crotonaldehyde, DL-glyceraldehyde and, notably, 4-hydroxy-3-methoxy benzaldehyde (vanillin).

Substrates and cofactors increases thermal stability of Cd36_03230p

Addition of NADP^+ (1.5 mM) resulted in a significant increase in the “melting temperature”, T_m , of Cd36_03230p as estimated by DSF (Table 2 and Fig. 3b). This suggests that this compound binds to, and stabilizes the protein. Long-chain aliphatic and aromatic aldehyde substrates (2 mM concentration) generally

Table 1 Steady state enzyme kinetics parameters for Cd36_03230p with various substrates^b

Substrate (CAS registry number)	k_{cat} (min^{-1})	$K_{0.5}$ (μM)	$k_{\text{cat}}/K_{0.5}$ ($\mu\text{M}^{-1} \text{min}$)	h
Cyclohexanecarboxyaldehyde (2043-61-0)	0.22 ± 0.03	48.0 ± 7.2	$(4.6 \pm 0.2) \times 10^{-3}$	2.9 ± 0.7
Benzaldehyde (100-52-7)	1.05 ± 0.14	42.8 ± 5.7	$(24.5 \pm 1.5) \times 10^{-3}$	3.2 ± 0.9
4-Hydroxybenzaldehyde (123-08-0)	^a			
Vanillin (121-33-5)	nd			

^a Although activity was observed with this compound, the kinetic data could not be fitted to Michaelis–Menten, cooperative or substrate inhibition kinetic models. ^b nd – no detectable turnover. Activity of Cd36_03230p was assayed using 100 mM potassium phosphate buffer (pH 7.3), 0.5 mM NADP^+ , varied concentrations of substrates (10–1200 μM) and 0.6 μM enzyme. Aldehydes were dissolved in 1.7% (v/v) DMSO as a solvent carrier. The final concentration of DMSO never exceeded 1% (v/v) in the assay. Values reported are those returned by non-linear fitting and are shown \pm the standard errors derived from this process.

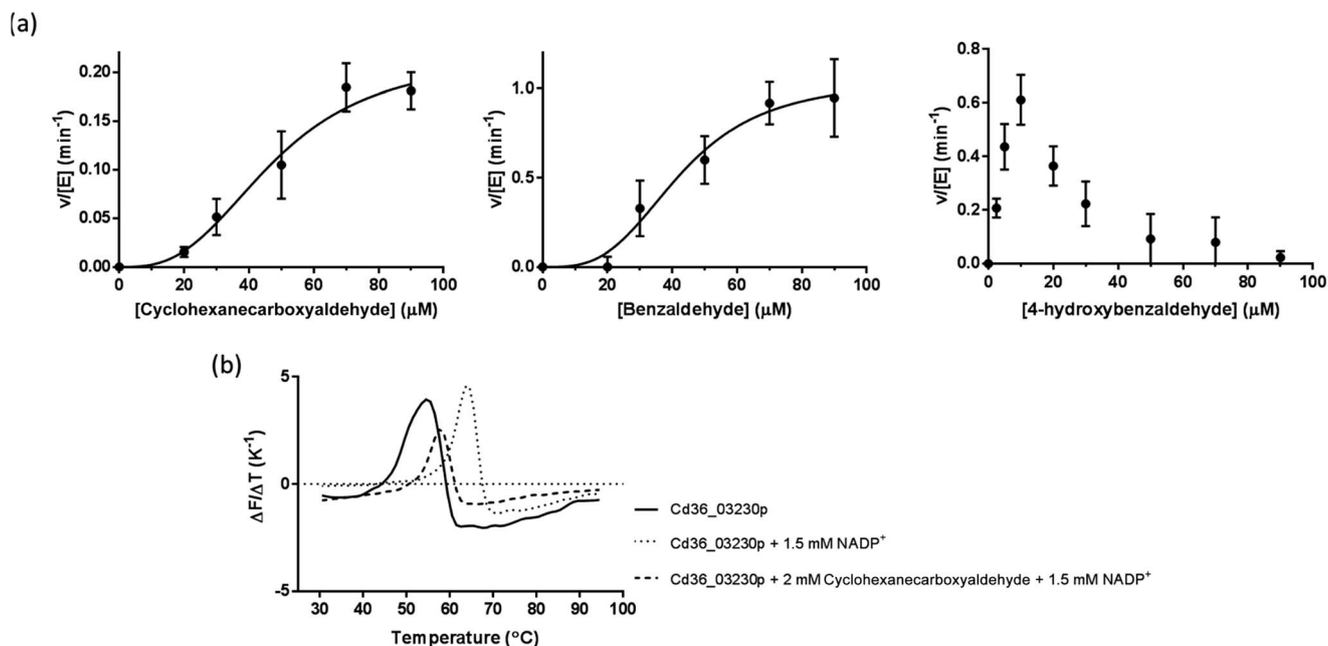


Fig. 3 Enzyme kinetics and thermal stability of Cd36_03230p. (a) The protein is an active aldehyde dehydrogenase with a limited range of substrates. Each point represents the mean of three determinations of the rate, the error bars the standard deviations of these means and the line the non-linear fit to the data. (b) A representative thermal denaturation experiment on Cd36_03230p (5 μM) in the presence of NADP^+ (1.5 mM) and cyclohexane aldehyde (2 mM).



Table 2 Thermal stability (T_m) of Cd36_03230p in the presence of various potential substrates

Substrate or cofactor	T_m (°C)
Untreated	53.5 ± 0.2
NADP ⁺	64.5 ± 0.0 ^a
Propionaldehyde	59.5 ± 0.0 ^a
Valeraldehyde	59.3 ± 0.1 ^a
Octanal	58.5 ± 0.6 ^a
Decanal	54.5 ± 0.7
Tridecanaldehyde	58.0 ± 0.3 ^a
Crotonaldehyde	59.0 ± 0.2 ^a
Cyclohexanecarboxyaldehyde	57.8 ± 0.5 ^a
Benzaldehyde	58.0 ± 0.4 ^a
4-Hydroxybenzaldehyde	58.8 ± 0.3 ^a
Vanillin	58.0 ± 0.3 ^a

^a Indicates a statistically significant difference ($p < 0.05$). Experiments with cofactors were compared to the appropriate untreated enzyme and experiments with cofactor and aldehyde were compared to the appropriate one with cofactor only. Aldehyde substrates were measured with 2 mM aldehyde and 1.5 mM NADP⁺.

reduced the thermal stability of the enzyme–NADP⁺ complex by ~4 °C (Table 2). This may indicate a slightly lower stability of the ternary enzyme–NADP⁺–aldehyde complex, perhaps resulting from increased overall flexibility in the protein. However, given that many of the compounds which affect the thermal stability are not substrates of the enzyme (and may, therefore, not interact with the protein), it is also possible that they cause a small destabilization through their general hydrophobic or chaotropic properties rather than through interaction at a specific site.⁴⁹

Conclusions

This study demonstrated that, despite being annotated as such, this enzyme has no detectable vanillin dehydrogenase activity. The data do show that the enzyme functions as an aldehyde dehydrogenase, with a strong preference for some cyclic and aromatic substrates. Our previous work suggested that a key residue in the active site influences the substrate specificity of yeast aldehyde dehydrogenases. In the case of *S. cerevisiae* Ald4p and Ald6p, the former has some activity to cyclic aldehyde substrates, but the latter does not.^{18,50,51} Ald6p has a bulky methionine residue (Met-177) which we hypothesized might sterically hinder the binding of cyclic substrates. In contrast, the structurally equivalent residue in Ald4p is Leu-196 and alteration of Met-177 in Ald6p to valine conferred some activity with cyclic aldehydes on this enzyme.¹⁸ In Cd36_03230p, the structurally equivalent residue is Ile-156. This provides further support for our hypothesis that a smaller hydrophobic residue at this position facilitates the binding of cyclic substrates. However, the lack of activity of Cd36_03230p with some cyclic substrates and all aliphatic substrates tested (in contrast with *S. cerevisiae* Ald4p and Ald6p which both act on aliphatic aldehydes) suggests that there are additional determinants of substrate specificity in these enzymes. Further studies are

required to elucidate these. The lack of activity with vanillin suggests that Cd36_03230p's putative annotation as a vanillin dehydrogenase is incorrect and should be changed. We suggest that cyclic/aromatic aldehyde dehydrogenase would be a more appropriate annotation.

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