

Zinc-selective inhibition of the promiscuous bacterial amide-hydrolase DapE: implications of metal heterogeneity for evolution and antibiotic drug design†‡

Cite this: *Metallomics*, 2014, 6, 88

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The development of resistance to virtually all current antibiotics makes the discovery of new antimicrobial compounds with novel protein targets an urgent challenge. The *dapE*-encoded *N*-succinyl-L,L-diaminopimelic acid desuccinylase (DapE) is an essential metallo-enzyme for growth and proliferation in many bacteria, acting in the desuccinylation of *N*-succinyl-L,L-diaminopimelic acid (SDAP) in a late stage of the anabolic pathway towards both lysine and a crucial building block of the peptidoglycan cell wall. L-Captopril, which has been shown to exhibit very promising inhibitory activity *in vitro* against DapE and has attractive drug-like properties, nevertheless does not target DapE in bacteria effectively. Here we show that L-captopril targets only the Zn²⁺-metallo-isoform of the enzyme, whereas the Mn²⁺-enzyme, which is also a physiologically relevant isoform in bacteria, is not inhibited. Our finding provides a rationale for the failure of this promising lead-compound to exhibit any significant antibiotic activity in bacteria and underlines the importance of addressing metallo-isoform heterogeneity in future drug design. Moreover, to our knowledge, this is the first example of metallo-isoform heterogeneity *in vivo* that provides an evolutionary advantage to bacteria upon drug-challenge.

Received 16th April 2013,
Accepted 8th August 2013

DOI: 10.1039/c3mt00125c

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Introduction

It has been estimated that about 25 000 people die in Europe alone every year from an infection due to common antibiotic-resistant bacteria, causing an economic burden in excess of 1.5 billion euro per year.¹ Similar alarming statistics are reported around the world, making the current inevitable rise of antibiotic-resistance to our limited subset of clinically useful drugs an issue of great concern.² It is therefore crucially important and urgent that new protein targets are identified and that their specific inhibitors are developed.

DapE, *N*-succinyl-L,L-diaminopimelic acid desuccinylase of the *meso*-diaminopimelate (mDAP)/lysine biosynthetic pathway of bacteria, has been identified as an attractive potential antibiotic target for two main reasons:³ (i) the growth of many bacteria is highly dependent on the biosynthesis of lysine and

diaminopimelic (DAP) acid, which are essential for protein synthesis and elaboration of their peptidoglycan cell wall, respectively,⁴ and (ii) both DAP and lysine are synthesized by the *dap* operon bacterial enzymes and no similar pathways take place in humans.

L-Captopril was identified as a low-micromolar inhibitor of the *H. influenzae* DapE enzyme *in vitro*.^{4a} Considering that L-captopril is a marketed drug against hypertension⁵ one may suppose that the compound already exhibits attractive pharmacokinetic properties. Moreover, considering the structural and functional conservation among DapE from various bacteria, one would expect that L-captopril may inhibit various bacterial homologues in a similar manner, thus potentially proving to be a broad-selectivity antibiotic. Disappointingly, however, L-captopril exhibited only very modest antimicrobial activity when tested in bacteria (including strains of *S. enterica* and *E. coli*). Moreover, this weak activity was revealed to be totally DapE-independent.⁶

The failure of a promising drug-like compound to exhibit any significant antimicrobial effect *in vivo* or to target the enzyme target identified *in vitro* is perplexing and acts as a sobering reminder of the difficulty of translating *in vitro* data to effects *in vivo*. In an attempt to overcome this initial setback, we aimed at understanding why L-captopril should fail to have any

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‡ Electronic supplementary information (ESI) available: Fig. S1. See DOI: 10.1039/c3mt00125c

significant antimicrobial potency in bacteria. Here, we present a simple answer to this conundrum, based on the surprising metal-selectivity of L-captopril and its inability to inhibit all microbiologically relevant metallo-isoforms. Further, our findings imply that heterogeneity in metal content of metallo-enzymes may confer selective advantages.

Materials and methods

Construction of C-terminus polyhistidine-tagged DapE (DapE-C6His)

A pET28b-based plasmid (Novagen) for DapE overexpression was constructed according to published methods.⁷ Briefly, the 1.2 kb *dapE* gene was PCR amplified from the plasmid pCM655dapE by using the primers from Microsynth, F-SEQpCM655 (5' GCGT ACGTCGCGACCGCGGACATGT 3') & R-pCM655XhoI (5' AGCCA AGCCTCGAGGGCGACGAGCTGTTC 3'), with *Pfu* turbo polymerase (Stratagene) and sub-cloned into pET28b plasmid (Novagen) at the restriction sites *Hind*III and *Xho*I (using restriction enzymes from Biolabs).

Expression and purification of recombinant DapE-C6His

The expression plasmid, pET28b-DapE, was transformed into strain BL21(DE3) (Stratagene); the pre-culture was made in LB-medium with 30 $\mu\text{g ml}^{-1}$ kanamycin and 2% glucose at 37 °C and the over-expression was induced with 1 mM isopropyl-1-thio- β -D-galactopyranoside (IPTG) when the OD₆₀₀ was between 0.5 and 0.8, in LB-medium containing 30 $\mu\text{g ml}^{-1}$ kanamycin at 20 °C throughout. The cells were harvested after 6 hours of IPTG induction and the cell pellet was frozen at -20 °C, which was then thawed on ice and re-suspended in lysis buffer (50 mM tricine pH 7.8, 0.5 M NaCl, 1 mM PMSF, 3 $\mu\text{g ml}^{-1}$ DNase I and 5 $\mu\text{g ml}^{-1}$ lysozyme) and the cell-lysis was carried out at 37 °C and 200 rpm for 1 hour. The soluble protein fraction was transferred into a fresh bottle at 4 °C after the centrifugation at high speed (9000g) for 30 minutes in a pre-cooled (4 °C) centrifuge.

The soluble protein fraction was filtered using a 0.45 μm filter and the loading buffer (50 mM tricine pH 7.8 & 0.5 M NaCl) and elution buffer (50 mM tricine, 0.5 M NaCl & 0.5 M imidazole and the final pH was adjusted to 7.8) were filtered using a 0.2 μm filter. Filtered DapE-C6His protein was purified on HiTrap 5 ml chelating HP columns (GE Healthcare) loaded with 0.1 M ZnCl₂ in 50 mM tricine pH 7.8 with an ÄKTA Purifier-10 from GE Life Sciences.

Removal of the labile metal from the purified protein solution was achieved by 3 step sequential dialysis: first dialysis against 10 l of 50 mM tricine pH 7.8 with 1 mM 1,10-phenanthroline, second dialysis against 10 l of 50 mM tricine pH 7.8 with 5 mM ethylenediaminetetraacetic acid (EDTA) and third dialysis with 50 mM de-metallated tricine pH 7.8. Each dialysis step was for 8 to 12 hours at 4 °C using the dialysis bags with an approximate molecular weight cut-off of 6 to 8 kDa (Spectrum Labs). Chelex-100 (Bio-Rad) chelating ion exchange resins were used for the de-metallation of the buffer, *i.e.* 50 mM tricine pH 7.8 used for dialysis.

Purified DapE protein from *S. enterica* exhibited a single band on SDS-PAGE, indicating a molecular weight of approximately 42 kDa. Protein concentrations were determined by using the Pierce BCA protein assay kit (Thermo Scientific). The yield of the purified DapE protein was about 1 mg L⁻¹ of bacterial culture.

Synthesis of SDAP

N-succinyl-L,L-diaminopimelic acid (SDAP) was synthesized in liquid phase through a standard coupling procedure between mono-*tert*-butyl succinate and enantiopure mono-*tert*-butyl protected L,L-diaminopimelic dimethyl ester (protected DAP), using *O*-benzotriazole-*N,N,N',N'*-tetramethyl-uronium-hexafluoro-phosphate (HBTU) as a coupling agent. The product of the coupling was then conveniently deprotected by standard cleavage of the protecting groups. DAP was previously synthesized according to published protocols *via* olefin cross-metathesis between enantiopure Boc-L-allylglycine-OMe and Cbz-L-vinylglycine-OMe, in yields above 70%.⁸

DapE kinetic assays

The enzymatic assays were carried out in triplicate in 50 mM tricine pH 7.8 at 37 °C, using the purified DapE-C6His to a final concentration of 1 ng μl^{-1} . The final substrate concentration of the native substrate (SDAP) was 20 mM in a reaction volume of 50 μl . Initially the DapE enzyme was pre-incubated with metals (0.2 mM ZnCl₂ & 0.5 mM MnCl₂ for the native substrate and 1 mM MnCl₂ for the promiscuous substrate) at 37 °C for an hour and then with the inhibitor for another 10 minutes. L-Captopril was purchased from Sigma-Aldrich. In all cases the reactions were initiated by adding the substrate, either SDAP or Asp-Leu (Shanghai FWD Chemicals). The enzymatic reaction was followed by measuring the hydrolysis of the peptide bond of the substrates.⁹ Readings were carried out every 2 minutes for about 3 hours at UV230nm in UV transparent 96-well plates (Greiner) using a TECAN Safire plate reader.

Growth assays

Bacterial growth assays were carried out with strain TN5911, a knockout strain for DapE and for several dipeptidases; therefore, mDAP (Bachem) has to be supplemented in both minimal and enriched media. An appropriate source of leucine and proline is also required in minimal medium and supplementation with lysine provides better growth (Prof. Charles G. Miller, personal communication). For DapE expression and selection assays, strain TN5911 was transformed with the plasmid pCM655/DapE to form strain TN5935.^{6,7}

The metal-dependent anti-microbial activity of L-captopril was monitored by measuring the growth curves of the TN5935 in LB-medium containing L-captopril (0, 1 mM, 5 mM, 25 mM, 100 mM & 300 mM) under native activity selection (no mDAP supplementation) and under no selection (supplemented with 1 mM mDAP) and with and without 0.35 mM metal (MnCl₂ or ZnCl₂) supplementation. All measurements were carried out at least in duplicate. The main cultures were grown and the OD₆₀₀ was measured in flat bottom 96-well plates (200 μl culture per well) in a plate reader (TECAN Safire) at 37 °C for every 16 minutes.

The main cultures were started by adding the pre-culture of OD₆₀₀ ~ 1 to an initial OD₆₀₀ between 0.1 and 0.2.

The effect of the metals on DapE enzyme activities in bacteria was monitored by measuring the growth curves of the TN5935 under no selection and under selection for native & promiscuous activities with and without 0.35 mM metal salt (MnCl₂ or ZnCl₂) supplementation. These growth curves were measured from the cultures of minimal medium (1% MgSO₄·7H₂O, 10% citric acid·H₂O, 50% K₂HPO₄, 17.5% NaH₂NH₄PO₄·4H₂O, 0.4% glucose, 0.4 mM proline, 0.4 mM lysine, 0.4 mM leucine or Asp-Leu, 1 mM IPTG and, if required, 1 mM mDAP) by following the OD₆₀₀ every 4 hours at 37 °C at 200 rpm. The cultures were started by adding the pre-culture of LB-medium of OD₆₀₀ ~ 1 to an initial OD₆₀₀ between 0.1 and 0.2, which corresponded in this case to 0.4 ml of overnight LB-medium culture in 100 ml of minimal medium in a 1 l Erlenmeyer flask. We carried out a control culture to confirm that at this dilution of rich LB-medium in minimal medium (250-fold dilution), there is insufficient carry-over of leucine to allow for any detectable growth of the *dapE* knockout strain (TN5911) when supplementing with mDAP.

Structural modelling

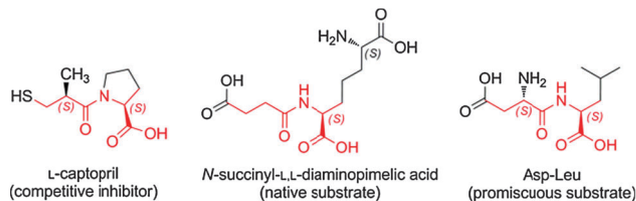
A homology model of the *Salmonella enterica* DapE model was built with SWISS-MODEL.¹⁰ The di-zinc *Haemophilus influenzae* X-ray structure [PDB: 3IC1] sharing 61% identity with *S. enterica* without water molecules was used as a modelling template.

The resulting model was energy minimized using the Yasara energy minimization server.¹¹ The conformation of residue Lys174 was replaced by an alternative rotamer in Maestro¹² to enlarge the ligand binding site. For the mono-zinc model, ZN2 was removed. The resulting structures were energy minimized using MacroModel.¹³ Subsequently, the L-captopril structure [PDB: X8Z] was docked into the active site of the mono- and di-zinc structures using SwissDock.¹⁴ Ligand conformations satisfying the constraints of metal-coordination that were imposed based on mechanistic considerations (see Results section) were found within the 25 and the 10 best ranked poses for the di-zinc and the mono-zinc models, respectively. These best-ranked models were submitted to the model archive (www.modelarchive.org) and are freely available for download (DOI: 10.5452/ma-a1nb6). All model figures were drawn with OpenStructure.¹⁵

Results

Metal substitution changes L-captopril inhibition *in vitro*

DapE was active as an SDAP desuccinylase *in vitro* both in its Zn²⁺ mono-metallated form (Zn/-) and as a Zn²⁺ binuclear (Zn-Zn) or mixed (Zn-Mn) form, with comparable V_{\max} (Scheme 1 and Table 1). As reported previously,⁷ no activity was detected for Asp-Leu hydrolysis with the Zn/- and Zn-Zn isoforms of the enzyme, whereas only the Zn-Mn isoform was active with this promiscuous substrate (in our hands, with k_{cat} 383 s⁻¹). These activity profiles correspond to the metallation status ascribed to the various enzyme preparations.



Scheme 1 L-Captopril inhibitor and substrates of DapE used in this study. Elements common to the three molecules are highlighted in red.

Table 1 DapE desuccinylation activity *in vitro*^a

DapE metallo-isoform	V_{\max} ($\mu\text{mol L}^{-1} \text{s}^{-1}$)	k_{cat} (s^{-1})
Zn/-	18.23	309
Zn-Zn	39.81	677
Zn-Mn	64.99	1104

^a V_{\max} and k_{cat} values of desuccinylase activity of different metallo-isoforms of recombinant DapE from *Salmonella enterica* typhimurium.

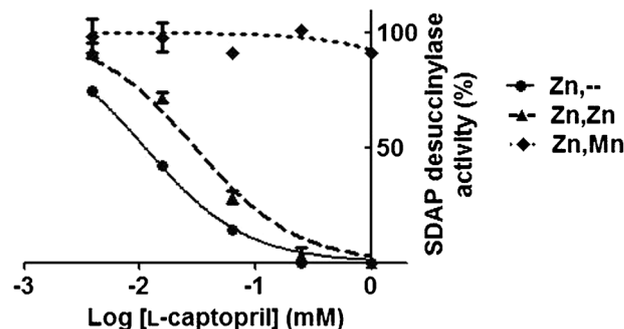


Fig. 1 Inhibition of desuccinylase activity of different DapE metallo-isoforms by L-captopril. Measurements were carried out in triplicate and error bars represent the standard deviation from the mean.

Addition of L-captopril *in vitro* inhibited the catalytic activity of the (Zn/-) or (Zn-Zn) metallated isoforms of DapE, with an IC₅₀ of 10 μM and 28 μM at 20 mM substrate, respectively (Fig. 1). However, no significant inhibition of the Zn-Mn isoform was found even at 1 mM L-captopril.

The substrate concentration at which half the maximum velocity was measured for SDAP hydrolysis (K_{50}) was similar for the Zn-Zn and Zn-Mn metalloisoforms (12 mM and 12.5 mM respectively). DapE from *Haemophilus influenzae* was reported in two independent studies^{9,16} to have comparable K_m for the Zn/-, Zn-Zn and Zn-Co metalloisoforms, which also supports the notion that the second metal only marginally affects the binding of the native substrate. Together, these results suggest that the metallation state at the second position influences the binding of both the inhibitor L-captopril and the promiscuous substrate Asp-Leu, but does not affect much the binding of the native substrate SDAP.

Manganese isoform is present *in vivo*

Traditionally, it has been considered that Zn²⁺ is the native and physiologically relevant metal of DapE. However, the enzyme presents also a promiscuous dipeptidase activity that is dependent on the presence of manganese ions (Mn²⁺), which can also be

selected *in vivo* in a leucine auxotroph strain using Asp-Leu as a substrate.⁷ Since this non-native dipeptidase activity occurs *in vitro* only when the labile, co-catalytic Zn^{2+} is replaced by Mn^{2+} to form a mixed Zn–Mn binuclear centre, it is surmised that the DapE-dependent promiscuous activity in bacterial cultures also arises from Mn^{2+} -metallo-isoforms of DapE *in vivo*.⁷

When selecting for the native desuccinylase activity in bacteria, the conditions in which no metal was added and where Zn^{2+} was supplemented showed similar growth; in contrast Zn^{2+} slowed down the growth of cell cultures when selecting for dipeptidase activity, which suggests that Zn^{2+} is deleterious for the promiscuous activity of DapE, but not for its native activity (Fig. 2). DapE was able to desuccinylate SDAP in its Zn-, Zn–Zn and Zn–Mn metallo-isoforms *in vitro*, albeit to different extents, with the following trends for k_{cat} : Zn–Mn > Zn–Zn > Zn- (Table 1). *S. enterica* grew appreciably better in Mn^{2+} -supplemented media when selected for either native or promiscuous activity (Fig. 2). Therefore, it appears that addition of Mn^{2+} promotes dipeptidase activity of DapE both *in vivo* and *in vitro*, whereas Zn^{2+} inhibits the dipeptidase activity, presumably by competing with Mn^{2+} for the labile metal-binding site of DapE.

We conclude from these observations that (i) metallation of DapE can be influenced by metal-supplementation with either Zn^{2+} or Mn^{2+} ; (ii) Mn^{2+} -metallation is growth-limiting in non-supplemented medium, particularly when selecting for the dipeptidase activity, and (iii) the Mn^{2+} isoform may be normally present *in vivo* but is not the only metallated species of DapE that occurs.

Zn^{2+} supplementation makes bacteria more susceptible to L-captopril inhibition compared to Mn^{2+}

Having concluded from growth profiles that metal supplementation of bacterial cultures can influence the state of metallation of DapE,

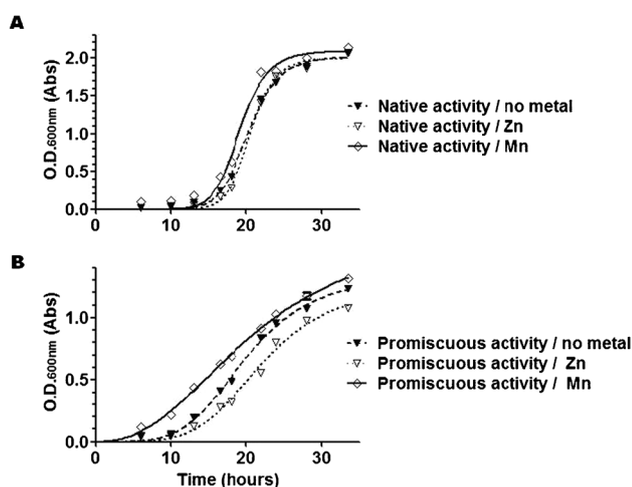


Fig. 2 Growth curves of bacterial cultures selected for native (desuccinylase) DapE activity (A) or promiscuous (dipeptidase) DapE activity (B) in the absence or presence of Zn^{2+} or Mn^{2+} supplementation. Note that Mn^{2+} accelerates the growth of the culture appreciably when selecting for both native and promiscuous activity. In contrast, Zn^{2+} slows down the growth of the culture appreciably when selecting for promiscuous activity but has little effect on proliferation when selecting for native activity. Plotted are fitted curves from means of duplicate cultures, with individual values comprised in the symbols. The values plotted are representative of two independent experiments with similar findings.

we next investigated whether metal-supplementation also affected susceptibility to L-captopril in bacterial cultures. We reasoned, since L-captopril targets only the Zn^{2+} -isoform of DapE *in vitro* and since metal supplementation can influence the metallation of DapE *in vivo*, that we may be able to influence the sensitivity of bacterial cultures to L-captopril through metallation of DapE.

First, we confirmed that L-captopril was toxic in a dose-dependent manner, shown by bacterial growth-curves in rich medium at different inhibitor concentrations (Fig. S1, ESI†). Second, in the growth conditions tested, in rich medium and 25 mM L-captopril, metal supplementation with either divalent Zn or Mn lowered L-captopril toxicity. The influence of metal supplementation on bacterial growth under these conditions was much larger in the presence of 25 mM L-captopril than in the absence of the drug (Fig. 3): Mn^{2+} -supplemented cultures clearly showed better growth as compared to Zn^{2+} supplemented or especially non-metal-supplemented cultures, which were more susceptible to L-captopril inhibition. The growth-profile in the presence of L-captopril was $Mn^{2+} > Zn^{2+} >$ no metal. Intriguingly, L-captopril inhibition of DapE *in vitro* has exactly the complementary activity profile: L-captopril inhibition $Mn^{2+} < Zn^{2+} <$ no metal.

Despite the metal-dependent effect of L-captopril, bacterial growth in the absence of selection for DapE activity (*i.e.* with mDAP supplementation) showed that the toxicity of this drug is largely independent of DapE activity (Fig. 3B), confirming our previous finding that this essential enzyme is not the main target of inhibition of L-captopril *in vivo* and suggesting that other metalloproteins may also be the main targets of L-captopril.⁶

We conclude that at least some of the targets of L-captopril are metal-dependent, *i.e.* metal-supplementation decreases sensitivity to L-captopril. The finding that Mn^{2+} supplementation

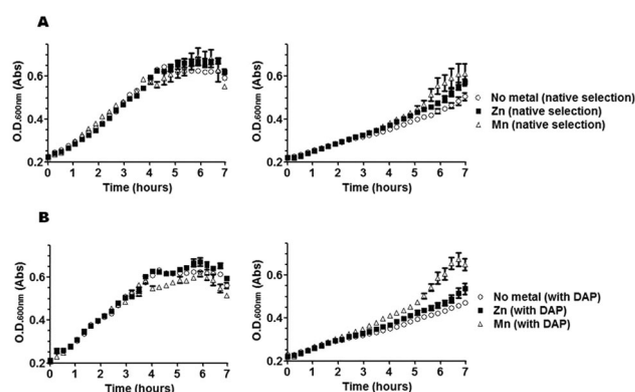


Fig. 3 Effect of metal supplementation on growth curves of bacterial cultures selected for DapE activity (A) or non-selected (B), in the absence of L-captopril (left) or in the presence of 25 mM L-captopril (right). Cultures were grown in LB broth either in the presence or absence of mDAP for selection of native (desuccinylase) activity of DapE. The effect of divalent Zn or Mn was tested and compared to the absence of metal supplementation (no metal). Note that L-captopril toxicity is dependent on metal-supplementation and that Mn^{2+} is more protective than Zn^{2+} , although the toxic effect of the drug is largely independent of DapE activity. Means of measurements from quadruplicate cultures are plotted with standard deviations shown as error bars.

is more protective than Zn^{2+} supplementation is consistent with the notion that L-captopril has a preference for Zn^{2+} -containing targets, at least some of which may also be functionally active with Mn^{2+} . We also surmise that the failure of L-captopril to exhibit any significant DapE-selective antibiotic activity in bacteria is due to the existence of sufficient amount of Zn–Mn isoform, which is not susceptible to L-captopril inhibition.

Model of L-captopril inhibition supports that thiol is the Zn^{2+} -binding group

We created a model of *S. enterica* DapE (Fig. 4) based on the high-resolution crystal structure of the homologous enzyme from *Haemophilus influenzae* reported by Nocek *et al.*¹⁷ The DapE enzyme is a dimer, each monomer of which contains two domains: a dimerization domain and a catalytic domain. The active site of the enzyme can contain either two or one zinc ions. The strongly bound Zn^{2+} is referred to as Zn1, whereas the

Zn^{2+} occupying the site of the labile metal (M2, which can be either Zn^{2+} or Mn^{2+}) is referred to as Zn2.

To generate the model of L-captopril binding to the active site, we imposed some constraints based on prior knowledge of DapE and L-captopril as a substrate analogue (Scheme 1). First, as in the suggested catalytic mechanism of DapE,¹⁷ we propose that the carbonyl group of the amide coordinates with the catalytic metal that acts as a Lewis acid (Zn1). Second, since captopril binds to the catalytic zinc of Angiotensin Converting Enzyme (ACE) through coordination by the sulfhydryl group⁵ and since we found experimentally that the nature of the co-catalytic metal (M2, either Zn^{2+} or Mn^{2+}) influences inhibition by L-captopril, we expect the sulfhydryl group to be oriented toward this second metal. Satisfyingly, docking-poses that fulfil these constraints were found within the 25 highest ranked poses (Fig. 4).

Divalent manganese behaves similar to magnesium in having a preference for oxygen ligands, although Mn^{2+} is more receptive of nitrogen ligands; in contrast, zinc prefers nitrogen and sulfur as ligands and, in general, a lower coordination number.¹⁸ Consequently, our model provides a reasonable and straightforward qualitative explanation for the metal-dependent selectivity of L-captopril inhibition of DapE: the presence of Mn^{2+} does not allow strong coordination by the sulfhydryl group, whereas either the absence of a co-catalytic metal or the presence of a second Zn^{2+} allows coordination with the sulfur liganding either Zn2 or Zn1, respectively.

Discussion

Implications for drug development

L-Captopril was the first marketed anti-hypertension drug, targeting the Zn-metalloprotease Angiotensin I Converting Enzyme (ACE) through coordination of the catalytic metal by a sulfhydryl group, with low nM affinity.⁵ However, L-captopril has also shown inhibitory activity towards a variety of other zinc metalloproteases, although this is typically several orders of magnitude weaker than with ACE.¹⁹ Recently, L-captopril was identified also as a low-micromolar inhibitor of *H. influenzae* DapE (IC₅₀ = 3.3 μ M). We found that L-captopril inhibited the Zn–Zn isoforms of *S. enterica* DapE with IC₅₀ values comparable to those of the *H. influenzae* enzyme (*i.e.* IC₅₀ of 10 μ M vs. 3.3 μ M, respectively), providing further evidence for the similarities in the structure and function of the active sites of DapE of different species.^{17,20} The low-micromolar affinity of L-captopril for the Zn/- and Zn–Zn isoforms of DapE could be seen as a reasonable starting point for further optimisation of this lead compound,^{3,4} akin to what succinyl-proline was for ACE.^{5,19} However, note that L-captopril at the concentrations at which it inhibits DapE might cause hypotension in normo-tensive individuals, which would warrant further development of its antibiotic properties. Moreover, at this low-micromolar level of affinity some off-targets are likely to occur, as indeed suggested by the poor activity of L-captopril⁶ and by the general protective effect of Zn^{2+} and especially Mn^{2+} supplementation (Fig. 3). We were nevertheless initially very surprised that we could detect

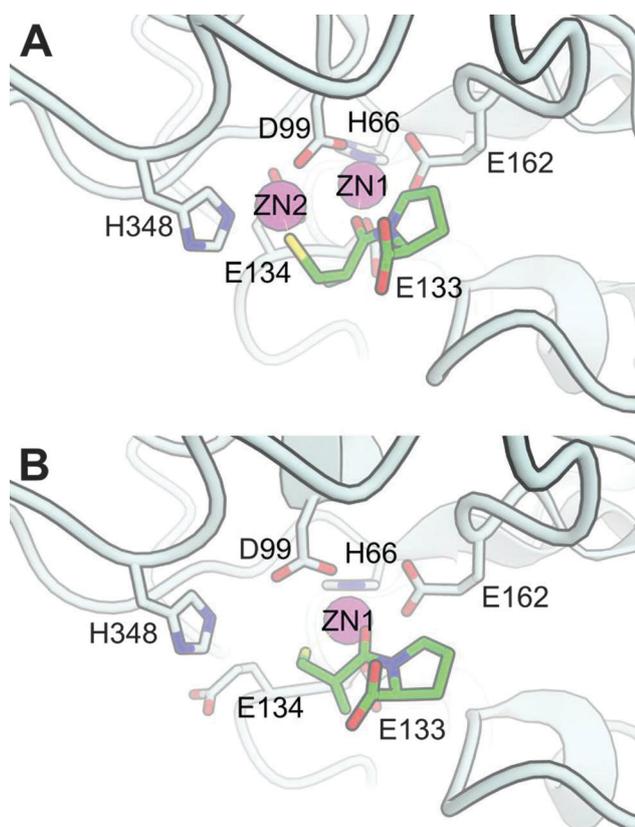


Fig. 4 Model of *S. enterica* DapE based on the crystal structure of *H. influenzae* DapE (PDB: 3IC1), with L-captopril docked to the (A) di-zinc and (B) mono-zinc catalytic metal center. Only the side-chains of important residues for catalysis are drawn and labelled, *i.e.* coordinating amino acids and the catalytic base (E134). L-Captopril is drawn in the stick-form with each type of atom colored differently: green carbon; blue nitrogen; red oxygen and yellow sulfur. Although coordination bonds with the ligand are not explicitly drawn, in both the mono- and di-zinc form the L-captopril carbonyl group of the amide coordinates with the Zn1, whereas the L-captopril thiol group can additionally coordinate either with Zn2 in the di-zinc structure (A) or with Zn1 in the mono-zinc structure, to form a 6-membered ring (B). These two models are freely available for download with DOI: 10.5452/ma-a1nb6 (www.modelarchive.org).

no evidence of L-captopril inhibiting DapE activity *in vivo* and that the compound had such a modest antimicrobial activity, despite low-micromolar affinity of this drug *in vitro* for an essential enzyme. We now forward the proposal that a crucial contribution to this low antibiotic activity is that L-captopril does not target one of the physiologically relevant isoforms of DapE, *i.e.* the Zn–Mn form of the enzyme.

It is generally acknowledged that any screen (or selection) should reflect the desired result as closely as possible, because ‘you get what you screen for’.²¹ Becker and colleagues^{4a} identified L-captopril as an inhibitor of DapE in a screen biased toward compounds containing zinc-binding groups, which included thiols, carboxylic acids, boronic acids, phosphonates and hydroxamates, and using the Zn-form of DapE of *H. influenzae* as a target *in vitro*. Consequently, a Zn-selective compound was identified with low micromolar affinity *in vitro*. Our discovery offers a plausible explanation for the failure of an apparently promising compound to show any DapE-targeted antimicrobial activity:⁶ the Zn–Mn mixed-metal isoform, which is not inhibited by L-captopril, is normally present *in bacteria*, as originally suggested by Broder and Miller⁷ judging by the presence of a selectable Mn²⁺-dependent promiscuous activity, which we confirmed here also by exploring bacterial growth upon metal-supplementation of cultures.

Exact knowledge of the metallation state of proteins *in vivo* is often thwarted by experimental difficulties.²² Moreover, metallation may exhibit strong dependence on the cellular context, differing for example in different media, cell-locations or in different species.²³ Several other examples have been reported where incomplete knowledge of the physiologically relevant metal has hindered drug development. For example, potent inhibitors of methionine aminopeptidase discovered *in vitro*, which were active against either Co²⁺ or Mn²⁺-isoforms of the enzyme, failed to have any of the expected antibacterial activity *in vivo*.²⁴ It was later suggested that Fe²⁺ is the native cofactor in *E. coli*, prompting a move either toward Fe²⁺-selective inhibitors that exhibited antimicrobial activity²⁵ or, in what may be called a “bet-hedging move” in the absence of a clear candidate, toward exploring selectivity against a plethora of different metallo-isoforms.²⁶ Similarly, peptide deformylase is another bacterial metalloenzyme and a promising antibiotic target²⁷ that was originally thought to be a Zn-metalloenzyme in *E. coli*²⁸ but was later shown instead to contain Fe²⁺, a metal that due to its tendency to oxidize in air leads to facile loss of enzyme activity.²⁹ However, the precise identity of the metal in peptide deformylase may be species-specific, since another bacterial pathogen, *Borrelia burgdorferi*, does use Zn²⁺ in this enzyme.³⁰ Despite considerable interest in peptide deformylase inhibitors in recent years^{27c} it appears that little attention has been paid to the issue of metallo-selectivity of drugs targeting this essential enzyme.

Implications for evolution

In general, the presence of different metal ions in metalloenzymes can maintain the overall structure of the active site, but provide alternative coordination interactions with substrates and inhibitors. It has already been pointed out that non-strict metal incorporation can be used as a facile way to create catalytic diversity.³¹ For

example, metal-heterogeneity has been proposed to have evolved in the carbonic anhydrase of marine diatoms that are challenged with low Zn-availability, thereby acquiring the capability of using either Zn²⁺ or Cd²⁺ in the active site of this essential enzyme.³² Indeed, we suggest that such metal-ambiguity and the resulting promiscuous behaviour *in vivo*, which may be considered a further example of “messiness in biology”,³³ is in fact a more general survival strategy, for example to avoid facile drug inhibition of an essential enzyme like DapE. To our knowledge, this may be the first example of metal-promiscuity providing a source of (non-genetic) diversity that confers an intrinsic survival advantage in defending against potential inhibitors, namely to the synthetic drug L-captopril, but also likely against other naturally occurring dipeptide-like compounds. Considering the importance of diversity as a driver of Darwinian evolution, we speculate that such metal-promiscuity may be much more common than hitherto thought, not only to provide substrate promiscuity in the development of new catalytic function, but also promiscuity for inhibitor-escape.

Conclusions

In summary, DapE in bacteria is heterogeneous, containing various metallated forms – mono-metallated with Zn²⁺ or binuclear with Zn²⁺ or Mn²⁺ – all of which are active as a desuccinylase, the essential and native function of the enzyme. We also conclude that the Zn–Mn binuclear isoform, which is very likely to be present *in vivo* as judged by bacterial selection of the promiscuous dipeptidase activity of DapE, is not inhibited by L-captopril, thus providing a mechanism of escape against this potential antibiotic. Such metal heterogeneity and resulting promiscuous properties may confer important selective advantages, including escape of inhibition, and therefore may have been selected for in evolution. Finally, the discovery of metal-selective inhibition of DapE that we describe here paves the way toward the discovery of compounds with improved antimicrobial properties. Our efforts are now directed toward finding novel inhibitors that target also the physiologically relevant Mn²⁺ isoform of DapE.

Abbreviations

mDAP	<i>meso</i> -Diaminopimelic acid
DAP	Diaminopimelic acid
SDAP	<i>N</i> -Succinyl-L,L-diaminopimelic acid
DapE	<i>N</i> -Succinyl-L,L-diaminopimelic acid desuccinylase
ACE	Angiotensin I converting enzyme

Acknowledgements

The authors would like to thank the financial support from the Swiss National Science Foundation (Grants 200021-116344 and CR22I3_143770), the State Secretariat for Education and Research (Grant C11.0054) and COST Actions TD0905 and CM0902. MC is particularly grateful for the support from the Holcim Foundation and GA from the Novartis Foundation. The

authors would also like to acknowledge Prof. Thomas R. Ward for general encouragement and for allowing the use of laboratory space and facilities, as well as thank Prof. Charles G. Miller (University of Illinois) for his generous donation of valuable research materials and insights.

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