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Structural Basis for Catalysis by the Mono- and Dimetalated Forms of the \textit{dapE}-Encoded \textit{N}-succinyl-l,l-Diaminopimelic Acid Desuccinylase☆

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Abstract

**Biosynthesis** of **lysine** and **meso**-diaminopimelic acid in **bacteria** provides essential components for **protein synthesis** and construction of the bacterial **peptidoglycan cell wall**. The **dapE operon** enzymes synthesize both **meso**-diaminopimelic acid and lysine and, therefore, represent **potential** targets for novel antibacterials. The **dapE**-encoded **N**-succinyl-**l,l**-diaminopimelic acid desuccinylase functions in a late step of the pathway and converts **N**-succinyl-**l,l**-diaminopimelic acid to **l,l**-diaminopimelic acid and succinate. Deletion of the **dapE gene** is lethal to **Helicobacter pylori** and **Mycobacterium smegmatis**, indicating that DapE's are essential for **cell growth and proliferation**. Since there are no similar pathways in **humans**, inhibitors that target DapE may have selective toxicity against only **bacteria**. A major limitation in developing antimicrobial agents that target DapE has been the lack of structural information. Herein, we report the high-resolution **X-ray crystal structures** of the DapE from **Haemophilus influenzae** with one and two **zinc** ions bound in the **active site**, respectively. These two forms show different activity. Based on these newly determined structures, we propose a revised **catalytic mechanism** of **peptide bond** cleavage by DapE enzymes. These structures provide important **insight** into catalytic mechanism of DapE enzymes as well as a structural foundation that is critical for the rational design of DapE inhibitors.

**Abbreviations**

- **mDAP**  
  *meso*-diaminopimelate
- **I,I-SDAP**  
  **N**-succinyl-**l,l**-diaminopimelic acid
- **CPG**₂  
  Carboxypeptidase from **Pseudomonas** sp. strain RS-16
- **AAP**  
  Leucine aminopeptidase from **Aeromonas proteolytica**
- **PDB**  
  Protein Data Bank

**Keywords**

- X-ray crystallography, zinc, hydrolases, mechanism, antimicrobials

**Introduction**

The **meso**-diaminopimelate (**mDAP**)/**lysine** biosynthetic pathway offers several potential antibacterial enzyme targets that have yet to be explored.¹, ², ³ One of the products of this pathway, lysine, is required in protein synthesis and is also used in the peptidoglycan layer of Gram-positive bacterial cell walls. A second product of this pathway, **mDAP**, is an essential component of the peptidoglycan cell wall for Gram-negative bacteria, providing a link between polysaccharide strands. It has been shown that deletion of the gene encoding one of the enzymes in this pathway, the **dapE**-encoded **N**-succinyl-**l,l**-diaminopimelic acid (**I,I-SDAP**) desuccinylase (DapE; EC 3.5.1.18), is lethal in **Helicobacter pylori** and **Mycobacterium smegmatis**. ⁴, ⁵ Notably, the **H. pylori dapE deletion mutant** was unable to grow in lysine-supplemented media, implying that lysine cannot be effectively imported. These results strongly suggest that the **mDAP/**lysine biosynthetic pathway is the **only** source for lysine and cannot be compensated by other pathways or import. Lysine is an essential amino acid that is not synthesized in humans; hence, it must be ingested as lysine or lysine-containing proteins. In contrast, most bacteria, plants, and algae can synthesize lysine and **mDAP** from aspartic acid.¹, ², ⁶ Since there are no similar biosynthetic pathways in mammals, including humans, inhibitors that target one or more of the enzymes in the **mDAP/**lysine pathway are hypothesized to exhibit selective toxicity against bacteria.¹

Genes encoding DapE's have been identified in several pathogenic Gram-positive and Gram-negative bacteria such as **Acinetobacter baumannii** (MDRAB), **Mycobacterium tuberculosis**, **Escherichia coli** (O157:H7), **Bordetella**
pertussis, Vibrio cholerae, Rickettsia prowazekii, Pseudomonas aeruginosa, Yersinia pestis, H. pylori, Haemophilus influenzae, Staphylococcus aureus (strain MRSA252), Enterococcus faecium, Salmonella enterica, and Streptococcus pneumoniae.4, 5, 7, 8, 9, 10 The fact that the DapE gene has been discovered in several multi-drug-resistant bacteria suggests that inhibitors of DapE enzymes may provide a new class of broad-spectrum antibiotics. Alignment of the DapE proteins listed above show a minimum of 49% sequence identity.11

Significantly, all DapE proteins characterized to date are medium-sized, dimeric enzymes (41.6 kDa/subunit) that require zinc ions for their activity.6, 12 The amino acid residues that function as metal ligands in the structurally characterized M28 family members, the carboxypeptidase from Pseudomonas sp. strain RS-16 (CPG2) and the leucine aminopeptidase from Aeromonas proteolytica (AAP).6, 13 are fully conserved in all DapE sequences.6, 12

Interestingly, it has been reported that the “as purified” DapE enzyme contains only one tightly bound Zn(II) ion and exhibits ~ 60% of its total activity, similar to AAP.6, 12 Thus, both metal ions seem to be required for full enzymatic activity, but their individual catalytic roles appear to differ markedly.

A major limitation in understanding the catalytic mechanism of DapE and in developing novel inhibitors that specifically can target DapE is the lack of knowledge about their structure and an active-site architecture.6, 18 The only X-ray crystal structure reported for any DapE enzyme is an apo form of the DapE from Neisseria meningitidis.19 The absence of metal ions in the structure makes it difficult to determine the spatial arrangement of the catalytically relevant residues that constitute the active site. While the current lack of structural data preclude definitive assignment of catalytically relevant residues, we have recently reported a three-dimensional homology model of the DapE from H. influenzae, using the crystal structure of the DapE from N. meningitidis as a template that exhibits ~ 54% identity to the DapE from H. influenzae.11 In an effort to clearly define the structure of DapE enzymes along with the catalytically relevant residues that constitute the active site, we have solved the 2.0- and 2.3-Å resolution structures of the mono- and dinuclear zinc DapE enzymes from H. influenzae, respectively. Now that active forms of DapE have been crystallographically characterized, the critical components of the active site including substrate binding can be elucidated.

Protein purification and X-ray structure of the DapE from H. influenzae

The recombinant DapE from H. influenzae was expressed and purified, as previously described, with minor modifications12 from a stock culture kindly provided by Professor John Blanchard.6 In order to obtain ultrapure protein suitable for crystallization, we further purified ~ 50 mg of DapE by loading it onto a Mono-Q column (5/50GL, GE Healthcare, Amersham Biosciences Corp., Piscataway, NJ, USA) that was pre-equilibrated with 10 mM Chelex-100-treated N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine buffer at pH 7.8. A flow rate of 0.5 mL min$^{-1}$ was used with a linear gradient of NaCl (0–0.25 M). Ultrapure DapE eluted as the first peak at ~ 0.15 M NaCl. The pure fractions were concentrated using a Centricon Plus 20 (Millipore Corp., Billerica, MA, USA). Purified DapE from H. influenzae exhibited a single band on SDS-PAGE, indicating an $M_r = 41,500$. Protein concentrations were determined from the absorbance at 280 nm using a molar absorptivity ($\varepsilon_{280} = 36,040$ M$^{-1}$ cm$^{-1}$), calculated using the method developed by Gill and von Hippel.20 The protein concentration determined using this molar absorptivity was in good agreement to that obtained using a Bradford assay. The concentration of DapE samples used for crystallization was ~ 15 mg mL$^{-1}$. Individual aliquots of purified DapE were stored in liquid nitrogen until needed.

DapE's catalyze the hydrolysis of l,l-SDAP, forming l,l-diaminopimelic acid and succinate.6, 12 Initial rates were fit directly to the Michaelis–Menten equation to obtain the catalytic constants $K_m$ and $k_{cat}$. The $k_{cat}$ and $K_m$ values for DapE were determined in triplicate by monitoring amide bond cleavage of l,l-SDAP10, 12, 21 at 225 nm in Chelex-100-treated 50 mM potassium phosphate buffer, pH 7.5, and found to be $140 \pm 10$ s$^{-1}$ and $730 \pm 15$ mM, respectively.12
Crystals of the dizinc form of DapE (ZnZn_DapE) were grown at 16 °C by vapor diffusion in hanging drops containing 1 mL of precipitant solution (1 M ammonium sulfate, 0.2 M NaCl, and 0.1 M Na acetate, pH 4.4) and 1 μL of 13 mg mL⁻¹ of DapE with three equivalents of zinc. The crystals grew within 2 weeks and reached a size of 0.1 mm × 0.1 mm × 0.05 mm. These crystals belonged to the primitive orthorhombic space group $P_{2_1}2_12_1$ with unit cell parameters $a = 44.7$ Å, $b = 95.7$ Å, and $c = 185.4$ Å. The asymmetric unit contains two molecules with a $V_m$ value of 2.4 Å (solvent content, 48%). In order to determine the structure of a monometalated form of DapE, we used a second batch of protein equilibrated with only one equivalent of zinc, and crystals were grown under the same conditions as described above. Crystals of the monozinc form of DapE (Zn_DapE) also grew within 10 days and reached a similar size with an identical morphology as crystals of ZnZn_DapE (space group $P_{2_1}2_12_1$ with unit cell parameters $a = 45.2$ Å, $b = 95.7$ Å, and $c = 181.2$ Å).

Prior to data collection, an X-ray fluorescence spectrum was recorded for both Zn_DapE and ZnZn_DapE, which identified the presence of Zn ions in the protein crystals. Data collection was carried out on the 19-ID beamline of the Structural Biology Center at the Advanced Photon Source according to the procedure described previously. Data to 2.0 and 2.3 Å were collected for Zn_DapE and ZnZn_DapE, respectively, at a wavelength of 0.9794 Å from the single crystals and were processed using HKL2000 (Table 1).

**Table 1. Data and refinement statistics**

<table>
<thead>
<tr>
<th>Data collection statistics</th>
<th>Zn_DapE</th>
<th>ZnZn_DapE</th>
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<tr>
<td>Space group</td>
<td>$P_{2_1}2_12_1$</td>
<td>$P_{2_1}2_12_1$</td>
</tr>
<tr>
<td>Unit cell parameters: $a$, $b$, $c$ (Å)</td>
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<td>44.7, 95.7, 185.4</td>
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<tr>
<td>Wavelength (Å)</td>
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<td>0.9794</td>
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<tr>
<td>Resolution (Å)</td>
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<td>$R_{merge}$ (%)</td>
<td>7.7 (69.2)</td>
<td>8.6 (56.7)</td>
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<tr>
<td>Completeness (%)</td>
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<td>95.9 (99.7)</td>
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<tr>
<td>$I/σ$</td>
<td>20.2 (2.5)</td>
<td>17.8(3.4)</td>
</tr>
</tbody>
</table>

**Phasing**

| Phasing | MR | MR |
| Resolution range (Å) | 30–3.0 | 35–3.0 |
| Molecular replacement statistics CC 0.5 $R_{fac}$ 0.46 | 0.5 | 0.42 |
| $R$-factor | 0.46 | 0.52 |

**Refinement statistics**

| Resolution range (Å) | 40.0–2.03 | 40.0–2.30 |
| $R_{cryst}$ (%)      | 20.2      | 19.4      |
| $R_{free}$ (%)       | 25.4      | 25.0      |
| Number of non-hydrogen atoms | 5718 | 5718 |
| Zn ions/SO₄ | 2/2 | 4/5 |
| Solvent           | 401      | 212      |

**rmsd from target values**

| Bond lengths (Å) | 0.017 | 0.015 |
| Bond angles (°)  | 1.57  | 1.43  |
### Average B-factors (Å²)

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<tr>
<td>Zn1/Zn2</td>
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### Ramachandran Plot (%)

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<td>100</td>
</tr>
<tr>
<td>Allowed</td>
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<td>0</td>
</tr>
<tr>
<td>Outliers</td>
<td>3ISZ</td>
<td>3IC1</td>
</tr>
</tbody>
</table>

The values in parentheses are for the highest-resolution shell.

1. $R_{merge} = \sum_h \sum_i |I_{h,i} - \langle I_{h,i} \rangle| / \sum_h \sum_i I_{h,i}$.
2. As defined by MolProbity.

The structure of Zn_DapE was determined by molecular replacement using the crystal structure of the DapE from *N. meningitidis* as a template [Protein Data Bank (PDB) ID: 1vgy], which exhibits ~ 54% identity with the DapE from *H. influenzae*, as described previously.11 Molecular replacement searches were completed with MOLREP (R-factor of 0.46%, correlation coefficient of 0.50) using the CCP4 suite.24, 25 Several rounds of rebuilding and readjusting using Coot and ARP/wARP were required to improve the initial model.26, 27 The final model was refined against all reflections using the program REFMAC 5.528 in the resolution range 30 to 2.0 Å except for 5% of the reflections, which were randomly selected and used to monitor $R_{free}$. The final refinement statistics are presented in Table 1.

The structure of ZnZn_DapE was determined by molecular replacement with MOLREP (R-factor of 0.43%, correlation coefficient of 0.52) using the catalytic domain of Zn_DapE in the first round and then the dimerization domain of Zn_DapE. Several cycles of model building and water picking with Coot followed by refinement with REFMAC 5.5 were undertaken to build a final structural model (Table 1). Analysis and validation of the structures were performed with the aid of MolProbity, Coot validation tools, and SSM and DALI servers.29, 30, 31 Figures were prepared using the program PyMOL.32

The crystal structures of Zn_DapE and ZnZn_DapE from *H. influenzae* were determined at 2.0 and 2.3 Å resolution, respectively. The final model for both enzymes includes two monomers in the asymmetric unit forming a dimer. The model of Zn_DapE contains 735 residues out of a possible 752 residues, 414 water molecules, 2 zinc ions, and 2 sulfate ions. The model of ZnZn_DapE contains 749 residues out of 752 residues, 4 zinc ions, 212 water molecules, 2 glycerol molecules, and 5 sulfate ions. The electron density maps obtained for both structures were of high quality and allowed reliable modeling of both monomers, except in the loop regions (residues 192–198 in chain A and 193–196 and 241–245 in chain B), which are disordered and were not included in the final model. The final models exhibited good crystallographic and geometric statistics and were refined against 2.0 Å data for Zn_DapE with a final $R_{work}$ of 20.2 and an $R_{free}$ of 25.4 and against 2.3 Å data for ZnZn_DapE to an $R_{work}/R_{free}$ of 19.4/25.0 (Table 1).

The DapE enzyme from *H. influenzae* is organized as a dimer and closely resembles the structure of the DapE from *N. meningitidis*. The search for structural homologues of DapE using the DALI and SSM programs30, 31 identified several closely related homologues with the apo form of DapE form *N. meningitidis* (PDB ID: 1vgy, Z-score = 47.8, rmsd = 2.031,19 acetylornithine deacetylase from *Bacteroides thetaiotaomicron* VPI-5482 (PDB ID: 3ct9, Z-score = 32.8, rmsd = 2.731),33 and the CPG2 from *Pseudomonas* sp. strain RS-16 (PDB ID: 1cg2, Z-score = 31.6, rmsd = 3.031)34 being the top three hits. Each subunit of the dimer consists of two functional domains: a large catalytic domain, which supplies the ligands for the zinc ions in the active site, and a smaller dimerization domain that contributes to dimer formation (Fig. 1). The domains are connected by a small hinge
region (residues 176–179 and 298–293) allowing movement of the dimerization domain with respect to the catalytic domain. The shape of the DapE dimer resembles a rotary style phone receiver with the larger catalytic domain being placed on the periphery of the dimer and separated by \( \sim 47 \) Å. The subunit topology is illustrated in Fig. 1a.

Fig. 1. Structure of DapE from *H. influenzae*. (a) Ribbon diagram showing the overall structure of the DapE dimer, with monomers shown in different colors (red and blue, and red and green). Individual domains are labeled and the secondary-structure elements are colored (blue and green, \( \beta \)-strand; red, \( \alpha \)-helices). (b) Diagram of the DapE monomer. The \( \alpha \)-helices are represented as cylinders, zinc ions are labeled in black (Zn1, the catalytic zinc) and gray (Zn2), and orange dots indicate a disordered loop that has not been modeled.

The dimerization domain of DapE consists of a 114-residue insertion between strand \( \beta 7 \) and helix \( \alpha 7 \) of the catalytic domain (Fig. 1) and comprises residues 180–292. The insertion folds into a two-layer \( \alpha + \beta \) sandwich fold (\( \beta 8 \), \( \alpha 5 \), \( \beta 9 \), \( \beta 10 \), and \( \alpha 6 \)). One layer is formed by the four \( \beta \)-strands arranged into an antiparallel \( \beta \)-sheet with an \( \sim 45^\circ \) twist across its length, while two helices sit on one site of the molecule forming a second layer. The secondary elements are organized into two \( \beta \alpha \beta \) motifs revealing a ferredoxin-like fold.

Active site of the DapE from *H. influenzae*

The catalytic domain is composed of residues 1–179 and 293–376. The core of the catalytic domain consists of an eight-stranded twisted \( \beta \)-sheet that is sandwiched between seven \( \alpha \)-helices. The mixed \( \beta \)-sheet is formed by two small antiparallel \( \beta \)-strands (\( \beta 1 \) and \( \beta 2 \)) followed by four centrally located parallel strands (\( \beta 5 \), \( \beta 3 \), \( \beta 6 \), and \( \beta 13 \)) as well as smaller parallel strands (\( \beta 7 \) and \( \beta 12 \)). The last two strands are rotated 180° and are facing in an opposite direction from the centrally located strands. The connecting helices are located below (\( \alpha 1 \), \( \alpha 2 \), \( \alpha 3 \), \( \alpha 9 \), and \( \alpha 7 \)) and above (\( \alpha 4 \) and \( \alpha 8 \)) the plane of the \( \beta \)-sheet. In addition to the central sheet (\( \beta 1 \), \( \beta 2 \), \( \beta 5 \), \( \beta 3 \), \( \beta 6 \), \( \beta 13 \), \( \beta 7 \), and \( \beta 12 \)), there is also a second small \( \beta \)-sheet located on the surface of the molecule. This \( \beta \)-sheet consists of two shorter antiparallel \( \beta \)-strands (\( \beta 4 \) and \( \beta 14 \)) (Fig. 1b). The zinc ion or ions are located near the C-terminal end of the catalytic domains at the surface of the protein (Fig. 1a and b).

The active-site cleft is located in the center of the catalytic domain above the centrally located parallel strands of the \( \beta \)-sheet (\( \beta 3 \), \( \beta 6 \), and \( \beta 13 \)) and is covered by the loops. The location and the architecture of the active site in ZnZn_DapE are strikingly similar to the dinuclear active sites of CPG\(_2\) and AAP (Fig. 2). The two zinc ions in ZnZn_DapE are 3.36 Å apart compared with 3.45 Å for AAP and 3.25 Å for CPG\(_2\). Like AAP and CPG\(_2\), each
of the zinc ions in ZnZn_DapE adopts a distorted tetrahedral or TBP geometry with the N\(^\varepsilon\) nitrogen of His67 for Zn1 and H349 for Zn2 along with a bridging water/hydroxide oxygen atom making up the axial positions of a potential TBP geometry. Identical with CPG\(_2\) and AAP, each zinc ion is coordinated by one imidazole group (H67 for Zn1 and H349 for Zn2) and the carboxylate oxygens of E163 (OE1 and the dangling oxygen OE2) for Zn1 and E135 for Zn2 (OE1 and the dangling oxygen OE2). Both zinc ions are bridged by D100 and a water/hydroxide. Interestingly, in the ZnZn_DapE structure, the Zn2 binding site exhibits only \(\sim 50\%\) occupancy. This structure confirms that the zinc ions in DapE form a \((\mu\text{-aquo})(\mu\text{-carboxylato})\text{dizinc(II)}\) core similar to AAP and CPG\(_2\) and are consistent with EXAFS data.\(^{16}\)

![Fig. 2. Close-up view showing the active site of ZnZn_DapE (a) and Zn_DapE (b) with 2Fo – Fc electron density map (blue) contoured at 1.0σ, zinc anomalous difference Fourier maps contoured at 4σ (magenta), and drawings of the active sites with distances displayed in angstroms. Zinc ions are labeled in black (Zn1, the catalytic zinc) and gray (Zn2). (c) Active site of carboxypeptidase G2 (PDB ID: 1cg2) and (d) close-up view of the active site of a bacterial leucyl aminopeptidase (PDB ID: 2DEA).]

Similar to AAP, it was shown that the as-purified DapE enzyme contains only one tightly bound zinc ion and exhibits \(\sim 60\%\) of its total activity in the presence of one zinc ion.\(^{12, 37}\) Because DapE’s can be activated with one zinc ion, it has been hypothesized that one divalent metal binding site is likely filled preferentially. This begs the following question: which metal ion binding site is filled first in the active site and which site is critical for
catalysis? Therefore, we attempted to obtain structural data for a mono-zinc form of DapE (Zn\textsubscript{DapE}) by limiting the amount of divalent metal ions present during the crystallization process (Fig. 2). The Zn(II) ion in Zn\textsubscript{DapE} resides in a distorted tetrahedral or TBP geometry with the N\text{ɛ} nitrogen of His\textsubscript{67} (2.1 Å) and the bridging water/hydroxide oxygen atom (2.1 Å) making up the axial positions of a potential TBP geometry. The remaining ligands are coordinated by the carboxylate oxygens of D100 (OD1, 2.0 Å) and E163 (OE, 2.0 Å, and the dangling oxygen OE\textsubscript{2}, 2.6 Å). Therefore, this structure confirms that the H67 site (Zn\textsubscript{1}) of the DapE active site is the high-affinity site and is occupied first and, hence, corresponds to the catalytic zinc. This observation is also consistent with ZnZn\textsubscript{DapE} structural data where we observed the same site being fully occupied and the second site being only partly occupied.

The structure of the monometalated form of DapE is only the second monometalated structure for any M28 family metalloprotease\textsuperscript{38} and is reminiscent of the monometalated structures of M24 family metalloproteases, namely, the aminopeptidase P from \textit{E. coli} and the methionine aminopeptidase from \textit{E. coli}.\textsuperscript{39, 40} Given that the vast majority of proteases that can bind two metal ions are active in the presence of one metal ion, the X-ray structure of the Zn\textsubscript{DapE} enzyme reported herein provides a structural model for the monometalated forms of these enzymes.

The structures of Zn\textsubscript{DapE} and ZnZn\textsubscript{DapE} along with the previously reported divalent metal binding studies\textsuperscript{12} provide insight into the observed metal binding properties of DapE as well as all M28 metallopeptidases.\textsuperscript{12} Based on our X-ray crystallographic data, metal binding to DapE occurs in a sequential fashion with the first zinc binding site (Zn\textsubscript{1}) bound to the H67 site of the active site and the second zinc ion binding to H349 (Zn\textsubscript{2}). EPR and electronic absorption data on the Co(II)-substituted forms of WT and mutant DapE enzymes also indicate very clearly that the first metal ion to bind to DapE resides in the H67 site of the active site whereas the second metal binding site corresponds to the H349 site of the active site,\textsuperscript{11, 12} consistent with our X-ray crystallographic data. These data are significant because substrate and also inhibitor zinc-binding groups have been hypothesized to bind to Zn\textsubscript{1}.\textsuperscript{11} Also of importance is that these data highlight the potential formation of heterodimetallic sites in DapE similar to AAP.\textsuperscript{41} The H67 residue in DapE corresponds to H97 in AAP while H349 corresponds to H256 in AAP. Based on the crystal structure of the butane boronic acid inhibited form of AAP,\textsuperscript{42} H97 (H67 in DapE) was proposed to function as a ligand in the second metal binding site. This apparent reversal of the zinc ion binding sites in DapE versus AAP may likely indicate that the catalytic roles of the active site zinc ions in DapE's are switched from that proposed for AAP. The latter suggestion is not necessarily surprising since AAP cleaves from the N-terminus while DapE cleaves the equivalent of a C-terminal carboxylate group.

Finally, several potentially important hydrogen-bonding interactions also exist in the active site of both Zn\textsubscript{DapE} and ZnZn\textsubscript{DapE}. Perhaps most notably, there is at least one interaction between the metal-bound water/hydroxide molecule and the carboxylate oxygen atoms of Glu\textsubscript{134}. Glu\textsubscript{134} was recently shown to be essential for catalysis and functions as a general acid/base.\textsuperscript{17} Glu\textsubscript{134} forms a strong hydrogen bond (\textasciitilde 2.8 Å) with the terminal metal-bound water/hydroxide in Zn\textsubscript{DapE} and also with the bridging water/hydroxide (\textasciitilde 2.9 Å) in ZnZn\textsubscript{DapE}. Other potentially important hydrogen-bonding interactions occur between the N\textsuperscript{5} proton of His\textsubscript{67} with a side-chain oxygen of Asp\textsubscript{69} forming an Asp–His–Zn triad that has been postulated to decrease the Lewis acidity of zinc ions\textsuperscript{43} and may further assist in facilitating the coordination of a double-bonded oxygen to Zn\textsubscript{2}. Similar arrangement and a fully conserved Asp–His–Zn triad are observed for both AAP and CPG\textsubscript{2} and are postulated to regulate the Lewis acidity of the zinc ion.\textsuperscript{36}

**Inhibitor design implications for DapE enzymes**

Inspection of the Zn\textsubscript{DapE} and ZnZn\textsubscript{DapE} structures, combined with surface analysis, reveals a smile-shaped cavity that extends along the catalytic domain and surrounds the active site (Fig. 3). This well-defined and negatively charged cavity is shaped from the top by strand \beta\textsubscript{12} and \alpha\textsubscript{8} and in the middle by the loop connecting these two elements. The bottom of the cavity is formed by loops (loop connecting \beta\textsubscript{strands} \beta\textsubscript{6} and \beta\textsubscript{7} and loop connecting \beta\textsubscript{5} and \alpha\textsubscript{4}) and \alpha\textsubscript{4}. Taking into account the linear character of the substrate, it is likely that the
substrate binds in an extended conformation along the smile-shaped groove with the peptide bond positioned over the active-site metal ions while the rest of the substrate could be further stabilized by the interaction of carboxyl groups of the substrate with the surrounding residues (Fig. 4). Assuming that the carbonyl moiety of the peptide bond binds to one or both metal ions in the active site, two residues, E134 and T325, likely form hydrogen bonds with the carbonyl group and/or the backbone amine group of l,l-SDAP. E134 had been shown to function as the general acid/base during catalysis, suggesting that T325, which is centrally positioned on the loop overhanging the active site, may play an important role in substrate recognition and transition-state stabilization. It is easy to envision this loop undergoing movements upon substrate binding.

Fig. 3. (a) Surface rendering of the ZnZn_DapE molecule (molecule A is shown) showing the charge distribution and depicting the smile-shaped active-site cavity. The surface charge distribution was determined using PyMOL. (b) Close-up view of the active-site cavity of ZnZn_DapE, molecule A, showing charged residues that may interact with the substrate. The zinc ions are represented as gray and black (catalytic zinc) spheres. Side chains of residues that may be involved in recognizing and binding the substrate are shown as orange sticks. Figures were prepared using PyMOL.
Fig. 4. Proposed catalytic reaction mechanisms of (a) monozinc DapE and (b) dizinc DapE.

DapE enzymes have been shown to exhibit strict specificity for the l,l-isoform of SDAP, indicating that the DapE active site is highly specific in both functional group recognition as well as stereochemistry. This specificity is built into the substrate binding pocket, which will minimize the nonspecific interactions of DapE with nonproductive inhibitors. Therefore, it is quite likely that substrate binding is further controlled by interactions of the substrate carboxylic groups with positively charged amino acid side chains. The potential positively charged residues that could interact with the carboxylic groups of the substrate are K175, R178, and N345 at one end and R258, R329, and K139 at the opposite end. The K175 side chain is within ~6 Å of the center of the active site; hence, it is easy to envision the ε-amino group of the lysine side chain within the appropriate distance to form a salt bridge interaction with a substrate carboxylate group and, thus, be involved in recognizing and binding substrate. On the opposite side, carboxylate groups of l,l-SDAP could also be further stabilized by an interaction with R258 and/or R329. In one of the monomers of the ZnZn-DapE structure, both of these residues form a charged dipole interaction with a sulfate ion, a possible mimic of the carboxylic group of
the substrate. R329 is centrally positioned in a positively charged pocket that it forms together with R258 and is further complimented by the N-terminus of α8. These data, in combination with previously reported inhibitor binding studies, indicate that DapE represents an excellent target for a highly specific drug that should have high efficacy and low toxicity.

Mechanistic implications
The X-ray structures of Zn_DapE and ZnZn_DapE presented herein provide a structural foundation for the corroboration of the proposed reaction mechanism of DapE. Analysis of structures and previously reported kinetic and spectroscopic studies on DapE enzymes allows us to propose a refined mechanism of catalysis for DapE's. Based on the recently proposed mechanism for AAP, the first step in catalysis for DapE's is likely recognition of the L,l-SDAP side chain by the smile-shaped cavity adjacent to the Zn1 site. Next, the peptide carbonyl oxygen of L,l-SDAP coordinates to Zn1, expanding its coordination number from four to five, activating it for nucleophilic attack. Deprotonation of the metal-bound water molecule by E134 to form a nucleophilic hydroxide moiety is consistent with the postulated pK₆ of the zinc-bound water molecule. Once the zinc-bound hydroxide is formed, it can attack the activated carbonyl carbon of the substrate, forming an η₁-μ₁-transition-state complex. Solvent kinetic isotope effect studies yielded an inverse isotope effect that was explained by the attack of a zinc-bound hydroxide on the amide carbonyl. E134 may provide a proton to the penultimate amino nitrogen, similar to that observed for AAP, returning it to its ionized state, thus facilitating product release. Once the products are released, a water molecule bridging the two metal ions is replaced. In the absence of a second metal ion, the catalytic mechanism does not likely change markedly as H349 is in position to assist in orienting the substrate properly in the active site through the formation of a hydrogen bond with a carboxylate side chain of the substrate, thereby stabilizing the transition-state intermediate reminiscent to proposals for the monometalated forms of AAP and the methionine aminopeptidase from E. coli. In the presence of a dinuclear site, the second metal ion likely coordinates either the peptide carbonyl oxygen in a bridging fashion or a carboxylate side chain of the substrate.

PDB accession numbers
The atomic coordinates and structure factor files for the structures of Zn_DapE and ZnZn_DapE from H. influenzae have been deposited in the Research Collaboratory for Structural Bioinformatics PDB with accession codes 3ISZ and 3IC1, respectively.

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**Notes**

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