

Specificity and Mechanism of *Acinetobacter baumannii* Nicotinamidase: Implications for Activation of the Front-Line Tuberculosis Drug Pyrazinamide**

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Nicotinamidase (EC 3.5.1.19) catalyzes hydrolysis of nicotinamide to nicotinic acid and ammonia, an important reaction in the NAD⁺ salvage pathway.^[1] This activity has a fortuitous medical benefit since the *Mycobacterium tuberculosis* enzyme converts the nicotinamide analogue prodrug pyrazinamide into the bacteriostatic pyrazinoic acid,^[2–4] hence the alternative name, pyrazinamidase (PncA). Pyrazinoic acid inhibits *M. tuberculosis* type I fatty acid synthase,^[5] represses mycolic acid biosynthesis, and appears to affect membrane energetics and acidification of the cytoplasm.^[4] It is active against semi-dormant tubercle *bacilli* and with rifampicin and isoniazid, forms the front-line tuberculosis treatment.^[2,3] Though studies of PncA have revealed aspects of its structure and biochemical activity^[6–9] there are no structural data on how the enzyme binds and processes physiological ligands. High-resolution crystal structures of *Acinetobacter baumannii* PncA (*AbPncA*) complexed with nicotinic acid and pyrazinoic acid now provide direct evidence for the interactions that govern the specificity and mechanism, and of how a valued anti-bacterial agent is activated.

Recombinant *AbPncA* was prepared, the dimeric, colorless enzyme was purified in high yield and its kinetic properties determined. With pyrazinamide as the substrate the following values were obtained; $K_M = 106.9 \mu\text{M}$, $V_{\text{max}} = 62.8 \text{ nmol min}^{-1}$, $k_{\text{cat}} = 3.1 \text{ min}^{-1}$, specific activity $132 \mu\text{M min}^{-1} \text{ mg}^{-1}$. These values are comparable to literature values, for example, the specific activity of *M. tuberculosis* PncA (*MtPncA*) with pyrazinamide is $82 \mu\text{M min}^{-1} \text{ mg}^{-1}$.^[8]

Two crystal forms (I and II) were obtained with nicotinic and pyrazinoic acid, respectively, and the structures determined. PncA is a divalent cation-dependent enzyme and activity has been reported with Fe²⁺, Mn²⁺,^[8] and Zn²⁺ ions.^[6] As expected, metal ions were observed in the structures. Inductively coupled plasma-atomic emission spectrometry (ICP-OES) identified that recombinant *AbPncA* contained

Fe²⁺ and Zn²⁺ ions in an approximate 1:1 ratio with a trace of Mn²⁺ present. However, anomalous dispersion measurements are consistent with a higher occupancy of Zn²⁺ at the active site and the crystallographic models contain that cation. We refer to Zn²⁺ ions in discussion but judge it likely that *AbPncA* functions in the presence of different divalent cations. (Experimental details, including enzyme activity and metal-ion identification, together with sequence alignments, and additional figures are given as Supporting Information Figure S1–S8).

Crystals were obtained in the presence of cacodylate buffer and form II shows dimethylarsinoyl-modified Cys159 in the active site, an artifact of crystallization (Supporting Information, Figure S1, S2). The steric hindrance of this modification precludes full occupancy of pyrazinoic acid such that the final refinement was performed with occupancy 0.8 for pyrazinoic acid, 0.2 for the modified cysteine. Crystal form I has two molecules, form II a single molecule in the asymmetric unit, respectively, with a root-mean-square deviation (r.m.s.d.) derived from least-squares fit of C α atoms of these three molecules of 0.2 Å. The structures and the interactions formed by ligands within the active sites are essentially identical and we concentrate on form I, a 1.65 Å resolution structure with full occupancy ligand (Supporting Information, Figure S3). About 60% of residues form elements of secondary structure, these are eight α -helices and nine β -strands (Figure 1, and Supporting Information, Figure S4). The core of the subunit is a parallel β -sheet of strands 1, 2, 5–9. Three helices ($\alpha 5$, $\alpha 6$, $\alpha 7$) lie on one side of the sheet, with $\alpha 2$ placed against the other. A subdomain is placed at

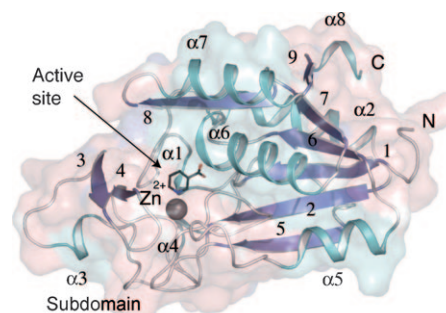


Figure 1. Ribbon diagram of the *AbPncA* monomer and location of the active site. The Zn²⁺ ion is shown as a gray sphere, nicotinic acid as a stick model: black C, blue N, red O. Helices (cyan) are labeled and β -strands (blue) numbered. The terminal positions of the polypeptide are labeled N and C. Molecular images prepared with PyMol.^[14]

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one end of the β -sheet and includes $\beta 3$ and $\beta 4$ and a single turn of helix, $\alpha 3$.

Gel filtration and analytical ultracentrifugation indicate that *AbPncA* is a stable dimer in solution, approximate mass 47 kDa. The asymmetric unit of form I is a dimer (Supporting Information, Figure S5), with an interface of 900 \AA^2 , which is about 10% of the surface area of a subunit. In form II the crystallographic twofold axis generates the same dimer. The interactions that stabilize the dimer mainly involve residues on $\alpha 5$ and $\beta 5$. The *AbPncA* subunit resembles orthologues from *Pyrococcus horikoshii* (*PhPncA*) and *Saccharomyces cerevisiae* (*ScPncA*).^[6,7] Superpositions give r.m.s.d. values of 1.2 \AA for the overlay of an *AbPncA* subunit on either *PhPncA* (44% sequence identity, 167 $\text{C}\alpha$ atoms) or *ScPncA* (33% identity, 191 $\text{C}\alpha$ atoms).

The *AbPncA* active site is between the core and the subdomain (Figure 1), and is formed by residues on $\beta 1$, $\beta 2$, the $\beta 4$ - $\alpha 4$ loop, $\beta 5$, and the $\beta 6$ - $\alpha 6$ turn. It is buried and completely occluded from solvent by four polypeptide segments; the loops linking $\alpha 3$ - $\beta 3$, $\beta 5$ - $\alpha 5$, and $\beta 8$ - $\alpha 9$ together with strand $\beta 4$. Numerous hydrophobic residues (discussed below) surround the active site and a gross conformational change is probably required to permit substrate binding or release of products.

The active site Cys159 is located at the N-terminus of $\alpha 6$ at one side of the active site with the Zn^{2+} ion positioned on the other side. The metal ion is held tightly in the *AbPncA* active site as shown by activity being retained in the presence of 10 mM EDTA (data not shown). Octahedral coordination of Zn^{2+} ion involves Asp54 OD2, His56 NE2, and His89 NE2, two water molecules and nicotinic acid N5 (Figure 2). Cation-ligand distances range from 2.11 to 2.28 \AA , consistent with data on Zn^{2+} ligand geometry.^[10] A network of hydrogen bonds position the coordinating groups. One water molecule forms hydrogen bonds with Ser62 OG and Asp121 OD2, the other with Asp54 OD1 and Asp121 OD1. The coordinating

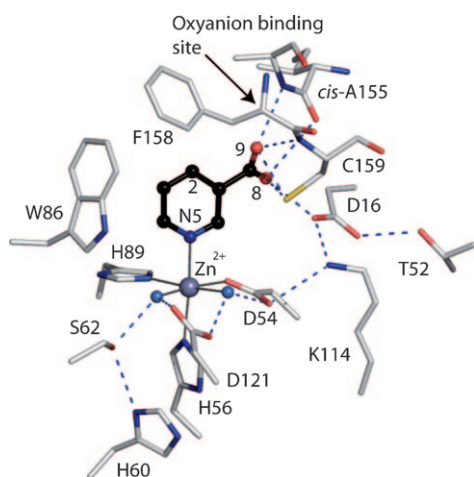


Figure 2. Active site of the nicotinic acid complex. The Zn^{2+} ion is a blue-gray sphere, thin lines mark coordination to amino acid side chains and two water molecules (marine spheres). Amino acids are in stick representation: C gray, N blue, O red, S yellow. Nicotinic acid is shown as a ball-and-stick model: C black, N blue, O red. Dashed lines represent potential hydrogen bonds.

His56 and His89 residues donate hydrogen bonds from ND1 to carbonyl groups of Gly115 and Pro87, respectively (data not shown). Nicotinic acid is tethered to the cation and positioned between five hydrophobic residues; Phe21, Leu27, Trp86, Tyr123, and Cys159 (Supporting Information, Figure S6). Trp86 NE1 and Tyr123 OH form a hydrogen bond to hold these residues in place over the ligand. A further six residues (Val29, Ile154, Ala155, Phe158, Ile184, and Leu186) stabilize the hydrophobic environment around the nicotinic acid and occlude the active site (Supporting Information, Figure S6). In the absence of a ligand, the Zn^{2+} coordination sphere is completed with a water as indicated in structures of *PhPncA* and *ScPncA*.^[6,7] Nicotinic acid O8 and O9 are 2.5 and 2.6 \AA , respectively, from Cys159 SG suggesting the presence of a bifurcated hydrogen bond. The next nearest functional group to Cys159 SG is Asp16 OD2 at a distance of 3.4 \AA . O9 accepts hydrogen bonds donated from main-chain amides of *cis*-Ala155 and Cys159. Interaction with the *cis*-Ala155 carbonyl group suggests that O8 is a hydroxy group and that protonation of Asp16 OD2 may facilitate a second hydrogen bond (Supporting Information, Figure S7). Alternatively, O8 as hydroxy group or if protonated may participate in a bifurcated hydrogen bond with Asp16 and Ala155. Asp16, together with Asp54 and Lys114 form a cluster of interacting hydrophilic residues on one side of the ligand-binding site. Lys114 NZ donates hydrogen bonds to Asp16 OD1, Asp54 OD2, (Figure 2) and the main-chain carbonyl group of Tyr123 (not shown). The close proximity of Lys114 to Asp16 is likely to influence the pK_a value. The Asp16 carboxylate and main-chain amide groups form hydrogen bonds with Thr52 OG1 (Supporting Information, Figure S7), a pairing strictly conserved in *PncA* (Supporting Information, Figure S8).

Amidation involves either acidic or basic hydrolysis and the structures of the enzyme-product complex suggest that basic hydrolysis applies in *PncA*. Acid hydrolysis would involve nucleophilic water attacking the carbon of a protonated amide. The hydrophobic environment on one side of the amide and close interactions with functional groups on the other renders it difficult to envisage how water could be placed to attack C7. Nicotinic acid O9 accepts hydrogen bonds from two amides so protonation at O9 would destabilize the complex due to the proximity of the amides.

A more likely mechanism is indicated by a strictly conserved and essential Cys159,^[6,8] on the polar side of the active site, ideally placed to attack the carbonyl carbon atom in a manner similar to that proposed for other enzymes, for example trypanothione synthetase amidase^[11] and the nitrilase enzyme superfamily.^[12] Nitrilases exploit a catalytic triad consisting of a reactive cysteine, a lysine, and a glutamate. The triad of *PncA* has a conservative difference with aspartate replacing glutamate.

We propose a four-stage mechanism (Figure 3). In stage I, substrate binds in the axial position displacing a water from the Zn^{2+} coordination sphere. The waters coordinating to equatorial sites of the metal ion are also held in position by hydrogen-bonding interactions to the enzyme (Figure 2), whereas the axial water lacks such a restraint. It is this water that vacates the coordination sphere as substrate binds.

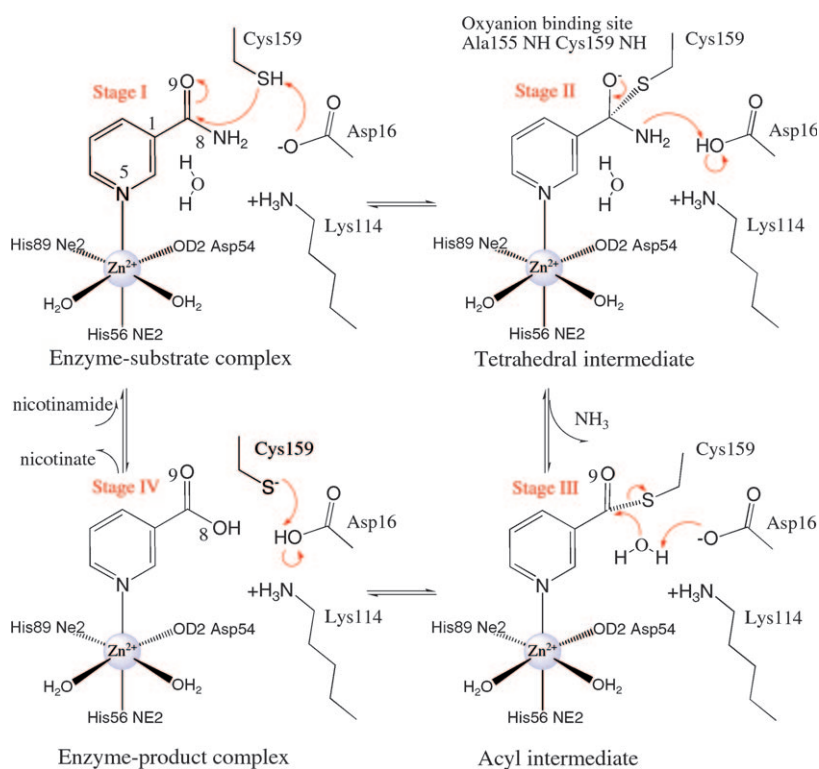


Figure 3. Proposed mechanism of nicotinamidase. see text for details.

This water may be prevented from exiting the active site by the hydrophobic lid covering the active site (Supporting Information, Figure S7). The displaced water, or an incoming water would preferentially bind to the hydrophilic side of the active site near Lys114. Proton abstraction by Asp16 would generate a Cys159 thiolate facilitating nucleophilic attack at C7. Two main-chain amides (*cis*-Ala155 and Cys159) form an oxyanion hole to support thiolate attack by stabilizing the resulting tetrahedral intermediate in a similar fashion to cysteine proteases.^[13] Proton donation from Asp16 to N8 of the substrate would promote C–N bond cleavage and release of NH₃ as the tetrahedral intermediate collapses in stage II to give an acyl intermediate. Water activation by Asp16, whereby the amino acid is protonated, generates a nucleophilic hydroxy group to attack the acyl intermediate producing nicotinic acid (or pyrazinoic acid) and a thiolate in stage III. In stage IV, products are released and the thiolate accepts a proton from Asp16 to regenerate a thiol.

A previously proposed mechanism invoked a Zn²⁺-coordinated hydroxy and an incoming water participating in catalysis.^[6] First a hydroxy group directly attacks the acyl intermediate to replace the amino group, and then another incoming water coordinates to the Zn²⁺ ion as a hydroxy with a proton removed by the nearby aspartate to enable nucleophilic attack on the acyl intermediate. This mechanism appears unduly complicated with two rounds of metal-ion-directed water activation and is incompatible with our new structures. Activation of water to hydroxy groups by the strong Lewis acid that is four-coordinate Zn²⁺ ion is a commonly invoked feature of zinc-dependent enzymes. In PncA, the metal ion is six-coordinate and this would reduce

the Lewis acid strength. In addition, as PncA catalysis is supported by different divalent cations this suggests that Lewis acid strength is less important than the structural role provided by octahedral coordination, precise placement of the substrate and an appropriate ligand exchange rate to support substrate binding with release of a water molecule. We therefore propose a simpler mechanism (Figure 3) with no requirement for the metal ion to activate water.

Recombinant *MtPncA* produced in *Escherichia coli* is reported to carry a mixture of Fe²⁺ and Mn²⁺ ions and reconstitution of apoenzyme with these ions resulted in almost full recovery of activity, surprisingly reconstitution with Zn²⁺ did not.^[8] Moreover, mutation of *MtPncA* His56 to alanine led to almost complete loss of metal-ion binding and enzyme activity. These observations were taken to imply that His56 was directly involved in metal-ion binding and that *MtPncA* binds ions in a different manner from *PhPncA*.^[8] *MtPncA* His56 corresponds to *AbPncA* His60, a residue that contributes significantly to the formation of the cation binding site by forming a hydrogen bond to Ser62 OG and positioning the serine to stabilize one of the waters that coordinates Zn²⁺ ion (Figure 2).

His60 also stabilizes the coordinating His56 by van der Waals interactions and this histidine pair is strictly conserved in PncA sequences. Indeed, 16 of the 20 amino acids involved in cation coordination, substrate recognition, and catalysis are strictly conserved between *AbPncA* and *MtPncA*, with a further three involving conservative substitutions (Supporting Information, Figure S8). This suggests that all PncA enzymes share metal-ion binding properties and mechanism of action. Structural data on recombinant enzyme and metal-ion analysis of native *MtPncA* would further clarify this.

Over 60 *M. tuberculosis* strains display pyrazinamide resistance owing to *pncA* gene mutations and more than 50% of these localize changes to three short polypeptide sections.^[4,8] In *AbPncA* these correspond to β1–α1, α3–β3 segments, and α6, near the active site and with residues important for substrate binding and catalysis. Of note is the strictly conserved Lys114 residue. Pyrazinamide-resistant mutants of *M. tuberculosis* containing a lysine–threonine change in this position have been identified.^[9] This change may disrupt the hydrophilic environment required to allow for the exploitation of water in the proposed mechanism and/or would fail to align Asp16 for catalysis.

In summary, the precise nature of how nicotinic acid, and pyrazinoic acid bind *AbPncA* casts doubt on a previously proposed mechanism, which was based on modeling substrate in a manner that did not include direct interaction with the Zn²⁺ ion. Our data reveal that substrate recognition involves interaction between the Zn²⁺ ion and the pyridyl nitrogen atom and that the position of functional groups has allowed us to propose a new mechanism for nicotinamidase activity and

one which likely applies to the activation of the anti-tuberculosis drug pyrazinamide.

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