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Crystal structures of the transpeptidase domain of the *Mycobacterium tuberculosis* penicillin-binding protein PonA1 reveal potential mechanisms of antibiotic resistance

Ekaterina V. Filippova¹,²,³, Karen J. Kieser⁴, Chi-Hao Luan²,⁵, Zdzislaw Wawrzak⁶, Olga Kiryukhina¹,²,³, Eric J. Rubin⁴,⁷ and Wayne F. Anderson¹,²,³

1 Department of Biochemistry and Molecular Genetics, Feinberg School of Medicine, Northwestern University, Chicago, IL, USA
2 Midwest Center for Structural Genomics (MCSG), Biosciences Division, Argonne National Laboratory, Argonne, IL, USA
3 Center for Structural Genomics of Infectious Diseases (CSGID), Feinberg School of Medicine, Northwestern University, Chicago, IL, USA
4 Department of Immunology and Infectious Disease, Harvard T.H. Chan School of Public Health, Boston, MA, USA
5 High Throughput Analysis Laboratory and Department of Molecular Biosciences, Northwestern University, Evanston, IL, USA
6 Life Science Collaborative Access Team, Synchrotron Research Center, Northwestern University, Evanston, IL, USA
7 Department of Microbiology and Immunobiology, Harvard Medical School, Boston, MA, USA

Keywords
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Correspondence
W. F. Anderson, Department of Biochemistry and Molecular Genetics, Feinberg School of Medicine, Northwestern University, Chicago, IL 60611, USA
Fax: +312 503 5349
Tel: +312 503 1697
E-mail: wf-anderson@northwestern.edu

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*Mycobacterium tuberculosis* is a human respiratory pathogen that causes the deadly disease tuberculosis. The rapid global spread of antibiotic-resistant *M. tuberculosis* makes tuberculosis infections difficult to treat. To overcome this problem new effective antimicrobial strategies are urgently needed. One promising target for new therapeutic approaches is PonA1, a class A penicillin-binding protein, which is required for maintaining physiological cell wall synthesis and cell shape during growth in mycobacteria. Here, crystal structures of the transpeptidase domain, the enzymatic domain responsible for penicillin binding, of PonA1 from *M. tuberculosis* in the inhibitor-free form and in complex with penicillin V are reported. We used site-directed mutagenesis, antibiotic profiling experiments, and fluorescence thermal shift assays to measure PonA1’s sensitivity to different classes of β-lactams. Structural comparison of the PonA1 apo-form and the antibiotic-bound form shows that binding of penicillin V induces conformational changes in the position of the loop β4-α3 surrounding the penicillin-binding site. We have also found that binding of different antibiotics including penicillin V positively impacts protein stability, while other tested β-lactams such as clavulanate or meropenem resulted in destabilization of PonA1. Our antibiotic profiling experiments indicate that the transpeptidase activity of PonA1 in both *M. tuberculosis* and *M. smegmatis* mediates tolerance to specific cell wall-targeting antibiotics, particularly to penicillin V and meropenem. Because *M. tuberculosis* is an important human pathogen, these structural data provide a template to design novel transpeptidase inhibitors to treat tuberculosis infections.

Database
Structural data are available in the PDB database under the accession numbers 5CRF and 5CXW.

Abbreviations
FTS, fluorescence thermal shift; PBP, penicillin-binding protein; TB, tuberculosis; TG-, transglycosylase domain mutant; TP-, transpeptidase domain mutant.
**Introduction**

*Mycobacterium tuberculosis* is a bacterial pathogen of the human respiratory system that primarily infects lungs but could also infect other parts of the body including kidney, spine, and brain. Tuberculosis (TB) infection is fatal for ~1.5 million people worldwide each year according to the World Health Organization and remains silent within 90% of the infected population [1]. Emergence of antibiotic-resistant *M. tuberculosis* bacteria has become a serious health problem during the last 40 years. In order to control and treat TB disease, the development of new effective drugs is urgently needed.

*Mycobacterium tuberculosis* has a complicated cell wall architecture compared to other antibiotic resistant bacteria [2–4]. The complexity of the cell wall is considered to be one of the reasons for the bacterium’s natural tolerance to antibiotics. One major cell wall component is peptidoglycan. The mutations of proteins involved in peptidoglycan synthesis in *M. tuberculosis* consequently lead to antibiotic resistance [5]. The peptidoglycan layer is formed by glycan chains of β-(1,4) linked N-acetylmuramic acid and N-acetylmuramic acid which are additionally crosslinked by 3,3 or 4,3 transpeptide bonds between short amino acid fragments of alanine, glutamate, and diaminopimelic acid residues. In mycobacteria, the 3,3 crosslinks are generated in equal abundance in all phases of bacterial growth by the 1,3,6-transpeptidases [6,7]. The classical 4,3 crosslinks are formed during the exponential phase of growth by the 1,3,6-transpeptidase activity of penicillin-binding proteins (PBPs) [8]. The 4,3 transpeptidase activity could be easily inhibited by penicillin or other β-lactam antibiotics by forming a stable covalent complex with the serine at the active site of the PBP [9]. PBPs have not traditionally been chosen as primary drug targets for *M. tuberculosis* because β-lactams are very susceptible to degradation by endogenous bacterial β-lactamases; however, recent reports indicate some level of efficacy against drug-sensitive and even drug-resistant *M. tuberculosis* strains [10]. A more recent study in *Escherichia coli* indicates that β-lactams do not only inhibit transpeptidase activity of PBPs but also induce degradation of the newly formed peptidoglycan chains, leading to systemic toxicity [11]. Therefore, investigating PBPs from *M. tuberculosis* may provide novel opportunities for the development of new antibiotics that would be resistant to β-lactamase cleavage and would simultaneously target multiple cellular proteins involved in peptidoglycan biosynthesis.

*Mycobacterium tuberculosis* possesses only two class A PBPs, PonA1 and PonA2, that contain both a transglycosylase domain involved in polymerization of the polyglycan chains and a transpeptidase domain, the enzymatic domain responsible for binding of penicillin [12,13]. Different studies indicate that PonA1 and PonA2 play a complex role in bacterial physiology and have an unusual hypersusceptibility to β-lactam antibiotics, suggesting that novel therapeutic development could optimally target these enzymes [14–17]. To understand the role of class A PBPs in mycobacterial peptidoglycan biology, we examined PonA1 from the *M. tuberculosis* strain H37Rv. PonA1 is a high molecular weight 71 kDa, two-domain protein that contains a noncleavable signal peptide at the N terminus. The N-terminal signal peptide of PonA1 is phosphorylated and was identified as a substrate for the serine-threonine protein kinase PknB [15,18]. The peptidoglycan transglycosylase domain is homologous to PBP1 from *E. coli* [19] and catalyzes the ligation of N-glycolyl muramic acid from the polyglycan sacculus to N-acetyl glucosamine from amphipathic peptidoglycan precursor molecules (lipid II). The C-terminal domain of PonA1 crosslinks the penultimate D-alanine to the D-meso-diaminopimelic acid between parallel peptidoglycan strands. PonA1 localizes to the poles and septa of *M. smegmatis* and is required for maintaining normal cell length in both *M. tuberculosis* and *M. smegmatis* [15,20,21]. Changes to the level of PonA1 protein result in a structurally abnormal cell shape, suggesting that PonA1 is critical for maintaining physiological cell wall synthesis and cell shape during growth. PonA1 regulates essential RipA peptidoglycan hydrolase activity and binds to the RipA endopeptidase domain at the end of PonA1’s C-terminal transpeptidase domain [20]. Additionally, recent work shows that PonA1 activity impacts antibiotic tolerance in mycobacteria [5]. Altering PonA1 transpeptidase or phosphorylation activity affects bacterial growth in the presence of the antibiotic teicoplanin, which targets peptidoglycan biosynthesis [15].

In this study, we describe the high-resolution crystal structure of the PonA1 transpeptidase domain from the *M. tuberculosis* virulent strain H37Rv in ligand-free form and in complex with penicillin V. We characterize the structural details and penicillin-binding site. Furthermore, using site-directed mutagenesis and antibiotic profiling, we provide evidence that PonA1’s transpeptidase activity in both *M. tuberculosis* and *M. smegmatis* mediates tolerance to different classes of β-lactams. Loss of PonA1’s transpeptidase activity renders cells more susceptible to β-lactams, especially to meropenem and penicillin V. Additionally, we tested the stability of PonA1 in the presence of different classes of β-lactams by fluorescence thermal shift.
(FTS) assays. We find that different antibiotics have different impact on the folding of PonA1. FTS data show that binding and formation of the acyl-enzyme by compounds like carbenicillin or penicillin V result in positive Tm shifts, while others like clavulanate or meropenem give negative Tm shifts, indicating that they induce a more destabilized conformation of PonA1. The structure of PonA1’s transpeptidase domain in complex with penicillin V provides a starting point to understand the mechanism underlying the development of antibiotic resistance in \textit{M. tuberculosi}s and to design novel transpeptidase inhibitors, which hold great potential for the treatment of TB disease.

**Results**

**Overall structure of PonA1’s transpeptidase domain**

Four PonA1 molecules in the asymmetric unit of the inhibitor-free form share a similar structure with an average root-mean-square deviation (RMSD) value of 0.2 Å calculated between the Cα atoms of their main chains. Gel filtration chromatography and analysis by the PISA server [22] suggest that the biological unit of PonA1 is a monomer (data not shown). Crystallized PonA1 molecules (residues 249–643) contain the transpeptidase domain and one small adjacent domain at the N terminus of the transpeptidase domain (Figs 1A and 2). The first 156 residues that form part of the N-terminal glycosyltransferase domain and 33 residues at the C terminus of PonA1 were not observed in the protein structure. These modifications could be due to protein degradation and/or structural disorder. In the structure, the PonA1 molecule has a unique unstructured C terminus that contains a proline-rich region. This region forms an exposed long hydrophobic tail (Fig. 1B), suggesting that it may be involved in the protein–protein interactions that have been suggested by previous studies [23].

The small adjacent N-terminal segment of the transpeptidase domain comprises two α-helices [α1n (250–268) and α2n (270–276)] and a three-stranded [β1n (280–284), β2n (480–486), and β3n (490–495)] antiparallel β-sheet (Figs 1A and 2). The PonA1 transpeptidase (or penicillin-binding domain) resembles other PBPs secondary structure: PBP1 of class A from \textit{Staphylococcus pneumoniae} (RMSD of 1.9 Å) [24,25], the transpeptidase domain of PBP2 from Neisseria gonorroeae (RMSD of 2.00 Å) [26], class A PBP1 from \textit{Pseudomonas aeruginosa} (RMSD of 1.65 Å) [27,28], and PBP4 from \textit{Listeria monocytogenes} (RMSD of 2.01 Å) [29]. A cleft in the middle of the domain contains the active site. The cleft divides the transpeptidase domain of PonA1 into two subdomains (Figs 1A and 2). One subdomain has α/β fold and contains an antiparallel β-sheet surrounded by α-helices in a following order: α1 (286–302), β1 (307–316), β2 (321–326), β1′ (341–343), β5′ (473–475), β6′ (503–505), α8 (520–525), β3 (537–545), β4 (550–560), β5 (563–572), and α9 (590–605). The second subdomain has an α-helical structure with six α-helices: α2 (349–358), α3 (391–397), α4 (399–410), α5

![Fig. 1. Structure of the transpeptidase domain of PonA1.](image)
(413–425), α6 (458–471), and α7 (507–519). Additionally, the second subdomain contains a long loop with three short β strands, β2’ (364–368), β3’ (371–373), and β4’ (376–378), that forms part of the inhibitor-binding pocket. The conformation of this loop is held by disulfide bond between Cys386 and Cys389 (Fig. 1A).

**Fluorescence thermal shift assays**

To investigate whether certain drugs may impact the stability of the protein, we determined thermal denaturation curves using FTS assays with different derivatives of β-lactam antibiotics. Effects of 23 antibiotics on the denaturation temperature of PonA1 were determined (Fig. 3). The data show that hetacillin (ΔT_m = 3.0 °C), carbenicillin (ΔT_m = 2.2 °C), dicloxacillin (ΔT_m = 1.9 °C), penicillin V (ΔT_m = 1.6 °C), or cefotaxime (ΔT_m = 1.6 °C) shifts the T_m of PonA1 by more than 1° in the positive direction, while nafcillin (ΔT_m = −2.0 °C), clavulanate (ΔT_m = −2.7 °C), and meropenem (ΔT_m = −3.8 °C) cause significant T_m shifts in the negative direction. The data shown are for PonA1 at 2.8 μM and the antibiotics at 5 μM.

The significant transition temperature shift at this low drug to protein ratio (1.8–1) indicates specific binding of these antibiotics to PonA1. Furthermore, carbenicillin gave rise to a similar positive T_m shift as penicillin V, cefotaxime, and dicloxacillin, while compounds such as clavulanate, meropenem, and nafcillin exhibited negative T_m shifts, indicating that these compounds make PonA1 less stable in these conditions.
While β-lactam compounds induced ~10 °C $T_m$ shift for other PBP proteins we studied (data not shown), the largest shift we observed for PonA1 was <3.5 °C.

**Catalytic site and PonA1–penicillin V interactions**

Similar to all known β-lactam-binding proteins, the PonA1 transpeptidase domain contains three conserved sequence motifs: SxxK (which contains the catalytic serine), SxN, and KTG at the penicillin-binding cleft [8,9]. The SxxK motif comprising residues Ser345, Ser346, Phe347, and Lys348 is located at the N terminus of the helix α2 and contains two important residues for catalysis in PBPs: Ser345 and Lys348. Ser345 is acylated by the substrate and situated at the center of the catalytic site. Lys348 is thought to enhance the nucleophilicity of the serine. The SxN motif comprising residues Ser398, Leu399, and Asn400 is located on the loop between helices α3 and α4. The KTG motif comprising residues Lys539, Thr540, and Gly541 is situated on the strand β3 of the PonA1 transpeptidase domain. In the inhibitor-free PonA1 structure, the Ser345 hydrogen bonds with Lys348, Glu382, and Thr540 and through a water molecule with Ser398. At the penicillin-binding site of the PonA1 structure, Lys348 hydrogen bonds with Asn400 from the SxN motif. We identify another hydrogen bond between Lys539 and Ser398 from the SxN motif. Despite the similarity in the position of the three conserved motifs within the active site, the rate of acylation of β-lactams varies considerably in different PBPs [30–33]. This suggests that the architecture of the antibiotic-binding pocket and adjacent loops plays an important role in the activity of different PBPs and likely influences PonA1’s interaction with inhibitors.
The high-resolution crystal structure of PonA1 in the penicillin V acylated form was determined and contained one protein molecule (residues 249–628) in the asymmetric unit. The part of the N-terminal glycosyltransferase domain (156 residues) and C terminus (51 residues) were missing in the structure due to protein degradation and/or disorder. We calculated RMSD values between the Ca atoms of PonA1 monomers in the apo-form and PonA1 monomer in the ligand-bound structure. The average RMSD value was 0.5 Å. Conformational changes with RMSD > 0.8 Å were observed in the position of the loops between β4'-α3 (loop with disulfide bridge), α5-α6, β3-β4, α5-α9 (in monomer C of the apo-form), and β3n-β6' (in monomer A of the apo-form) as well as in the position of the small adjacent domain at the N terminus of the transpeptidase domain (Fig. 1C). Another change that was observed was in the position of the C-terminal proline-rich tail in all monomers of PonA1 (Fig. 1C). To investigate if packing interactions between adjacent monomers in the crystal lattice were responsible for conformational differences in the structures, we analyzed their crystal contacts. In the crystal lattice, PonA1 monomers were very tightly packed with Matthews coefficients \( V_m = 1.81 \) and \( V_m = 2.17 \) for the inhibitor-free form and penicillin V-bound form, respectively. The contacts between the PonA1 monomers and symmetry-related monomers in ligand-free and penicillin V-bound structures with Van der Waals interactions < 4.6 Å were calculated with the program CONTACT (ccp4 support program [34]). The analysis indicated that all monomers have different contact areas with symmetry-related monomers. Most of the conformational changes including the N-terminal adjacent domain and C-terminal proline-rich tail have contacts with symmetry equivalent monomers in one or both structures. However, the position of the loop between β4'-α3 surrounding the active site was not associated with crystal packing and was most likely affected by reaction with the penicillin V (Fig. 4A). The disposition of the loop β4'-α3 in the PonA1 monomers has the largest RMSD value ~ 3 Å (calculated average value). In two monomers (B and D) of the apo-form structure, this loop is partially disordered due to its mobility.

At the binding cleft, penicillin V forms an acyl–enzyme complex through the covalent bond between the OG atom of Ser345 and the C6 atom of the β-lactam ring (Fig. 4A and 4B). The carboxyl group of the thiazolidine ring forms hydrogen bonds with OG1 atoms of Thr540 and Thr542 located on the β3 strand, and through a water molecule with the nitrogen of the main chain and the OG atom of the side chain of Ser589 on the loop β5-α9 (Fig. 4C). The N7 atom of the thiazolidine ring of penicillin V hydrogen bonds with the OG atom of Ser398 located on helix α3. The N4 and O18 atoms of the acyl side chain of penicillin V interact with the O atom of the main chain of Thr542 of β3 and the ND2 atom of the side chain of Asn400 of α4, respectively. Other interactions occur between the O1 and O18 atoms of penicillin V and Thr379 of β4', Ser381 of B4', and Ser398 of α4 residues through water molecules (Fig. 4C). Therefore, structural analysis of PonA1 in an antibiotic-bound and free form suggests that positioning of the R group of penicillin V at the active site is accompanied by conformational change of the loop β4'-α3 that widened the penicillin-binding pocket (Fig. 4A).
Antibiotic profiling experiments

PonA1’s interaction with small molecules occurs in the inhibitor cleft, which includes its active site serine. Because of the importance of PonA1’s active site in cell wall biosynthesis and cell growth, we hypothesized that PonA1’s active site serine plays a key role in the protein’s interactions with small molecules during cell growth. To test the importance of PonA1’s active site for interaction with antibiotics, we generated M. tuberculosis and M. smegmatis mutants of the transpeptidase active site. Coupling these genetic mutants with several derivatives of β-lactams, we assessed the efficacy of particular PBP-targeting antibiotics in M. tuberculosis and M. smegmatis (Fig. 5A). For both M. tuberculosis and M. smegmatis, loss of PonA1’s transpeptidase activity rendered cells more susceptible to β-lactams, specifically to carbenicillin and meropenem (Fig. 5A). It is possible that the observed difference in antibiotic tolerance may be due to altered effective antibiotic concentration for the remaining PBPs in the cell (because transpeptidase mutant PonA1 no longer covalently binds transpeptidase inhibitors). However, our previous work demonstrated that M. tuberculosis cells that lack PonA1 and PonA2 exhibit differential susceptibility to cell wall-targeting antibiotics, including transpeptidase inhibitors. Thus, although these proteins are homologous, their distinct

![Fig. 5. In vitro antibiotic efficacy experiments. (A) The indicated Mycobacterium tuberculosis H37Rv strains were grown in the presence of carbenicillin and clavulinate (left panel) or meropenem (right panel) at the indicated concentrations. Bacterial survival in the presence of antibiotic was calculated in comparison to untreated bacterial cells. Wild-type PonA1, wt; TG-mutant PonA1, TG-; TP-mutant PonA1, TP-; double TG- and TP-mutant PonA1, TG-TP-. Data were collected in duplicate for each strain and are representative of two experiments. Error bars represent SD. The asterisk indicates *P*-value of < 0.05 and was determined by an unpaired, two-tailed t-test. (B) M. smegmatis mc2155 strains expressing wild-type PonA1 (wt, nonoutlined squares) or transpeptidase mutant PonA1 (TP-, outlined circles) were grown in the presence of penicillin V and clavulinate (left panel) or meropenem (right panel) at the indicated concentrations. Untreated control cells are shown in gray for both strains. Data were collected in triplicate (for carbenicillin data) or duplicate (for penicillin V data) for each strain and are representative of two experiments. Error bars represent SEM.](image-url)
physiological roles in mycobacterial cell wall biology result in different susceptibility to cell wall-active drugs [35].

Interestingly, cell growth plateaued at different times when *M. smegmatis* cells were treated with penicillin V compared with meropenem. The PonA1 transpeptidase-inactivated cells exhibit a substantial delay in growth at a range of penicillin V concentrations when compared to wild-type control (Fig. 5B). At 1 μg·mL⁻¹, the TP cells never recover. These data suggest that cells lacking PonA1’s transpeptidase activity grow much less robustly in the presence of penicillin V or meropenem.

We observed that when we treat cells that encode an allele of ponA1 wherein both enzymatic domains are inactivated (Fig. 5A, TG-TP-) with meropenem, the survival rate is almost the same as the single TG-mutant. This may correlate with other cell biology data that show the expression of a TP allele is important for normal peptidoglycan and how these 4,3 and 3,3 crosslinkings in mycobacterial peptidoglycan and how these 4,3 and 3,3 crosslinking enzymes work in mycobacteria. Because mycobacteria have more 3,3 crosslinks than 4,3 crosslinks, we may observe this glycan strand toxicity more in the presence of 3,3 inhibitors (such as meropenem), unlike the situation in *E. coli*.

**Discussion**

In this study, we characterize the crystal structure of the transpeptidase domain of PonA1, which affects antibiotic tolerance in mycobacteria. The structural comparison of inhibitor-free and inhibitor-bound states indicates that binding of penicillin V induces conformational changes of the loop β4'-α3 leading to a widening of the penicillin-binding pocket. In the structure, penicillin V forms a network of interactions with PonA1 residues of highly conserved motifs (SxK, SxN, and KTG) in known β-lactam PBPs. These findings highlight why penicillins are such effective inhibitors of this class of enzymes.

Our antibiotic profiling experiments suggest that PonA1’s transpeptidase activity is important for normal cellular tolerance of transpeptidase inhibitors, especially to penicillin V and meropenem. We also observed that cells that lack PonA1’s transpeptidase activity grown in the presence of penicillin V exhibit a substantial delay in growth and develop resistance in 25 h. This may suggest that homologous peptidoglycan synthesizing enzymes are more insensitive to penicillin V take over PonA1’s function to promote new peptidoglycan synthesis or that mutations occurred that alter the PBP binding affinity of penicillin V. Moreover, our data indicate that meropenem and penicillin V affect cell growth differently. This may be correlated with the ability of these classes of compounds to target 4,3 crosslinking enzymes vs. 3,3 crosslinking enzymes. For example, meropenem is thought to target both the enzyme groups, whereas carbenicillin or penicillin V should target just the 4,3 crosslinking enzymes.

In addition, the FTS assays show that different derivatives of β-lactam antibiotics have different effects on PonA1 protein stability. The presence of meropenem, clavulanate, or nafcillin induces unstable conformations of the protein, while carbenicillin, penicillin V, metacillin, dicloxacillin have positive effects on protein stability. There is a possibility that binding of meropenem, clavulanate, or nafcillin increases more disordered conformations of the protein. The initial noncovalent complexes should have favorable interactions. When the covalent complex is formed and the serine is acylated, the interactions with product change. The covalent adducts with the acylated serine make more unfavorable interactions at the protein active site that destabilize the structure. We also suggest that, when compared to acylation with penicillin V, acylation with meropenem may affect the formation of the disulfide bond, which could lead to a more unstable conformation of the PonA1 transpeptidase domain. We found that binding of penicillin V induced conformational changes in the loop β4'-α3 with the two cysteines (C386 and C389). The changes in the transpeptidase domain consequently may disturb contacts between PonA1 and its interactors, such as the endopeptidase RipA, which are important for normal peptidoglycan metabolism [23]. It has been shown that the C-terminal 259 residues of the transpeptidase domain of PonA1, where the last 55 residues form an unstructured proline-rich tail, are required for interactions with the 25 C-terminal residues of RipA thus contributing to peptidoglycan remodeling processes [20]. Our combined data suggest that meropenem could be a useful model for future drug design to target mycobacterial PBPs. Alternatively, TB treatment could be served by further optimization of meropenem to improve its efficacy and render it even less susceptible to beta-lactamase cleavage [10].

In conclusion, because *M. tuberculosis* is an important human pathogen, these data provide clear molecular details toward understanding the mechanism. 

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underlying the development of penicillin resistance. These structural data provide a template to design novel transpeptidase inhibitors and improve already known drugs to treat TB disease.

**Experimental procedures**

**Cloning**

The gene encoding PonA1 (NCBI accession code YP_177687.1) containing residues 92–676 excluding N-terminal signal peptide was amplified from the genomic DNA of *M. tuberculosis* strain H37Rv using PCR forward and reverse primers: 5'-TACTTCAATCCAATGCCAAC AACCTGTTCGGCGGCGAT and 5'-TTATCCACTTCC AATGTTCAGGCGTGAGGAGCCGAC, respectively. The product was cloned into the pMCSG73 vector, which incorporates a T7 promoter, TVMV-cleavable N-terminal NusA tag, TEV-cleavable N-terminal hexa-histidine tag, and StrepII tag developed at Midwest Center of Structural Genomics (www.mcsg.anl.gov). The vector was transformed into *E. coli* BL21(DE3) Magic cells.

**Expression and purification**

PonA1 fusion protein with NusA was expressed in 4 L of High Yield selenomethionine M9 medium (Mediclon Inc., Shanghai, China) containing 100 mg L⁻¹ ampicillin and 50 mg L⁻¹ kanamycin. BL21-Magic cells were overgrown at 37 °C until OD₆₀₀ reached 0.6–0.8 and were induced with 5 mM isopropyl β-D-thiogalactoside (IPTG) overnight at 22 °C. Cells were harvested by centrifugation at 10 876 g for 10 min and resuspended in lysis buffer pH 8.3 containing 0.25 M sodium chloride, 0.1 M ammonium sulfate, 43.6 mM sodium phosphate dibasic, 3.25 mM citric acid, 10% glycerol, 1 mM tris-(2-chloroethyl)-phosphate (TCEP), 5 mM imidazole, and 0.08% n-dodecyl β-D-maltoside (DDM). The cells were then sonicated and the cleared lysate was prepared by centrifugation at 36 000 g for 40 min at 4 °C. The NusA fusion protein in pMCSG73 constructs is removed before purification by TVMV protease cleavage. TVMV protease was expressed the same way as PonA1 fusion protein and cleared lysate was obtained. The two cleared lysates were mixed together in 5 : 1 ratio (PonA1–NusA:TVMV protease), 5 mM β-mercaptoethanol was added, and the mixture was incubated overnight at 4 °C. The protein was purified by Ni-affinity chromatography followed by gel filtration. Loading buffers (10 mM Tris/HCl pH 8.3, 0.5 M sodium chloride, 1 mM TCEP) containing 25 mM and 0.5 mM imidazole, respectively, were used for washing and elution. The purity of acquired protein was examined with SDS/PAGE and Coomassie blue staining. SDS/PAGE revealed that the protein showed a tendency to degrade. An additional chromatographic step was performed to improve purity. Protein was loaded on a Ni column, washed with loading buffer containing 0.1 mM DDM, 1 M sodium chloride, 1 mM TCEP, and eluted with 0.5 M imidazole in loading buffer with 0.5 M sodium chloride. It was then dialyzed against buffer containing 10 mM Tris/HCl pH 8.3, 0.5 M sodium chloride, and 1 mM TCEP. The hexa-histidine tag was cleaved overnight at 4 °C by recombinant TEV protease having a noncleavable histidine tag. PonA1 was separated from TEV protease by passing through the Ni column. Purified PonA1 was concentrated to 16 mg·mL⁻¹ in the loading buffer defined above for crystallization.

**Crystallization and data collection**

Crystallization trays of selenomethionine-labeled PonA1 were set up on Corning 96-well sitting-drop plates using the sitting-drop vapor-diffusion method at 19 °C. One microlitre of protein solution (concentration 8 mg·mL⁻¹) was mixed with 1 μL of reservoir solution from the crystal screens Classics, Classics II, JCSG+, PACT, PEGs, and PEGs II (Qiagen Inc., Valencia, CA, USA). Prior to cocryocrystallization with antibiotic, PonA1 was incubated with 5 mM penicillin V for 30 min. Crystals of PonA1 in inhibitor-free form were obtained with a reservoir solution containing 0.2 M lithium chloride, 0.1 M HEPES buffer pH 7.0, and 20% (w/v) PEG6000. The cocryocrystals of PonA1 with penicillin V were obtained with a reservoir solution containing 0.1 M HEPES buffer pH 7.5 and 25% (w/v) PEG3350. Before data collection, cocryocrystals of PonA1 with antibiotic were soaked with 10 mM penicillin V for 10 min in 5 μL of well solution. All crystals were flash cooled in liquid nitrogen using reservoir solution as a cryo-protectant. Low temperature (100 K) X-ray diffraction data were collected at LS-CAT 21ID-G and 21ID-F beamlines of the Advanced Photon Source (Argonne National Laboratory, Argonne, IL, USA) from crystals of PonA1 in inhibitor-free form and in the presence of penicillin V at 1.8 and 1.75 A resolutions, respectively. The crystal space groups and cell parameters are summarized in Table 1.

**Structure determination and refinement**

The structure of the PonA1 inhibitor-free form was determined using the single wavelength anomalous dispersion method (SAD) and the AutoSol Wizard from the phenoix program suite [36]. AutoSol contains hyss [37], solve/re-solve [38], xtriage, and phenoix.refine scripts to determine Se positions, generate experimental phases, build and refine model structure. The structure of PonA1 in complex with penicillin V was determined by molecular replacement using Phaser [39]. The structure of the inhibitor-free form was taken as the starting model. The structure of the PonA1 inhibitor-bound form was processed in HKL-3000 [40]. Structures were refined in refmac [41] and manually built in COOT [42]. Water molecules were placed using ARP/WARP.
Table 1. Data collection, structure determination, and refinement statistics.

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<tr>
<td>a, b, c (Å)</td>
<td>46.0, 333.3, 47.5</td>
<td>47.3, 60.6, 133.9</td>
</tr>
<tr>
<td>α, β, γ (°)</td>
<td>90, 108.4, 90</td>
<td>90, 90, 90</td>
</tr>
<tr>
<td>Matthews coefficient</td>
<td>1.81</td>
<td>2.17</td>
</tr>
<tr>
<td>Solvent content (%)</td>
<td>31.9</td>
<td>43.2</td>
</tr>
<tr>
<td><strong>Data collection</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>96.8 (97.6)</td>
<td>99.5 (99)</td>
</tr>
<tr>
<td>No. of unique reflections</td>
<td>120754</td>
<td>39656</td>
</tr>
<tr>
<td>I/σ(I)</td>
<td>30.3 (2.4)</td>
<td>23.5 (3.1)</td>
</tr>
<tr>
<td>Rmerge (%)</td>
<td>0.07 (0.5)</td>
<td>0.09 (0.6)</td>
</tr>
<tr>
<td>CC1/2</td>
<td>0.99 (0.79)</td>
<td>0.99 (0.67)</td>
</tr>
<tr>
<td>Redundancy</td>
<td>4.9 (4.1)</td>
<td>7.2 (7.1)</td>
</tr>
<tr>
<td>Wilson B-factor (Å²)</td>
<td>43.9</td>
<td>16.3</td>
</tr>
<tr>
<td><strong>Refinement</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R (%)/Rfree (%)</td>
<td>16.1/20.5</td>
<td>14.7/18.9</td>
</tr>
<tr>
<td>RMSD bond length (Å)</td>
<td>0.014</td>
<td>0.016</td>
</tr>
<tr>
<td>RMSD bond angle (°)</td>
<td>1.6</td>
<td>1.7</td>
</tr>
<tr>
<td>Average B value (Å²)</td>
<td>31.9</td>
<td>20.1</td>
</tr>
<tr>
<td>Number of molecules in AU</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>No. of atoms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Protein</td>
<td>11251</td>
<td>2834</td>
</tr>
<tr>
<td>Inhibitor/other</td>
<td>–25</td>
<td>24/21</td>
</tr>
<tr>
<td>Ligands</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water molecules</td>
<td>964</td>
<td>458</td>
</tr>
<tr>
<td>Ramachandran analysisb</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Favored (%)/n</td>
<td>95.4/</td>
<td>97.2/</td>
</tr>
<tr>
<td>Allowed (%)/n</td>
<td>4.2/43</td>
<td>2.8/28</td>
</tr>
<tr>
<td>Outlier (%)/n</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

a Data for the highest resolution shell are given in parentheses. The abbreviations RMSD and AU stand for root-mean-square deviation and asymmetric unit, respectively.  
b Defined by validation program MOLPROBITY.

[43] and manually checked in COOT [42]. The quality of structures was checked using the PDB ADIT validation server (http://deposit.pdb.org/validate/) and MOLPROBITY [44]. Refinement and validation statistics are summarized in Table 1. The figures were generated with PYMOL [45], CCP4MG [46], LIGPLOT+ [47], and ESPRIT 3.0 [48].

PDB accession codes

Atomic coordinates and structure factors of PonA1 in inhibitor-free form and in penicillin V-bound form are deposited into the Protein Data Bank [49] with accession codes 5CRF and 5CXW, respectively.

Fluorescence thermal shift assay

Fluorescence thermal shift assays were conducted in 384-well plate format with an assay volume of 10 μL. Concentrated PonA1 protein sample and 5000× Sypro Orange (Invitrogen, Carlsbad, CA, USA) were diluted and mixed in a Heps buffer (100 mM HEPES, 150 mM NaCl, pH7.5), resulting in protein assay concentration of 2.8 μM and 5× Sypro Orange. After adding the protein Sypro Orange mix to a PCR plate, antibiotics from 10 mM DMSO stocks of the spectrum collection (MicroSource Discovery Systems, Gaylordsville, CT, USA) were added in nanoliter volumes using a Labcyte Echo550 acoustic liquid transfer robot for testing. The assay plates were mixed with a plate shaker, sealed with optical clear plastic seal, and centrifuged. Thermal scanning coupled with fluorescence detection was performed on a Bio-Rad CFX384 qPCR machine at 1.5 °C-min⁻¹ from 10 °C to 80 °C. Data analysis was performed using the in-house software excelFTS which uses IDBS Xlfit for fitting the fluorescence data to a Boltzmann function to determine the melting temperature Tm and other thermal transition parameters. The experiments were performed in triplicate.

Site-directed mutagenesis

Transpeptidase active site mutants were generated as previously described [15]. The M. smegmatis mutant is S468/469A (using the appropriate start site for the enzyme, see [15])—both serines in the canonical SSxK motif were mutated to alanine to ensure that activity would be abolished. For M. tuberculosis, the mutations are S487/488A. The mutations were generated in single copy with vectors that integrate into the L5 phage site in the genome (in M. smegmatis, pMC1s TetO MSMEG_6900-FLAG; or in M. tuberculosis pMC1s TetO RV0050-FLAG) in background strains where endogenous ponA1 was deleted. The FLAG epitope does not compromise PonA1 function [15], as this form of the protein fully complements bacterial growth by OD assay.

Antibiotic profiling experiments

The single copy mutant strains described above were used for all experiments. Isogenic wild-type controls (wherein wild-type ponA1 is integrated at the same L5 phage site and endogenous ponA1 is deleted) were included for all experiments, along with untreated cells. For M. tuberculosis experiments, log phase M. tuberculosis H37Rv strains were back-diluted to a calculated OD600 of 0.01 and grown in 7H9 + OADC complete medium ± antibiotic for 6 days in...
24-well plates. Cells were grown at 37 °C with shaking. Each strain was cultured in triplicate. Control wells contained no antibiotic. Optical density of each well was measured after 6 days. Triplicate wells treated with antibiotic were averaged and then normalized to the average of triplicate untreated wells to calculate the percent survival in the presence of antibiotic. Carbenicillin (1 μg·mL⁻¹) was cotreated with clavulanate (0.5 μg·mL⁻¹), Meropenem was used at 0.15625 μg·mL⁻¹. For *M. smegmatis* experiments, log phase *M. smegmatis* mc²155 strains were back-diluted to a calculated OD600 of 0.01 and grown in 7H9 + ADC complete medium ± antibiotic for 48 h in 96-well plates. Cells were grown at 37 °C with shaking. Control wells contained no antibiotic. Optical density of each well was measured every 30 min. All data were plotted without normalization to untreated control strains. Cells were treated with penicillin V at 1, 0.5, and 0.25 μg·mL⁻¹ in the presence of 5 μg·mL⁻¹ clavulanate (cultured in triplicate) or meropenem at 2.5, 1.25, and 0.625 μg·mL⁻¹ (cultured in duplicate).

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**Conflict of interest**

The authors have no conflict of interest to declare.

**Author contributions**

EVF, KJK, CHL, EJR, and WFA planned experiments; EVF, KJK, CHL, ZW, and OK performed experiments; EVF, KJK, CHL, EJR, and WFA analyzed data; EVF, KJK, CHL, OK, EJR, and WFA wrote the paper.

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