A piperidinol-containing molecule is active against *Mycobacterium tuberculosis* by inhibiting the mycolic acid flippase activity of MmpL3

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*Mycobacterium tuberculosis*, the causative agent of tuberculosis, remains a major human pathogen, and current treatment options to combat this disease are under threat because of the emergence of multidrug-resistant and extensively drug-resistant tuberculosis. High-throughput whole-cell screening of an extensive compound library has recently identified a piperidinol-containing molecule, PIPD1, as a potent lead compound against *M. tuberculosis*. Herein, we show that PIPD1 and related analogs exert *in vitro* bactericidal activity against the *M. tuberculosis* strain mc^**6**30 and also against a panel of multidrug-resistant and extensively drug-resistant clinical isolates of *M. tuberculosis*, suggesting that PIPD1’s mode of action differs from those of most first- and second-line anti-tubercular drugs. Selection and DNA sequencing of PIPD1-resistant mycobacterial mutants revealed the presence of single-nucleotide polymorphisms in *mmpL3*, encoding an inner membrane–associated mycolic acid flippase in *M. tuberculosis*. Results from functional assays with spheroplasts derived from a *M. smegmatis* strain lacking the endogenous *mmpL3* gene but harboring the *M. tuberculosis* *mmpL3* homolog indicated that PIPD1 inhibits the MmpL3-driven translocation of trehalose monomycolate across the inner membrane without altering the proton motive force. Using a predictive structural model of MmpL3 from *M. tuberculosis*, docking studies revealed a PIPD1-binding cavity recently found to accommodate different inhibitors in *M. smegmatis* MmpL3. In conclusion, our findings have uncovered bactericidal activity of a new chemical scaffold. Its anti-tubercular activity is mediated by direct inhibition of the flippase activity of MmpL3 rather than by inhibition of the inner membrane proton motive force, significantly advancing our understanding of MmpL3-targeted inhibition in mycobacteria.

With 10 million new cases and 1.6 million deaths in 2017, tuberculosis (TB)^4 in *M. tuberculosis*, a resilient microorganism that can persist through long courses of antibiotics and years of dormancy within the host. Moreover, the emergence of multidrug-resistant (MDR) and extensively drug-resistant (XDR) TB largely contributes to the difficulty in curing TB patients (1). The standard chemotherapeutic treatments remain very complicated and challenging, substantiated by the slow rate of growth of the bacilli and the existence of a thick and waxy drug-impermeable cell envelope (2). This latter structure consists of a complex skeleton where peptidoglycan, arabinogalactan, and mycolic acids are covalently linked together, and in which noncovalently associated lipids are interspersed (3). The mycolic acid portion of the envelope (the mycomembrane) is composed of very long fatty acids (C70 – 90) that are either covalently attached to the arabinan moiety of the arabinogalactan polymer or found esterified to trehalose as trehalose monomycolate (TMM) or trehalose dimycolate (TDM). From a drug discovery perspective, the pathway involving the enzymatic steps responsible for mycolic acid biosynthesis and/or cell wall attachment of mycolic acids is of particular interest, supported by the mode of action of several important anti-TB drugs, including isoniazid, ethionamide, delamanid, or ethambutol (4–6). In this context, new chemical entities that kill *M. tuberculosis* have been discovered in recent years.
including inhibitors of the Ag85 complex catalyzing the transfer of mycolic acids to arabinogalactan (7, 8) or inhibitors of the mycolic acid transporter MmpL3, which flips TMM across the inner membrane (9).

MmpL3 has been identified in multiple high-throughput whole-cell screens as the putative target of multiple anti-TB compounds during the last few years, thereby representing one of the most promising pharmacological targets being investigated (10). In addition, the essentiality of mmpL3 for M. tuberculosis growth in vitro and for establishing infection in human macrophages or in mice has been demonstrated using conditional knockdown mutants (11, 12). A large panoply of chemical entities with different scaffolds have been reported to target MmpL3, not only in M. tuberculosis but also in nontuberculous mycobacteria, such as Mycobacterium abscessus, opening a new field in the inhibition of mycolic acid translocation (13–23). In most cases, the anti-TB activities of these compounds were believed to act by targeting MmpL3 because of the isolation of spontaneous resistant mutants harboring altered mmpL3 alleles. MmpL3 inhibitors are associated with a decrease in TDM biosynthesis and accumulation of its direct TMM precursor, as a consequence of ineffective transport of TMM. In addition, several inhibitors decrease the intracellular ATP concentration and inhibit the proton motive force (PMF) in the inner membrane by perturbing the membrane potential (24, 25). It therefore remains possible that several small molecules may indirectly modulate MmpL3 activity at least by dissipating the inner membrane PMF, which is critical to MmpL3 lipid transport activity (26). So far, using elegant spheroplast-based functional assays, only two molecules, including BM212 and AU1235, have in fact been shown to inhibit MmpL3-mediated TMM flipping across the IM (27). Several putative MmpL3 inhibitors also display synergistic interactions with other anti-TB drugs, further increasing interest in this new pharmacological target (28).

We recently performed a phenotypic screen against M. abscessus using a library of a 177 confirmed chemical series arising from a known set of potent nontoxic anti-TB hits (29), which led to the discovery of a new piperidinol-based compound, termed PIPD1, exhibiting potent bactericidal activity against clinical M. abscessus strains in vitro and in infected macrophages and zebrafish (17). Whole-genome sequencing of multiple PIPD1-resistant M. abscessus mutants identified several mutations in MAB_4508, encoding a protein homologous to MmpL3. In addition, biochemical analyses proved that although de novo synthesis of mycolic acids remained unchanged by PIPD1 treatment, the compound strongly impaired TMM transport (17). However, it is not clear whether PIPD1 can directly inhibit MmpL3 activity or whether it dissipates the inner membrane PMF, which in turn would affect the biological activity of MmpL3. In addition, whether PIPD1 also targets MmpL3 in M. tuberculosis remains unknown. Herein, we conducted a thorough study to characterize the activity of PIPD1 and its mode of action in M. tuberculosis through the combination of genetic and biochemical approaches.

Results

**PIPD1 and related analogs are highly active compounds against M. tuberculosis in vitro**

GSK1985270A or 4-(4-chloro-3-(trifluoromethyl)phenyl)-1-(2-methylbenzyl)piperidin-4-ol, hereafter designated PIPD1, was originally identified as a new class of MmpL3 inhibitor active against M. abscessus (17) via selection from a library of 177 potent hits against M. tuberculosis (29). However, the activity and mode of action of PIPD1 and related analogs in M. tuberculosis have not been investigated yet. Herein, we determined the MIC₉₀ of PIPD1 and 26 structural analogs against M. tuberculosis mc² 6230 (Table S1). Although all compounds were active against M. tuberculosis, they could be classified into two categories based on their MIC values: highly active compounds sharing MIC ≤ 0.65 μM, including PIPD1 and five analogs (FMD88, FMD6, FMD93, FMD65, and FMD10) and moderately or poorly active compounds sharing 1.1 < MIC < 39 μM (all remaining compounds). The potential toxicity of the 11 most active compounds was determined on Vero cells using resazurin as a cell survival indicator. In general, most compounds, including PIPD1, exhibited low cytotoxicity with IC₅₀ values ranging from 13 to 130 μM. Both the IC₅₀ and the selectivity index corresponding to the IC₅₀/MIC₉₀ ratio are indicated in Table S1. Apart from FMD96, FMD91, FMD15, and FMD6, the selectivity index values are in the 100–207 range, particularly attractive for guiding future structure–activity relationship studies.

The presence of a substituent on one of the ortho positions in ring B is critical for the antitubercular activity of the compounds and appears to depend on the size of the substituent. FMD88, the most active compound of the PIPD1 analog series, contains an iodine atom at the ortho position in ring B (Fig. S1). Interestingly, the van der Waals radius of an iodine atom (1.98 Å) is very close to the radius of the methyl group in PIPD1 (2 Å). We noticed a decrease of the biological activity as a function of the decrease in the van der Waals radii of the substituents present on ortho position in ring B; substitutions with a bromine in FMD93 (1.85 Å), a chlorine in FMD65 (1.75 Å), a hydroxyl group in FMD10 (1.51 Å), or a hydrogen atom in FMD15 resulted in a 2–4-fold decrease in the anti-TB activity, relative to PIPD1 (Table S1 and Fig. S1). Addition of another substituent on the second ortho position in ring B slightly decreases the activity (FMD94 and FMD96 compared with FMD65 and PIPD1, respectively). In addition, single substitutions on meta and para positions are less favorable as evidenced by the reduced activity of FMD63 (meta position of the methyl group in ring B) or FMD66 (para position of the methyl group on ring B) relative to PIPD1 (Table S1 and Fig. S1). There was no obvious correlation between the biological activities of PIPD1 analogs and the electronic effects of substituents present on aromatic rings. Furthermore, the presence of a sterically hindered substituent at these positions in ring B appears detrimental for the biological activity, as judged by the presence of an antracene substituent at these positions in ring B (FMD32, FMD33, FMD99, FMD102, and FMD103).

**PIPD1 inhibits MmpL3 in M. tuberculosis**
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**Figure 1. Dose-dependent kill kinetic profiles of M. tuberculosis mc²6230 (A) and mc²6230 R4 (B) exposed to PIPD1.** The parental strain mc²6230 and the spontaneous resistant strain mc²6230 R4 (I292T substitution in MmpL3) were grown in Middlebrook 7H9 broth supplemented with OADC and pantothenic acid and exposed to increasing concentrations of PIPD1. MIC₉₀ and MIC₉₉ were determined by the microdilution method in 96-well plates. MIC₉₀ is defined as the minimal drug concentration that virtually inhibits completely bacterial growth in 7H9 broth medium, whereas the MIC₉₉ represents the drug concentration that kills 99% of the bacteria present in the inoculum after 2 weeks of incubation at 37 °C. Bacterial loads at the beginning and at the end of the experiment were determined by CFU counting on 7H10 containing OADC and pantothenate after 3 weeks of incubation at 37 °C. The results are from three independent experiments.

**Table 1**

| Strains          | MIC₉₀ | MIC₉₉ | MIC₉₀  | MIC₉₉  | MIC₉₀  |
|------------------|------|------|-------|-------|-------|
| mc²6230          | 0.125| 0.25 | 0.0625| 0.125 | 0.0625|
| mc²6230 L567P    | 1    | 1    | 0.5   | 1     | 0.0625|
| mc²6230 I292T    | 4    | 4    | 0.5   | 1     | 0.0625|
| mc²6230 Y252D    | 8    | 8    | 0.125 | 0.25  | 0.0625|

**PIPD1 is bactericidal against M. tuberculosis and retains activity against MDR and XDR isolates**

The high activity against *M. tuberculosis* and lack of toxicity toward eukaryotic cells, PIPD1 was chosen for all subsequent analyses. Dose- and time-dependent killing of *M. tuberculosis* mc²6230 by PIPD1 was examined by exposing culture over 7 days across a 0.25–8 μg/ml dose-response curve and enumerating the CFU (Fig. 1A). At a concentration of 0.25 μg/ml (corresponding to 2× MIC), PIPD1 partially inhibited growth. At 1 μg/ml, PIPD1 was bactericidal, consistent with its minimal bactericidal concentration (MIC₉₀ of 0.25 μg/ml) (Table 1). This effect was further increased at higher concentrations (4 and 8 μg/ml) with an ~5-log reduction of CFU at day 7, relative to untreated control (Fig. 1A). Overall, these results suggest that PIPD1 bactericidal activity is dose- and time-dependent.

Next, we assessed the activity of PIPD1 against H37Rv as well as a panel of MDR and XDR clinical isolates using the MGIT system. The drug susceptibility profile clearly indicates that all strains were equally sensitive to 0.5–1 μg/ml PIPD1, even isolates resistant to all first-line drugs (isoniazid, rifampicin, ethambutol, and pyrazinamide) and resistant to second intent drugs such as moxifloxacin, amikacin, and protoniamide (Table 2). This strongly suggests that PIPD1 is acting on pathways that are different from those inhibited by first- and second-line anti-tubercular drugs.

**PIPD1-resistant strains harbor mutations in mmpL3**

- It was previously shown, in *M. abscessus*, PIPD1 likely targets the mycolic acid transporter MmpL3. However, it is not clear if PIPD1 acts directly or indirectly (17). To investigate whether MmpL3 is the target of this inhibitor in *M. tuberculosis*, spontaneous resistant mutants were isolated by plating *M. tuberculosis* cells on 7H10 plates containing either PIPD1 (at 2×, 4×, and 8× MIC) or the FMD88 derivative (at 16× MIC). In total, 49 individual mutants were obtained, and their resistance profiles to PIPD1 were confirmed, as indicated by their MIC upshifts (Table 3). All mutants exhibited a 4–32-fold increase in resistance to PIPD1 compared with the parental strain. In all mutants, PCR amplicons covering the entire *mmpL3* gene were generated and subjected to DNA sequencing. The presence of SNPs was confirmed in all 49 mutants, leading to 10 different amino acid substitutions (Table 3). Unexpectedly, there were important variations in the frequency of the various SNPs, with the A1945C substitution (resulting in M649I replacement) accounting for 69% of the mutations identified (frequency of 2.7 × 10⁻⁶). The Y252D mutation was the second most represented mutation found in 18% of the resistors. All other point mutations were isolated in one to three strains. The Y252D and I292T mutations led to the highest resistance levels (32- and 16-fold, respectively).

- To determine whether point mutations in MmpL3 affect also the bactericidal activity of PIPD1, the kill kinetic profile of mc²6230 R₄, carrying the I292T substitution, was compared with the one of the parental strain mc²6230. Fig. 1B shows that...
this mutation abrogates the growth inhibitory effect of the drug at 1 \( \mu \text{g/ml} \) but not at high concentrations (8 \( \mu \text{g/ml} \)). This is strengthened by the fact that mc26230 \( R_4 \) exhibits a higher MBC value than the parental strain (Table 1). Similarly, mc26230 \( R_5 \) and mc26230 \( R_7 \), carrying the L567P and Y252D mutations, respectively, exhibited higher MIC and MBC values, suggesting that these mutations may affect the binding site of PIPD1, in silico docking of the inhibitor was performed on the \( M. tuberculosis \) MmpL3 homology model (Fig. 2B), assuming that the inhibitor binds into the same cavity that has been shown to accommodate four different inhibitors in the crystal structures of MmpL3 from \( M. smegmatis \) (30). This prediction highlights the occurrence of seventeen residues forming the binding pocket of PIPD1, comprising 12 residues (Ile-248, Asp-251, Asp-640, and Tyr-641) merly identified the conserved Asp-251, Asp-640, and Tyr-641 residues in all MmpL members and demonstrated their crucial role for MmpL3 activity by translocating protons through the transmembrane domain (34), an observation subsequently confirmed by the elucidation of the MmpL3 crystal structure (30). Altogether, these results support the view that PIPD1 acts by blocking the proton relay, leading to TMM transport inhibition and mycobacterial death. In addition, four residues of the six-

**Localization of the mutations conferring PIPD1 resistance on a structural MmpL3 model**

To get structural insights into the mechanism conferring resistance to PIPD1, the various mutations identified in the spontaneous resistant mutants were mapped on an \( M. tuberculosis \) MmpL3 three-dimensional homology model that was generated using the crystal structure of \( M. smegmatis \) MmpL3 as a template (30). MmpL3 is a transporter comprising 12 transmembrane helices (30–33). The nine substituted residues are found in seven different transmembrane helices (TMs): P209A is on TM3, Y252D and V257L are on TM4, V285A/G and I292T are on TM5, L567P is on TM8, M649L is on TM10, V681I is on TM11, and M704T is on TM10 (Fig. 2A). To further assess whether these mutations may affect the binding site of PIPD1, in silico docking of the inhibitor was performed on the \( M. tuberculosis \) MmpL3 homology model (Fig. 2B), assuming that the inhibitor binds into the same cavity that has been shown to accommodate four different inhibitors in the crystal structures of MmpL3 from \( M. smegmatis \) (30). This prediction highlights the occurrence of seventeen residues forming the binding pocket of PIPD1, comprising 12 residues (Ile-248, Asp-251, Asp-640, and Tyr-641) merly identified the conserved Asp-251, Asp-640, and Tyr-641 residues in all MmpL members and demonstrated their crucial role for MmpL3 activity by translocating protons through the transmembrane domain (34), an observation subsequently confirmed by the elucidation of the MmpL3 crystal structure (30). Altogether, these results support the view that PIPD1 acts by blocking the proton relay, leading to TMM transport inhibition and mycobacterial death. In addition, four residues of the six-

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PIP1 inhibits MmpL3 in M. tuberculosis

Figure 2. Localization of mutations conferring resistance to PIP1 on a M. tuberculosis MmpL3 predictive structural model. A, mapping of the various mutations identified in spontaneous resistors selected against PIP1 on a M. tuberculosis MmpL3 three-dimensional homology model. Only the transmembrane helices are depicted. The mutated residues are shown as sticks, and the mutations are indicated in the colored boxes. All the TM helices are in gray, excepting those carrying mutations, notably TM3 (blue), TM4 (magenta), TM5 (wheat), TM8 (red), TM10 (pink), TM11 (pale green), and TM12 (green). PIP1 is shown as yellow sticks. B, the figure depicts the residues forming the proposed PIP1-binding cavity. Residues are in the same color code as in A. The four residues interacting with PIP1 and found mutated in the PIP1 resistant strains are in indicated in colored boxes. PIP1 was docked in the model and the best docking pose is represented and corresponds to a binding energy of ΔG° = −7.6 kcal/mol. The black dashed lines indicate hydrogen bonds. C, comparison of the PIP1-binding sites in MmpL3 from M. abscessus (green), M. smegmatis (orange), and M. tuberculosis (blue).

PIP1 targets MmpL3 and inhibits its flippase activity

The primary function of MmpL3 involves exporting TMM across the plasma membrane, allowing the lipid synthesized in the cytoplasm to reach the mycomembrane (17, 21, 27, 32, 35). Although all of the data acquired so far are consistent with the idea that PIP1 inhibits MmpL3, it remains unknown whether PIP1 directly affects MmpL3 activity. To evaluate whether PIP1 inhibits TMM flipping, we exploited recently developed TMM topological assays (27), based on the production of spheroplasts, which are devoid of the outer membrane and cell wall but retain an intact inner membrane. In this study, spheroplasts were derived from a genetically engineered M. smegmatis strain in which the endogenous mmpL3 gene has been deleted and mmpL3 from M. tuberculosis has been inserted in the chromosomal attB site (∆mmpL3 attB::mmpL3-tb) (Fig. S2). The MmpL3 flippase activity in spheroplasts can be assessed by...
monitoring surface exposure of 6-azido modified TMM. In this method, 6-azido-trehalose was fed to spheroplasts and incorporated into 6-azido-TMM in the cytoplasmic membrane (27, 36). 6-Azido-TMM can be labeled via bioorthogonal chemistry with alkyne-containing biotin (sDIBO-biotin) to generate biotin-conjugated TMM, and the surface-exposed fraction of these molecules can be selectively detected by streptavidin–Alexa 488 (Fig. 3A). The well-characterized MmpL3 inhibitor, 1,5-diarylpyrrole BM212 was included as a positive control (13). 4× MIC PIPD1-treated spheroplasts showed significantly lower fluorescence on the surface compared with the control cells treated with DMSO, suggesting a strong inhibition of MmpL3 flippase activity. The reduction of cell surface fluorescence by PIPD1 is comparable with that by 4× MIC BM212 (Fig. 3, B and C), which has previously been shown to inhibit TMM flipping (27).

In another method to assess the TMM flipping activity of MmpL3, specific lipases that can only access mycolic acids that have been flipped to the outer leaflet on the inner membrane were added (27) (Fig. 4A). Cellular lipids were labeled with [14C]acetate, and the accessibility of the newly formed radioactive TMM molecules to the lipase lysin B (LysB) was determined in spheroplasts incubated with PIPD1, followed by extraction of the lipids and subsequent resolution by TLC. We found that PIPD1 significantly reduced TMM accessibility to LysB in a weak dose-dependent manner (Fig. 4B). When the spheroplasts were treated with 4× MIC of PIPD1, the percentage of TMM accessible to LysB was reduced to a level comparable with the "no drug treatment" control.
Figure 4. PIPD1 reduces TMM accessibility to LysB in spheroplasts expressing WT MmpL3-tb, but not in spheroplasts expressing MmpL3I292T-tb. A, schematic representation illustrating the inner membrane (IM) of the spheroplast and the MmpL3-mediated mycolic acid translocation. Newly synthesized TMM is flipped across the IM, released from IM where it becomes accessible to LysB (left panel). In the presence of MmpL3 inhibitors, it is expected that, because of inhibition of the flippase activity, the TMM level accessible to LysB is reduced (right panel). B and C, representative TLC analyses of 14C-labeled lipids newly synthesized in the presence of indicated concentrations of PIPD1 and extracted from M. smegmatis ΔmmpL3 attB::mmpL3-tb (WT) (B) or ΔmmpL3 attB::mmpL3T875C-tb (encoding MmpL3I292T-tb) (C) spheroplasts following treatment with or without purified LysB. DMSO was used to dissolve PIPD1 and thus served as the negative control. BM212 was used as a positive control. Equal amounts of radioactivity were spotted for each sample. The developing solvent system comprises chloroform-methanol-water (30:8:1, v:v:v). D, graphical plot showing the effects of PIPD1 on the amounts of LysB-accessible TMMs in spheroplasts expressing WT MmpL3-tb (WT) or MmpL3T875C-tb (I292T). The percentage of TMMs accessible to LysB is given by the difference in TMM levels between samples with or without LysB treatment, normalized against the level in control samples without LysB treatment. TMM levels in each sample were quantified as a fraction of total mycolates (TMM + mycolic acid (MA)). Average percentages and standard deviations from three biological replicates are plotted. For Student’s t test, NS, not significant; *, p < 0.05; **, p < 0.01 compared with the respective DMSO control.

To further address whether PIPD1 inhibits TMM flipping by targeting MmpL3, we sought to examine LysB accessibility of TMMs in M. smegmatis spheroplasts expressing mutated mmpL3 alleles that conferred to M. tuberculosis PIPD1 resistance. Consistent with the MIC results in M. tuberculosis, introducing mmpL3T754G-tb (encoding MmpL3Y252D-tb) or mmpL3T875C-tb (encoding MmpL3I292T-tb) as the only copy of mmpL3 into M. smegmatis increased resistance against PIPD1 by at least 8-fold (Table S2). Using the LysB assay, we demonstrate that TMM surface exposure was unaffected by PIPD1.
treatment in spheroplasts expressing MmpL3_{1292T-tb} (Fig. 4, C and D), indicating that this single mutation prevents MmpL3 from being inhibited by PIPD1. We conclude that MmpL3 is the target of PIPD1 in M. tuberculosis.

Because the activity of MmpL3 relies on the PMF of the inner membrane, we next investigated whether PIPD1 may interfere with the membrane potential (Δψ) and/or the proton gradient (ΔpH). Δψ was monitored by following the fluorescence intensity variations of the 3,3′-dipropylthiodicarbocyanine (DiSC3(5)) dye in ΔmmpL3 attB::mmpL3-tb spheroplasts upon the addition of 2× and 4× MIC of PIPD1. Similar to BM212, PIPD1 failed to show variations in the intensity of fluorescence, whereas the addition of valinomycin + KCl led to a pronounced increase in fluorescence caused by Δψ disruption (Fig. S4), suggesting that PIPD1 does not affect the membrane potential. We next investigated the potential effect of PIPD1 on the ΔpH across the inner membrane of ΔmmpL3 attB::mmpL3-tb spheroplasts using a pH-sensitive fluorescent dye BCECF. In contrast to CCCP (at 2 and 5 μM), the addition of PIPD1 did not disrupt the proton gradient across the inner membrane (Table S3). Together, these results suggest that PIPD1 blocks TMM translocation without affecting the PMF of the inner membrane but through direct inhibition of MmpL3, likely by blocking the translocation of protons through the transmembrane domain.

**Discussion**

The discovery efforts focused on selecting new drugs are critical for the long-term control of TB, particularly against MDR and XDR TB. In this context, new classes of compounds active against M. tuberculosis and inhibiting new target pathways have been identified in recent years (37). In this study, we describe the potent activity of PIPD1 against M. tuberculosis drug-sensitive and -resistant strains. The parental molecule, along with a panel of structural analogs, exhibited low MICs and high selectivity indices and elicited resistant mutations in mmpL3. Analysis of the mmpL3 mutations, causing various levels of resistance to PIPD1, shows similarities with previously identified inhibitors as well as unique amino acid replacements. In particular, the L567P mutation in TM8 can also be found in the strains resistant to HC2091, SQ109, and TBL140 (14, 19). That the PIPD1-resistant strains carrying the L567P, I92T, and Y252D substitutions were cross-resistant to the adamantyl urea AU1235 (21) further supports the view that PIPD1 targets MmpL3, as shown earlier in M. abscessus (17). The fact that the MIC values of PIPD1 (0.125 μg/ml) were very similar against M. tuberculosis and M. abscessus prompted us to advance that the mechanism of action and/or binding site would be conserved in both species. Despite an extensive screening leading to the selection of 49 individual spontaneous resistant mutants and the identification of 10 different amino acid substitutions, these mutations were not structurally clustered on an M. tuberculosis MmpL3 three-dimensional predictive model. However, computational docking revealed that PIPD1 is able to occupy the cavity within the proton translocation channel of this M. tuberculosis MmpL3 model. Because the same cavity has recently been shown to accommodate other small molecule inhibitors in the crystal structure of M. smegmatis MmpL3 (30), we believe that PIPD1 likely binds MmpL3 directly. However, full confirmation of this binding mode awaits the experimentally determined 3D structure of the MmpL3–PIPD1 complex.

MmpL3 carries an essential function in mycobacteria by exporting mycolic acids to the mycomembrane (21, 35). Consistent with these findings, conditionally regulated knockdown of mmpL3 demonstrated requirement of MmpL3 function for replication and viability of M. tuberculosis, both under laboratory growth conditions and during infection in mice (11, 12). Furthermore, depletion of MmpL3 rendered M. tuberculosis hypersusceptible to MmpL3 inhibitors (11). Recent work demonstrated that MmpL3 is the flippase catalyzing translocation of TMM from the inner leaflet to the outer leaflet in the inner membrane in mycobacteria (27). TMM is the precursor of the mycolyl-arabinogalactan-epitidoglycan skeleton and TDM, which are major components of the mycomembrane resulting from the action of the Ag85 complex (8, 38, 39). Although we believe that PIPD1 binds to MmpL3, this binding may not necessarily translate to inhibition of its activity. A striking example is SQ109, which has been shown to bind and co-crystallize with MmpL3 (30) but fails to inhibit the TMM flipping activity of MmpL3 in mycobacterial spheroplasts (27). We show here that PIPD1 targets MmpL3 and reduces its flippase activity in a M. smegmatis mutant lacking the endogenous mmpL3 gene but carrying the M. tuberculosis homologous gene, without affecting the inner membrane PMF, thus providing new functional knowledge that cannot be gleaned from binding/structural studies alone. MmpL3 inhibition by PIPD1 is very likely to result in the cessation of the mycolyl-arabinogalactan-peptidoglycan and TDM synthesis, as reported previously in M. abscessus (17). In a recent study, Shetty et al. (40) showed that the E11 acetamide analog targets MmpL3 by an indirect mechanism involving disruption of the inner membrane PMF utilized by MmpL3 for TMM translocation. This contrasts with PIPD1, behaving like BM212, altering neither the membrane potential (Δψ) nor the proton gradient (ΔpH). This is particularly important because it has been proposed that several MmpL3 inhibitors may act indirectly by targeting the PMF, which drives proton translocation for MmpL3 lipid translocation (26) and that spontaneous resistance mutants in MmpL3 may be compensatory, thereby obscuring possible alternative targets, as demonstrated for the tetrahydroprazolopyrimidine THPP1 (41, 42). Therefore, combined together, our genetic, biochemical, and docking studies support a scenario whereby PIPD1 directly interacts with MmpL3 and inhibits its function.

This adds PIPD1 to the growing list of MmpL3 inhibitor classes (BM212, adamantly ureas and indolecarboxamides) that all bind to the same site within the proton translocation channel of the protein whether or not they trigger additional effects on the PMF that potentiates their activity (30, 43).

Although most of the PIPD1 analogs evaluated in this study are equally or less potent than the parental PIPD1 molecule against M. tuberculosis, one of them (FMD88) showed a slightly increased activity. This demonstrates that simple structural optimization of the PIPD1 scaffold is possible and may lead to the second generation of derived drug candidates for the potential treatment of MDR/XDR TB as mycolic acid transporter inhibitors. The structure–activity relationship study provided here will guide fur-
ther optimization of the parental scaffold followed by subsequent testing of these compounds in animal models to strengthen the development pipeline of this new class of antitubercular agents.

The exquisite vulnerability of MmpL3 at all stages of the infection establishes this transporter as an attractive new target with the potential to improve and shorten current drug-susceptible and drug-resistant TB chemotherapies. This led to studies showing that the indolecarboxamides and adamantyl urea MmpL3 inhibitors act also synergistically with rifampin, bedaquiline, clofazimine, and β-lactams, offering the perspective of reducing the duration of TB treatments (28). Therefore, it remains to be investigated whether PIPD1 or its related analogs may also potentiate the activity of other anti-TB drugs.

**Experimental procedures**

**Synthesis of PIPD1 analogs**

A library of 26 PIPD1 analogs was prepared following a methodology reported elsewhere. AU1235 was purchased from MedChemTronica (Sweden). All compounds were dissolved in DMSO.

**Bacterial strains and growth conditions**

*M. tuberculosis* mc²6230 (44) was grown at 37 °C in Middlebrook 7H9 supplemented with 0.05% Tween 80, 10% oleic-albumin-dextrose-catalase enrichment (7H9 OADC/T) and 24 μg/ml pantothenic acid.

**Drug susceptibility testing**

Exponentially growing *M. tuberculosis* cultures were diluted in 7H9OADC/T containing 24 μg/ml pantothenic acid to an A₆₀₀ = 0.01. The bacteria were then seeded in 100-μl volumes in all the wells of a 96-well plate except the first column of wells, which contained 200 μl of the bacterial suspension. Compounds (stock concentration at 10 mg/ml) were directly added to the wells of the first column so as to achieve a concentration of 128 or 64 μg/ml. 2-fold serial dilutions were then carried out by transferring 100 μl of the bacterial suspension from the first column of wells to the second column, mixing, and repeating this procedure for each consecutive column. The plates were then placed in sealed plastic bags and incubated at 37 °C for 7–10 days. The MIC is defined as the lowest concentration of compound inhibiting growth of 99% of the bacteria present in the inoculum, judged by visual inspection of the plates. The MBC was obtained after CFU enumeration and corresponds to the concentration for which less than 1% of bacteria survived after 7 days of treatment as compared with the number of viable bacteria in the original inoculum.

In addition, phenotypic drug susceptibility testing (pDST) for PIPD1 was performed on clinical MTBC isolates (non-MDR, n = 3; MDR, n = 3; XDR, n = 2) using the BACTEC MGIT 960 instrument (Becton Dickinson, NJ). In all experiments, strain H37Rv was tested in parallel. Procedures to test PIPD1 were adopted from the standard protocol for commercially available first- and second-line compounds. In brief, suitable dilutions of a stock solution containing PIPD1 (100 mg/ml in DMSO) were added to MGIT tubes containing growth medium and BBL OADC supplement (Becton Dickinson) to result in final drug concentrations of 0.0625 μg/ml to 2 μg/ml. MGIT tubes were each inoculated with 0.5 ml of a fresh bacterial inoculum suspension prepared from a liquid subculture. Tubes were mixed well and loaded into the MGIT instrument. Growth was monitored by automated fluorometric detection of oxygen consumption and compared with drug-free control tubes containing a 1:100 dilution of the investigated culture suspension. Incubation was stopped when the growth control reached a growth unit (GU) value of 400. At that point, GU values of drug-containing tubes were retrieved from the MGIT instrument software and results were interpreted as follows. If the GU of the drug-containing tube was more than 100 when the GU of the growth control was 400, the result was defined as resistant. If the GU value was equal to or less than 100, the result was considered susceptible.

**Time-kill assay**

*M. tuberculosis* liquid cultures were exposed to increasing drug concentrations and plated on Middlebrook 7H10OADC containing 24 μg/ml pantothenic acid at a daily basis for 7 days. CFU were enumerated after 2–3 weeks of incubation at 37 °C.

**Selection of spontaneous resistant mutants and identification of point mutations in MmpL3**

Exponentially growing *M. tuberculosis* cultures were plated on 7H10OADC containing pantothenate and 0.25, 0.5, or 1 μg/ml PIPD1. After 2–3 weeks of incubation at 37 °C, single colonies were selected and grown in a liquid medium and individually subjected to MIC determination and scored for resistance to PIPD1. Identification of SNPs in *mmpL3* in the resistant strains was done by PCR amplification using primers *mmpL3f*’-GAT CGA TAT CCT TCG CCA AAC CGA AAG TAG-3’ and *mmpL3r* 5’-AGT TAA GCT TCT AAT CGC GGT GAA CCA ACT-3’ to produce a 3520-bp amplicon and subsequent sequencing.

**Generation of *M. smegmatis***

To generate this strain in *M. smegmatis* mc²155, an integrative vector pJE402 was first used to introduce a WT *mmpL3-msm* allele together with a kanR cassette at the attB site. Next, the chromosomal *mmpL3* was deleted by two-step homologous recombination as described earlier (45). A new pJE402 plasmid harboring *mmpL3-msm* and a hygR cassette was electroporated into the strain to replace the *kanR* with *hygR*; the resulting *M. smegmatis* mutant *mmpL3* attB::*hygR-mmpL3-msm* strain allows other *mmpL3* alleles to be introduced into the *attB* site via allelic replacement. To generate *M. smegmatis* Δ*mmpL3* attB::*hygR-mmpL3-msm* strain allows other *mmpL3* alleles to be introduced into the *attB* site via allelic replacement. To generate *M. smegmatis* Δ*mmpL3* attB::*mmpL3*-tb, another pJE402 plasmid harboring *mmpL3*-tb and a *kanR* cassette was electroporated into the Δ*mmpL3* attB::*hygR-mmpL3-msm* strain. In the transformsants sensitive to Hyg and resistant to Kan, the WT *mmpL3-msm* allele and the *hygR* cassette were successfully replaced by *mmpL3*-tb and *kanR*, and proper replacement was confirmed by PCR and sequencing. The same latter steps were used for the generation of Δ*mmpL3* attB::*mmpL3*₇₇₅₆C-tb (encoding MmpL3₇₇₅₆₋₂₂₅ autoloads), and Δ*mmpL3* attB::*mmpL3*₉₂₇₅C-tb (encoding MmpL3₉₂₇₅₋₂₂₅ autoloads).
**Spheroplast formation**

*M. smegmatis* mc²155 ΔmmpL3 attB::mmpL3-tb cells were converted to spheroplasts, as described previously (27). Briefly, *M. smegmatis* ΔmmpL3 attB::mmpL3-tb was inoculated into tryptic soy broth (TSB) with 0.05% Tween 80, followed by incubation at 37 °C and 180 rpm until the A₆₀₀ reached ~1.0. Glycine was then added into the culture to a final concentration of 1.2% (w/v), followed by continued incubation at 37 °C and 180 rpm for 20–24 h. The glycine-treated cells were harvested by centrifugation at 4,000 g, washed with 1× SMM buffer (final pH 6.8, 0.5 M sucrose and 20 mM maleate buffer at pH 6.6), and resuspended in TSB-SMM (TSB containing 1× SMM buffer) by gentle pipetting. Filter-sterilized solutions of lysozyme (10 mg/ml) and glycine (20%, w/v) were added to final concentrations of 50 μg/ml and 1.2% (w/v), respectively. Spheroplast formation was complete after incubation at 37 °C and 120 rpm for another 20–24 h. The final mixture was washed with 1× SMM buffer and filtered through a sterile cell strainer (20 μm; BD Falcon) to eliminate clumps of cells.

**Assessing TMM accessibility to degradation by purified LysB in spheroplasts**

The LysB-accessibility assay was conducted essentially as previously described (27). Briefly, *M. smegmatis* ΔmmpL3 attB::mmpL3-tb spheroplasts were metabolically labeled with sodium [1-¹⁴C]acetate for 2 h, followed by addition of purified LysB for 30 min at 37 °C. Lipids were extracted directly after the 30 min incubation. Where indicated, putative MmpL3 inhibitors were added 10 min before addition of [¹⁴C]acetate.

**Assessing 6-azido-TMM surface display in spheroplasts**

The 6-azido-TMM surface display assay was conducted essentially as previously described, although the sDIBO-biotin probe was used instead of the discontinued DIBO-biotin probe (27). Briefly, *M. smegmatis* ΔmmpL3 attB::mmpL3-tb spheroplasts were metabolically labeled with 6-azido-trehalose for 2 h at 37 °C to synthesize 6-azido-TMM, which then reacted with Click-IT Biotin sDIBO alkyne to generate biotin-TMM. Surface-exposed biotin-TMM were detected on spheroplasts using Alexa Fluor 488–conjugated streptavidin and visualized by fluorescence microscopy. Use of sDIBO-biotin, instead of DIBO-biotin as in (27), gave higher nonspecific background labeling, even in the negative control where 6-azido-trehalose was not added. Where indicated, putative MmpL3 inhibitors were added 15 min before addition of 6-azido-trehalose and included in all wash buffers. 15 μg/ml chloramphenicol was added to prevent interference from the activity of newly synthesized and exported Ag85.

**Determination of intracellular pH using the BCECF-AM dye**

The effects of inhibitors on ΔpH were determined using the BCECF-AM dye. Briefly, *M. smegmatis* ΔmmpL3 attB::mmpL3-tb spheroplasts (A₆₀₀ = 0.8) in 1× SMM buffers at various pH levels (6.0–8.0) were incubated with 20 μM BCECF-AM at 37 °C for 30 min in the presence of 20 μM nigericin. Nigericin is a proton uncoupler and serves to allow equilibration of protons (pH) across the membrane. The buffers also contained 100 mM KCl to ensure a steady membrane potential (Δψ). Fluorescence emission (λₑₓ 525 nm) intensities of intracellular BCECF were measured following excitation at λₑₓ 488 and 440 nm in a SPECTRAmax 250 microplate spectrophotometer equipped with SOFTmax PRO software (Molecular Devices). The ratio of fluorescence emission intensities at these two excitation wavelengths, or the fluorescence excitation profile (λₑₓ 488 nm/λₑₓ 440 nm), is pH-dependent. Average (λₑₓ 488 nm/λₑₓ 440 nm) values were obtained from technical triplicates and plotted against pH to obtain a linear standard curve. To test the effects of putative MmpL3 inhibitors on intracellular pH (and hence ΔpH), spheroplasts (A₆₀₀ = 0.8) in 1× SMM buffer at pH 6.8 were pretreated with indicated concentrations of PIPD1, and incubated at 37 °C for 30 min. BM212 was used as a negative control, whereas CCCP was used as a positive control. 20 μM BCECF-AM was then added to the samples, and incubation was continued for 30 min before fluorescence measurements. Fluorescence excitation profiles (λₑₓ 488 nm/λₑₓ 440 nm) of BCECF for each condition were averaged (across three technical replicates) and calibrated against the standard curve.

**Membrane potential measurements in M. smegmatis ΔmmpL3 attB::mmpL3-tb spheroplasts**

The effects of inhibitors on Δψ were determined using the membrane potential-sensitive DiSC3(5) dye (47). DiSC3(5) binds to energized membranes and becomes quenched. When Δψ is disrupted, the dye leaves the membrane, resulting in an increase in fluorescence. 1.5 ml *M. smegmatis* ΔmmpL3 attB::mmpL3-tb spheroplasts (A₆₀₀ = 0.8) were used in 1× SMM buffer containing 10 mM glucose and 1 mM nigericin (added to remove the effects of ΔpH). DiSC3(5) was then added to samples to get a final concentration of 5 μM and equilibrated for 10 min at room temperature. From this point, fluorescence was continuously monitored with a SPECTRAmax 250 microplate spectrophotometer equipped with SOFTmax PRO software (Molecular Devices), employing an excitation wavelength of 643 nm and an emission wavelength of 666 nm. The effect of PIPD1 on Δψ was measured by monitoring an increase in fluorescence when these compounds were added at specific time points at indicated concentrations. BM212 and valinomycin (with potassium chloride) were used as negative and positive controls, respectively.

**3D modeling of MmpL3 and docking studies**

The MmpL3 three-dimensional homology models of *M. tuberculosis* (Rv0206) and *M. abscessus* (MAB_4508) were generated using the Phyre2 server (48) and the crystal structure of *M. smegmatis* MmpL3 as a template (PDB code: 6aajj) (30). The model geometry was further optimized using the Phenix package (49). In silico docking of PIPD1 was performed with the PyRx software (50) running AutoDock Vina (51) and using the MmpL3 homology models (*M. tuberculosis* and *M. abscessus*) or the crystal structure of *M. smegmatis* as receptors and the following grid parameters.
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center X = 10.5, Y = 6.1, and Z = 18.2 and dimensions (Å) X = 16.1, Y = 15.3, and Z = 22.1.

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