Pyridylpiperazine-based allosteric inhibitors of RND-type multidrug efflux pumps

Coline Plé1,6, Heng-Keat Tam2,5,6, Anais Vieira Da Cruz3, Nina Compagne3, Juan-Carlos Jiménez-Castellanos1, Reinke T. Müller2, Elizabeth Pradel1, Wuen Ee Foong2, Giuliano Malloci4, Alexia Ballée3, Moritz A. Kirchner2, Parisa Moshfegh3, Adrien Herledan3, Andrea Herrmann2, Benoit Deprez3, Nicolas Willand3, Attilio Vittorio Vargiu4, Klaas M. Pos2,7, Marion Flipo3,7 & Ruben C. Hartkoorn1,7

Efflux transporters of the RND family confer resistance to multiple antibiotics in Gram-negative bacteria. Here, we identify and chemically optimize pyridylpiperazine-based compounds that potentiate antibiotic activity in E. coli through inhibition of its primary RND transporter, AcrAB-TolC. Characterisation of resistant E. coli mutants and structural biology analyses indicate that the compounds bind to a unique site on the transmembrane domain of the AcrB L protomer, lined by key catalytic residues involved in proton relay. Molecular dynamics simulations suggest that the inhibitors access this binding pocket from the cytoplasm via a channel exclusively present in the AcrB L protomer. Thus, our work unveils a class of allosteric efflux-pump inhibitors that likely act by preventing the functional catalytic cycle of the RND pump.
Infections by multidrug-resistant (MDR) Gram-negative bacteria are a major threat to global healthcare and are currently classified as a critical priority for R&D by the World Health Organization. Among many antibiotic-specific resistance mechanisms, active antibiotic extrusion by multidrug efflux pumps is a major contributor to both intrinsic (basal) and acquired multiple drug resistance phenotypes in Gram-negative bacteria. In particular, efflux pumps belonging to the Resistance Nodulation division (RND) superfamily extrude a plethora of chemically diverse antibiotic molecules and represent a major barrier to antibiotic efficacy and drug development.

RND-type efflux pumps span the entire Gram-negative envelope and are composed of a proton motive force-driven inner membrane transporter (e.g. AcrB), a periplasmic adaptor protein (e.g. AcrA), and an outer membrane channel (e.g. TolC). Homotrimeric AcrB is comprised of three protomers that cycle interdependently through the loose (L), tight (T), and open (O) conformational states during H⁺/drug antiport catalysis 1-12. Efflux pump substrates enter AcrB via dedicated entrance tunnels toward the proximal access pocket (AP) in the L protomer or toward the more distal deep binding pocket (DBP) exclusively present in the T protomer. Substrate binding to this DBP leads to structural rearrangements that propagate into the transmembrane (TM) region to allow for protonation of titratable side chains (D407 and D408) located deeply inside the TM domain 13. The electrostatic changes upon protonation lead to the transition from the T state to the O state, resulting in the collapse of the entrance tunnels and the DBP. Concurrently, the substrate leaves the DBP via a newly formed exit tunnel in the AcrB O protomer and is transported through the AcrA-TolC channel across the outer membrane. The AcrB O protomer subsequently resets to the L state through a proton motive force-dependent H⁺-release from the protonated Asp pair into the cytoplasm 5,9,12,14,15.

Considering the critical role of RND pumps in innate and acquired antibiotic resistance in Gram-negative bacteria, there have been considerable efforts to discover and develop efflux pump inhibitors (EPI). Characterized EPIs include Phe-Arg β-naphthylamide (PAβN) and 1-(1-naphthylmethyl)-piperazine (NMP), which inhibit RND-efflux by binding and blocking the DBP or competitively preventing antibiotic binding to this site. More specific allosteric EPIs such as pyridopyrimidines (such as D13-9001), and pyranopyridines (the MBX series) bind to a hydrophobic pocket (aka “hydrophobic trap”) proximal to the DBP of E. coli AcrB 16-23, halting the catalytic efflux cycle by blocking the T to O transition, while also likely competing with drug binding in the DBP 22,24.

In this work, we sought to discover a novel chemical scaffold able to boost antibiotic activity in E. coli by inhibiting the AcrAB-TolC efflux pump. In line with this goal, a pyridylpiperazine-based molecule was discovered by phenotypic screening and was validated to act specifically on AcrB. Medicinal chemistry uncovered more potent inhibitors and structural biology showed the location of the pyridylpiperazine binding pocket in the AcrB TM domain. Our findings bring to light a novel class of EPIs that act in a previously unexploited pocket, which likely allosterically prevents protomer cycling and the drug efflux process. Together, the presented work offers an alternative prospect in the fight against antibiotic resistance of Gram-negative bacteria.

**Results**

**Identification of new EPIs.** To identify EPIs, a phenotypic assay was established to screen a chemical library of 1280 fragments (Table S1). The molecules selected in this chemical library (originating from both commercially available molecules and unique in-house synthesized compounds) were enriched with C-sp³, spiro carbon atoms and halogen atoms 25,26. Additionally, they fulfilled the commonly accepted “rule of three” (MW < 300 g/mol, cLogP < 3, number of H-bond donors ≤ 3, number of H-bond acceptors ≤ 3) 27. The EPI screening assay searched for molecules that synergized the activity of a sub-MIC (Minimal Inhibitory Concentration) dose of the antibiotic pyridomycin 28, a particularly good substrate of the AcrAB-TolC efflux pump (MIC for E. coli BW25113 is 12.5–25 µg/mL whereas for E. coli BW25113 ΔacrB or ΔtolC mutants it is 0.78 µg/mL). Screening of the fragment library at 300 µM identified a one-hit molecule, BDM73185 (1), which prevented E. coli BW25113 growth in the presence of sub-MIC pyridomycin (5 µg/mL) but exhibited no antibiotic activity by itself. BDM73185 (1) is a small soluble molecule (MW: 265 g/mol, PBS solubility > 900 µM) with a pyridine core, two electron-withdrawing substituents (Cl and CF₃) in positions 3 and 5 and a basic piperazine moiety in position 2 (Table 1). In validation experiments, BDM73185 (1) (300 µM) boosted the antibiotic activity of a panel of AcrAB-TolC substrates such as chloramphenicol (8-fold), pyridomycin (4-fold), tetracycline (4-fold), erythromycin and chloroxacin (2-fold), but did not alter the activity of non-AcrB substrates streptomycin and kanamycin (Table S2). In addition, BDM73185 (1) did not mediate antibiotic boosting in E. coli ΔacrB, ΔacrA, or ΔtolC mutants (Table S3).

**Spontaneous AcrB point mutations confer BDM73185 resistance.** To validate AcrAB-TolC as the BDM73185 (1) target, spontaneous E. coli mutants resistant to BDM73185 (1)-mediated antibiotic boosting were selected, without changing the basal antibiotic susceptibility in the absence of the inhibitor. Selection of E. coli BW25113 on solid media containing 600 µM BDM73185 (1) and different sub-inhibitory concentrations of erythromycin (1.25, 2.5, or 5 µg/mL) resulted in the selection of 2, 1, and 1 resistant colony, respectively (frequency of resistance ~1.5 × 10⁶). The four selected resistant isolates showed no change in basal susceptibility to either erythromycin or pyridomycin (confirming no change in the basal activity of the AcrAB-TolC efflux pump), and this antibiotic susceptibility was no longer boosted by BDM73185 (1) (Table S4). Targeted Sanger sequencing found that all four resistant isolates carried a non-synonymous mutation of acrB coding for proximal residues in the 5th TM helix (TM5) of AcrB (3 mutants with t1348c [S450P], and one with g1336c [A446P]). Whole-genome sequencing and variant analysis confirmed that no other mutations were selected in these BDM73185 (1) resistant mutants, implying that the resistance phenotype selected was linked solely to an acrB point mutation. These mutations in TM5 are spatially distant from the periplasmic AcrB regions targeted by known EPIs such as PAβN 29, MBX series 19 and NMP 30. Concordantly, the canonical AcrB inhibitors PAβN (15 µg/mL) and NMP (250 µg/mL) were found to still boost pyridomycin antibiotic activity in the BDM73185 (1) resistant strains (Table S5). As an additional validation, a reverse genetics approach by recombineering was used to specifically reintroduce wildtype acrB, acrB t1348c [S450P], or acrB g1336c [A446P] into its natural chromosomal locus in E. coli ΔacrB, and these engineered strains showed the same BDM73185 (1) resistance phenotype as the selected strains (Table S6). Together, these data imply that the identified acrB mutations are solely responsible for the BDM73185 (1) resistance phenotype.

**Medicinal chemistry.** While BDM73185 (1) was the cornerstone in discovering this class of EPIs, it lacked potency and was unable to revert pyridomycin efficacy to that of E. coli ΔacrB and E. coli ΔtolC. Therefore, structure-activity relationships (SAR) around
the pyridylpiperazine hit were investigated to generate more potent analogues. This SAR study interrogated the role of the piperazine ring, the trifluoromethyl group and the chlorine atom of BDM73185 (1) (Table 1).

SAR analysis of the piperazine ring showed that the basic nitrogen was necessary for pyridomycin boosting activity in *E. coli*, as its replacement by oxygen (2) or a carbon atom (3) resulted in inactive compounds (EC$_{90}$ > 500 µM). Addition of a methyl group on the basic nitrogen (4) led to a 6-fold loss in potency (EC$_{90}$ = 375 ± 125 µM) suggesting a possible steric hindrance.

Replacement of the hydrophobic trifluoromethyl group with a more polar methoxy group (6, EC$_{90}$ = 94 ± 32 µM) caused a slight

**Table 1 Biological activities of compounds 1-12.**

| Compound | A   | R1  | R2  | R3  | MIC$_{90}$ (µM)$^a$ | EC$_{90}$ (µM)$^b$ | EPI antibiotic activity | EPI antibiotic boosting activity |
|----------|-----|-----|-----|-----|---------------------|---------------------|-------------------------|-----------------------------|
| 1 (BDM73185) | NH  | Cl  | CF$_3$ | H  | > 500               | 62 ± 21             |                         |                             |
| 2        | O   | Cl  | CF$_3$ | H  | > 500               | > 500               |                         |                             |
| 3        | CH$_2$ | Cl  | CF$_3$ | H  | > 500               | > 500               |                         |                             |
| 4        | N-CH$_3$ | Cl  | CF$_3$ | H  | > 500               | 375 ± 125           |                         |                             |
| 5        | NH.HCl | Cl  | CH$_3$ | H  | > 500               | 62                  |                         |                             |
| 6        | NH.HCl | Cl  | OCH$_3$ | H  | > 250               | 94 ± 32             |                         |                             |
| 7        | NH.HCl | Cl  | Br   | H  | > 500               | 32                  |                         |                             |
| 8 (BDM88832) | NH  | Cl  | I    | H  | > 500               | 12 ± 4              |                         |                             |
| 9 (BDM88855) | NH  | Cl  | 5  | 6  | > 250               | 3.4 ± 0.7           |                         |                             |
| 9' (BDM88855.HCl) | NH.HCl | Cl  | 5  | 6  | > 250               | 3.6 ± 0.5           |                         |                             |
| 10       | NH.HCl | H   | 5  | 6  | > 250               | > 250               |                         |                             |
| 11       | NH.HCl | Br  | 5  | 6  | > 250               | 1.5 ± 0.5           |                         |                             |
| 12       | NH.HCl | I   | 5  | 6  | > 250               | 3 ± 1               |                         |                             |

$^a$MIC$_{90}$ represents the minimal inhibitory concentration of tested compounds that prevent 90% of *E. coli* BW25113 growth as measured by resazurin reduction.

$^b$EC$_{90}$ represents the effective concentration of tested compounds that prevents the growth of *E. coli* BW25113 in the presence of 8 µg/mL pyridomycin as measured by resazurin reduction (the MIC$_{90}$ of pyridomycin alone is 12.5–25 µg/mL). Data are the result of at least two biological replicates and are presented as mean values ± SEM. Source data are provided as a Source Data file.
decrease in activity. In contrast, replacement with a hydrophobic methyl group (5, EC₉₀ = 62 µM) did not impact compound potency, suggesting that hydrophobicity is an important criterion for activity. This was confirmed by the improved potency observed with the introduction of bromine (7, EC₉₀ = 32 µM) or an iodine atom (BDM88832 (8), EC₉₀ = 12 µM) instead of the trifluoromethyl group. It was therefore decided to replace the 5-iodopyridine with a quinoline ring (BDM88855 (9) and BDM88855.HCl (9’)), leading to a 15-fold more potent compound than the initial hit. Finally, removal of the chlorine atom (R1 group) of the quinoline ring significantly decreased activity (10, EC₉₀ > 250 µM), whereas its replacement by other halogen atoms was well tolerated (11 and 12, EC₉₀ = 1.5 µM and 3 µM, respectively). None of the compounds showed antibacterial activity by themselves at the highest concentration tested. Overall, this SAR study led to the identification of fragment-sized pyridylpiperazine derivatives (compounds 9, 11, and 12) 15 to 30 times more potent than the initial hit. Both BDM73185 (1) and BDM88855.HCl (9’) did not induce apoptosis (CC₉₀ > 100 µM) in cytoxicity assays on BALB/3T3 cells following 24 and 48 h of exposure and were therefore considered non-cytotoxic in vitro.

The spectrum of antibiotic boosting by BDM88855. The activity of a large panel of antibiotics was determined on wild-type *E. coli* in the absence and presence of 100 µM BDM88855.HCl (9’). The antibiotic activity was also determined on *E. coli ΔacrAB*, which served as a proxy for full AcrB inhibition, helping clearly define the AcrB antibiotic substrates (Table 2). Data confirmed that all the AcrB substrates antibiotics were efficiently boosted by BDM88855.HCl (9’) in wild-type *E. coli*, to levels similar to that observed for *E. coli ΔacrAB*. Parallel experiments evaluating this antibiotic boosting by BDM88855.HCl (9’) in the pyridylpiperazine resistant *E. coli* isolates (with AcrB mutations S450P or A446P), showed these isolates to be cross-resistant to BDM88855.HCl (9’), confirming that these analogues of the initial hit retained activity on AcrB (Table S7).

Checkerboard assay to evaluate antibiotic boosting. To compare the efficacy of BDM88855.HCl (9’) and PAßN, their concentration-dependent boosting of pyridomycin and oxacillin activity was determined using a checkerboard assay. Data show a concentration-dependent modulation of antibiotic activity by BDM88855.HCl (9’), with boosting observed from 3.1–6.25 µM, reaching maximal inhibition at 100–200 µM (Fig. 1). This inhibition was largely shifted in the pyridylpiperazine resistant *E. coli* isolate (carrying the S450P AcrB substitution), confirming its resistance to BDM88855.HCl (9’). In parallel assays, PAßN inhibition of pyridomycin and oxacillin started from 6.25–12.5 µM and was similar in both the wild-type and pyridylpiperazine resistant *E. coli* isolate. With oxacillin, PAßN was

| Antibiotic        | *E. coli BW25113* | *E. coli BW25113 ΔacrAB* |
|-------------------|------------------|-------------------------|
|                   | MIC (µg/mL)      | MIC (µg/mL)             |
| Oxacillin         | >100             | 3.0                     |
| Linezol           | 100              | 3.1 ± 0.07              |
| Novobiocin        | >100             | 12                      |
| Fusid acid        | >100             | 12                      |
| Pyridomycin       | 9 ± 3            | 0.75 ± 0.25             |
| Chloramphenicol   | 4.5 ± 1.5        | 0.5                     |
| Erythromycin      | 18.5 ± 6.5       | 4.0 ± 2.0               |
| Ciprofloxacin     | 0.052 ± 0.014    | 0.013 ± 0.004           |
| Piperacillin      | 0.39             | 0.10                    |
| Tetracycline      | 0.75 ± 0.25      | 0.25                    |
| Triclosan         | 0.045 ± 0.005    | 0.026 ± 0.018           |
| Ampicillin        | 4.5 ± 1.5        | 2.5 ± 0.5               |
| Ceftazidime       | 0.12             | 0.25                    |
| Streptomycin      | 3.13 ± 0.005     | 6.3                     |
| Aztreonam         | 0.060            | 0.09 ± 0.03             |
| Gentamicin        | 0.59 ± 0.2       | 1.60                    |

As a comparison, the antibiotic susceptibility of *E. coli* lacking AcrAB is also indicated.

BDM88855.HCl (9’) on its own has no impact on *E. coli* growth up to the highest concentration tested (250 µM). Data are the mean ± SEM of two independent biological replicates (SEM is not indicated when all replicates showed the same MIC). Source data are provided as a Source Data file.
Confirmation of direct inhibition of AcrB by pyridylpiperazines. To reinforce that BDM88855·HCl (9’) acts directly on AcrB, its impact was evaluated on the immediate bacterial accumulation of AcrB substrate; berberine, and biophysically, on the stabilization of the inhibitor/AcrB complex using thermal shift assay (TSA). Accordingly, fluorescence uptake assays showed that BDM88855·HCl (9’) increased berberine accumulation in E. coli expressing wild-type acrB in a concentration-dependent manner, while bacteria producing the AcrB variants A446P and S450P were de facto unaffected by the inhibitor (Fig. S1). Likewise, TSAs confirmed a concentration-dependent increment of the melting temperature of the purified wild-type AcrB, but not of the AcrB (A446P) variant (Fig. S2), indicating inhibitor-mediated stabilization of the protein/inhibitor complex for wild-type AcrB and not for the resistant mutant. Of note is that the concentration-dependent boosting of antibiotic activity, berberine accumulation and protein stabilization by TSA are congruent. Together, these results reinforce that BDM88855·HCl (9’) directly binds and inhibits AcrB.

Structural confirmation of pyridylpiperazine binding site. As evidenced by the selection of natural resistant isolates described above, we anticipated inhibitor binding at the TM domain, in contrast to the previously structurally described inhibitors targeting the periplasmic porter domain19,20. To confirm this, we determined the co-crystal structure of wildtype AcrB with BDM88855 (9) at 2.6 Å resolution, and the inactive AcrB R971A variant13 with BDM88855 (9) and BDM88832 (8) at 2.5 and 2.8 Å resolution (Table S8). Non-proteinaceous electron density between TM4, TM5 and TM10 showed that BDM88855 (9) and BDM88832 (8) bind specifically to the TM domain of the L protomer (Fig. 2, Fig. S3). Analysis of the anomalous signals originating from the iodine atom of BDM88832 (8) or from the brominated pyridylpiperazine inhibitor (compound 11) indicated the location of the halogen atoms (Fig. S3) and confirmed the binding of BDM88832 (8) and BDM88855 (9) at this position in the TM domain, with minimal halogen and hydrogen bond length deviations between the interaction of the latter inhibitor in wildtype and the R971A variant (Fig. S3d).

Superimposition of the unliganded AcrB L protomers31 (PDB: 4DX5) and inhibitor-bound structures reveals a root mean square deviation (RMSD) (Cα) of 0.89–1.05 Å over 1024 residues (Table S9), with the most obvious deviations at the TM domain (Fig. 2b, Table S9). Here, the TM helices show subtle differences in their conformation at the cytoplasmic side, leading to an expansion of the narrow channel between the D407, D408, and K940 residues of the proton relay site and the cytoplasm, thus allowing for the accommodation of BDM88855 (9) (Fig. 2b). The quinoline moiety of the inhibitor interacts with surrounding hydrophobic residues, including V411, L442, I445, and F948 (Fig. 2d, Fig. S3a, Table S10). A subtle shift of TM10 permits the formation of a halogen bond32 between the chlorine atom and carbonyl oxygen of main chain K940. The piperazine ring is mainly surrounded by residues involved in the proton relay network (D407, D408, and K940), as well as L449, and the carbonyl oxygen of the main chain L404 (Fig. 2d, Fig. S3a). The

![Fig. 2 Structure of AcrB in complex with BDM88855 (9). a Side view of the AcrB trimer comprising the L (marine), T (yellow), and O (red) protomers. BDM88855 (9) (green) binds to the TM domain of the L protomer. The boundaries of the inner membrane (IM) are indicated for the L protomer. b Side view on the TM domain of the L protomer. The apo-structure (dark blue) and the BDM88855 (9·stick representation) near the proton relay triad D407, D408, and K940. c Cytoplasmic side view into the TM domain of the AcrB L protomer (marine) and the location of BDM88855 (9·stick representation) nested between TM helices 4, 5, and 10. d Enlarged view of the inhibitor binding site showing interacting residues (marine sticks) in less or equal than 4 Å distance from BDM88855 (9).](image-url)
carboxyl moiety of D408 and carbonyl moiety of L404 are re-orientated compared to the apo-structure to allow the formation of a salt bridge with the protonated secondary nitrogen atom (\(-\text{NH}_2^+\)) of the piperazine moiety (Fig. 2d). Furthermore, the D407 and K940 side chains are re-orientated to accommodate the piperazine moiety of BDM88855 (8), (normally, K940 occupies this area in the unliganded L protomer) (Fig. S4). This re-orientation disrupts the H-bond network between S481, D407, D408, K940, and T978\(^13\), while maintaining H-bonding between S481 and D408 in the BDM88855 (9) co-structure (Fig. S5). The presence of a halogen atom in position 3 of the quinoline ring is required for inhibitor activity and correlates with the formation of a halogen bond to the main chain carbonyl oxygen of K940. In addition, the formation of a salt bridge between the charged piperazine ring and the carbonyl side chain of D408 reinforces the importance of the basic nitrogen of the piperazine ring for activity. Therefore, the inhibitor likely interferes with the proton relay site at the TM domain of AcrB and appears to lock the L protomer in this state.

**Structure-function validation of the BDM88855 binding site.** Drug susceptibility assays show that the AcrB variants I438A, I445A, I943A, and L944A, conferred increased susceptibility against all AcrB substrates tested (Fig. 2d, Table S10, Fig. S6/S7), indicating that these residues may be involved in drug transport and/or proton binding/coupling. AcrB variants V411A, L442A, Glu947A conferred wild-type like susceptibilities, in the absence and presence of BDM88855 (9) (except linezolid in case of the L442A variant) (Table S10), indicating that these substitutions result in a loss of inhibitor affinity. The alkyl side chains of V411, L442, and Glu947 orient towards the quinoline moiety of the inhibitor resulting in van der Waals interactions (Fig. S3a), so substitution with alanine (with a shorter side chain) most likely leads to a loss of inhibitor interactions and affinity. In contrast to the mutations that prevent AcrB inhibition, the functionally active F948A variant was hypersensitive to inhibition by BDM88855 (9) compared to wildtype AcrB (Table S10, Fig. S7). As F948 resides at the cytoplasmic rim of the TM domain, proximal to the inhibitor binding site, the Ala-substitution may facilitate inhibitor access from the cytoplasm towards its binding site.

**Putative pyridylpiperazine uptake into AcrB.** The location of the key residues at the cytoplasmic rim of the TM domain (Fig. 3), and the presence of water accessible channels from the cytoplasmic side of this region in the L state\(^3\), suggest that the inhibitor gains access to its binding pocket via the cytoplasmic side. To assess this hypothesis, we performed all-atom molecular dynamics (MD) simulations\(^33\) of the interaction between one hundred molecules of BDM88832 (8) and wild-type AcrB (Fig. S9). Computational data confirmed the tendency of one or more inhibitor molecules to persistently reside in a channel linking the cytoplasm and the BDM88832 (8) binding site, as inferred from the crystal structure in the L protomer of AcrB (Fig. 3). A lower density of compounds was seen for both the T and O protomers. BDM88832 (8) interacted mostly with residues V411, L442, E947, and F948 (Fig. S10) as it travelled from the cytoplasm towards the binding pocket. A persistent interaction between the inhibitor and F948 could rationalize the observed hypersensitive F948A phenotype in comparison to wild-type AcrB. Overall, these results validate the AcrB/inhibitor co-structures and provide strong evidence that pyridylpiperazine inhibitors access AcrB from the cytoplasm rather than the periplasm.

**Discussion**

In Gram-negative bacteria, RND efflux pumps play a central role in both innate and acquired antibiotic resistance by extruding a plethora of antimicrobial compounds. Therefore, the discovery and development of potent efflux pump inhibitors would allow for improved antibiotic efficacy and drug repurposing\(^34,35\). In addition, preventing antibiotic efflux would greatly improve the success rate of drug discovery programs, where RND-pump compound extrusion is the origin for low hit rates (New drugs for bad bugs\(^36\)). Some RND-efflux pump inhibitors have already been discovered, including PAβN and derivatives, pyrido-pyrimidine (such as D13-9001), and the pyranopyridine derivatives (MBX series). PAβN is a broad spectrum competitive inhibitor of the DBP of RND pumps (but also targets the outer membrane LPS layer\(^37\)), and was recently shown to perturb the functional dynamics of AcrB\(^38\).

Both the pyrido-pyrimidine and pyranopyridine inhibitors bind to a specific “hydrophobic trap” in the DBP in the T protomer of asymmetric AcrB (Fig. 4), either preventing substrate binding or inhibiting the T to O transition required for the functional catalytic cycle of the RND pump\(^19,20\). As the DBP is a lipophilic pocket, medicinal chemistry for optimizing the drug-like properties of hit inhibitor molecules can be challenging. To facilitate downstream medicinal chemistry, in this...
work we opted to screen a selected library of small hydrophilic fragments. Due to their physicochemical properties, it was anticipated that more hydrophilic fragments might bind to alternative (non-DBP) sites of AcrB.

A breakthrough in this work was the selection of natural E. coli point mutants specifically resistant towards the pyridylpiperazine-based EPIs (at a low frequency). These mutations were located in the AcrB TM domain, suggesting a hitherto undiscovered mechanism of efflux pump inhibition, a finding that was consolidated by structural biology data. Additional mutagenesis data suggest that these EPIs act directly on the AcrAB-ToIC efflux pump, rather than on its transcriptional/translational regulation or on other putative targets.

The functional state of the AcrB trimer comprises three protomers in different conformational states, the L, T, and O protomers. From the crystal co-structure, it is apparent that the pyridylpiperazine EPIs bind uniquely to the L protomer of AcrB. MD simulations also predicted pyridylpiperazine inhibitors to preferentially interact with the cytosolic surface of the L protomer. Specifically, simulations indicated that compounds enter AcrB in the L protomer state via a cytoplasmic-open access channel leading to the inhibitor binding pocket. As the L protomer TM domain is in an inward open conformation, this entry pathway is more accessible than in the T or the O protomer, which are in an outward open (inward closed) or occluded conformation, respectively.

The elucidated crystal structure of the pyridylpiperazine binding pocket in AcrB points to two possible mechanisms by which these EPIs inhibit AcrB function. Firstly, by binding deep in the TM domain of the L protomer, the pyridylpiperazine may sterically prevent the L to T transition, a vital process involving major structural movement in the TM domain from an inward open to an outward open conformation. An alternative possible mechanism of action is that the pyridylpiperazine impedes AcrB from using the proton motive force as its energy source for substrate translocation. From the identified inhibitor binding pocket, the protonated piperazine moiety of the pyridylpiperazine forms a critical salt bridge with the side chain of D408. D408, together with D407, K940, and R971 are essential catalytic residues of the AcrB proton relay, and the interaction of the pyridylpiperazine with D408 likely interferes with the H+-translocation pathway. Interestingly, a similar mechanism of action has been proposed for a number of anti-tuberculosis drugs targeting the Gram-positive RND-type mycolic acid efflux pump, MmpL3.36

The elucidated crystal structure of the pyridylpiperazine series as a novel class of efflux pump inhibitors that act by binding a previously unexploited pocket in the TM region of the L protomer of AcrB. Our data suggest that occupancy of this conserved pocket prevents the functional catalytic cycle of this RND pump resulting in enhanced drug susceptibility. Together, these lead molecules provide a promising basis for optimization into potent drug-like EPIs for in vivo efficacy and clinical development to disarm Gram-negative bacteria.

**Methods**

**Strains, media, and antibiotics.** *Escherichia coli* BW25113 (CGSC 7636) and its derivatives *E. coli ΔtolC* (JW5503-1, ΔtolC732::kan, CGSC 11430), *E. coli ΔacrA* (JW0452-3, ΔacrA748::kan, CGSC 11843) and *E. coli ΔacrB* (JW0451-2, ΔacrB747::kan, CGSC 8609) were obtained from *E. coli* Genetic Stock Center (CGSC, New Haven, Connecticut) and originated from the Keio Collection.14 Bacterial were regularly cultured on LB broth (BD, DIFCO) at 37°C. All bacterial antibiotic susceptibility tests were performed in cation-adjusted Mueller-Hinton broth (CAMHB; BD Difco) at 37°C. Commercially available molecules, including antibiotics and efflux pump inhibitors, were purchased from various vendors, including Sigma–Aldrich, Carbosynth Limited, Fisher Scientific, Euromedex, and details can be provided on request. Pyridomycin was extracted and purified from *Dactylosporangium fulvum*, as previously described.

**Screening library.** To screen for potential EPIs, a chemical library of 1280 compounds (at 100 mM in DMSO) were transferred (in duplicate) by acoustical technology (Echo® 550, Labcyte Inc) to a destination 384-well plate (150 nL/well, the...
**Chemical synthesis** can be found in the supplementary methods section and structural validation of the compounds can be found in the source data files.

**Liquid Mic determination.** The minimum inhibitory concentration (MIC) determination was performed in 96-well format in order to confirm screening results, define EPI activity on bacterial mutants, and evaluate the activity of improved EPIs. Briefly, bacterial cultures were diluted to OD<sub>600</sub> of 0.001 in CAMHB and spiked with either EPI (to determine the shift in antibiotic MIC, typically 500 µM BDM73185 (1) or 100 µM BDM88855.HCI (9)) or with AcrAB-ToCA substrates (to determine the activity of EPI typically 8 µg/mL pyrromycin). These bacterial/compound mixes were then transferred to a 96-well plate (100 µL well) and a dose-response of the compounds of interest was added by serial dilutions. Following culture (5 h, 37 °C), strain viability was evaluated using the resazurin reduction assay, and measured by fluorescence (POLARstar Omega, BMG Labtech: Ex= 530 nm Em: 590 nm). EC<sub>90</sub> was determined based on the ratio of the apoptotic cell fluorescence (POLARstar Omega, BMG Labtech: Ex= 530 nm Em: 590 nm). EC<sub>90</sub> was determined based on the ratio of the apoptotic cell fluorescence. For compounds with a 0.001, spiked with DMSO or 100 µM BDM88855.HCI (9) and added to the destination plate (50 µL per well). Viability was determined as described above in 5 h co-culture.

**Cytotoxicity.** The cytotoxicity of compounds BDM73185 (1) and BDM88855.HCI (9) was determined using BALB/c3T3 cells using live imaging following both Hoechst 33342 and NucView 488 Caspase-3 staining. Briefly, BALB/c3T3 cells were seeded in 384-well plates. Four hours later, compounds with a concentration of 80 nM BDM73185 (1) or a concentration of 100 µM to the culture medium, as well as Hoechst 33342 and NucView 488 Caspase-3 substrate. 24 h and 48 h after compounds addition, live imaging was performed using an In Cell Analyzer 6000 (GE Healthcare). The cytotoxicity was defined based on the ratio of the apoptotic cell population (NucView staining) and the total population (Hoechst staining) using Columbus software. Compounds were tested in triplicate. Carfilzomib (499, 249,142, 61, and 30.6 nM) was used as a positive control in this assay.

**Isolation of BDM73185 (1)-resistant mutants.** The selection of E. coli BDM73185 (1)-resistant clones was performed by plating 50 µL of concentrated log-phase BW25113 culture at OD<sub>600</sub> = 8 onto CAMHB agar containing either erythromycin alone (1.25, 2, 5, 10, 20, or 40 µg/mL) or erythromycin with 600 µM BDM73185 (1). Plates were incubated at 37 °C until the appearance of resistant colonies (1–2 days). Clones were picked and grown in CAMHB without selective pressure, before confirmation of antibiotic susceptibility (erythromycin, pyromycin, and linezolid) in the absence and presence of BDM73185 (1) at 300 µM.

**gDNA extraction and acrB variant analysis.** E. coli BW25113 wild-type and the four BDM73185 (1)-resistant clones were grown overnight in CAMHB medium at 37 °C. Cells (1 mL) were collected by centrifugation, re-suspended in Phosphate Buffered saline (500 µL) and were broken by bead beating using the MP Biomedicals<sup>324</sup> Instrument FastPrep<sup>24-34</sup> (MP Biomedicals, Santa Ana, California). Lyssates were collected and incubated with 4.5 µL of 20 mg/mL RNaseA (PureLink<sup>®</sup> RNase A, Thermofisher scientific) for 10 min at room temperature. DNA was extracted as per standard phenol-chloroform-isooamyl alcohol (25:24:1) procedure (Thermofisher scientific). Genomic DNA and purity were checked by Denovix DS-11 Series (Denovix, Wilmington, Delaware), and acrB sequenced by Sanger sequencing (Genoscreen, France).

**Whole-genome sequencing and variant analysis.** For whole-genome sequencing, genomic DNA from bacteria was subjected to MicrobesNG (http://www.microbesng.com, Birmingham, UK) for library preparation (Nextera protocol) and whole-genome sequencing on an Illumina sequencer (Novaseq6000). Trimmed reads (depicted at NCBI BioProject ID: PRJNA764862) were initially processed using PRINSEQ-lite PERL script (PRINSEQ-lite version 0.20.4<sup>44</sup>) to remove low-quality data with the following parameters (-min_len 50 -min_qual 30 -trim_left 0 -trim_right 0 -trim_min 0 -nonindup). Reads were then aligned to the E. coli BW25113 reference genome (NCBI accession number CP009973) with BWA-MEM version 0.7.17-r118843<sup>4</sup>. The output SAM alignment files were converted to BAM files and sorted using SAMtools version 1.1v with default parameters<sup>14</sup>. SNPs and Indels were called using GATK HaplotypeCaller Version 4.2.0<sup>45</sup>. Identified variants were finally annotated using snpsEff version 4.3<sup>46</sup>.

**Construction of E. coli BW25113 acrB chromosomal point mutants.** To generate unmarked single point mutant clones, an E. coli BW25113 AcrB mutant was generated (chromosomal reconstitution) using lambda-Red recombination with either wildtype acrB, acrB S450P, or acrB A446P. Briefly, unmarked BW25113 AcrB was generated from E. coli ΔacrB (BW0451-2, ΔacrB::kanR) strain by replacement of the FRT-Kan-FRT recombinase cassette using lambda-Red recombination transformation with PFLP2<sup>47</sup>. A resulting AmpR KmR mutant was streaked on LB-5% sucrose agar plates to cure the bacteria of PFLP2. This unmarked E. coli BW25113 ΔacrB strain was then transformed with the inducible Red-recombinase plasmid; pEP136<sup>48</sup> and AmpR resistant clones were selected at 30 °C. Arabinose induced ΔacrB electroporation was preformed at 30 °C and transformed with three different PCR products containing either wildtype acrB, acrB S450P, or acrB A446P. Specifically, these 3.9 kb PCR products were generated by amplifying pyridomycin (primers 5′-ATACATCCAGGCGGTATCCT-3′ and 5′-GGCGACGCGATTAAAAGCACGTT-3′) from wild-type E. coli, or the two selected plasmids<sup>48</sup> and lincomycin, and linezolid) in the absence and presence of BDM73185 (1) (50 µL per well), and a dose-response of the compounds of interest was added by serial dilutions. Following culture (5 h, 37 °C), strain viability was evaluated using the resazurin reduction assay, and measured by fluorescence (POLARstar Omega, BMG Labtech: Ex= 530 nm Em: 590 nm). EC<sub>90</sub> was determined based on the ratio of the apoptotic cell fluorescence. For compounds with a 0.001, spiked with DMSO or 100 µM BDM88855.HCI (9) and added to the destination plate (50 µL per well). Viability was determined as described above in 5 h co-culture.

**Whole-cell accumulation assay.** Semi-quantitative accumulation assays were carried out to further characterize the impairment of substrate efflux in the presence of BDM88855.HCI (9). The accumulation of the pump substrate berberine by E. coli BW25113 ΔacrB/pET24a_ (WT, D407N, A446P or S450P) leaky expression system was monitored over time in the presence of different BW25113 ΔacrB (9) concentrations.

**AcrB** was assayed according to previous experiments<sup>20</sup> with some modifications. In brief, chemically competent E. coli BW25113 AcrB cells were transformed with the corresponding pET24a_ aacrB variants WT, D407N, A446P, or S450P, plated on LB, 50 mg/mL kanamycin (LB-K50) agar and incubated overnight at 37 °C. Colonies (1-2 mm) were picked, transferred to a 96-well plate (1 mL per well), and a dose-response of the compounds of interest was added by serial dilutions. Following culture (5 h, 37 °C), strain viability was evaluated using the resazurin reduction assay, and measured by fluorescence (POLARstar Omega, BMG Labtech: Ex= 530 nm Em: 590 nm). EC<sub>90</sub> was determined based on the ratio of the apoptotic cell fluorescence. For compounds with a 0.001, spiked with DMSO or 100 µM BDM88855.HCI (9) and added to the destination plate (50 µL per well). Viability was determined as described above in 5 h co-culture.

**Isolation of BDM73185 (1)-resistant mutants.** The selection of E. coli BDM73185 (1)-resistant clones was performed by plating 50 µL of concentrated log-phase BW25113 culture at OD<sub>600</sub> = 8 onto CAMHB agar containing either erythromycin alone (1.25, 2, 5, 10, 20, or 40 µg/mL) or erythromycin with 600 µM BDM73185 (1). Plates were incubated at 37 °C until the appearance of resistant colonies (1–2 days). Clones were picked and grown in CAMHB without selective pressure, before confirmation of antibiotic susceptibility (erythromycin, pyromycin, and linezolid) in the absence and presence of BDM73185 (1) at 300 µM. Brie
30 min. The protease inhibitor PMSF (200 µM, final) was added directly before cell disruption with a pressure cell homogenizer (Stansted Fluid power Ltd.), 2× at times of ~15 min using a cell disruption plate (in 800 mL PCW, 20% w/v stock) and 12 mM imidazole (1 M stock, pH 7.5) in a final volume of 12 mL for at least 30 min under mild sample rotation. Buffer P1 was used for volume adjustment. After ultra-centrifugation (160,000 × g, 30 min) the supernatant was rotated with 400 mL (1 M) suspension. The protein was eluted with 4 mL P1, 400 mM imidazole in a further volume of 4 mL P1. Eluates were concentrated in an Amicon Ultra-100 kDa spin concentrator to 500 µL, applied and incubated on a Cytiva Superose 6 increase 10/300 GL size exclusion column for at least 30 min. The resin was washed in a gravity flow column with each 15 CV (6 mL) of P1 buffer (20 mM Tris pH 7.5, 300 mM NaCl, 0.02% DDM) supplement with (i) 20 mM imidazole and (ii) 80 mM imidazole. The protein was diluted to 0.1 mg/mL for TSA analysis.

Drug agar plate assay. A colony of E. coli BW25113 ΔacrB harbouring pET24acrB-His encoding wild-type (WT) or variant AcrB were grown overnight in LB medium containing 50 μg/mL kanamycin at 37 °C. Dilution of the cultures to OD600 = 10−4 was performed and 4 μL of each diluted culture were spotted on LB agar plates containing 50 μg/mL kanamycin, supplied drugs (12.5 μg/mL dicyclohexyl, 12.5 μg/mL fusicid acid, 12 μg/mL lincosolid, or 80 μg/mL TIP). Plates were incubated at 37 °C for 14–16 h. Agar plates were imaged by ImageQuant TL (GE Healthcare Biosciences AB, Uppsala, Sweden). The intensity of the cell growth (Table S10) was quantified by ImageJ 1.52o software. The cell growth of WT at first dilution is set to 1. The intensity of cell growth at each dilution higher than 10% of cell growth of WT at first dilution is considered as cell growth. To calculate the relative cell growth of each mutant, the cell growth of each mutant is normalized with cell growth of WT after deduction of the cell growth of cells harbouring the inactive AcrB ΔacrB(T) variant (cell growth = 0). E.g. if the cell growth of harbouring AcrB WT is 4 dilution steps, the fourth dilution step is set to 100% growth. Growth of cells harboring AcrB variants lower than 75% of the growth of cells containing AcrB WT is considered significantly affected if p < 0.005 or slightly affected if p < 0.05. Growth of cells harbouring AcrB variants in between 75–85% of growth of cells harbouring AcrB-WT is considered as slightly affected if p < 0.005, p value was calculated by two-sided Student’s t-test (Table S10).

Overproduction and purification of DARPinS. A single colony of E. coli XLI-1 Blue cells harbouring pQE30-DARP16,61 was inoculated in LB liquid medium supplemented with 50 μg/mL kanamycin at 37 °C and cultivated overnight. Overnight cultures were inoculated into a fresh LB liquid medium supplemented with antibiotic as described above and grown until OD600 = 0.8 for 3 h. Cell membranes were resuspended in Buffer B (20 mM Tris-HCl buffer at pH 7.5, 150 mM NaCl, 20 mM Imidazole and 10% Glycerol and solubilized with 1% dodecyl maltoside (D-97002-C, DDM, Glycos) at 4 °C for 1 h. Solubilized membranes were cleared at 160,000 × g for 30 min and the supernatant was loaded onto a HisTrap HP Ni2+ affinity column (1 mL bed volume, GE Healthcare). The column was washed with Buffer B supplemented with 0.2 M NaCl, 0.6 M and 90% EtOH for 30 min. The protein was eluted with Buffer D (20 mM Tris-HCl buffer at pH 7.5, 150 mM NaCl, 220 mM imidazole, 10% glycerol and 0.02% DDM), for 30 min before setting up crystallization, AcrB was mixed with DARPinS in a molar ratio of 2:1 and incubated on ice for 1 h. DARPinS were mixed by size exclusion chromatography (Superose 6, GE Healthcare) with buffer containing 20 mM Tris pH 7.5, 150 mM NaCl, 0.03% DDM and 0.05% DDAO (Anatrace). Co-crystallization of AcrB/DARPin variants with various inhibitors were obtained by hanging drop crystallization within 1–2 weeks. Briefly, BDM88832 (8) or BDM88859 (9) (6–8 mM) were mixed with a protein solution (12 mg/mL final concentration), and incubated on ice for 30 min. Subsequently, all the protein solutions were centrifuged at 13,000 × g for 10 min at 4 °C to remove the insoluble materials, before setting up the crystallization. The reservoir contained 50 mM NaCl, 10% (w/v) PEG4000 and 10–120 mM ammonium sulfate. Crystals were harvested after 2 months and cryo-protected by serial transfer of crystals into reservoir solution supplemented with increasing of glycerol concentration to 28% before flash-cooling in liquid nitrogen.

Diffraction data collection and refinement. Data were collected on beamline PROXIMA 2 A ( Soleil Synchrotron, Saint Aubin, France) using an Eiger detector (Dectris), respectively, indexed and integrated with XDS65. All the structural models were iteratively built-in COOT66 with a structural model (PDB ID: 2BM1)67 and refined with REFMACS in CCP4i v768. The structure was validated with MolProbity v4.59. Polder electron density maps were calculated by phenix.polder57. All figures were generated by Pymol (Schrödinger LLC).
tool of AMBER and atomic charges derived from an electrostatic potential distribution calculated with Gaussian as in ref. Each system was first subjected to a multi-step structural relaxation via a combination of steepest descent (5000 steps) and conjugate gradient (up to 25000 steps) methods using the pmemd program implemented in AMBER18, as described previously 39,38,36,60–68. The systems were then heated from 0 to 310 K in two steps: (i) from 0 to 100 K in 1 ns under constant-volume conditions, with harmonic restraints (k = 1 kcal mol\(^{-1}\) A\(^{-2}\)) applied to the heavy atoms of both the protein and the lipids; (ii) from 100 to 310 K in 5 ns under constant pressure (set to a value of 1 atm) with restraints on the heavy atoms of the protein and on the z coordinates of the phosphorous atoms of the lipids, so as to allow membrane rearrangement during heating. As a final step before production runs, a series of 20 equilibration steps, each of 500 ps in duration (total 10 ns) and with restraints on the protein coordinates, were performed to equilibrate the simulation. These equilibration steps were carried out under isotropic pressure scaling using the Berendsen thermostat, whereas a Langevin thermostat (with a collision frequency of 1 ps\(^{-1}\)) was used to maintain a constant temperature. Finally, five MD simulations of 2 μs each were performed on an isothermal-isobaric ensemble. A time step of 2 fs was used for all runs before production, while the latter runs were carried out with a time step of 4 fs after hydrogen mass repartitioning 69. R–H bonds lengths were constrained using the SHAKE algorithm, and coordinates were saved every 500 ps. The Particle mesh Ewald algorithm was used to evaluate long-range electrostatic forces with a non-bonded cut-off of 9 Å.

**Post-processing of the MD trajectories.** MD trajectories were analyzed using either in-house tcl and bash scripts or the cpptraj tool of AMBER18. Hydration of the TM region of the protein (including the residues lining the binding sites of BDM88832 (8) and BDM88855 (9)) was estimated by calculating the average density values along the cumulative trajectory, using the grid tool of AMBER18. Densities were normalized to the value of bulk water (1 g/mL). Figures were prepared using xmgrace (https://plasma-gate.weizmann.ac.il/Grace/) and VMD 1.9.4 (23).

**Reporting summary** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability** Atomic coordinates and structure factors reported in this paper have been deposited in the Protein Data Bank under accession numbers 7OKU (BDM88855 inhibitor bound to the transmembrane domain of AcrB), 70UL (BDM88832 inhibitor bound to the transmembrane domain of AcrB-R971A), 70UM (BDM88855 inhibitor bound to the transmembrane domain of AcrB-R971A). Atomic coordinates that were used and support the findings of this study are available in the Protein Data Bank under accession numbers 4DX5, 4DX7, 4U96, and 5JMN. Whole-genome sequencing data (fastq files) of the multidrug transporter AcrB were deposited in GenBank (BioProject ID: PRJNA764862). Source data are provided with this paper.

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**Author contributions**

C.P., H.K.T., A.V.D.C., N.C., J.C.J.C., W.E.F., G.M., E.P., A.V.V., R.T.M., K.M.P., M.F., and R.C.H. designed experiments. C.P. and Ad.H. performed EPI screening and C.P. performed selection and characterization of resistant isolates. N.W. and M.F. designed the FLP-based system and B.D. advised on compounds design. A.V.D.C., N.C., A.R., and P.M. performed the chemical synthesis. C.P. and E.P. generated genetic knockouts. I.C.J.C. performed checkerboard assays. An.H. performed protein production, membrane preparation, and western blot experiments. K.H.T., W.E.F., M.A.K., and An.H. performed the protein purification, crystallization, data collection, and biochemical/microbiological assay. K.H.T., W.E.F., and K.M.P. determined crystal structures. R.T.M. conducted berberine accumulation and thermal shift assays. G.M. contributed compounds of BDM compounds. A.V.V. performed MD simulations and analysed simulation data. All authors contributed to data analysis. C.P., H.K.T., A.V.D.C., N.C., W.E.F., N.W., G.M., A.V.V., K.M.P., M.F., and R.C.H. wrote the paper. K.M.P., M.F., and R.C.H. obtained funding for this work.

**Competing interests**

C.P., H.K.T., A.V.D.C., N.C., J.C.J.C., R.T.M., K.M.P., M.F., and R.C.H. are inventors on patent application covering the EPI described in this manuscript. The remaining authors declare no competing interests.

**Additional information**

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**Correspondence** and requests for materials should be addressed to Klaas M. Pos, Marion Flipo or Ruben C. Hartkooij.

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