Development of a Novel Antimicrobial Screening System Targeting the Pyoverdine-Mediated Iron Acquisition System and Xenobiotic Efflux Pumps

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Academic Editor: Peter J. Rutledge

Received: 9 February 2015 / Accepted: 23 April 2015 / Published: 29 April 2015

Abstract: The iron acquisition systems in Pseudomonas aeruginosa are inducible in response to low-iron conditions and important for growth of this organism under iron limitation. OprM is the essential outer membrane subunit of the MexAB-OprM xenobiotic efflux pump. We designed and constructed a new model antimicrobial screening system targeting both the iron-uptake system and xenobiotic efflux pumps. The oprM gene was placed immediately downstream of the ferri-pyoverdine receptor gene, fpvA, in the host lacking chromosomal oprM and the expression of oprM was monitored by an antibiotic susceptibility test under iron depleted and replete conditions. The recombinant cells showed wild-type susceptibility to pump substrate antibiotics, e.g., aztreonam, under iron limitation.
and became supersusceptible to them under iron repletion, suggesting that expression of oprM is under control of the iron acquisition system. Upon screening of a chemical library comprising 2952 compounds using this strain, a compound—ethyl 2-(1-acetylpiperidine-4-carboxamido)-4,5,6,7-tetrahydrobenzo[b] thiophene-3-carboxylate—was found to enhance the efficacy of aztreonam under iron limitation, suggesting that the compound inhibits either the iron acquisition system or the MexAB-OprM efflux pump. This compound was subsequently found to inhibit the growth of wild-type cells in the presence of sublethal amounts of aztreonam, regardless of the presence or absence of dipyridyl, an iron-chelator. The compound was eventually identified to block the function of the MexAB-OprM efflux pump, showing the validity of this new method.

Keywords: antimicrobials; iron acquisition systems; efflux pumps; Pseudomonas aeruginosa

1. Introduction

Infections caused by multidrug resistant bacteria are a serious problem in hospitals and a major public health concern these days [1,2]. The discovery of new antimicrobial agents to combat such pathogens is an urgent and crucial concern [3–5]. One of the approaches may be to search for essential metabolic targets through genome analyses and subsequently perform target-based high-throughput in vitro screenings of novel lead compounds [6]. However, the agents that act upon essential metabolic targets inevitably select resistant mutants due to their strong selective pressure [7,8]. Thus, alternative approaches may be needed to tackle these issues. One such approach could be to find agents which interfere with the action of bacterial virulence factors. If anything, virulence factors are less likely to exert selective pressure on the development of resistant strains since they are virtually secondary [9–11]. Thus, means to control bacterial virulence open a window to search for new types of antimicrobial agents.

The iron acquisition system of bacteria is an important virulence factor for the establishment of infection [12,13]. Bacterial cells generally require micromolar orders of free iron to support their growth [12]. However, the human body sequesters iron molecules through high-affinity iron-binding proteins such as transferrin, lactoferrin, and ferritin, lowering the free-iron level to about $10^{-18}$ M or less [13,14]. Bacterial cells cannot grow under these iron concentrations [12–14]. To overcome this difficulty, bacterial cells elaborate specific iron acquisition systems under the iron-limited environment inducing high affinity iron chelator molecules and their transport systems [13,15]. The iron acquisition system in Pseudomonas aeruginosa, for instance, works as follows: in iron-limiting environments, the cells secrete high affinity iron chelator molecules, e.g., pyoverdine, which trap iron molecules and the iron-pyoverdine complex binds with the outer membrane receptor, FpvA. This signal may be transmitted to the inner membrane protein anti-σ factor, FpvR, and the interaction of the iron-pyoverdine-FpvA complex with the periplasmic domain of FpvR releases bound extracytoplasmic σ factors, PvdS and FpvI, promoting the transcription of the genes encoding pyoverdine biosynthetic proteins, its receptor FpvA, a virulent factor ToxA, and a protease PrpL [15–17]. In addition, another global regulator, Fur, negatively controls the transcription of many genes involved in the iron utilization. In iron limiting environments, extracytoplasmic σ factors such as fpvI and pvdS mediate Fur action resulting in activation...
of toxA and prpL [18]. The iron-uptake systems in bacteria are tightly regulated to an optimum level, since excessive intracellular iron molecules may cause the generation of oxygen radicals, which are highly toxic to the cells under aerobic conditions [12]. On the other hand, iron deficiency results in restriction of bacterial growth [12,13]. Thus, blocking the iron acquisition signal transduction circuits would be a potential means to hamper bacterial growth.

The protein PvdN, which is essential for the biogenesis of pyoverdine, is translocated to the periplasmic space across the inner membrane via the twin-arginine protein translocation machinery (Tat) [19]. Hence, blocking of the Tat protein-secretion system results in an inability to produce pyoverdine, leading to iron deficiency in P. aeruginosa under iron limited conditions.

The MexAB-OprM xenobiotic efflux pump consists of the inner membrane transporter MexB, the outer membrane duct protein OprM, and the periplasmic clump protein MexA [20]. The OprM protein functions as a duct in several other efflux-pump systems [20–22]. The pump is intrinsically expressed in the wild-type strain of P. aeruginosa rendering the cells resistant to multiple classes of antibiotics. The cells lacking the pump protein became supersusceptible to the pump substrate antibiotics, such as aztreonam, chloramphenicol and others [20]. Therefore, pump inhibitors represent powerful auxiliary agents to potentiate antibiotics. The aim of this study was to construct a model antimicrobial screening system targeting the iron acquisition systems, the Tat protein secretion system and/or the multidrug efflux pump(s) in P. aeruginosa. Then, using this newly developed system, a chemical library was screened.

2. Results

2.1. Experimental Design for Screening of Inhibitors toward Pyoverdine-Mediated Iron-Acquisition System and Xenobiotic Efflux Pumps

Pseudomonas aeruginosa senses low iron environments and responds to them by inducing a high-affinity ferric iron chelator, pyoverdine, and its outer membrane receptor, FpvA [16,23]. If the gene coding for the outer membrane subunit of the MexAB-OprM efflux pump, oprM, is placed immediately downstream site of the fpvA gene of the oprM-deficient cell, expression of oprM will be under control of the iron acquisition system. Thus, expression of the iron acquisition system may be monitored as the resistance to the pump substrate antibiotics (Figure 1). Alternatively, pump inhibitors render the cells supersusceptible to the substrate antibiotics. Using this system, potential inhibitors of the iron-uptake system or the pump inhibitor may be screened by a simple phenotypic change of the test strain from wild-type level susceptibility to supersusceptibility in the presence of sublethal concentrations of a MexAB-OprM substrate antibiotic, e.g., aztreonam, under iron-depletion conditions (Figure 1). The reason to choose OprM subunit is that: (i) the protein collaborates with several xenobiotic exporter systems [21] and (ii) OprM-deficient cells are more susceptible to the substrate antibiotics than mexA- or mexB-deficient cells [20]. Thus, OprM serves as a pivotal efflux pump subunit suitable for a highly sensitive screening system. We constructed such a recombinant P. aeruginosa strain as described in the Experimental Section and the strain was designated as TNP091.
Figure 1. Experimental design of the screening system. The oprM gene is placed immediately downstream of the fpvA gene of the oprM-deficient cell by homologous recombination. As a result, the reporter strain shows susceptibility and resistance (wild-type susceptibility) to a MexAB-OprM efflux pump substrate antibiotic (pentagons) under iron-replete (A) and iron-depleted (B) conditions, respectively, since the expression of the oprM gene is under the control of the fpvA gene.

2.2. Antibiotic Susceptibility of the Test Strain Depends on the Extracellular Iron Levels

We next assessed the impact of the extracellular iron levels on the antibiotic susceptibility of TNP091 (Table 1). The MICs of aztreonam in TNP091 under iron-replete and depleted (in the presence of dipyridyl) conditions were 0.25 and 2.0 µg/mL, respectively, and the values were comparable to those for TNP072 lacking the oprM gene and TNP072 harboring pOPRM1, respectively (Table 1).

| Strain            | Plasmid | Mic (µg/mL) |                       |                       |
|-------------------|---------|-------------|-----------------------|-----------------------|
|                   |         | Aztreonam   | Chloramphenicol       |                       |
|                   |         | Fe-replete  | Fe-depleted           | Fe-replete  | Fe-depleted |
| PAO4290 (parent)  | -       | 4.0         | 4.0                   | 32.0                  | 32.0        |
| TNP072 (∆oprM)    | -       | 0.25        | 0.25                  | 2.0                   | 2.0         |
| TNP072 (∆oprM)    | pOPRM1  | 4.0         | 4.0                   | 32.0                  | 32.0        |
| TNP091 (fpvA::oprM)| -        | 0.25        | 2.0                   | 4.0                   | 16.0        |
| TNP092 (fpvA::oprM ∆pvdS) | - | 0.25        | 0.5                   | 4.0                   | 4.0         |
| TNP092 (fpvA::oprM ∆pvdS) | pPvdS | 4.0         | 4.0                   | 32.0                  | 32.0        |

MICs were determined by the agar dilution method. Mueller-Hinton medium were prepared with or without 0.5 mM dipyridyl.

The MICs of chloramphenicol in TNP091 under iron-replete and depleted environments were 4.0 and 16 µg/mL, respectively. MICs of aztreonam and chloramphenicol in PAO4290, TNP072, and TNP072 (pOPRM1) were unchanged irrespective of the extracellular iron levels (Table 1). The wild-type susceptibility in TNP091 in the presence of dipyridyl was interpreted to mean that TNP091 cells sensed a low iron level in the milieu and this signal was conveyed to the fpvA promoter via an iron-responsive signal transduction pathway and induced the oprM gene expression. We analyzed the level of OprM
expression in the presence and absence of 0.5 mM dipyridyl by the western-blotting method using an anti-OprM antibody as a probe (Figure 2). Although TNP091 produced undetectable levels of OprM under the iron-replete conditions (Figure 2, lane 3), a significant level of OprM was detected under the iron-depleted conditions (Figure 2, lane 7), confirming that OprM was induced in response to the iron-depletion.

**Figure 2.** Western blot analysis of the cells grown under iron-replete and iron-depleted conditions. Cells were grown in the absence (lanes 1 to 4) or presence (lanes 5 to 9) of 0.5 mM dipyridyl and the whole cell lysate was subjected to SDS-polyacrylamide gel (10%) electrophoresis followed by visualization with an anti-OprM antibody. The position of OprM is indicated by an arrowhead. Lanes 1 and 5, PAO4290 (wild-type); lanes 2 and 6, TNP072 (∆oprM); lanes 3 and 7, TNP091 (∆oprM, fpvA::oprM); lanes 4 and 8, TNP092 (∆oprM, fpvA::oprM, ∆pvdS); lane 9, TNP092 (pPvdS).

### 2.3. The pvdS Mutation Prevents Induction of the oprM Expression

The above results suggested that antibiotic resistance in TNP091 in the presence of dipyridyl is most likely attributable to the induction of OprM through an iron responsive signal transduction system. To validate this assumption, we disrupted the *pvdS* gene, activation of which is essential for the induction of the *fpvA* gene and hence the *oprM* gene too [16]. First, we assessed the impact of the *pvdS* disruption on the pyoverdine production. The culture supernatant of a wild-type strain PAO4290, an *oprM*-deficient mutant TNP072, and the test strain TNP091 showed typical pyoverdine absorption spectra (Figure 3A–C), whereas that of the *pvdS* deletion mutant TNP092 produced an undetectable level of pyoverdine (Figure 3D), confirming that the iron acquisition cascade did not function at all in TNP092. The result is consistent with the previous report [16].

We next determined the MICs of antibiotics in TNP092 (Table 1). The MICs of aztreonam and chloramphenicol in TNP092 were 0.5 µg/mL and 4.0 µg/mL, respectively, under the iron-depleted conditions. The values were comparable with those obtained under the iron-replete conditions (Table 1). Consistent with this, TNP092 produced only a trace level of OprM under an iron-limiting condition (Figure 2, lane 8). To set back the *pvdS* gene into TNP092, the *pvdS* gene was cloned on a shuttle vector pMWB67HE [24] under control of *lac* promoter, pPvdS, and this plasmid was introduced into TNP092. The impact of the plasmid-borne *pvdS* expression on drug susceptibility of the transformant TNP092 (pPvdS) was tested. Under iron-depleted conditions, the cells recovered the wild-type susceptibility to both aztreonam and chloramphenicol (Table 1). A western blotting experiment revealed that the production of the OprM protein was restored in TNP092 (pPvdS) (Figure 2, lane 9). All these data supported an assumption that the wild-type susceptibility in TNP091 is attributable to the induction of OprM under iron-depleted conditions.
Figure 3. Absorption spectra of the culture supernatant. Cells were grown in casamino acid medium overnight at 37 °C under aerobic conditions and removed by the centrifugation at 13,700×g for 10 min. The clear supernatant fractions were scanned with a spectrophotometer. (A), PAO4290 (wild-type); (B), TNP072 (ΔoprM); (C), TNP091 (ΔoprM, fpvA::oprM); (D), TNP092 (ΔoprM, fpvA::oprM, ΔpvdS).

2.4. Screening of Inhibitors of the Iron-Acquisition System or the Efflux Pump in a Chemical Library

A structurally diverse chemical compound library (n = 2952) from ChemBridge was screened for inhibitors of the iron-acquisition system or a xenobiotic extrusion pump, MexAB-OprM, taking advantage of the phenotypic change of the TNP091 reporter. As expected, TNP091 showed an undetectable level of growth in the presence of 1.0 µg/mL of aztreonam in the absence of dipyridyl (Figure 4B). On the other hand, in the presence of dipyridyl, the TNP091 cells grew on the antibiotic-impregnated plate (Figure 4A). Among the chemicals in the library, one of them, ethyl 2-(1-acetyl-piperidine-4-carboxamido)-4,5,6,7-tetrahydrobenzo[b]thiophene-3-carboxylate (Figure 4C), showed a dose dependent growth inhibition of TNP091 in the presence of both aztreonam and dipyridyl (Figure 4A, see clear spots). This compound was designated as YH001 for convenience.
Figure 4. Effect of ethyl 2-(1-acetylpiperidine-4-carboxamido)-4,5,6,7-tetrahydrobenzo-[b]thiophene-3-carboxylate (YH001) on the growth of the test strain TNP091 (ΔoprM, fpvA::oprM) and chemical structure of YH001. (A) Liquefied Mueller-Hinton agar medium was mixed with the TNP091 cells and 1.0 µg/mL of aztreonam in the presence of 0.5 mM dipyridyl, and was solidified. Two µL of YH001 solution was spotted on the agar surface and the plate was incubated for 24 h at 37 °C. Concentrations of YH001 used were; 1, 10 mM; 2, 5 mM; 3, 2.5 mM; and 4, solvent control DMSO only. (B) Mueller-Hinton agar was mixed with the TNP091 cells and 1.0 µg/mL of aztreonam and was solidified. Two µL of YH001 solution was spotted on the agar surface and the plates were incubated for 24 h at 37 °C. 1, 2, 3, and 4 are the same as above. (C) Chemical structure of YH001.

2.5. Characterization of the Hit-Compound

Since this screening system employed a functional MexAB-OprM efflux pump as the reporter, it is possible that the hit-compounds inhibited either any one of the iron-responsive signal transduction pathway, the efflux pump activity or the growth of the cells by non-specific cytotoxicity. To identify the target of YH001, the growth-inhibitory activity of this compound in the presence of 2.0 µg/mL of aztreonam was tested in the wild-type strain PAO4290 (MIC of aztreonam 4.0 µg/mL) in the presence and absence of 0.5 mM dipyridyl. If YH001 blocked only the iron-responsive signal transduction pathway, MexAB-OprM may be expressed from their chromosomal genes irrespective of the presence or absence of dipyridyl rendering the cells resistant to the antibiotics. However, this was not the case (Figure 5A,B).

PAO4290 was unable to grow in the presence of 2.0 µg/mL of aztreonam and YH001, regardless of the presence of dipyridyl, suggesting a possibility that compound YH001 acted on the efflux pump itself. To assure this assumption, a strain producing an elevated level of MexAB-OprM, TNP30#1 (MIC of aztreonam 25 µg/mL) was similarly tested in the presence of 8 µg/mL of aztreonam. The growth of the cells was inhibited in a similar manner to above, confirming that YH001 directly acts on the efflux pump (Figure 5C,D). As a control, we took a known pump inhibitor molecule, L-phenyl-alanyl-L-arginyl-β-naphthylamide (PAβN) [25], and determined a synergistic effect with aztreonam in PAO4290 and TNP091. The results confirmed that this inhibitor exhibited similar results to YH001 shown in
Figures 4 and 5 (data not shown). To see whether compound YH001 exerts non-specific cytotoxic activity, strains TNP091, PAO4290 and TNP30#1 were incubated on the aztreonam-free medium in the presence of an appropriate concentration of YH001. The result showed no detectable growth inhibitory activity (data not shown). Consistently, MIC of YH001 in TNP30#1 appeared >320 µg/mL. Taken together, all the evidence suggests that YH001 acts on the efflux pump.

**Figure 5.** Effect of YH001 on the growth of *P. aeruginosa* producing different levels of the efflux pump in the presence of antibiotics. (A,B), a wild type strain PAO4290 producing a basal level of MexAB-OprM pump. (C,D), a strain producing an elevated level of MexAB-OprM pump, TNP30#1. Agar plates were prepared as described in the legend to Figure 4 impregnated with the following: (A) dipyridyl (0.5 mM) and aztreonam (2.0 µg/mL), (B) aztreonam (2.0 µg/mL), (C) dipyridyl (0.5 mM) and aztreonam (8.0 µg/mL), and (D) aztreonam (8.0 µg/mL). Concentrations of YH001 used were; 1, 10 mM; 2, 5 mM; 3, 2.5 mM; 4, solvent control DMSO only.

Efflux pump inhibitors are known to potentiate efficacy of the efflux-pump-substrate antibiotics [26]. This possibility was tested by checkerboard assay with PAO4290 and TNP30#1. The MICs of aztreonam and chloramphenicol in PAO4290 decreased to 1/2 and 1/4 of the original MICs, respectively, in the presence of 20 µg/mL of YH001 (Figure 6A). However, that of imipenem was unchanged in the presence of even 80 µg/mL of YH001. This is attributable to the fact that imipenem is a poor substrate of the MexAB-OprM efflux pump [20]. Hence, imipenem served as an excellent control compound. Similar tendencies were observed in the pump-overproducing strain, TNP30#1 (Figure 6B). In addition, YH001 did not exert a synergistic effect with chloramphenicol in TNP072 (ΔoprM) and TNP076 (ΔmexA ΔmexB ΔoprM) as expected (data not shown). Thus, it became evident that YH001 inhibits the activity of the MexAB-OprM efflux pump in a dose dependent manner.
Figure 6. Synergistic effect of YH001 with antibiotics. Potentiation of the antibiotic effect with YH001 was assayed by the checkerboard method. The MIC values were expressed relative to that in the YH001-free medium. (A) *P. aeruginosa* wild type strain PAO4290. MICs of aztreonam, chloramphenicol, and imipenem in PAO4290 are 2.0, 16 and 4.0 µg/mL, respectively. (B) *P. aeruginosa* mutant TNP30#1 producing an elevated level of MexAB-OprM. MICs of aztreonam, chloramphenicol, and imipenem in TNP30#1 are 32, 64 and 4.0 µg/mL, respectively. Symbols: circles, aztreonam; triangles, chloramphenicol; and squares, imipenem.

Since the inhibitor YH001 acts on MexAB-OprM efflux pump of *P. aeruginosa*, we asked whether this compound exerts similar activity on the major multiantibiotic efflux pump of *E. coli*, AcrAB-TolC, which is a homologue of MexAB-OprM. The growth of the wild-type *E. coli* strain BW25113 was inhibited in the presence of sublethal levels of chloramphenicol and YH001 (discernible spots in Figure 7A). Since the presence of YH001 and a sublethal concentration of chloramphenicol exerted no discernible effect on the growth of *acrb*-deficient mutant JW0451 (Figure 7B), HY001 appeared to inhibit AcrAB-TolC efflux pump as well. However, the activity of YH001 toward AcrAB-TolC seems much weaker than to MexAB-OprM. To be sure that YH001 acts on MexAB-OprM in the *E. coli* cells, the checkerboard assay was carried out using aztreonam and erythromycin in the mutant lacking *acrb* and *tolC*. The MICs of aztreonam and erythromycin were lowered to 1/2 to 1/4 in the presence of 2.0 to 32 µg/mL of YH001 (Figure 7C) compared with the value without the inhibitor, indicating clearly that the inhibitor acts directly on the MexAB-OprM efflux pump, even in the *E. coli* host.

3. Discussion

We have designed and constructed a new model antimicrobial compound screening system targeting the iron-acquisition system and the multidrug efflux pump(s) in *Pseudomonas aeruginosa*. 
Figure 7. Effect of YH001 on the growth of E. coli strains in the presence of sub-lethal concentrations of chloramphenicol and synergistic effects of YH001 with antibiotics. 

(A) Effect of YH001 on the wild-type E. coli strain BW25113; (B) Effect of YH001 on acrB-deficient mutant E. coli JW0451 (ΔacrB). Agar plates were prepared mixing with the indicated concentrations of chloramphenicol and the respective bacterial cells as described in the legend to Figure 4. Two µL of YH001 solution was spotted on the surface of the agar medium and the plates were incubated at 37 °C for 24 h. (C) Synergistic effects of YH001 with antibiotics in the E. coli MG1655ΔBC (ΔacrB, ΔtolC) and E. coli MG1655ΔBC/pABM harboring a plasmid carrying the mexAB-oprM genes. MICs of antibiotics were determined by the checkerboard method. The MIC values were expressed relative to that in the YH001-free medium. MICs of aztreonam and erythromycin in the transformants are 1.0 and 32 µg/mL, respectively. Symbols: open circles, aztreonam; open triangles, erythromycin.

The compounds obtained from the screening by this system may have either any one of the following modes of action: (i) if the inhibitor-compound blocked the pyoverdine-synthesizing pathway, iron-pyoverdine complex may not be formed and, therefore, a σ factor, PvdS, may be kept associated with an anti-σ factor, FpvR. Thus, the test strain will be aztreonam supersusceptible; (ii) if the inhibitor blocked the function of MexAB-OprM efflux pump, the test strain will be aztreonam supersusceptible irrespective of the iron-acquisition system; (iii) if the inhibitor blocked the Tat secretion machinery, biogenesis of pyoverdine may be blocked because a periplasmic protein, PvdN, cannot be translocated.
to its destination [19] rendering the cells supersusceptible to aztreonam. In fact, the MICs of aztreonam and chloramphenicol in the tatC-deficient derivative of TNP091 dropped four-fold each compared with that in the parent cell, TNP091, under iron-depleted conditions. Since lines of evidence are accumulating that the Tat system secretes several other important virulence factors [27], blocking of this machinery likely reduces the pathogenesis of the bacteria; (iv) if the inhibitor blocked the Fur and Fur-box-mediated iron acquisition pathway, the cells will be killed, since the Fur protein is assumed to be essential for survival of P. aeruginosa [15].

The primary aim of the experiment was to design a new antimicrobial screening system and to construct a strain suitable for the purpose. A model system we chose was to screen potential inhibitors that block the iron acquisition systems. Advantages of using inhibitors found by this screening system over classical antibiotics may be as follows: (i) since the compounds reduce the virulence of the pathogenic bacteria but may not kill them, they are less likely to select resistant mutants; (ii) it is expected that an efflux pump inhibitor will potentiate the activity of classical antibiotics. This assumption was validated by the fact that YH001 lowered the MICs of aztreonam and chloramphenicol (Figure 6A,B).

As mentioned above, this screening system could select inhibitors of RND-type xenobiotic efflux pumps. The reason why OprM was chosen was as follows: OprM functions as a common xenobiotic discharge duct for the pump units of MexAB, MexXY, MexCD and others [20–22]. Therefore, if the inhibitors blocked the function of OprM, it is potentially possible to block the function of multiple species of xenobiotic efflux pumps rendering the cells susceptible to a wide variety of substrate antibiotics. To test such a possibility, the MIC of gentamicin, which is an excellent substrate of the MexXY pump, was determined in the presence of YH001. However, the MIC remained unchanged in mexXY-overexpressing strains in the presence and absence of YH001 (data not shown). Therefore, it is most likely that inhibitor YH001 specifically blocks the function of the MexAB pump but not the OprM exit channel. In addition, however, this compound appeared to have an inhibitory activity toward the major antibiotic efflux pump AcrAB-ToIC in E. coli leaving a possibility that a broad spectrum inhibitor, once selected, can be extended to use in a wide range of other bacterial species. A sole drawback of this system would be that the inhibitor must be selected by negative means. However, patient and stubborn searches may open a way to combat multiantibiotic resistant pathogens.

4. Experimental Section

4.1. Bacterial Strains, Plasmids, and Growth Conditions

The bacterial strains and plasmids used are listed in Table 2. Cells were grown aerobically in L-broth containing 10 g of tryptone, 5 g of yeast extract, and 5 g of NaCl per liter (pH 7.2) at 37 °C. For the pyoverdine assay, cells were grown in medium containing 5 g per liter of casamino acids, 5 mM K2HPO4, and 1 mM MgSO4 under aeration at 37 °C overnight. Antibiotics used were aztreonam (Sigma-Aldrich, St. Louis, MO, USA), chloramphenicol (Wako Pure Chemicals, Tokyo, Japan), imipenem (Sigma-Aldrich) and erythromycin (Sigma-Aldrich). To generate iron-depleted conditions, dipyridyl (Sigma-Aldrich) to a final concentration of 0.5 mM was added to Mueller-Hinton agar medium.
### Table 2. Bacterial strains and plasmids.

| Strains or Plasmids       | Relevant Property                                              | Source or Reference |
|---------------------------|-----------------------------------------------------------------|---------------------|
| **Strain**                | **Relevant Property**                                           | **Source or Reference** |
| *Escherichia coli* JM109  | recA1, endA1, gyrA96, thi, hsdR17(rk/cmk1), e14 (mcrA+), supE44, relA1, Δ(lac-proAB)/F[traD36, proAB’, lacF’, lacZΔM15] | Laboratory strain   |
| *Escherichia coli* S17-1  | RP4-2-Tc::Mu-Km::Tn7, pro, res-, mod'                           | [28]                |
| *Escherichia coli* MG1655ΔABC | ΔacrB, ΔtoIC                                                      | [29]                |
| *Escherichia coli* BW25113 | Wild type, parent of JW0451                                     | [30]                |
| *Escherichia coli* JW0451  | ΔacrB derived from BW25113                                      | [30]                |
| *Pseudomonas aeruginosa* PAO1 | wild type                                                        | Laboratory strain   |
| *Pseudomonas aeruginosa* PAO4290 | leu-10, argF10, aph-9004, FP-                                   | Matsumoto collection|
| *Pseudomonas aeruginosa* TNP072 | PAO4290 ΔoprM::Tc’                                              | [20]                |
| *Pseudomonas aeruginosa* TNP091 | TNP072 fpvA::oprM                                              | This study          |
| *Pseudomonas aeruginosa* TNP092 | TNP072 fpvA::oprM, ΔpvdS                                         | This study          |
| *Pseudomonas aeruginosa* TNP30#1 | PAO4290-derived multiantibiotic resistant mutant overexpressing | [31]                |
|                           | mexAB-OprM genes                                                 |                     |
| **Plasmid**               | **Relevant Property**                                           | **Source or Reference** |
| pK19mobsacB               | pK19 derivative carrying RP4 mob region and Bacillus subtilis sacB gene, Km’ | [32]                |
| pK18mob                   | pK18 derivative carrying RP4 mob region, Km’                     | [32]                |
| pMMB67EH                  | Broad host range vector, Ap’, IncQ                               | [24]                |
| pMMB67HE                  | Broad host range vector, Ap’, IncQ                               | [24]                |
| pK19FpvOprM               | pK19mobsacB derivative carrying 3’-end of fpvA region and its downstream region in which oprM gene is inserted | This study          |
| pK18mobPvdS               | pK18mob derivative caring internal region of the pvdS gene       | This study          |
| pOPRM1                    | pMMB67EH derivative carrying wild type oprM gene, Ap’            | [21]                |
| pPvdS                     | pMMB67EH derivative carrying the pvdS gene                       | This study          |
| pABM                      | pMMB67HE derivative carrying the mexA-mexB-oprM genes            | [29]                |

_Abbreviations:_ KM, kanamycin; TC, tetracycline; Ap, ampicillin.

### 4.2. Construction of the Recombinant Strain

To insert the oprM gene immediate downstream site of the fpvA gene in the chromosome of TNP072, a suicide vector was constructed as follows. First, a DNA fragment containing downstream region of fpvA was amplified using a pair of primers, FpvAdown-F and FpvAdown-R (Table 3), using the chromosomal DNA of PAO1 as the template and TaKaRa PrimeStar HS (TAKARA, Otsu, Japan) according to the manufacturer’s instructions. The resulting DNA fragments were digested with SmaI and EcoRI and subcloned into pK19mobsacB [32] pretreated with SmaI, EcoRI, and bacterial alkaline phosphatase (TOYOBO, Osaka, Japan). Next, a fragment containing oprM was amplified using primers, OprM-F and OprM-R (Table 3), as above, and the XbaI/SmaI-treated fragment was inserted into the
above-mentioned recombinant plasmid pretreated with XbaI and SmaI. Lastly, the DNA fragment containing the 3'-end of fpvA gene (1428 bp) was amplified with primers, FpvA-F and FpvA-R (Table 3), as above, and the HindIII/XbaI-treated fragment was inserted into the HindIII and XbaI sites of the above oprM-bearing plasmid. The resulting suicide vector pK19FvpAOpM was then transferred from E. coli S17-1 to P. aeruginosa TNP072 by the filter mating method [20]. Transconjugants were selected on L-agar medium containing 12.5 µg/mL of kanamycin and 12.5 µg/mL of tetracycline at 37 °C overnight. Subsequently, the resulting transconjugants were counterselected on L-agar medium containing 10% (w/v) sucrose at 37 °C overnight. The chromosomal region of the fpvA gene in the resulting kanamycin-susceptible clones was analyzed by PCR using forward and reverse primers, FpvA-F and FpvAdown-R, respectively, to confirm that oprM was properly inserted into the downstream site of the fpvA gene. The strain was designated as TNP091.

To disrupt the pvdS gene in TNP091 cells, we first amplified internal region (419 bp) of the pvdS gene by Takara Ex Taq (TAKARA) using a primer set, PvdS-F and PvdS-R (Table 3), and the chromosomal DNA from PAO1 as the template. The resulting DNA fragments were treated with EcoRI and XbaI, and was ligated to pK18mob [32] pretreated with EcoRI, XbaI, and bacterial alkaline phosphatase, to yield pK18mobPvdS. Next, pK18mobPvdS in E. coli S17-1 was transferred to TNP091 by the conjugation as described above and transconjugants were selected on the medium containing 12.5 µg/mL of kanamycin and 12.5 µg/mL of tetracycline. The chromosomal region contiguous to pvdS was examined by PCR to confirm the insertional inactivation.

Table 3. Primers.

| Primer         | Nucleotide sequence * | Tag        |
|---------------|-----------------------|------------|
| FpvAdown-F    | 5′-GGCCCCGGGGCCCGACTGCAGAAAAC-3′ | SmaI       |
| FpvAdown-R    | 5′-CGAATTCACCGCAACTGTTGGGATAC-3′ | EcoRI      |
| OprM-F        | 5′-TGTCTAGACAAAGCAGCCGCTGCCGTCGTC-3′ | XbaI       |
| OprM-R        | 5′-TGACCCCCGAGCGCTGCGGATGCTCCTGGGACAGTT-3′ | SmaI       |
| FpvA-F        | 5′-ATTAAAGCTTAATCCCGACACCATGCTTAC-3′ | HindIII    |
| FpvA-R        | 5′-GATCTAGATCAGAAGCAGCCGCTGCCGTCGTC-3′ | XbaI       |
| PvdS-F        | 5′-TGAAATCCACCGATGCCGATAGCTGTTGGGATAC-3′ | EcoRI      |
| PvdS-R        | 5′-TGCTTACATGAAGCAGTTGGGACAGTGTT-3′ | XbaI       |

*a Restriction site tags are underlined.

4.3. Expression of OprM under Iron-Replete and Iron-Depleted Conditions

Fully-grown TNP091 cells in Mueller Hinton medium were diluted with a 100-fold volume of the same medium in the presence or absence of 0.5 mM dipyridyl, an iron chelator, and were continued to grow until a late-log phase. The cells were harvested by centrifugation at 9000× g for 10 min, washed once with phosphate-buffered saline (pH, 7.2), suspended in the same solution and the cell-free lysate was subjected to the western blot analysis.

4.4. Screening of a Chemical Library

The test strain TNP091 was grown in Mueller-Hinton broth at 37 °C overnight and was diluted a 100-fold with pre-warmed Mueller-Hinton agar containing 1.0 µg/mL of aztreonam and 0.5 mM dipyridyl. The
agar medium was solidified in a Petri dish. Two microliters each of agents from a small molecule chemical library (ChemBridge Co., San Diego, CA, USA) dissolved in dimethyl sulfoxide (10 mM or 0.1 mM) were placed on the surface of the solid agar medium and the plates were incubated at 37 °C overnight. The compound, which inhibited the growth of the cells, was scored at 24 h of incubation.

4.5. Assay for the Potentiation of Antibiotic Activity by a Hit-Compound

Potentiation of the antibiotic activity by a hit-compound in the \( P. \ aeruginosa \) cells was assayed by the checkerboard method. A test strain TNP091 was grown in 2 mL of Mueller-Hinton broth containing 0.4% of KNO\(_3\) (w/v) at 37 °C overnight by standing culture. Five microliters of a 1000-fold diluted cell suspension (1–3 \( \times \) \( 10^4 \) cells/inoculum) were added into 2 mL of fresh Mueller-Hinton broth containing a 2-fold serially diluted antibiotic solution, to that various amounts of the compound to be tested were added. After 20 to 24 h of standing culture at 37 °C, growth of the test strain was scored by reading absorbance at 660 nm. Potentiation of the antibiotic activity by the hit-compound was also assayed in \( Escherichia \ coli \) MG1655 \( \Delta\)acrB\(\Delta\)tolC producing plasmid-borne MexAB-OprM by the microdilution method in a 96-well plate as described previously [29]. The synergistic effect was calculated by dividing an MIC of antibiotic in the presence of the hit-compound by that without the compound.

4.6. Other Techniques

The MIC of antibiotics was determined by the agar dilution method using Mueller-Hinton agar (Becton, Dickinson and Company, Sparks, MD, USA) [33]. Pyoverdine was quantified by measuring the absorption spectra from 380 nm to 500 nm of the culture supernatants [33]. SDS-PAGE was carried out by the method of Laemmli [34]. To determine the expression levels of OprM, the whole cell lysates were subjected to SDS-PAGE followed by blotting onto a polyvinylidene fluoride membrane (Millipore, Billerica, MA, USA). OprM was visualized by probing with an anti-OprM antibody using an alkaline phosphatase-conjugated secondary antibody [20]. The protein concentration was quantified by the method of Lowry \textit{et al.} [35]. The nucleotide sequence was determined by the dideoxy chain termination method [36].

Acknowledgments

We are grateful to Y. Ito for allowing us to use the chemical library and S. Kuwahara for helpful discussion. This study was supported in part by the Adaptable and Seamless Technology Transfer Program through Target-driven R&D from the Japan Science and Technology Agency.

Author Contributions

Kazuki S. and Hiroshi Y. designed and performed research; Kenichi U., Keiji A., Yoshimi M., Hideaki M., Tasuke A., and Emiko I. performed research and analyzed the data; Taiji N. and Hiroshi Y. wrote the paper. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.
1. Boucher, H.W.; Talbot, G.H.; Bradley, J.S.; Edwards, J.E.; Gilbert, D.; Rice, L.B.; Scheld, M.; Spellberg, B.; Bartlett, J. Bad bugs, no drugs: No ESKAPE! An update from the Infectious Diseases Society of America. *Clin. Infect. Dis.* 2009, 48, 1–12.

2. Taubes, G. The bacteria fight back. *Science* 2008, 321, 356–361.

3. Hunter, P. Where next for antibiotics? The immune system and the nature of pathogenicity are providing vital clues in the fight against antibiotic-resistant bacteria. *EMBO Rep.* 2012, 13, 680–683.

4. Bush, K.; Courvalin, P.; Dantas, G.; Davies, J.; Eisenstein, B.; Huovinen, P.; Jacoby, G.A.; Kishony, R.; Kreiswirth, B.N.; Kutter, E.; et al. Tackling antibiotic resistance. *Nat. Rev. Microbiol.* 2011, 9, 894–896.

5. Fischbach, M.A.; Walsh, C.T. Antibiotics for emerging pathogens. *Science* 2009, 325, 1089–1093.

6. Payne, D.J.; Gwynn, M.N.; Holmes, D.J.; Pompliano, D.L. Drugs for bad bugs: Confronting the challenges of antibacterial discovery. *Nat. Rev. Drug Discov.* 2007, 6, 29–40.

7. Taylor, P.L.; Wright, G.D. Novel approaches to discovery of antibacterial agents. *Anim. Health Res. Rev.* 2008, 9, 237–246.

8. Gwynn, M.N.; Portnoy, A.; Rittenhouse, S.F.; Payne, D.J. Challenges of antibacterial discovery revisited. *Ann. N. Y. Acad. Sci.* 2010, 1213, 5–19.

9. Rasko, D.A.; Sperandio, V. Anti-virulence strategies to combat bacteria-mediated disease. *Nat. Rev. Drug Discov.* 2010, 9, 117–128.

10. Baron, C. Antivirulence drugs to target bacterial secretion systems. *Curr. Opin. Microbiol.* 2010, 13, 100–105.

11. Alekshun, M.N.; Levy, S.B. Targeting virulence to prevent infection: To kill or not to kill? *Drug Discov. Today Ther. Strategies* 2004, 1, 483–489.

12. Andrews, S.C.; Robinson, A.K.; Rodriguez-Quinones, F. Bacterial iron homeostasis. *FEMS Microbiol. Rev.* 2003, 27, 215–237.

13. Miethke, M.; Marahiel, M.A. Siderophore-based iron acquisition and pathogen control. *Microbiol Mol. Biol. Rev.* 2007, 71, 413–451.

14. Drakesmith, H.; Prentice, A.M. Hepcidin and the iron-infection axis. *Science* 2012, 338, 768–772.

15. Cornelis, P. Iron uptake and metabolism in pseudomonads. *Appl. Microbiol. Biotechnol.* 2010, 86, 1637–1645.

16. Beare, P.A.; For, R.J.; Martin, L.W.; Lamont, I.L. Siderophore-mediated cell signalling in *Pseudomonas aeruginosa*: Divergent pathways regulate virulence factor production and siderophore receptor synthesis. *Mol. Microbiol.* 2003, 47, 195–207.

17. Lamont, I.L.; Beare, P.A.; Ochsner, U.; Vasil, A.I.; Vasil, M.L. Siderophore-mediated signaling regulates virulence factor production in *Pseudomonas aeruginosa*. *Proc. Natl. Acad. Sci. USA* 2002, 99, 7072–7077.

18. Cornelis, P.; Matthijs, S.; Van Oeffelen, L. Iron uptake regulation in *Pseudomonas aeruginosa*. *Biometals* 2009, 22, 15–22.

19. Vouhoux, R.; Filloux, A.; Schalk, I.J. Pyoverdine-mediated iron uptake in *Pseudomonas aeruginosa*: The Tat system is required for PvdN but not for FpvA transport. *J. Bacteriol.* 2006, 188, 3317–3323.
20. Yoneyama, H.; Ocaktan, A.; Tsuda, M.; Nakae, T. The role of mex-gene products in antibiotic extrusion in *Pseudomonas aeruginosa*. Biochem. Biophys. Res. Commun. 1997, 233, 611–618.

21. Yoneyama, H.; Ocaktan, A.; Gotoh, N.; Nishino, T.; Nakae, T. Subunit swapping in the Mex-extrusion pumps in *Pseudomonas aeruginosa*. Biochem. Biophys. Res. Commun. 1998, 244, 898–902.

22. Aires, J.R.; Kohler, T.; Nikaido, H.; Plesiat, P. Involvement of an active efflux system in the natural resistance of *Pseudomonas aeruginosa* to aminoglycosides. Antimicrob. Agents Chemother. 1999, 43, 2624–2628.

23. Poole, K.; Neshat, S.; Heinrichs, D. Pyoverdine-mediated iron transport in *Pseudomonas aeruginosa*: Involvement of a high-molecular-mass outer membrane protein. FEMS Microbiol. Lett. 1991, 62, 1–5.

24. Furste, J.P.; Pansegrau, W.; Frank, R.; Blocker, H.; Scholz, P.; Bagdasarian, M.; Lanka, E. Molecular cloning of the plasmid RP4 primase region in a multi-host-range tacP expression vector. Gene 1986, 48, 119–131.

25. Piddock, L.J.; Garvey, M.I.; Rahman, M.M.; Gibbons, S. Natural and synthetic compounds such as trimethoprim behave as inhibitors of efflux in Gram-negative bacteria. J. Antimicrob. Chemother. 2010, 65, 1215–1223.

26. Stavri, M.; Piddock, L.J.; Gibbons, S. Bacterial efflux pump inhibitors from natural sources. J. Antimicrob. Chemother. 2007, 59, 1247–1260.

27. Ochsner, U.A.; Snyder, A.; Vasil, A.I.; Vasil, M.L. Effects of the twin-arginine translocase on secretion of virulence factors, stress response, and pathogenesis. Proc. Natl. Acad. Sci. USA 2002, 99, 8312–8317.

28. Simon, R.; Priefer, U.; Puhler, A. A Broad Host Range Mobilization System for In Vivo Genetic Engineering: Transposon Mutagenesis in Gram Negative Bacteria. Nat. Biotech. 1983, 1, 784–791.

29. Matsumoto, Y.; Hayama, K.; Sakakihara, S.; Nishino, K.; Noji, H.; Iino, R.; Yamaguchi, A. Evaluation of multidrug efflux pump inhibitors by a new method using microfluidic channels. PLoS ONE 2011, 6, e18547.

30. Baba, T.; Ara, T.; Hasegawa, M.; Takai, Y.; Okumura, Y.; Baba, M.; Datsenko, K.A.; Tomita, M.; Wanner, B.L.; Mori, H. Construction of *Escherichia coli* K-12 in-frame, single-gene knockout mutants: The Keio collection. Mol. Syst. Biol. 2006, 2, 2006.0008.

31. Saito, K.; Yoneyama, H.; Nakae, T. *nalB*-type mutations causing the overexpression of the MexAB-OprM efflux pump are located in the *mexR* gene of the *Pseudomonas aeruginosa* chromosome. FEMS Microbiol. Lett. 1999, 179, 67–72.

32. Schaefer, A.; Tauch, A.; Jager, W.; Kalinowski, J.; Thierbach, G.; Puhler, A. Small mobilizable multi-purpose cloning vectors derived from the *Escherichia coli* plasmids pK18 and pK19: Selection of defined deletions in the chromosome of *Corynebacterium glutamicum*. Gene 1994, 145, 69–73.

33. Akiba, K.; Ando, T.; Isogai, E.; Nakae, T.; Yoneyama, H. Tat pathway-mediated translocation of the Sec pathway substrate OprM, an outer membrane subunit of the resistance nodulation division xenobiotic extrusion pumps, in *Pseudomonas aeruginosa*. Chemotherapy 2013, 59, 129–137.

34. Laemmli, U.K. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature 1970, 227, 680–685.
35. Lowry, O.H.; Rosebrough, N.J.; Farr, A.L.; Randall, R.J. Protein measurement with the Folin phenol reagent. *J. Biol. Chem.* **1951**, *193*, 265–275.
36. Sanger, F.; Nicklen, S.; Coulson, A.R. DNA sequencing with chain-terminating inhibitors. *Proc. Natl. Acad. Sci. USA* **1977**, *74*, 5463–5467.

Sample Availability: Not available.

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