Thiamine pyrophosphate (TPP) is an essential cofactor for various pivotal cellular processes in all living organisms, including bacteria. Thiamine biosynthesis occurs in bacteria but not in humans; therefore, the enzymes in this pathway are attractive targets for antibiotic development. Among these enzymes, thiamine monophosphate kinase (ThiK) catalyzes the final step of this pathway, phosphorylating thiamine monophosphate to produce TPP. Here, we extensively investigated ThiK in *Pseudomonas aeruginosa*, a major pathogen responsible for hospital-acquired infections. We demonstrate that thiK deletion abolishes not only thiamine biosynthesis but also thiamine salvage capability and results in growth defects of the ΔthiK strain even in the presence of thiamine derivatives, except for TPP. Most importantly, the pathogenesis of the ΔthiK strain was markedly attenuated, compared with that of WT cells, with lower inflammatory cytokine induction and 103–105-fold decreased bacterial loads in an in vivo infection model in which the intracellular TPP level was in the submicromolar range. To validate *P. aeruginosa* ThiK (PaThiK) as a drug target, we further characterized its biochemical properties, determining a $V_{\text{max}}$ of $4.0 \pm 0.2$ nmol min$^{-1}$ and $K_{\text{m}}$ values of $111 \pm 8$ and $8.0 \pm 3.5$ μM for ATP and thiamine monophosphate, respectively. An *in vitro* small-molecule screening assay identified PaThiK inhibitors including WAY213613, a noncompetitive inhibitor with a $K_{\text{i}}$ value of 13.4 ± 2.3 μM and potential antibacterial activity against *P. aeruginosa*. These comprehensive biological and biochemical results indicate that PaThiK represents a potential drug target for the development of an augmented repertoire of antibiotics against *P. aeruginosa*.

*Pseudomonas aeruginosa* is a Gram-negative opportunistic pathogen associated with a wide range of acute and chronic infections of various body sites, including the urinary tract, skin, and respiratory tract (1). Patients with compromised immune defenses due to underlying diseases such as cancer or HIV infection or with severe burns, cystic fibrosis, bronchiectasis, or chronic obstructive pulmonary disease are particularly susceptible to *P. aeruginosa* infection (2–7). The mainstay of treatment for *P. aeruginosa* infection is antibiotics. However, the expression of multiple efflux pumps, reduced permeability of the outer membrane, the capacity to form biofilms, and the presence of persisters have rendered *P. aeruginosa* intrinsically resistant to many antibiotics (8, 9). Given this natural resistance, excessive use of antibiotics is required to treat *P. aeruginosa* infections, accelerating the development of drug-resistant strains. Indeed, multidrug-resistant and extensively drug-resistant *P. aeruginosa* strains are now prevalent worldwide, with the emergence of pan-drug-resistant strains threatening global public health (10–13). The World Health Organization recently listed drug-resistant *P. aeruginosa* as a priority pathogen necessitating urgent action to develop novel antibiotics to overcome current antibiotic resistance (14).

Thiamine is a crucial molecule in all living organisms, from microorganisms to mammals. Therefore, thiamine metabolism has attracted growing attention in the development of various drugs, including antibiotics (15–19). The physiologically active form of thiamine, thiamine pyrophosphate (TPP), plays important roles as a cofactor in various essential cellular processes, including carbohydrate, lipid, and amino acid metabolism (20–22). Microorganisms such as bacteria and fungi, as well as plants, produce TPP via *de novo* biosynthetic pathways that mammals lack (23, 24). The TPP biosynthetic pathway of bacteria involves the separate biosynthesis of thiazole and pyrimidine moieties, which are joined to form thiamine monophosphate (TMP) in a reaction catalyzed by thiamine phosphate synthase (ThiE) (25–27). Thiamine monophosphate kinase (ThiK) catalyzes the final step of the pathway by phosphorylating TMP to TPP, the biologically active form of the cofactor (28, 29). In addition to the TPP biosynthetic pathway, bacteria are capable of salvaging thiamine from exogenous sources to generate TPP. Some bacteria, such as *Bacillus subtilis*, take up exogenous thiamine and convert it to TPP in a one-step reaction catalyzed by thiamine pyrophosphate kinase (TPK), similar to the mammalian thiamine salvage pathway (30–33). Other bacteria, including *Escherichia coli*, first convert thiamine to TMP by thiamine kinase (ThiK) and then subsequently generate TPP by adding one additional phosphate to TMP through ThiL, the enzyme in the main TPP biosynthetic pathway (30, 34). Because TPP is indispensable for bacterial survival and humans lack the TPP biosynthetic pathway, enzymes involved in bacterial TPP biosynthesis are potential targets in the development of new antibiotics. ThiK is of particular interest,

This article contains supporting information.

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considering its key role in thiamine metabolism. Nevertheless, few studies, involving a limited number of bacterial species, have been conducted on ThiL, and it has never been validated as a target for antibacterial agents (28, 29, 35, 36).

In this work, we constructed a clean thiL deletion mutant of 
P. aeruginosa and investigated the impact of thiL deficiency on bacterial survival and in vivo pathogenesis. We also biochemically characterized 
P. aeruginosa ThiL and identified small molecules that inhibit the enzyme by using an optimized luminescent kinase assay. To the best of our knowledge, this is the first work to demonstrate the role of 
P. aeruginosa ThiL in bacterial physiology and pathogenesis and the first to validate ThiL as a new target for drug development, providing comprehensive biochemical characterization of ThiL and identifying its inhibitors.

**Results**

**Role of ThiL in 
P. aeruginosa thiamine metabolism**

To investigate the physiological role of ThiL in 
P. aeruginosa, we constructed an in-frame deletion mutant of the thiL gene in 
P. aeruginosa PAO1 by two-step allele exchange (37–39). TPP was supplied via selection medium for the last step of mutant generation, to avoid the loss of bacterial viability due to impaired thiamine biosynthesis. Deletion of thiL in the bacterial genome was confirmed by amplification of the thiL flanking region (Fig. S1). Phenotypic analysis of the ∆thiL mutant confirmed that deletion of thiL is lethal to 
P. aeruginosa unless TPP is exogenously provided in the medium (Fig. 1). Complementation with the plasmid expressing thiL (pthiL) relieved the growth defect of the ∆thiL mutant caused by TPP depletion, suggesting that ThiL is essential for TPP biosynthesis in 
P. aeruginosa (see Fig. 2A).

The thiamine salvage pathway is another way to generate TPP in many organisms, including bacteria and even mammals, which cannot synthesize TPP de novo. Two types of direct thiamine salvage pathways have been identified in bacteria, namely, one-step pyrophosphorylation of thiamine to TPP by thiamine pyrophosphokinase (TPK/ThiN) and two steps of subsequent monophosphorylation of thiamine to TMP and then TPP by ThiK and ThiL, respectively. As the thiamine salvage pathway in 
P. aeruginosa has yet to be characterized and no thiamine transporter genes have been identified in this species (26), we first tested whether the bacteria are capable of producing TPP using extracellular thiamine. We found that the thiE mutant in which de novo TPP synthesis is impaired was able to grow in the presence of extracellular thiamine, TMP, and TPP, indicating that 
P. aeruginosa can take up thiamine and has a thiamine salvage pathway (Fig. 3). Unlike the thiE mutant, the ∆thiL strain exhibited a growth defect even with exogenously provided thiamine and TMP (Fig. 2). This result suggests that ThiL is involved in not only de novo TPP synthesis but also the thiamine salvage pathway in 
P. aeruginosa, indicating that ThiL plays a critical role in 
P. aeruginosa thiamine metabolism.
**Role of ThiL in the virulence of P. aeruginosa in vivo**

Next, we sought to determine the role of ThiL in the virulence of *P. aeruginosa* in vivo. C57BL/6 mice were infected intranasally with 2 × 10⁷ CFU of the PAO1, Δ*thiL*, or thiL-complemented strains. During the first 20 h, all infected mice exhibited decreased movement, compared with uninfected mice. Histopathological analysis of the lung tissues at 20 h postinfection also suggested that, unlike uninfected control mice, all infected mice showed a distinct inflammatory response composed mainly of peribronchial and alveolar neutrophilic infiltrates, with neutrophilic consolidation and smooth muscle hyperplasia in the arterioles of the lung tissues (Fig. 4). Although mice infected with the Δ*thiL* strain exhibited a slightly lower degree of lung neutrophil infiltration (8.2 ± 6.3%) than mice infected with the PAO1 strain (11 ± 4%), the difference was not significant, suggesting that the Δ*thiL* mutant is capable of recruiting host immune cells and causing lung inflammation similar to the PAO1 strain (Fig. S2). Interestingly, when the average bacterial load in the left lobe of the lungs was measured at 20 h postinfection (*n* = 5 for each group), over 1,000 times more bacteria were found in mice infected with the PAO1 strain (1.6 × 10⁸ CFU) and thiL-complemented strain (1.6 × 10⁸ CFU), compared with mice infected with the Δ*thiL* strain (6.7 × 10⁴ CFU) (Fig. 5A). In addition, the survival rates of the infected mice clearly demonstrated the attenuated virulence of the Δ*thiL* strain, compared with the PAO1 and thiL-complemented strains. Over 50% of mice infected with the PAO1 and thiL-complemented strains died by 24 h postinfection, and the rest of the infected mice were dead by 48 h postinfection (Fig. 5B). In contrast, uninfected control mice and mice infected with the Δ*thiL* mutant survived as long as 72 h postinfection (Fig. 5B).

Because there was a discrepancy between the results of the lung histology and survival analyses, we further investigated pathogenesis in the infected mice by measuring the host immune response in the blood. Levels of three proinflammatory cytokines, IL-6, IL-8, and tumor necrosis factor α (TNFα), were measured in the blood of infected mice at 20 h postinfection. The level of macrophage inflammatory protein-2 (MIP-2) was measured as a murine counterpart of IL-8. Levels of all three tested cytokines were significantly higher in PAO1-infected mice than in Δ*thiL*-infected mice (Fig. 6, A, B, and C). Similarly, the splenic bacterial count in mice infected with the PAO1 strain (6 × 10⁵ CFU/organ) was 4 orders of magnitude higher than that in Δ*thiL*-infected mice (2 × 10⁴ CFU/organ) (Fig. 6D). Taken together, these results suggest that overall virulence is attenuated in the Δ*thiL* strain due to decreased severity of bacteremia and sepsis, ultimately leading to survival of the mice, demonstrating that ThiL is essential for the full virulence of *P. aeruginosa*.

**Biochemical characterization of P. aeruginosa ThiL**

Despite its importance in bacterial physiology and virulence, to our knowledge *P. aeruginosa* ThiL (PaThiL) has never been studied before, at either the molecular or biochemical level. For biochemical characterization of the enzyme, the *P. aeruginosa thiL* gene (PA4051) was PCR amplified and cloned into pET28b. The recombinant PaThiL was purified as a ~37-kDa His-tagged protein using a nickel-nitrilotriacetic acid affinity matrix, with >95% purity (Fig. S3). We found that purified PaThiL was highly unstable without desalting, exhibiting 80% and 56% activity loss within 24 h at 4 °C and −210 °C, respectively (Fig. S4). A further gradual decrease in activity was observed over time. Therefore, purified PaThiL was frozen and stored in liquid nitrogen (−210 °C) after immediate desalting, and activity was retained for over 1 month without significant loss (Fig. S4).

Previous studies of ThiL used a HPLC-based assay or a coupled assay using apo-carboxylase to assess ThiL activity (28, 36, 40). Because both types of assays are quite inconvenient and inefficient, we adapted a luminescent kinase assay that detects ATP consumption and optimized it to determine PaThiL enzymatic kinetics (see Experimental Procedures). The final assay was established to analyze the activity of 10 μg of PaThiL in a reaction buffer containing 0.05 mM TMP, 0.05 mM ATP, 50 mM Tris-HCl (pH 8.0), 5 mM MgCl₂, and 350 mM KCl. The *Kₘ* values for TMP and ATP, as well as the *Vₘₐₓ* value, were calculated based on Michaelis-Menten and Lineeweaver-Burk plots of PaThiL activity in reactions containing...
several different concentrations of ATP and TMP. The $V_{\text{max}}$ value of PaThiL was 4.0 ± 0.2 nmol·min$^{-1}$. The $K_m$ values for ATP and TMP were 111 ± 8 µM and 8.0 ± 3.5 µM, respectively, with a random bi-bi mechanism (Fig. 7). Similar ranges of $K_m$ values were previously reported for partially purified E. coli ThiL (270 and 1.1 µM for ATP and TMP, respectively) (36). Further characterization of PaThiL revealed that the enzyme phosphorylates oxythiamine monophosphate (a TMP analogue) with a $K_m$ value of 15.2 ± 2.0 µM (Fig. S5). Thiamine, as well as other thiamine analogues, including oxythiamine,
pyrithiamine, and amprolium, were not phosphorylated by PaThiL, indicating that prior acquisition of monophosphate is a requirement for PaThiL substrates (Fig. S6). TPP was not further phosphorylated by ThiL, suggesting that PaThiL does not contribute to the generation of intracellular thiamine triphosphate.

Validation of PaThiL as a new antibacterial target

Gram-negative bacteria such as *P. aeruginosa* are notorious for their multidrug and pan-drug resistance, which has created an urgent need for new classes of antibiotics. Our *in vivo* virulence study of the ΔthiL mutant indicated that ThiL is a promising novel candidate for an antibacterial molecule, as humans have no PaThiL homologue. In order to validate the druggability of PaThiL, 2,800 commercially available compounds were screened for inhibitory activity against PaThiL. The screening was conducted using a slight modification of our established *in vitro* PaThiL assay. The screening assay was initiated upon addition of 5 μg of purified PaThiL to a reaction mixture containing 10 μM TMP and 100 μM test compound; ThiL activity was measured by monitoring ATP consumption during a 10-min reaction. WAY213613 (Tocriscreen) and 5-hydroxyindolacetic acid (Lopac) were among several compounds that inhibited PaThiL. Further biochemical studies revealed that WAY213613 was a noncompetitive inhibitor of TMP for PaThiL, with a $K_i$ value of $13.4 \pm 2.3 \mu M$, whereas 5-hydroxyindolacetic acid was an uncompetitive inhibitor that exhibited a $K_i$ value of $114 \pm 27 \mu M$ (Fig. 8). Additional binding mode analyses of each compound with the homology modeling-derived 3D structure of PaThiL suggested energetically favorable interactions with PaThiL (data not shown).

We next tested whether the identified PaThiL inhibitors exhibited antibacterial activity. Because the outer membrane of *P. aeruginosa* can impede the entry of drugs into the cell and thereby mask potential antibacterial activity of PaThiL inhibitors, we also tested the antibacterial activity with a low concentration of colistin (0.5 μg/ml), which can increase cell permeability by disrupting the bacterial outer membrane while not affecting bacterial viability (Fig. S7 and Table S1). Although 5-hydroxyindolacetic acid did not exhibit antibacterial activity at 100 μM even in the presence of colistin, we found that 100 μM WAY213613 completely stopped bacterial growth not by itself but in the presence of colistin (Fig. 9). These results suggest that PaThiL is a valid target for development of new antibacterial therapeutics, demonstrating the feasibility of identification of small molecules that inhibit PaThiL.

**Discussion**

TPP-dependent enzymes, including transketolase, pyruvate dehydrogenase, 2-oxoglutarate dehydrogenase, and 1-deoxy-d-
xylulose-5-phosphate synthase, catalyze essential cellular reactions in bacteria, from central metabolism to biosynthesis of amino acids, cofactors, and lipids (21). Recently, TPP-dependent pyruvate and α-ketoglutarate dehydrogenase complexes in *P. aeruginosa* were found to intracellularly reduce phenazine derivatives such as pyocyanin (a pseudomonal toxin), thus
contributing to iron acquisition and redox homeostasis of the bacteria in addition to their designated roles in central metabolism (41). This evidence indicates that TPP deficiency can have a substantial effect on the metabolic network of bacteria, thus making TPP metabolism a very attractive target for the development of new antibiotics. Indeed, inhibitors of ThiE exhibit antibacterial activity against Mycobacterium tuberculosis, a notorious human pathogen (42). Because M. tuberculosis does not harbor the genes for the thiamine salvage pathway and transporters, inhibition of ThiE can effectively deplete the TPP pool by impairing TPP biosynthesis, the only source of TPP in the bacteria, ultimately causing bacterial death.

However, inducing the death of many pathogenic bacteria via TPP deficiency is hampered by several challenges, particularly in vivo; 1) complete deficiency of TPP requires concurrent inhibition of both thiamine salvage and TPP biosynthesis, and 2) levels of thiamine derivatives in the host, specifically TPP, should not be sufficient for directly promoting bacterial survival. In this work, we addressed these two challenges by investigating ThiL in P. aeruginosa. We demonstrated that ThiL is the key enzyme for both de novo TPP synthesis and thiamine salvage in P. aeruginosa by revealing an in vitro growth defect of the thiL mutant, thus indicating a possibility to deplete the TPP pool in P. aeruginosa by targeting ThiL (Fig. 1).

Although thiamine transporter genes have not been identified in P. aeruginosa, the growth of thiE and thiL mutants in the presence of thiamine, TMP, or TPP suggests the existence of transporters that import thiamine derivatives in P. aeruginosa (26) (Figs. 2 and 3). In E. coli, intracellular TPP concentrations were measured at 150–300 μM under optimal growth conditions, which are presumably TPP levels for normal bacterial growth (43, 44). Consistent with this notion, when we determined the extracellular TPP concentration necessary to relieve the growth defect of the thiE mutant in which the TPP biosynthetic pathway is impaired, we found that a minimum TPP concentration of 25 μM was required to detect any degree of thiE mutant growth and a concentration over 200 μM was required to fully support the growth of the thiE mutant (Fig. 3). This result suggests that, if P. aeruginosa relies solely on exogenous TPP for growth, then the concentration should be higher than 25 μM. Reported TPP levels in mouse blood range from 0.3 to 1.2 μM, well below the TPP concentration sufficient to support growth of the thiL mutant (45, 46). Consistent with these data, we found that the thiL mutant exhibited markedly reduced pathogenicity in a mouse infection model, compared with the WT strain (Figs. 5 and 6). TPP levels in healthy human blood (116–138 nM) are even lower than in mouse blood, suggesting that ThiL is a viable therapeutic target in the treatment of Pseudomonas infections (47, 48).

We also validated PaThiL as a suitable target for new antibiotic agents by conducting an intensive biochemical characterization followed by identification of inhibitors of PaThiL. Considering the high intracellular concentrations of ThiL substrates (ATP, ~1.8 mM; TMP, ~12.5 μM) and product (TPP, ~0.3 mM) in bacteria (44), noncompetitive ThiL inhibitors might be more beneficial than competitive inhibitors that probably require a very high affinity for ThiL. Interestingly, we found that WAY213613, a known inhibitor of the EAAT2 glutamate transporter, could inhibit PaThiL in a noncompetitive manner, with a K_i value of 13.4 ± 2.3 μM. When we tested antibacterial activity, WAY213613 was able to inhibit the growth of P. aeruginosa, although the permeability of the bacterial outer membrane had to be increased to detect the activity (Fig. 9).

Antibiotic-resistant bacteria such as pan-drug-resistant P. aeruginosa pose an immediate threat to global public health and highlight an urgent need for new antibiotics and targets. In this work, we extensively characterized PaThiL and demonstrated its roles in bacterial physiology and pathogenesis. Although this study focused on PaThiL, this enzyme is expected to play similar roles in other bacteria that have a thiamine salvage pathway composed of ThiK and ThiL, such as E. coli and Salmonella species (26). Taken together, the results of this work demonstrate that ThiL is a suitable therapeutic target for the development of new drugs to treat not only P. aeruginosa infections but potentially many other bacterial infections as well.

**Experimental procedures**

**Bacterial strains, culture media, and chemicals**

All strains and plasmids used in this study are listed in Table 1. Bacteria were cultured in Luria-Bertani (LB) broth at 37°C

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**Figure 9. Growth kinetics of PAO1 in antibacterial effect tests with WAY213613 or 5-hydroxyindolacetic acid.** No antibacterial activities were observed with 100 μM WAY213613 (A) or 5-hydroxyindolacetic acid (B). A bacterial inhibition effect for WAY213613 (100 μM) was observed with addition of a sublethal dose of colistin (0.5 μg/ml) (A). 5-Hydroxyindolacetic acid (100 μM) did not show an antibacterial effect with addition of colistin (0.5 μg/ml) (B). Control groups were treated with no compound in LB medium or LB medium with colistin (0.5 μg/ml). Error bars are means ± S.D. of triplicate experimental groups.
ThiL is essential for pseudomonal thiamine metabolism

**Table 1**

<table>
<thead>
<tr>
<th>Strains, plasmids, and primers used in this study</th>
<th>Description</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strains</strong></td>
<td>P. aeruginosa PAO1</td>
<td>E. coli BL21(DE3)</td>
</tr>
<tr>
<td></td>
<td>PW7731</td>
<td>ΔthiL</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
<td>pET28b+</td>
<td>E. coli expression vector, Km&lt;sup&gt;R&lt;/sup&gt;</td>
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<tr>
<td></td>
<td>pCVD442-Gm</td>
<td>sacB suicide vector from pUM24, Gm&lt;sup&gt;R&lt;/sup&gt;</td>
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<td></td>
<td>pBSP II SK&lt;sup&gt;–&lt;/sup&gt;</td>
<td>Broad-host-range expression vector, Amp&lt;sup&gt;R&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>pET28&lt;sup&gt;b+&lt;/sup&gt;</td>
<td>PAO1 thiL gene inserted in pET28&lt;sup&gt;b+&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>pTHI</td>
<td>PAO1 thiL gene inserted in pBSP II SK&lt;sup&gt;–&lt;/sup&gt;</td>
</tr>
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<td><strong>Primers</strong></td>
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<tr>
<td></td>
<td></td>
<td>Reverse, ATTAATCGAGCTCTCAGCGCTTGCCGCC</td>
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<td></td>
<td>nusB (PA4052)</td>
<td>Forward, GTTTAAGCTTTCAGTCACGTTGGGTTCCGAAATGTTGG</td>
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<td>Reverse, GGGATCCATATGGGTGAGTTCGAGCTGATCCGCC</td>
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<tr>
<td></td>
<td>pggA (PA4050)</td>
<td>Forward, GGATCCCATATGGGTGAGTTCGAGCTGATCCGCC</td>
</tr>
</tbody>
</table>

<sup>a</sup>Km<sup>R</sup>, kanamycin resistance; Gm<sup>R</sup>, gentamycin resistance; Amp<sup>R</sup>, ampicillin resistance.

<sup>b</sup>Restriction sites are underlined.

with agitation. The thiE transposon mutant (PW7731) was cultured in minimal medium supplemented with TMP for normal growth. LB powder was dissolved in purified water and autoclaved at 121 °C for 15 min. As minimal medium, M9 salt powder was dissolved in purified water and autoclaved at 121 °C for 15 min. Sterile MgSO<sub>4</sub> (2 mM) and CaCl<sub>2</sub> (0.1 mM) were added to M9 solution for complete minimal medium. Supplements for the media, including glucose (20 mM), thiamine, TMP, and TPP, were sterilized by filtration through a 0.22-μm PVDF membrane (Millipore, Burlington, MA) before addition. Agar (1.5%) was added to the minimal medium to prepare solid growth plates. Kanamycin (50 mg/ml) and ampicillin (100 mg/ml) were dissolved in purified water and sterilized by filtration through a 0.22-μm PVDF membrane. Antibiotic stock solutions were stored at −20 °C. Isopropyl β-D-1-thiogalactopyranoside (IPTG) (1 mM) was dissolved in purified water and filtered through a 0.22-μm filter. All chemicals were purchased from Sigma-Aldrich (St. Louis, MO), except for ATP and IPTG, which were purchased from Thermo Fisher Scientific (Waltham, MA).

**Growth of the thiE transposon mutant (PW7731) with thiamine, TMP, or TPP**

M9 glucose was supplemented with thiamine (50 μM), TMP (50 μM), or TPP (100 μM), and the medium was applied to PW7731 with an initial OD<sub>600</sub> of 0.02. After a 16-h incubation with agitation, absorbance (OD<sub>600</sub>) was measured for each growth condition. The TPP salvage concentration was determined by 2-fold serial dilution of TPP in a range from 0.02 to 400 μM. Each concentration of TPP was applied to PW7731 with an initial OD<sub>600</sub> of 0.0001, which was prepared by 1/500 dilution of the bacteria at an OD<sub>600</sub> of 0.05, and the mutant was incubated in 37 °C for 16 h. The absorbance (OD<sub>600</sub>) was measured with a multilabel plate reader.

**Overexpression and isolation of ThiL**

The gene encoding thiamine monophosphate kinase (thiL) was amplified by PCR analysis using the primers listed in Table 1 and was inserted into pET28<sup>b+</sup> (Novagen, Madison, WI), a vector for the cloning and expression of recombinant proteins with His tags, at restriction sites between NdeI and HindIII. The plasmid was transformed into E. coli BL21 cells, which were selected on medium containing kanamycin (50 μg/ml). IPTG (1 mM) was added to the mid-log-phase culture to induce protein expression, and the cells were incubated for 2.5 h at 37 °C with agitation. The cells were centrifuged at 4,000 × g, and the resulting pellet was resuspended in 5 ml of 100 mM Tris-HCl (pH 8.0) supplemented with a protease inhibitor (Roche, Mannheim, Germany). The suspended cells were then sonicated using a Bioruptor sonication system (Diagenode, Denville, NJ) and centrifuged at 14,000 × g. ThiL protein in the supernatant was purified by affinity chromatography (GE Healthcare, Freiburg, Germany). The collected protein was desalted using a 30kDa Centrifugal Filter Unit (Millipore) and centrifuged at 14,000 × g. ThiL protein was added to the reaction buffer to initiate the kinase reaction. After 10 min, the amount of ATP was determined using a luminescence assay (Promega, Madison, WI). ATP hydrolysis in the reaction buffer without TMP was less than 1% in 10 min. Km and V<sub>max</sub> values were determined for TMP and ATP using the ThiL assay with TMP and ATP concentrations ranging from 0.78 to 50 μM and from 25 to 100 μM, respectively, in 2-fold dilution steps. The assay data were fit to Michaelis-Menten or Lineweaver-Burk functions using the random bistrubute module schema.

**ThiL assay**

The ThiL assay was established based on previous protocols (28, 40). The assay reaction buffer contained 0.05 mM TMP, 0.05 mM ATP, 50 mM Tris-HCl (pH 8.0), 5 mM MgCl<sub>2</sub>, and 350 mM KCl. A total of 10 μg ThiL protein was added to the reaction buffer to initiate the kinase reaction. After 10 min, the amount of ATP was determined using a luminescence assay (Promega, Madison, WI). ATP hydrolysis in the reaction buffer without TMP was less than 1% in 10 min. Km and V<sub>max</sub> values were determined for TMP and ATP using the ThiL assay with TMP and ATP concentrations ranging from 0.78 to 50 μM and from 25 to 100 μM, respectively, in 2-fold dilution steps. The assay data were fit to Michaelis-Menten or Lineweaver-Burk functions using the random bistrubute module in Sigma.
ThiL is essential for pseudomonal thiamine metabolism

Plot v. 14 (Systat Software Inc., San Jose, CA) to obtain the $K_m$ and $V_{max}$ values. To determine the $K_v$ of ThiL inhibitors, ThiL assays were carried out with WAY213613 (0–66.7 μM) or 5-hydroxyindolacetic acid (0–100 μM). The activity rates of ThiL were fit to Michaelis-Menten or Lineweaver-Burk functions using the single-substrate/single-inhibitor module in SigmaPlot.

Bacterial growth kinetics with ThiL inhibitors

For measurement of PAO1 growth kinetics with ThiL inhibitors, 100 μM WAY213613 or 5-hydroxyindolacetic acid was applied to PAO1 with or without colistin at 0.5 μg/ml. The growth kinetic assays were carried out at 37°C in 96-well plates with a final OD$_{600}$ of 0.0001. Absorbance values (OD$_{600}$) were measured every 20 min for 13 h with a multilabel plate reader (SpectraMax M5; Molecular Devices, San Jose, CA). A checkerboard assay was also performed for the combination of WAY213613 and colistin in the ranges of 0–200 μM and 0–20 μM, respectively. The MIC of each antimicrobial agent alone and in combination was defined as the lowest concentration that inhibited visible growth of PAO1. The fractional inhibitory concentration (FIC) index was calculated by the formula FIC = (MIC of WAY213613 in combination/MIC of WAY213613 alone) + (MIC of colistin in combination/MIC of colistin alone), as described previously (49, 50).

Construction of thiL deletion mutants via allelic exchange

To construct a P. aeruginosa mutant lacking thiL, allelic exchange was carried out according to previous studies (37, 39). Briefly, both upstream (nudB; 480 bp) and downstream (ppgA; 516 bp) genes flanking thiL were amplified by PCR analysis using the primers listed in Table 1. Both ends of the amplified genes carried specific restriction sites (Table 1). The downstream gene and pCVD442-Gm, a suicide vector, were double-digested with SacI and SmaI and then ligated. Consecutively, the upstream gene and pCVD442-Gm-ppgA were double-digested with SpH1 and SmaI and then ligated to obtain the suicide vector carrying the amplified genes. The generated vector was electroporated into E. coli SM10 λpir to facilitate conjugation into P. aeruginosa. Transconjugants were selected on medium containing gentamicin (100 μg/ml), and allele exchange was induced by incubation in medium containing sucrose (6%). The deletion mutant was selected on LB medium supplemented with 1 mM TPP, and deletion of thiL was confirmed by PCR analysis.

Generation of the complemented strain of the thiL deletion mutant

To generate a complemented strain of the P. aeruginosa ΔthiL mutant, the amplified thiL gene was double-digested with BamHI and HindIII and inserted into pBSP II SK(−) to create pthiL. The constructed plasmid was electroporated into competent cells of the PAO1 ΔthiL strain using a previously reported method (51). Briefly, the bacterial culture was washed twice and suspended with 300 μl sucrose at room temperature to generate electrocompetent cells. Next, 1 μg pthiL was mixed with 100 μl electrocompetent cells, and the pulse was applied to the mixture (P. aeruginosa 2.5-kV setting; Bio-Rad). LB medium was then immediately added to the mixture and incubated for 1 h at 37°C. Transformed cell were selected on LB agar plates supplemented with ampicillin (0.5 mg/ml). Complementation was confirmed by PCR analysis and bacterial growth without TPP supplementation.

P. aeruginosa acute murine infection model

Inbred 6-week-old female C57BL/6 mice were purchased from Orient Bio (Seongnam, Korea) and maintained in our animal facility, which cared for the animals in accordance with the institutional guidelines. All mice were subjected to a 1-week adjustment period prior to the experiment. The protocol was approved by the Animal Care and Ethics Committee of Institut Pasteur Korea (Approval number IPK-17014).

The acute infection model was established based on previously published methods (52). Bacteria were grown to mid-log phase and washed three times with Dulbecco’s PBS (DPBS). Mice were anesthetized with a mixture of ketamine and xylazine and inoculated intranasally with 2 × 10^7 CFU bacteria suspended in 20 μl DPBS. At 20 h after inoculation, mice were sacrificed using isoflurane, and both lungs and the spleen were harvested. The left lung and spleen were homogenized in PBS and then plated separately onto LB agar plates for WT and complementation strains; the LB agar was supplemented with TPP for the ΔthiL strain. The plates were incubated for 20 h at 37°C, and then the viable bacterial cells were counted to determine the number of bacteria in the organs. The cranial lobe of the lungs was placed in 10% formalin for histopathological analysis by H&E staining. For histopathological analysis, 4-μm-thick sections of fixed lungs were blindly analyzed for cellular infiltration using Image-Pro Plus v.4.5 (Media Cybernetics, Rockville, MD). Briefly, degrees of neutrophil infiltration were assessed as percentages of inflammatory lesions per total area in microscopic images (magnification, ×40) of the lung sections. Another set of mice (n = 5) were subjected to body weight measurements and assessments of clinical signs of infection, and results were recorded until 72 h after inoculation. At the end of the experiment, the mice were euthanized using isoflurane.

Determination of proinflammatory cytokine plasma concentrations

At the time of sacrifice, an average of 0.7 ml of whole blood was withdrawn from each mouse by cardiac puncture and collected in heparin-treated tubes. The blood was immediately centrifuged at 2,000 × g for 20 min to obtain plasma. The isolated plasma was fast-frozen on dry ice and stored at −80°C. To measure cytokine concentrations, we used Quantikine ELISA kits for IL-6, TNFα, MIP-2, and IL-1β (R & D Systems, Minneapolis, MN). Each assay was performed according to the manufacturer’s instructions.
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**Data availability**

All data are contained within the article and the supporting information.

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**Conflict of interest**—The authors declare that they have no conflicts of interests with the contents of this article.

**Abbreviations**—The abbreviations used are: TPP, thiamine pyrophosphate; ThiL, thiamine monophosphate kinase; ThiE, thiamine phosphate synthase; ThiK, thiamine kinase; TMP, thiamine monophosphate; TPK, thiamine pyrophosphate kinase; TNFα, tumor necrosis factor α; MIP-2, macrophage inflammatory protein 2; PaThiL, *Pseudomonas aeruginosa* thiamine monophosphate kinase; LB, Luria-Bertani; IPTG, isopropyl β-d-thiogalactopyranoside; DPBS, Dulbecco’s phosphate-buffered saline.

**References**

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