Computationally designed pyocyanin demethylase acts synergistically with tobramycin to kill recalcitrant *Pseudomonas aeruginosa* biofilms

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*Pseudomonas aeruginosa* is an opportunistic human pathogen that develops difficult-to-treat biofilms in immunocompromised individuals, cystic fibrosis patients, and in chronic wounds. *P. aeruginosa* has an arsenal of physiological attributes that enable it to evade standard antibiotic treatments, particularly in the context of biofilms where it grows slowly and becomes tolerant to many drugs. One of its survival strategies involves the production of the redox-active phenazine, pyocyanin, which promotes biofilm development. We previously identified an enzyme, PodA, that demethylated pyocyanin and disrupted *P. aeruginosa* biofilm development in vitro. Here, we asked if this protein could be used as a potential therapeutic for *P. aeruginosa* infections together with tobramycin, an antibiotic typically used in the clinic. A major roadblock to answering this question was the poor yield and instability of wild-type PodA purified from standard *Escherichia coli* overexpression systems. We hypothesized that the insufficient yields were due to poor packing within PodA’s obligatory homotrimeric interfaces. We therefore applied the protein design algorithm, AffiLib, to optimize the symmetric core of this interface, resulting in a design that incorporated five mutations leading to a 20-fold increase in protein yield from heterologous expression and purification and a substantial increase in stability to environmental conditions. The addition of the designed PodA with tobramycin led to increased killing of *P. aeruginosa* cultures under oxic and hypoxic conditions in both the planktonic and biofilm states. This study highlights the potential for targeting extracellular metabolites to assist the control of *P. aeruginosa* biofilms that tolerate conventional antibiotic treatment.

*Pseudomonas aeruginosa* | biofilms | antibiotic tolerance | pyocyanin | AffiLib

The opportunistic human pathogen, *Pseudomonas aeruginosa*, is able to form biofilms that are notoriously problematic and difficult to treat. Biofilms are multicellular structures that are characterized by adherence of microorganisms to each other and to surfaces via a secreted polymeric matrix. *P. aeruginosa* species are known to form biofilms within host tissues (e.g., diabetic foot ulcers and the lungs of cystic fibrosis patients) and on medical devices (e.g., implants and catheters) (1). While the cells within a biofilm are metabolically active, they can tolerate high concentrations of antibiotics (2), making infections difficult to clear. For this reason, studying how *P. aeruginosa* cells sustain their metabolism in biofilms and how biofilms can be disrupted is of particular interest.

During *P. aeruginosa* biofilm development, cells produce redox-active metabolites known as phenazines. One such phenazine, pyocyanin (PYO), is blue in color and can be found in pus from wound infections (3, 4) and in lung infection sputum samples (5). PYO confers a competitive advantage for *P. aeruginosa* by contributing to iron acquisition (6), anaerobic energy conservation (7), signaling (8), and biofilm development (9). It has also been shown that *P. aeruginosa* cells lacking the ability to synthesize phenazines (10), and PYO in particular (11), are more sensitive to conventional antibiotics when grown both planktonically and in the biofilm state; the mechanisms underpinning these phenomena are beginning to be elucidated (12). Accordingly, we reasoned that enzymatically degrading PYO within *P. aeruginosa* biofilms might improve conventional drug treatment, representing a strategy for biofilm control.

Previous work in our laboratory identified and isolated a *Mycobacterium fortuitum* enzyme (PodA) that demethylated pyocyanin to another phenazine derivative, 1-hydroxy-phenazine (SI Appendix, Fig. S1A) (13):

Pyocyanin<sub>red</sub> + H<sub>2</sub>O $\xrightarrow{PodA}$ 1 − OH − Phenazine<sub>red</sub> + CH<sub>3</sub>O

Consistent with the requirement of PYO for *P. aeruginosa* biofilm formation and metabolic maintenance (14, 15), addition of this enzyme inhibited biofilm development (13). Motivated by these preliminary observations, we sought to further explore PodA’s therapeutic potential under in vitro conditions that were more relevant to chronic infections, such as those that characterize the mucus-filled lungs of individuals living with cystic fibrosis (16). While our preliminary studies indicated that PodA is an effective *P. aeruginosa* biofilm disruptor (13), its purification through standard heterologous overexpression provided insufficient yield to enable physiological studies to reveal a novel enzyme’s therapeutic potential.

**Significance**

*Pseudomonas aeruginosa* is a major cause of hospital-acquired infections due to its formation of biofilms that are highly tolerant to antibiotics. Conventional drugs often fail to kill slowly growing biofilms because they do not target the mechanisms that sustain cells in this state; alternative biofilm control strategies are thus urgently needed. One way in which *P. aeruginosa* builds robust biofilms is through the production of redox-active phenazines such as pyocyanin. We identified an enzyme that degrades pyocyanin but were stymied in studying its potential to combat biofilms due to its poor expression yield. Here we show how protein design can stabilize the enzyme to improve purification yields, enabling physiological studies to reveal a novel enzyme’s therapeutic potential.


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studies, posing a major barrier to exploring its potential synergy with clinical antibiotics. 

There are many factors that contribute to protein stability, ranging from exposure of hydrophobic surfaces (17), entropy of folding (18), cellular turnover by proteases (19, 20), potential production of misfolded or aggregated protein (21), and the atomic contacts that stabilize the native state (22). Overexpressing a protein in a heterologous system may also expose incompatibilities between the host’s protein-synthesis machinery and the target protein (21). Attempts to produce stable protein variants through phylogenetic analyses or the application of in vitro evolution have proven very successful. Nevertheless, these conventional strategies are especially difficult to apply to enzymes like PodA, which require time-consuming protein-production protocols that are impossible to implement in medium or high-throughput screening. By contrast, in recent years, computational design methods that use a combination of phylogenetic analysis and atomistic design calculations have exhibited much higher accuracy than conventional optimization strategies (23), enabling the improvement of protein stability, expressibility (24), and activity (25, 26) through the experimental screening of only a few dozen designs.

The goals of this study were three-fold: first, to increase the yield and stability of PodA through the AffiLib protein design method (25); second, with sufficient protein in hand, to characterize the enzymatic activity of designed PodA under a variety of conditions; third, to perform Michaelis-Menten kinetics of PodA variants to determine the optimal conditions for activity.

![Graphs and tables showing specific activity, buffer, salt, and viscosity sweeps for WT PodA and PodA variants, as well as Michaelis-Menten kinetics for PodA8 and PodA10.](https://doi.org/10.1073/pnas.2022012118)
clinically relevant conditions; third, to determine whether the addition of PodA (i.e., PYO depletion) could increase efficacy of tobramycin, one of the most commonly used antibiotics to treat chronic *P. aeruginosa* infections, in different growth media. Our results demonstrate the potential of PodA to assist the treatment of chronic infections. Due to increasing rates of multidrug resistance and *P. aeruginosa* biofilms’ extreme tolerance to antibiotics, it is crucial that we find alternative mechanisms for treating such infections.

**Results**

**PodA Activity Is Stable Across Conditions Relevant to Cystic Fibrosis Sputum.** To assess PodA’s potential for therapeutic relevance in the context of human chronic infections, such as those in the cystic fibrosis (CF) airways, we began by quantifying PodA activity and determining its kinetic parameters under a spectrum of physiologically relevant variables (i.e., pH, viscosity, and sodium and potassium concentrations). As done previously to facilitate PodA purification (13), we excluded a predicted N-terminal transmembrane domain, resulting in a wild-type (WT) variant called WT PodA	extsubscript{30–162}. Moving forward, all purified PodA proteins maintained this deletion. WT PodA	extsubscript{30–162} denaturation activity was measured using a spectrophotometric assay that monitors the absorbance of PYO over time. Using a variety of buffers to analyze activity from pH 5.5 to 10.0, we found that WT PodA	extsubscript{30–162} had optimal activity at pH 6 (Fig. 1A). Activity of WT PodA	extsubscript{30–162} below pH 5.5 was not established due to the pKₐ of PYO (pKₐ = 4.9), which changes the chemical and therefore absorptive properties of the substrate. The pH of CF airway surface liquid is acidic and is thought to increase viscosity of CF mucus by influencing mucin electrostatic interactions (27). Using a reaction condition with a pH of 6, a slight increase in viscosity with ethylene glycol or mucin electrostatic interactions (27). Using a reaction condition and is thought to increase viscosity of CF mucus by influencing ties of the substrate. The pH of CF airway surface liquid is acidic 0.018, a

yields (typically with respect to the enzyme

sodium in cystic fibrosis sputum are 66 mM (Na⁺) a n d1 5m M

0.24 activity compared to controls, as previously seen (13). A variant tors the absorbance of PYO over time. Using a variety of buffers to

stabilize PodA in the hopes of efficiently overcoming this

screening such as in vitro evolution or deep mutational scanning. Because PodA is a homotrimer, all modeling and design simulations used symmetric sequence, backbone, and rigid-body sampling. Furthermore, because the available crystallographic structure of PodA shows that each monomer binds a Ca²⁺ ion (SI Appendix, Fig. S2), we disallowed any change in amino acid identity or conformation at the ion-ligation sites. The designs were ranked according to their computed energies, and the top 35 designs were visually inspected. From those, we chose to test a diverse set of 10 mutants that exhibited improved core packing or hydrogen-bonding interactions across the obligatory interfaces (Fig. 2A–D).

Induction studies showed that a majority of the PodA variants were shuttled into inclusion bodies (SI Appendix, Fig. S3). A protocol was developed (Materials and Methods) to denature and purify WT PodA	extsubscript{30–162} from inclusion bodies, followed by a refolding procedure. Before moving forward using the inclusion body purification exclusively for the designed variants, we verified that WT PodA	extsubscript{30–162} activity and kinetics were not altered after denaturing and refolding the enzyme. Soluble fraction WT PodA	extsubscript{30–162} had a catalytic efficiency of 1.05 ± 0.093 × 10⁻³ M⁻¹ s⁻¹ and WT PodA	extsubscript{30–162} purified from inclusion bodies of the same prep had a catalytic efficiency of 1.14 ± 0.25 × 10⁻³ M⁻¹ s⁻¹.

Remarkably, eight PodA designs exhibited increased protein yields relative to the wild-type protein when purified from inclusion bodies (Fig. 2E). This is consistent with our hypothesis that the trimeric interfaces in the wild-type PodA were unstable and that improving packing and interchain contacts at these interfaces might improve protein production yields. Of these, eight, wild-type levels of specific activity were maintained for PodA8 and PodA10 (Fig. 1B). We compared the kinetics of these proteins and found no change in catalytic efficiency between PodA	extsubscript{10} and PodA8 and a slight increase in efficiency for PodA10 (Fig. 1C). Due to the slightly increased efficiency combined with a higher yield, we decided to use PodA10 to study its effects on *P. aeruginosa*. As in Fig. 1L with WT PodA	extsubscript{30–162}, we analyzed PodA10 activity across a broad range of buffers, salts, and viscosity agents. Unlike WT PodA	extsubscript{30–162}, PodA10 maintained activity over a large range of pH, with no observable preference between a pH of 5.5 and 9 (Fig. 1D), indicating that the design was also more stable to environmental conditions than the wild-type enzyme. PodA10 activity was not altered with addition of salts, and a slight advantage was seen with addition of ethylene glycol (Fig. 2D), similar to WT PodA	extsubscript{30–162}. Due to the high thermal stability of PodA, we were unable to complete stability assays that rely on monitoring protein unfolding over a temperature gradient. We therefore analyzed the enzyme’s kinetic stability by boiling WT PodA	extsubscript{30–162} and PodA10 and measuring activity of each protein after specified boiling times. After 30 and 60 min of boiling, PodA10 had significantly higher activity compared to WT PodA	extsubscript{30–162} (SI Appendix, Fig. S4). Thus, the optimized intersubunit interactions in the designed PodA10 resulted in significantly higher *Escherichia coli*
expression levels and kinetic stability compared to the parental enzyme. We also noted that several designed mutations in PodA8 and PodA10 were not prevalent in a multiple-sequence alignment of PodA homologs (SI Appendix, Fig. S5), indicating the importance of the atomistic calculations to design success.

Addition of PodA10 to P. aeruginosa Cultures Enhances Planktonic Killing and Blocks Biofilm Development. Prior to testing the impact of PodA10 on biofilms, we sought to determine its effects on P. aeruginosa grown planktonically. To ensure that its activity and kinetics would not be compromised in physiologically relevant conditions, the yield of trimer interface designs purified from inclusion bodies per liter of overexpression culture is shown in Table E. Designed Mutations. Fig. 2. Structure-based design of stabilized PodA variants. (A) PodA forms a symmetric trimer. Each monomer is colored differently, and positions that were subject to design are labeled with their mutation in PodA10 and marked by spheres. Ca²⁺ ions are colored orange. (B–D) Selected mutations implemented by AffiLib and their structural context. Colors for each monomer (green, pink, blue) are as in A. (B) Mutation Ala87Val (sticks) increases core packing in a hydrophobic region. (C) Mutation Ile73Thr improves surface polarity. (D) Mutation Ala53Asn form an interfacial hydrogen bond with a backbone carbonyl. (E) Yield of trimer interface designs purified from inclusion bodies per liter of overexpression culture.

Fig. 3. Effects of PodA10 on planktonic viability and crude biofilm development. Cell viability counts of P. aeruginosa liquid media synergy experiments with PodA10 (0 or 5 μM) and tobramycin (concentrations indicated on the x axis) for LB (A) and SCFM (B). Data represent the mean of biological triplicates with error bars representing the SEM. NG, No Growth. (C) Crystal violet biofilm assay. P. aeruginosa cells were inoculated in a 96-well polystyrene plate containing minimal medium with arginine (40 mM) and no PodA, WT PodA, PodA10, or inactive PodA10, followed by incubation at 37 °C. After 24 h of growth, planktonic cells were washed away, and attached cells were stained with crystal violet. Cells were destained with an acetic acid and methanol mixture, after which biomass was quantified based on crystal violet absorbance at 550 nm. Error bars represent SEM. For all graphs, an unpaired t test was used to calculate P values between treatments, ns, not significant; *P < 0.05; ****P < 0.0001. For both panels, PodA10inactive represents the variant PodA10H121A,E154A,Y156A.
growth medium, we tested PodA10 PYO demethylation activity in lysogeny broth (LB) and synthetic cystic fibrosis medium (SCFM) (16). We saw no change in catalytic efficiency in either medium compared to experiments performed in buffer (SI Appendix, Fig. S6), allowing us to move forward with studying PodA10 effects on \textit{P. aeruginosa}. We proceeded to ask whether PodA is synergistic with tobramycin (an antibiotic commonly used to treat \textit{P. aeruginosa} infections in the clinic). Tobramycin is an aminoglycoside that targets the ribosome, and its efficacy requires a sufficient proton motive force for entry (29, 30). The minimum inhibitory concentration (50\%) for tobramycin on \textit{Pseudomonas} ranges from 1 to 3 \( \mu \text{g} \cdot \text{mL}^{-1} \) (31), with the target tobramycin concentration in sputum of cystic fibrosis infections being \( \sim 130 \mu \text{g} \cdot \text{mL}^{-1} \) (32).

To assess whether PodA10 is synergistic with tobramycin, we grew planktonic cultures of \textit{P. aeruginosa} strain PA14 to high cell viability...
densities in LB or SCFM over a range of concentrations of PodA10 (0 to 20 μM) and tobramycin (0 to 100 μg·mL⁻¹). Cells were incubated statically. After incubation, cells were resuspended and reinoculated into fresh medium. Lag times were calculated for each well, and we hypothesize that increased lag times reflect increased cell death from the treatment. Based on the results of these screens, we identified synergistic concentrations of PodA and tobramycin to be analyzed in triplicate. A minimum of 10 mg of PodA was needed to complete preliminary experiments in triplicate, followed by targeted experiments in biological triplicate with technical triplicate in LB and SCFM. To perform these studies, we would have had to purify protein from over 30 L of culture for WT PodA30–162 but, due to the improvements in yield observed for PodA10, we were able to achieve our goals using only 1.6 L of culture—a substantial improvement in experimental efficiency.

In LB, addition of PodA10 (>1 μM) together with tobramycin increased lag times by 1.5 to 2.0 h compared to PodA (−) controls (SI Appendix, Fig. S7). To verify that lag times were due to increased cell death, cells were plated to count colony forming units (CFUs). PodA10 (5 μM) alone did not increase cell death, but when PodA10 was added with tobramycin, cell viability greatly decreased compared to no protein or inactive PodA10 controls (Fig. 3A). The same experiments were repeated in SCFM, a minimal medium whose composition is based upon that of splash media (31) from cystic fibrosis patients (16) and which elicits a similar response from P. aeruginosa grown in vitro compared to the in vivo CF lung environment (33). While slight differences were seen in lag times when PodA10 was added with tobramycin, the effects were not as dramatic as seen in our experiments with LB (SI Appendix, Fig. S7). However, when plating the same SCFM cultures to calculate viability via CFUs, there was a significant decrease in survival when PodA10 (5 μM) was added with tobramycin compared to no protein or inactive PodA10 controls (Fig. 3B). We speculate that the inconsistent lag times observed in our experiments (SI Appendix, Fig. S7) were due to cells adjusting from being treated in minimal defined medium (SCFM) and then grown in rich medium (LB), an observation that is common in microbiological growth experiments. This indicates that direct cell counts are a better test for analyzing PodA10 and tobramycin synergistic killing.

Having established that PodA10 enhances killing of planktonic P. aeruginosa, we wanted to verify that this variant could also inhibit biofilm formation as shown previously for WT PodA30–162 (13). For our first biofilm test, we used the biofilm crystal violet microtiter plate assay to measure the amount of biomass attached to microtiter plate wells after a defined period of time (34). Incubation for 24 h with addition of PodA10 from the beginning of the assay led to a biofilm attachment defect that was comparable to that achieved by WT PodA30–162: addition of inactive PodA10 had no effect on attachment (Fig. 3C).

As a final test of PodA effects being attributable to PYO removal, Pseudomonas cells that could not synthesize PYO (pH2M::aacC) were subjected to the same experimental setup as in Fig. 3. For liquid cultures, there were no significant differences when PodA10 was added in combination with tobramycin, compared to the tobramycin-only controls (SI Appendix, Fig. S8). Addition of WT PodA30–162/PodA10, or inactive PodA, did not impact attachment of a PYO-deficient Pseudomonas strain compared to no protein controls (SI Appendix, Fig. S8). These results establish that the decreased viability observed for the addition of PodA10 (Fig. 3) is specifically due to the elimination of PYO from the cultures.

**Addition of PodA10 to Aggregate Biofilms Together with Conventional Antibiotics Leads to Synergistic Killing.** While the crystal violet assay permits biofilm phenotypes to be rapidly screened, it only provides a crude measurement of a biofilm defect related to biofilm development and attachment. To better understand how PodA10 impacts mature biofilms and their viability, we turned to the agar block biofilm assay (ABBA), which was designed to enable the experimental study (13, 35) of the type of aggregate biofilms that characterize human chronic infections (36). Hundreds of distinct biofilm aggregates grow in a single ABBA, allowing for simultaneous observation of biofilms of different sizes. Due to the ABBA being low throughput, we chose to test a single tobramycin concentration (50 μg·mL⁻¹) with PodA10 (5 μM) in LB and SCFM, as these concentrations showed the most dramatic synergistic effects in liquid culture (Fig. 3 and SI Appendix, Fig. S7). Cells were inoculated into the agar and allowed to grow into aggregates for 24 h (LB) or 12 h (SCFM), after which PodA10 and tobramycin were added simultaneously and allowed to incubate for 12 h (LB) or 8 h (SCFM) (Fig. 4A). Addition of PodA10 and tobramycin for LB and SCFM was administered at the latest time point for which an oxygen gradient was present (24 h for LB; 12 h for SCFM) (SI Appendix, Fig. S9). This ensured that biofilms were at their most mature state, while still exhibiting metabolic stratification. After treatment, samples were stained with propidium iodide (PI), a fluorescent DNA-binding dye that is incapable of passing through the membrane of highly energized cells and imaged. Generally, higher PI staining indicates lower cell viability due to the compromised membranes of these cells. Aggregates were analyzed via confocal microscopy throughout the depth of the ABBA, with each aggregate representing a distinctly growing Pseudomonas biofilm.

Incubation of LB-grown aggregates with tobramycin and PodA10 led to PI staining intensity that was significantly greater than PodA10 or tobramycin alone (Fig. 4B). To quantify this visual effect, a three-dimensional mask was generated around each aggregate, and the average PI fluorescence of each aggregate was measured as a function of depth from the top of the agar block (Fig. 4C). In LB, the average PI intensity with PodA10 and tobramycin was higher than the PI intensity of tobramycin alone. To ensure higher intensities were not due to differing aggregate sizes, we plotted the average volume of each aggregate over depth and saw no significant differences in volumes between treatments (SI Appendix, Fig. S10). Plotting the aggregate volume vs. PI intensity of each aggregate further showed that PI staining was highest in the PodA10 + tobramycin treated samples (SI Appendix, Fig. S10). These staining results correlated with cell viability, which was determined by homogenizing ABBAs and plating in LB. ABBAs treated in minimal defined medium (SCFM) and then grown in LB showed the lowest viable cell counts compared to ABBAs incubated with tobramycin alone or tobramycin and inactive PodA10 (Fig. 4D).

For the LB ABBA, the largest differences in PI staining between tobramycin and tobramycin + PodA10 (Fig. 4 C and F) were between 0 and 300 μm, a region previously characterized as theoxic/hypoxic zone within the agar (13), with oxygen respiration contributing to increased tobramycin efficacy (29). To confirm this, parallel experiments were set up identically, and oxygen profiles were measured using microelectrodes. We observed an oxygen gradient through the depth of the LB ABBA that correlated with the viability patterns observed (Fig. 4C and SI Appendix, Fig. S9). Importantly, the oxygen gradient changed over time: at 12 h, no oxygen was present at a depth of ∼300 μm (SI Appendix, Fig. S9); however, by 24 h, oxygen was again detected at the lower depths. We speculate that these dynamics reflect a change in the limiting nutrient at these different time points: first oxygen, then carbon. Once carbon is depleted, oxygen concentrations rise again due to diffusion and a lack of cellular consumption.

At smaller scales, an oxygen gradient also exists within each single aggregate, due to cellular oxygen consumption at the periphery outpacing O₂ diffusion toward the center of the aggregate (37). Additionally, phenazine gradients have been shown to exist within biofilms, with PYO localized to the periphery (38), reflecting the requirement for molecular oxygen in PYO biosynthesis. Analyzing data from Fig. 4B in an XY plane rather...
than an XZ plane, we find a staining pattern consistent with such gradients: untreated aggregates grown in LB show an outer ring of PI staining that is abolished by addition of PodA10 (SI Appendix, Fig. S11).

Similar synergistic effects between PodA10 and tobramycin treatment were observed for biofilm aggregates grown in SCFM. Qualitative differences in PI staining trends were also seen in this medium (Fig. 4E), with the PI intensity of the PodA10 + tobramycin treatment being higher than tobramycin alone (Fig. 4F). This difference was similar to that seen in LB and was also manifested in the viability counts (Fig. 4G). Microelectrode measurements of 12-h SCFM ABBAs revealed an oxygen gradient decline over depth, mirroring the higher PI staining intensity throughout the upper depths of the ABBA in the presence of tobramycin (Fig. 4F).

Discussion

As a first step toward exploring PodA’s therapeutic potential, we sought to increase its yield during heterologous overexpression. The AffiLib design approach that we utilized was recently applied to the obligatory interfaces in the variable domains of several antibodies, leading to substantial increases in production yields and thermal resistance (28). Applying AffiLib to PodA successfully increased the production yields 20-fold by optimizing contact interaction with tobramycin produced interfaces. This enabled us to study the physiological effects of PodA on planktonic and biofilm-grown P. aeruginosa cells under conditions relevant to human infections, revealing significant synergistic killing with the commonly used clinical antibiotic tobramycin.

The phenazine PYO is beneficial to P. aeruginosa biofilm development and metabolic activity (14, 15), and cells that cannot synthesize phenazines have reduced rates of survival in anoxic conditions in the absence of alternative terminal electron acceptors (9, 39). Phenazines must be in an oxidized form to act as electron acceptors, so we wondered whether the conversion of PYOox to 1-OH-PHZred via PodA would be detrimental to P. aeruginosa metabolic vitality within the anoxic zone. Because an oxygen gradient was present throughout the ABBA aggregates, we could examine this effect in deeper regions (>300 μm) of the ABBA. Interestingly, PodA addition alone did not significantly change PI staining at depth (>300 μm), where oxygen is consumed more rapidly; the possibility of PI staining at this depth is limited to the upper depths of the ABBA (Fig. S12).

As expected, treatment with PodA10 + tobramycin showed an outer ring of PI staining characteristic of these cells surviving by cycling phenazines (7), the membrane potential of cells at this depth of the ABBA system is sufficiently low that PI can be taken up as efficiently by metabolically active cells as by dead cells, thus obscuring a viability readout; and/or 3) PYO is not the primary phenazine sustaining metabolic activity in the anoxic zones; rather, another phenazine, such as phenazine-1-carboxamide [which can also sustain anaerobic survival (40) and localizes to the interior of large anoxic colony biofilms grown on 1% tryptone (38)], may maintain viability under these conditions.

In contrast to the anoxic zone in the ABBA, PodA addition in combination with tobramycin produced a striking effect in theoxic/hypoxic zones of the ABBA assay in LB and SCFM (Fig. 4). Additionally, PodA-only treatments in 24-h LB ABBA showed a lower PI staining pattern in the oxic region. Importantly, while PYO is beneficial to cells when they are electron-donor replete but oxidant-limited, PYO is toxic to P. aeruginosa under regimes where electron donors are limited but oxidants are replete, such as in theoxic/hypoxic zone of older cultures, where PYO can react with oxygen and cause oxidative stress (41). This fact, combined with the knowledge that PYO is maximally concentrated in the outer regions of large colony biofilms grown on 1% tryptone (38), suggests the following explanation of the effects of PodA on biofilm viability in our 24-h LB ABBA experiments: For untreated cells that are electron donor-limited, PYO causes localized toxicity at the oxic interface of biofilm aggregates. Addition of PodA alone removes PYO from oxic regions, leading to greater metabolic activity of aggregates in these zones, as well as in the outer rings of large aggregates (reflected by lower PI staining). In contrast, when PodA is added together with tobramycin, this enhancement of metabolic activity is a double-edged sword. Greater metabolic activity—specifically, having an inner membrane that is sufficiently energized to be able to take up tobramycin—has been shown to be correlated with greater tobramycin susceptibility (29). Accordingly, in LB and SCFM ABBA regions that have both oxygen and PYO and sufficient carbon, the depletion of PYO by PodA sensitizes these cells to tobramycin treatment. In the LB ABBA, oxygen was consumed by 300 μm, leading to metabolic restriction after 12 h (SI Appendix, Fig. S9), which correlated with no significant differences in PI staining between nontreated and tobramycin-treated samples at these lower depths (Fig. 4C).

It is likely that the differences in effects observed for PodA treatment (in the absence of tobramycin) in LB and SCFM simply reflect differences in timing of treatment and media composition, with varying electron donor:electron acceptor abundance. Notably, SCFM has 3 mM glucose and 9 mM lactate as a carbon source, in contrast to LB that has only amino acids and purines/pyrimidines as carbon sources (42). It is therefore possible that, in the oxic zone at 12 h, cells in SCFM are more protected from PYO toxicity than in LB at 24 h due to a higher electron donor:electron acceptor ratio (41). Finally, the ratios and production of different phenazines (e.g., PYO, PCA, PCN) in P. aeruginosa are known to vary depending on the carbon source in the growth medium (43); such variation could also contribute to the differences that we observed in PodA effects between our media.

Overall, this study demonstrates the utility of protein design methods such as AffiLib to optimize the yield of proteins of interest such as PodA. The significant improvement of protein production yields by optimizing the oligomeric interfaces of PodA and by experimentally testing only 10 designs suggests that other obligatory complexes could similarly be stabilized; this may be quite significant given that a majority of natural proteins form homo-oligomers (44). The availability of a functionally expressible design enabled physiological studies that would have been practically inaccessible otherwise. Because PYO and other phenazines permit P. aeruginosa to adopt metabolic strategies that lead to evasion of antibiotic treatments (10–12), it stands to reason that removal of these metabolites might offer an attractive therapeutic approach. Our results showing enhancement of synergistic killing by a designed PodA and tobramycin provide further motivation and the means to explore PodA’s potential to be used as a biologic therapeutic for treating chronic P. aeruginosa infections. Future experiments will include studying the effects of PodA10 and tobramycin in animal models to determine treatment outcomes.

Materials and Methods

Bacterial Strains, Culture Media, and Chemicals. Strains used in this study are listed in SI Appendix, Table S1. E. coli BL21 (DE3) (43) was used for protein overexpression and E. coli DH5α (New England Biolabs) was used for plasmid construction. All E. coli strains were grown at 37°C in LB (Difco) or Terrific Broth (TB, Difco). Ampicillin for culturing E. coli was used at 100 μg mL⁻¹. P. aeruginosa strain UCBPP-PA14 (hereafter P. aeruginosa) was used for growth analyses and cultivated on LB. The following chemicals were purchased from Sigma-Aldrich unless otherwise noted: glycerol (VWR), Hepes (Gold BioTechnology), and sodium chloride (Fisher Scientific).

Plasmid Construction. All plasmids used in this work are listed in SI Appendix, Table S2. Primers were synthesized by Integrated DNA Technologies (IDT) and are listed in SI Appendix, Table S3. For heterologous protein expression, the gene coding for PodA was amplified from M. fortuitum strain.
ATCC 8641 using Phusion High Fidelity DNA Polymerase (ThermoFisher Scientific) per manufacturer’s instructions and cloned into an isopropyl-β-D-thiogalactoside (IPTG)-inducible vector pTEV16 (NEB), using Type II Restriction BspQI cloning (47). The resulting plasmid was referred to as pPodA1. Catalytic PodA variants were utilized from a previous study (13). For designed proteins, gBlocks were synthesized from IDT to code for relevant amino acid substitutions, with 5’ and 3’ base pairs corresponding to the multiple cloning site (MCS) of pTEV16. One set of primers was used to amplify each gBlock and another set to amplify pTEV16. PCR constructs were isolated using the Monarch PCR & DNA Cleanup Kit (NEB). Genes were then cloned into pTEV16 using Gibson Assembly Master Mix (NEB) per manufacturer’s protocol. The resulting plasmids (pPodA2 to pPodA11) are listed in Table S2.

Protein Overproduction and Purification of Soluble and Inclusion Body Fractions. Plasmids coding for proteins of interest were transformed into E. coli BL21 (DE3). Overnight cultures (10 mL) of transformants were sub-cultured (1% [vol/vol]) into TSB (10 L for WT PodA and 1 L for PodA variants) containing ampicillin. Cultures were grown at 37 °C with shaking to an optical density of 0.4 (OD600nm), and plasmid expression was induced with IPTG (50 μM) and shaken overnight at 16 °C. Cells were harvested by centrifugation at 5,000 g for 10 min in a Beckman Coulter Avanti J-20 XIO refrigerated centrifuge using a JLA-8.1000 rotor at 4 °C. Cell pellets were stored at -80 °C until use.

For purification, cell pellets were thawed and resuspended in 30 mL buffer A (Hepes [50 mM, pH 7.5], NaCl [500 mM], and imidazole [20 mM]) containing lysozyme (1 mg mL−1). DNase (5 μg mL−1), and protease inhibitor (phenylmethylsulfonyl fluoride, 0.5 mM). Cells were lysed by four passages through an Avemist Emulsiflex C3 (ATA Scientific) at 15,000 psi. Due to difficulties filtering lysate in subsequent steps, additional DNase (5 μg mL−1) was added before filtration. Clarified lysate was applied to a pre-equilibrated 5-mL HisTrap FF column (GE Healthcare) using a 25 mL, 200 mM buffer and incubated in a PCR thermocycler at 100 °C for time periods listed in Table S1. Volume was increased to 20 mL with Hepes (50 mM, pH 7.0), NaCl (500 mM), 30% glycerol (vol/vol), and shaken overnight at 16 °C. Using an AKTA FPLC, lysate was applied to a pre-equilibrated 5-mL HisTrap FF column (Sigma-Aldrich) after which the column was washed with 10 column volumes (CV) of buffer A, 7 CV of 8% buffer B (Hepes [50 mM, pH 7.5], NaCl [500 mM], and imidazole [500 mM]), and a gradient to 100% buffer B over 10 CV (50 mM Hepes [pH 7.5], NaCl [500 mM], and imidazole [500 mM]). Fractions containing PodA were combined and cleaved for 3 h at 25 °C with tobacco etch virus (TEV) protease (1:50 mg:mg ratio of TEV to PodA). The combined and cleaved for 3 h at 25 °C with TEV protease (1:50 mg:mg ratio of TEV to PodA). The flow through was confirmed by SDS/PAGE. Cleaved PodA protein was then dialyzed for storage at 4 °C in Hepes (50 mM, pH 7.5), NaCl (250 mM), and 30% glycerol (vol/vol). Finally, cleaved PodA was stored at −80 °C until use.

Protein Design Calculations. Using the PodA structure as a starting point (Protein Data Bank ID 5K21), we selected 13 positions in the homotrimeric interfaces for design: S3A1u, 64Val, 67Met, 73Ile, 87Ala, 91Thr, 92Asn, 99Met, 112Gil, 118Leu, 129Ala, 134Thr, and 141Lys. As previously described (25, 26), we generated a Position Specific Scoring Matrix (PSSM) using the default parameters and calculated the tolerated sequence identities at each of the 13 positions with PSSM cutoffs ≥ 2 and Rosetta ΔΔG < 6 Rosetta energy units. Next, we enumerated, modeled, and refined in Rosetta all the possible combinations of mutations allowed by the tolerated sequence space that differed from the WT protein by 3 to 5 mutations (a total of 118,424 combinations of mutations). During all Rosetta modeling and design calculations, the structure was modeled with the Ca2+ ions, and their ligating residues were unchanged and subject to C3 symmetry constraints. We then ranked the designs according to Rosetta energy and clustered them, retaining the low-energy designs that exhibited at least two mutations relative to another. We selected 10 for experimental screening out of the top 35 by visual inspection. The command lines and RosettaScripts (49) used to run the symmetric refinement, mutational scanning, and design trajectories are provided in supplemental files refine_auto.xml and filter_scan_auto.xml and mutate_auto.xml. The PSSM file used in the design calculations is available as file 5K21_pssm.txt, and the results of the mutational scanning calculations are provided as refsite_ddg6.txt.

A web-accessible version of RosettaLib (for nonsymmetric design) is available at http://siftlib.weizmann.ac.il for academic users.

Synthesis of PYO. PYO was synthesized from phenazine methosulfate using a protocol described previously (50). Deviations included eliminating purification of PYO via thin-layer chromatography plates and utilizing dichloromethane in the place of chloroform. PYO was analyzed for purity via high-performance liquid chromatography (HPLC) and by analysis and detection of 5-methylsulfonate, a reporter group used to determine the extent of PYO (4130 M−1 cm−1) in combination with Beer’s Law (A = εc).

Specific Activity and Kinetics of PodA. To determine specific activity and kinetic parameters of PodA for PYO, a continuous spectrophotometric assay monitoring the absorbance of PYO (690 nm) was utilized. Briefly, assays were performed at 25 °C in 100 μL reaction volumes in 96-well plates, and reactions were monitored at 690 nm over 10 min. Reaction mixtures contained phosphate buffer (50 mM, pH 6.0), ethylene glycol (1%, vol/vol), protein (3 μM), and substrate (for specific activity, 100 μM; for kinetics, varying). For each reaction, 10 μL of 25 mM IPTG solution was added to each reaction and incubated in a PCR thermocycler at 100 °C for time periods listed in Table S1. Reactions were cooled to room temperature for 1 h and diluted (3 μL) into 100 μL reaction volumes. Data were acquired using the SpectraMax M3 Microplate Reader ( Molecular Devices) using the Soft Max Pro software every 10 s over 10 min. Path lengths for each well were calculated using Soft Max Pro endpoint readings, and slopes were corrected for path lengths of 1 cm.

Specific activity was calculated from the slope of the linear range (ΔOD690 min−1) using Beer’s Law (A = εc) with a path length (l) of 1 cm and the molar extinction coefficient (ε) of PYO (4,130 M−1 cm−1). Knowing the absorption (A), this equation was solved for c, giving specific activity in nmol·min−1·mg−1 of PodA. For kinetic parameters, graphs of initial velocity (Vmax) versus substrate concentration (S) were plotted using Prism v8 (GraphPad). The Michaelis–Menten kinetics model was used to determine Km and Vmax. The Michaelis–Menten kinetics model was used to determine km and kcat of PodA. The turnover number (kcat) was determined using the following equation: kcat = kcat[E], where [E] is the concentration of PodA added. All spectrophotometric assays mentioned above were completed three times, each in technical triplicate with a representative dataset shown. Error bars represent SD as calculated by Prism v8 (GraphPad). SD for kcat and kcat were calculated using standard formulas for propagation of error.

P. aeruginosa PodA and Tobramycin Liquid Synergy Experiments. For planktonic synergy experiments, starter cultures were grown 24 h at 37 °C shaking in LB or SCFM supplemented with 30 μM FeSO4. Stationary phase cultures

8 of 10 | PNAS
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Computationally designed pyocyanin demethylase acts synergistically with tobramycin to kill recalcitrant Pseudomonas aeruginosa biofilms
VanDriese et al.
(usually between OD_{500} 4 and 6) were aliquoted (100 μL) into 96-well flat bottom wells (VWR). Tobramycin and PodA were diluted into stock concentrations in vol/vol antibiotic stock solution (biotic and 2 μL PodA per well). Tobramycin was made fresh daily and was diluted in water, and PodA was diluted in Hepes (50 mM, pH 7.5). Plates were incubated in a humid Tupperware chamber lined with wet paper towel at 37 °C for 8 h. After incubation, wells were pipetted to resuspend, and each well was diluted 1:10 into fresh LB. From that mixture, 10 μL of diluted cells were pipetted into 90 μL fresh LB in 96-well flat bottom plates. Light mineral oil (65 μL undiluted, Sigma-Aldrich) was pipetted onto the top of cultures to prevent dehydration and allow for oxygen diffusion. Microtiter plates were incubated at 37 °C inside a temperature-controlled chamber of a BioTek Synergy 4 plate reader. Plates were continuously shaken (medium setting) and time points (OD_{500}) were taken for 24 h every 30 min. Data were plotted using Prism v8. Growth studies were completed three times with a representation from one experiment shown. For concentrations that led to an increase in lag times, the same experiment was set up but, rather than an outgrowth step, cells were continuously shaken (medium setting) and time points (OD_{500}) were done with a 1:10 dilution for a final OD of 0.1. Molten LB agar (1% v/v) was added with cells simultaneously and was present during growth and biofilm development. Planktonic cells were removed, and attached cells were quantified with crystal violet staining (125 μL, 0.1% v/v), followed by washes with water and destaining with an acetic acid (30%, vol/vol) mixture in water. Crystal violet was measured using a SpectraMax M3 Microplate Reader ( Molecular Devices) at an absorbance of 550 nm. Details of this procedure were published previously (34).

**Crystal Violet Assay.** *P. aeruginosa* grown overnight in LB were inoculated (1.5%, vol/vol of OD_{500} = 5 culture) and grown for 24 h at 37 °C in 96-well round bottom plates in minimal medium (100 μL) with arginine (40 mM) as the sole carbon source. PodA (1 μM) was added with cells simultaneously and was present during growth and biofilm development. Planktonic cells were removed, and attached cells were quantified with crystal violet staining (125 μL, 0.1% v/v, made by mixing premade 2× SCFM + 30 mM FeSO_{4} and 2% agar) was incubated at 44 °C, after which 10 μL of OD 0.1 culture was added and mixed into 1 mL of agar for a final starting OD of 0.001. A portion of the mixture (200 μL) was pipetted into a well of an eight-well glass chamber slide for microscopy (Thermo Fisher Scientific #155409) or into a 2-mL Eppendorf tube for cell viability experiments. LB ABBAs were incubated at 37 °C in a humid chamber and were in- cubated in a 37 °C sand bath during oxygen probe measurements. Oxygen concentrations were measured using a Clark-type amperometric electrode with a 10-μm tip diameter, which was connected to an amplifier of a multimeter (Unisense). A two-step calibration was performed using an oxygen-free solution (0.1 M NaOH, 0.1 M sodium ascorbate) and an oxygen satu- rated LB or SCFM solution (with 1% v/vol salinity). The agar surface was found by decreasing the tip depth by 25 μm until oxygen measurements declined by 2 μmol·L^{-1}. The oxygen sensor was positioned 100 μm above the agar interface, and data were collected in 25-μm steps for a total of 700 μm. Technical triplicate measurements were made at each depth, with each measurement taking 3 s with 2 s between measurements. Oxygen profile data are averages from one experiment of technical triplicates of biologic triplicates with each experiment done in triplicate on different days. Data were collected using SensorTrace Pro-3.1.3 software and were plotted using Prism v8 (GraphPad).

**Data Availability.** All study data are included in the article and/or supporting information.

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VanDrisse et al.

Computationally designed pyocyanin dimethylate acts synergistically with tobramycin to kill recalcitrant *Pseudomonas aeruginosa* biofilms

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**Oxygen Probe Measurements.** ABBAs were inoculated and set up identically as described in *P. aeruginosa* PodA and Tobramycin ABBA Synergy Experiments. ABBAs were incubated at 37 °C in a humid chamber and were incubated in a 37 °C sand bath during oxygen probe measurements. Oxygen concentrations were measured using a Clark-type amperometric electrode with a 10-μm tip diameter, which was connected to an amplifier of a multimeter (Unisense). A two-step calibration was performed using an oxygen-free solution (0.1 M NaOH, 0.1 M sodium ascorbate) and an oxygen saturated LB or SCFM solution (with 1% v/vol salinity). The agar surface was found by decreasing the tip depth by 25 μm until oxygen measurements declined by 2 μmol·L^{-1}. The oxygen sensor was positioned 100 μm above the agar interface, and data were collected in 25-μm steps for a total of 700 μm. Technical triplicate measurements were made at each depth, with each measurement taking 3 s with 2 s between measurements. Oxygen profile data are averages from one experiment of technical triplicates of biologic triplicates with each experiment done in triplicate on different days. Data were collected using SensorTrace Pro-3.1.3 software and were plotted using Prism v8 (GraphPad).

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