Genome-scale modeling of *Pseudomonas aeruginosa* PA14 unveils its broad metabolic capabilities and role of metabolism in drug potentiation

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**Abstract**

*P. aeruginosa* is an opportunist human pathogen that is one of the leading causes of hospital-acquired infections. We have developed an updated genome-scale model (GEM) of *Pseudomonas aeruginosa* PA14 for systems-study of the pathogen. We used both automated and semi-manual approaches to reconstruct and curate the model. After an extensive literature research, we added organism-specific reactions (e.g., phenazine transport and redox metabolism, cofactor metabolism, carnitine metabolism, oxalate production, etc.) to the model. This effort led to a highly curated, three-compartment, and mass-and-charge balanced BiGG model of PA14 that contains 1509 genes, 1779 metabolic reactions and 1151 unique metabolites. The model (iSD1509) has the largest genome coverage of *P. aeruginosa* PA14 to date with 424 more genes than the previous model (iPau1129). It is also the most accurate with prediction accuracies as high as 92.4% (for gene essentiality) and 93.5% (for substrate utilization). The model simulates growth in both aerobic and anaerobic conditions. It predicts the biosynthesis of the virulence factor phenazine as a process for the pathogen to grow in low-oxygen environment. Further, a mechanism for the overproduction of a drug susceptibility biomarker (gluconate) can be elucidated by the principles of optimal growth. Finally, the model also simulates drug activity potentiation and protection by fumarate and glyoxylate, respectively, and provides mechanistic explanations for these processes. Overall, iSD1509 can be utilized to decipher the metabolic mechanisms associated with virulence and antibiotic susceptibility of *P. aeruginosa* PA14 to aid in the development of effective intervention strategies.

**Introduction**

*Pseudomonas aeruginosa* is a Gram-negative proteobacterium that is metabolically versatile and an opportunistic human pathogen. It is a leading cause of nosocomial infections\(^1\),\(^2\). One of the well-known *P. aeruginosa* infection sites is in the lungs of cystic fibrosis (CF) patients. Such infections can lead to high morbidity and mortality...
The ability to resist multiple drugs (including aminoglycosides, quinolones and β-lactams), synthesize virulence factors (e.g., phenazines, proteases, lysins, exotoxins, etc.), and produce biofilms allows *P. aeruginosa* to infect and colonize its host. The pathogenicity of *P. aeruginosa* and the host response can differ between strains. *P. aeruginosa* PA14 is a hypervirulent strain and belongs to the most common clonal group. A study in 2004 first published its genome, which is highly similar to *P. aeruginosa* PA01’s genome, but carries two additional pathogenicity islands that contribute significantly to the virulence of PA14. Therefore, the investigation of PA14 has gained particular interest in the research community.

Genome-scale metabolic models (GEMs) provide a reliable tool for systems study of bacteria. GEMs can be advantageous in investigating pathogens because identifying potential intervention strategies can be challenging due to the wide range of genetic mutations and metabolic targets, and niche-specific alteration of metabolic processes. GEMs have been utilized for systems investigation of pathogenic species such as *Acinetobacter baumannii*, *Klebsiella pneumoniae*, *Mycobacterium tuberculosis*, *Vibrio vulnificus* and *P. aeruginosa*. For the development of GEMs, first a reconstruction of the metabolic pathways of the organism of interest is required, which can then be converted to a mathematical format that can be analyzed using constraint-based modeling and flux balance analysis approaches. A GEM of *Pseudomonas aeruginosa* PA14 (referred to as iPau1129) was published in 2017 in a study investigating the association between growth and virulence-linked pathways. Several missing metabolic capabilities and standardization considerations in iPau1129 motivated us to reconstruct an updated model for PA14. For instance, iPau1129 does not possess multiple terminal oxidases, which are present in *P. aeruginosa*. Likewise, phenazine-dependent redox reactions including reactive oxygen species (ROS) formation were not included in iPau1129. We also sought to use human-interpretable metabolite and reaction identifiers standardized across a large repository of models—in particular, the Biochemical Genetic and Genomic (BiGG) standardization and modeling platform. iPau1129 was constructed using ModelSEED identifiers without cross-referencing other databases.

In this study, we developed a reconstruction of PA14 strain using BiGG identifiers with 424 additional unique genes compared with iPau1129. The model (iSD1509) demonstrated high prediction accuracy, and can also simulate growth in anaerobic environment. Furthermore, using the limed-FBA approach, we extended iSD1509 to investigate the effect of phenazine biosynthesis on the biomass production at various levels of oxygen availability. Finally, we also demonstrate the utility of iSD1509 in studying antibiotic resistance in *P. aeruginosa*.

### Materials and Methods

#### Model simulations

For the simulation of the model, we performed flux balance analysis (FBA) on conditions reflecting the media used in a particular experimental study (SI Appendix, Table S1). The FBA simulations were performed using COBRApy package (v. 0.18.1). In most simulations, we used the aerobic biomass reaction as the objective function. For anaerobic growth predictions and comparison with aerobic growth rates, we changed the objective function to the anaerobic biomass reaction. If the biomass flux was computed to less than $10^{-5}$ h$^{-1}$, “no growth” was assigned. For the computation of ubiquinone yield and growth rate comparison in aerobic and anaerobic environment, parsimonious FBA (pFBA) was applied.

For the computation of gluconate production, both FBA and flux variability analysis (FVA) (with loopless method and fraction of optimum set to 1) were applied. All
simulations were performed in M9 medium. Experimental data to validate gluconate production were obtained from [31], where measurements for RpoN mutant and wild-type were performed in M9 media. Gluconate production for mutants besides RpoN were measured by [31] in SFCM media. However, for consistency and to facilitate interpretation, we simulated all mutants in M9 medium (a minimal medium) and found that these simulations were consistent with experiments in SFCM media (a rich medium).

To compare the wild-type and mutant fluxes, we applied a heuristic approach. First, any zero flux was replaced by 0.01. To compute gluconate yield, flux of gluconate secretion reaction was divided by that of glucose exchange reaction. For FBA simulations, the reaction fluxes or yields of the mutant were divided by those of the wildtype. For FVA fluxes, an average flux of each reaction was calculated by taking the mean of minimum and maximum flux values for both mutant and wildtype. Then, the mutant average reaction flux values were normalized by those of the wildtype.

**Superoxide leakage** We reconstructed the superoxide leakage by creating a net reaction composed of two: (1) normal reaction, in which water is produced as a by-product of redox metabolism (0.5 \( \text{o}_2 \text{e} + \text{pyoh}_2 \text{e} \to \text{h}_2\text{e} + \text{pyo}_2 \text{e} \)), and (2) leaky reaction in which superoxide is produced as a by-product (2.0 \( \text{o}_2 \text{e} + \text{pyoh}_2 \text{e} \to 2.0 \text{h}_2\text{e} + 2.0 \text{o}_2\text{e} + \text{pyo}_2 \text{e} \)). We estimated the ROS leakage stoichiometric coefficient (\( \kappa \)) by taking into account the changes in the amount of pyocyanin (which can lead to superoxide production) and hydrogen peroxide (a product of superoxide dismutation) produced after 24 hours of growth [24]. According to this method,

\[
\kappa = \frac{1}{\Delta \text{OD}_{\text{phenazine}} / \Delta \text{OD}_{\text{hydrogen peroxide}}}
\]

Then, we multiplied the stoichiometric coefficients of reaction (2) by \( \kappa \) and those of reaction (1) by 1-\( \kappa \) before adding up the reactions to generate a net reaction. The value of \( \kappa \) was estimated to be 0.29. Therefore, the net reaction used in the model is: 0.935 \( \text{o}_2 \text{e} + \text{pyoh}_2 \text{e} \to 0.58 \text{h}_2\text{e} + 0.71 \text{h}_2\text{e} + 0.58 \text{o}_2 \text{e} + \text{pyo}_2 \text{e} \).

**Addition of Unknown Reactions** For the addition of unknown reactions derived from the literature, we first checked for mass-and-charge balance. Then, we entered the reactions in the eQuilibrator program (v. 0.4.1) [32] in Python (v. 3.7.7). The parameters- pH = 7.5, ionic strength = 0.25 M, temperature = 25°C, control of magnesium ion (pMg) = 3 were used. Physiological concentrations (aqueous reactants at 1 mM) were assumed for the calculation of Gibbs free energy of transformation (\( \Delta G^m \)). Reactions with \( \Delta G^m < 0 \) were added to the model.

**Results**

**Model reconstruction and validation**

The reconstruction of iSD1509 was developed using the pipeline outlined (SI appendix, Fig. S1 and Extended Results). We compared the reconstructions of iSD1509 and iPau1129. The iSD1509 model contains considerably higher number of genes, reactions, and metabolites than iPau1129 (Fig. 1A). It possesses 424 more unique genes than in iPau1129. When we compared the top KEGG pathways between the two reconstructions, the number of genes in almost every pathway was higher in iSD1509 than in iPau1129 (Fig. 1B). The current MEMOTE (v. 0.12.0) score of iSD1509 is 88% (SI Appendix, Fig. S2).
Next, we validated iSD1509 by performing two tests: 1) substrate utilization in minimal media, and 2) gene essentiality. We used the same datasets utilized for validating iPau1129 [17].

The model can simulate growth on LB, SCFM (Synthetic Cystic Fibrosis Medium), and minimal media containing individual substrates (87 compounds). On the minimal medium, the model predicts substrate utilization with an accuracy of 94.3%. This is a significant improvement compared to the iPau1129 model (80.5%). The predicted growth rate on the glucose minimal medium (0.89 h⁻¹) is within the range of experimentally determined growth rates for \( P. \) aeruginosa [33]. Next, we compared the metabolic flux analysis (MFA) results with the predicted FBA values for growth on glucose minimal medium. Due to the unavailability of MFA data for PA14 and since both PA14 and PAO1 strains are highly similar [5], we used the MFA data for PAO1 [34]. FBA predictions and MFA results were highly correlated (Pearson correlation coefficient: 0.91, \( p \)-value<0.001) (SI Appendix, Fig. S3).

For gene essentiality validation, we performed predictions in LB medium. For the genes common between iSD1509 and iPau1129 (1,085 genes), iSD1509 retained a similar accuracy (iSD1509: 90.5% vs. iPau1129: 91.0%) by achieving a >6% higher recall (SI Appendix, Fig. S4A). An increase in 1% accuracy was achieved if all the genes in iSD1509 were considered (SI Appendix, Fig. S4B).

### Predictive assessment of iSD1509

Next, we assessed the model using new datasets. For the gene essentiality data, 321 core essential genes identified in a recent transposon insertion sequence study were used [35]. Since the core essential genes were identified in three different media conditions—LB, SCFM and glucose minimal (M9) medium, we computed gene essentiality on all three conditions. For LB, SCFM, and M9 minimal medium, overall accuracies were 92.4%, 92.3%, and 90.2%, respectively. Precision ranged between 51.2% (M9) and 64.0% (LB), and recall ranged from 57.2% (LB) to 57.9% (M9 and SFCM) (Fig. 1C). Our model also demonstrated higher prediction accuracy than iPau1129 in LB medium for the common core essential genes (SI Appendix, Fig. S4C).

We assessed substrate utilization using the data published in Dunphy et al. [36]. We simulated the model for growth on 131 minimal media containing different substrates. On eight substrates, the two experimental data sets ([17] vs. [36]) disagreed on \( P. \) aeruginosa’s ability to grow. Hence, we excluded these eight substrates when assessing our model. On 123 substrates, the model simulated growth with a high accuracy of 93.5% (Fig. 1D).

### Anaerobic growth

\( P. \) aeruginosa can utilize nitrate or nitrite as terminal acceptors for growth in the absence of oxygen. When we first simulated the model in anaerobic condition, it could not simulate growth because the two biomass constituents- ubiquinone-9 (UQ9) and thiamine diphosphate (THMPP) could not be produced.

For anaerobic UQ9 production, a recent study [37] demonstrated that UQ9 can be produced by an alternate pathway within the UQ9 biosynthesis chain. The biosynthetic proteins are shared between aerobic and anaerobic pathways except for the ones that catalyze the three reactions associated with hydroxylation of intermediate metabolites (Fig. 2A). In an anaerobic environment, an alternative oxidizing molecule (proposed to be prephenate) could be involved [38]. Therefore, we added prephenate-based hypothetical reactions as anaerobic alternatives to the aforementioned three hydroxylation reactions. These reactions are mass-and-charge balanced, and their G°'s
Figure 1. Genome-scale reconstruction of *Pseudomonas aeruginosa* PA14, iSD1509 and its predictive assessment. (A) When compared with iPau1129, the latest model iSD1509 has considerable increase in the reaction, gene, and metabolite content. (B) The top KEGG pathways between both reconstructions were compared, and iSD1509 evidently has higher gene content in almost every top KEGG pathway. (C) iSD1509 was then used for predicting core essential genes determined in a separate study [35]. The model prediction accuracy was found to be >90% in three different media conditions indicating a highly predictive model. (D) Using iSD1509, carbon substrate utilization was predicted for a new set of data [36]. The prediction accuracy was computed to be 93.5%.
were computed using eQuilibrator (v. 0.4.1) [32] to determine their feasibility. For anaerobic THMPP production, no alternative pathway could be identified in *P. aeruginosa* by either literature search or annotation-based methods. Hence, we removed this metabolite and adjusted the biomass reaction creating an anaerobic biomass reaction.

Following the changes, we could simulate anaerobic growth in LB medium. We then compared the growth rates between aerobic and anaerobic conditions by applying pFBA [29] and using an anaerobic biomass reaction as the objective function. We observed that the biomass production is lower under the anaerobic conditions than in the aerobic condition. (Fig. 2B). Studies have suggested that the energy yield of nitrate is lower than that of oxygen in *Pseudomonas* strains [39]. We also computed the UQ9 production in aerobic and anaerobic conditions. For this analysis, the reactions pertaining to ubiquinone-8 (UQ8) were turned off, and the flux-sum was computed for UQ9 which was then divided by the growth rate. The model predicted that the UQ9 production is slightly higher in the nitrate-supplemented medium than in the aerobic condition, which has been experimentally demonstrated (Fig. 2B [37]). Our model also showed that the UQ9 yield in nitrite-supplemented medium is considerably low (2.63 x 10^{-4} mmol gDW^{-1}), but the prediction has not been experimentally validated.

**Production of pyocyanin in *P. aeruginosa* can be simulated by extended *i*SD1509**

We next used *i*SD1509 to investigate phenazine production in *P. aeruginosa* PA14. *Pseudomonas* can produce and secrete phenazines (e.g., pyocyanin), which possess redox properties and can cycle in and out of the cell. Usually, they are reduced within the cell, then they go to the extracellular space to get oxidized (by donating electrons to acceptors such as oxygen), and finally get transported back in the cell to complete a redox cycle [23, 40]. Studies have demonstrated that phenazine production is stimulated by low oxygen tension [41]. Further, phenazines are involved in the survival of *P. aeruginosa* in biofilm environment in which oxygen is limiting [23, 40].

To simulate the production of pyocyanin, which can act as a cofactor [42], we applied limed-FBA on *i*SD1509 to account for the dilution of this cofactor [27]. Since the cofactor is not part of the biomass reaction in *i*SD1509, it does not dilute as the model simulates growth. Hence, the model cannot predict the synthesis of pyocyanin as it is regenerated as part of the cofactor cycle (SI Appendix, Fig. S5A). Using *i*SD1509 limed, the oxygen-dependent synthesis of pyocyanin could be simulated (SI Appendix, Fig. S5A).

We utilized *i*SD1509 limed to study the effect of oxygen availability and phenazine production on the growth rate of *P. aeruginosa*. For the simulation, the oxygen import flux was constrained over a range (0.5 mmol gDW^{-1} h^{-1} to 10 mmol gDW^{-1} h^{-1}). For each of those flux constraints, pyocyanin synthesis flux was also constrained over a range (0.000088 mmol gDW^{-1} h^{-1} to maximum value (0.00088 mmol gDW^{-1} h^{-1} computed for 0.5 mmol gDW^{-1} h^{-1} oxygen import flux)). For each value of oxygen import flux and pyocyanin biosynthesis flux, the growth rate was computed. With this analysis, we observed that for lower oxygen import flux (less oxygen available to the cell), the effect of pyocyanin production on the growth rate is pronounced. In contrast, when more oxygen is available to the cell, pyocyanin synthesis does not considerably contribute to the biomass production (Fig. 3 and SI Appendix, Fig. S5B).

The *i*SD1509 limed can mechanistically explain the production of pyocyanin at low oxygen availability. In such environments, *Pseudomonas* is forced to divert resources
Figure 2. Our model predicts anaerobic growth of *P. aeruginosa* along with higher production of ubiquinone in anaerobic condition than in aerobic one. (A) The proposed pathway for ubiquinone production in both aerobic and anaerobic conditions share same enzymes except for the hydroxylation reactions [37, 38]. (B) The model simulates growth rates (green bars) in media with different terminal electron acceptors including nitrate and nitrite. Likewise, the model predicts that the ubiquinone (UQ9) yield (blue bars) in nitrate is slightly higher compared to that in aerobic conditions which agrees with the experimental data [37]. In nitrite, the UQ9 yield is extremely low (2.63 x 10^{-4} mmol gDW^{-1}).

towards phenazine production to counteract the imbalance in the intracellular redox state caused by the reduced availability of oxygen, which has been demonstrated experimentally [23]. Therefore, phenazines are crucial to the survival of the pathogen at low oxygen availability (e.g., within biofilms). We can, hence, use iSD1509limed to identify potential intervention strategies that target phenazine redox cycle (SI Appendix, Fig. S5C).
Figure 3. The limed-FBA model can predict the biomass production dependence on pyocyanin biosynthesis and oxygen availability. Simulations were performed over a wide range of oxygen uptake (from extracellular to periplasm) and pyocyanin biosynthesis flux constraints by optimizing for biomass production. For oxygen-limited condition (lower oxygen import flux), the effect of pyocyanin production on the growth rate is more profound compared to that in more oxygen-rich condition (higher oxygen import flux).
Identification of potential mutants that overproduce gluconate

The rpoN mutant is a significant gluconate producer. The loss of function mutation in rpoN is common among the clinical isolates in CF patients [43, 44]. Behrends et al. proposed that gluconate production is positively (weak but significant) correlated to reduced antibiotic susceptibility. Further, the investigators demonstrated that only one indirect target of RpoN, 6-phosphogluconate dehydratase (6PGDH, PA14_29210) could replicate the gluconate overproduction phenotype of the rpoN mutant [31].

We used iSD1509 to recapitulate the results of the study [31] on the glucose minimal (M9) medium. Of the knockouts of the thirteen indirect targets of RpoN, the FBA simulations accurately predicted that only the deletion of 6PGDH (edd gene) leads to a significant increase in the gluconate production compared to the wildtype (Fig. 4A). The model also correctly showed that the growth of the edd mutant is considerably affected. The simulations provided insights into other mutants such as deletion of gluconate symporter gene (PA14_34630) leading to increased flux through glucose transport reaction (GLCabcpp) (Fig. 4A).

Next, we performed FVA simulations to determine whether gluconate production in edd mutant was a requirement for the optimal growth. The gluconate secretion (GLCNtex) reaction flux range (absolute value) in the mutant (4.72 mmol gDW\(^{-1}\) h\(^{-1}\)) is fixed at a higher rate than in the wildtype (0 mmol gDW\(^{-1}\) h\(^{-1}\)) suggesting that for the optimal growth of the mutant, gluconate is forced to be transported out of the cell.

Glyoxylate shunt decreases the TCA flux leading to lower respiration rate and lower proton motive force

Meylan et al. [45] showed that the addition of fumarate along with tobramycin leads to greater drug uptake and activity whereas glyoxylate protects the cells from tobramycin lethality. The study showed that the higher tobramycin uptake and activity in fumarate-containing medium is due to greater proton motive force (PMF) and increased respiration rate caused by higher flux through the TCA cycle, respectively. Likewise, in glyoxylate-containing medium, glyoxylate, by directly inhibiting α-ketoglutarate dehydrogenase, diverts the flux away from TCA cycle towards the glyoxylate shunt leading to reduced TCA cycle activity. We used iSD1509 to test an alternative, i.e., whether the law of optimal growth can explain drug protection by glyoxylate supplementation. Furthermore, we used iSD1509 to study the pathway utilization differences between the two metabolite supplementations.

We simulated iSD1509 on minimal (M9) medium containing glyoxylate or fumarate, and low amounts of citrate by optimizing for the biomass production. Since glyoxylate was not predicted to be a growth-inducing substrate in the substrate utilization assessment step, an artificial glyoxylate uptake reaction was added. We performed a flux sampling analysis by constraining the biomass flux to 90% of the estimated growth rate (by FBA) on both media. For proton flux calculations, we computed the flux-sum of periplasmic proton. Then, we compared the median fluxes of the reactions pertaining to TCA cycle and glyoxylate cycle between the two media conditions. The model predictions agreed with Meylan et al. [45] that the glyoxylate shunt indeed drives the flux away from TCA cycle in glyoxylate-containing medium. Likewise, the simulations also indicate that the glyoxylate flux is diverted towards the reactions catalyzed by malic enzymes, which recycle the necessary cofactors— NADH and NADPH. Unlike the Meylan study, the model predicted higher flux activity through pyruvate dehydrogenase reaction in the glyoxylate minimal medium. Moreover, the production of oxalate and glycolate were not confirmed by the model predictions. Instead, the glyoxylate flux was diverted towards the reaction catalyzed by glyoxylate carboligase. We also observed that the oxygen uptake rate and proton flux-sum were
Figure 4. Our model can accurately predict gluconate production in the mutants of the genes regulated by RpoN. Thirteen genes regulated by RpoN were in silico knocked out in this study in order to recapitulate the results of experimental study [31]. (A) FBA simulations were able to accurately predict that only edd mutant (PA14_22910) produces considerable amount of gluconate compared to the wildtype. Furthermore, the decrease in the growth rate of the mutant was also recapitulated in this study. For calculations, zero flux values were first replaced with 0.01. The fluxes of the reactions catalyzed by glucose dehydrogenase (GLCDpp and GLCDpp_q9) have been averaged (column GLCDpp_avg). For gluconate yield (mmol gluconate per mmol glucose, last column), gluconate secretion flux was divided by glucose exchange flux. Then, mutant reaction fluxes/yield were divided by respective wildtype reaction fluxes/yield. (B) To further characterize the edd mutant, FVA simulations were carried out in order to examine the flux range of desired reactions in both wildtype and mutant. Gluconate excretion is required by the mutant for optimal growth in the given condition. Furthermore, the glucose flux in the mutant is divided between glucose transport and glucose dehydrogenase reactions whereas in the wildtype, majority of the flux is channeled towards the Entner-Doudoroff pathway through glucose dehydrogenase reaction (SI appendix, Fig. S4). The values (after the reaction names) are derived by dividing the average reaction fluxes of the edd mutant by those of the wildtype. To avoid division by zeroes, all the zeroes were replaced by 0.01.
Figure 5. The iSD1509 model predicts that the glyoxylate shunt causes lower flux through the TCA cycle leading to lower oxygen uptake rate and PMF. Flux sampling analysis was performed in minimal media containing fumarate and glyoxylate to identify the flux distributions of TCA cycle reactions. According to the analysis, the TCA cycle is significantly upregulated in fumarate-supplemented medium compared to the glyoxylate one. The asterisk for reversible succinyl-CoA synthetase reaction is to note that the reaction flux is negative. In glyoxylate, the flux from acetyl-CoA is shunted towards the glyoxylate cycle as shown in the figure. The flux goes through the reactions catalyzed by malic enzymes to regenerate the cofactors—NADH and NADPH. This leads to a reduced TCA flux, decreased oxygen uptake rate, and PMF. All the sampling results (boxes with flux distributions where green: glyoxylate and blue: fumarate) are for reaction fluxes except for proton for which flux-sum yield was computed from the sampling data. Arrow colors indicate median fluxes being higher in one condition versus the other such that green: glyoxylate, blue: fumarate, black and bold: similar, and black and dotted: low flux in both conditions.

higher in the fumarate- than in the glyoxylate-supplemented medium, which led to increased drug uptake and activity in fumarate treatment (Fig. 5). Therefore, the law of optimal growth could also explain drug potentiation and drug protection by fumarate and glyoxylate supplementation, respectively.

Conclusion

Pseudomonas aeruginosa is an extensively studied organism in association with human infections especially in CF lungs. Since it is known to possess multiple mechanisms of drug resistance (e.g., lipopolysaccharide modification, overexpression of efflux pumps) and an ability to survive within the biofilm environment, eradication of Pseudomonas infections can be challenging. Even though P. aeruginosa PAO1, a laboratory strain, has been widely investigated, and a copious amount of data is available for the organism, such progress has not been made for other P. aeruginosa
organisms including the hypervirulent strain PA14. Using a predictive model such as iSD1509, which contains organism-specific knowledge, we can not only bridge the gaps but can also provide a platform to design experiments and strategies to combat infections caused by PA14 strain.

In this study, we have demonstrated that iSD1509 is a highly predictive and reliable model with accuracies of 90.2-93.4% and 93.5% for gene essentiality and substrate utilization predictions, respectively. Further, guided by the model simulations, we predicted that the biosynthesis of ubiquinone-9 is required for the anaerobic growth of PA14, and a recent study [37] corroborated this hypothesis. The model also predicted that the survival of P. aeruginosa in low oxygen environment (such as within biofilms) is possible due to phenazine production and provided mechanistic insight into their biosynthesis.

The model successfully identified the mutant that produced higher amounts of gluconate than the wildtype among thirteen candidates. Using iSD1509, more computational experiments can be designed to identify other mutants that overproduce gluconate. Likewise, other possible biomarkers of drug resistance can be predicted. Furthermore, using multi-strain modeling approaches [46], models of various P. aeruginosa isolates can be simulated to compare biomarker production simultaneously.

Finally, we used iSD1509 to recapitulate the potentiation of an antibiotic by metabolite supplementation [45]. The model correctly differentiated between metabolites that increased drug lethality versus those that did not and offered mechanistic explanations for these responses. Namely, supplementing glyoxylate diverted flux away from the TCA cycle, lowering the respiration rate that consequently led to lower drug activity. Supplementing the medium with fumarate caused higher oxygen uptake rate due to higher TCA cycle activity which leads to higher drug activity [45]. Likewise, PMF (and ultimately drug uptake) was higher in the fumarate-containing medium than in the glyoxylate-containing medium. These results suggest that our model can be used to design antimicrobial strategies based on metabolic mechanisms, including metabolite supplementation, against P. aeruginosa.

In conclusion, this model can be seen as a computational platform to design experiments targeting P. aeruginosa metabolism at various growth states including within the biofilm environment. Likewise, novel computational targets of P. aeruginosa (i.e., pyocyanin redox cycle) can now be investigated to identify the best strategy to inhibit the growth of the pathogen. Overall, we expect the model to significantly accelerate our understanding of P. aeruginosa to combat the associated infections.

Supporting Information Appendix (SI)

The materials and methods are detailed in SI Appendix, including reconstruction of iSD1509, simulations using the model, which were performed in COBRApy package (v. 0.18.1) [28].

Data availability

The model and sample codes are available at https://github.com/dahalsanzeev/iSD1509-M. All other study data are included in the article and/or supporting information.
Acknowledgments

This work was supported by Queen’s University and the Natural Sciences and Engineering Research Council of Canada (NSERC) [RGPIN-2020-06325]. AR and AD received support from the German Center for Infection Research (grant number 8020708703) and the Cluster of Excellence CMFI (Controlling Microbes to Fight Infections), project number EXC-2124/05.037_0, funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) under Germany’s Excellence Strategy – EXC 20124 – 390838134.
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Extended Results

Initial reconstruction and manual refinement

We used CarveMe [47] to create a draft reconstruction that contained 2196 reactions and 1482 genes. Following this, we added the biomass reaction from the previous SEED model (iPau1129) [17]. We gap-filled our model in LB rich medium to produce all the biomass constituents by adding the reactions from iPau1129 or from other BiGG models.

Next, we curated our model through a semi-automated approach. We chose seventeen different BiGG models to perform a bidirectional blast hit for each of the reactions present in the model (Table S2). We then manually checked the reactions for any discrepancy in the gene-protein-reaction (GPR) associations made by CarveMe and by our approach. Any discrepancy was resolved using the knowledge from other databases (e.g., IMG, KEGG). Reactions removed from the model are provided (Dataset S1).

Using carbon substrate essentiality [17] and gene essentiality data [48], we iteratively filled more knowledge gaps either through annotation- or literature-derived information. A list of reactions that were added to the model is provided (Dataset S2). We also utilized gene essentiality data to fix GPR associations. Likewise, biomass reaction was also modified during this process to better reflect the gene essentiality data.

We next added any extra genes in iPau1129 not present in our model. This process led to the addition of reactions related to alginate production, rhamnolipid production, pyochelin production, etc. Only 44 genes from iPau1129 are missing in iSD1509 (Dataset S3). Finally, we checked and corrected any mass-and-charge imbalance, first by applying the knowledge from well-curated BiGG models (iJN1462 [49] and iML1515 [50]), which lead to only 129 metabolites that required additional manual curation. The formula and charges of those metabolites were changed either using information from metabolite databases (BiGG, MetaNetX, PubChem or ModelSeed) or manually by mass-and-charge balancing the reactions (Dataset S4).

Literature derived curation and addition of reactions

We curated the knowledge derived from literature to add to the reconstruction to make the model more strain-specific. The reactions pertaining to multiple terminal oxidases, reactive oxygen species elimination, n-alkane degradation, phenazine-associated metabolism, oxalate production, rubredoxin-based metabolism and H+-translocating NADH:quinone oxidoreductases were added. This led to the development of a model possessing 1509 genes. The model (referred to as iSD1509) contains 2023 total reactions and 1151 unique metabolites and possesses three compartments.

We analyzed the additional genes in iSD1509 by enrichment in both KEGG and COG categories (Fig. S6). In KEGG, notable enriched pathways were oxidative phosphorylation, sulfur metabolism, folate biosynthesis, glyoxylate and dicarboxylate metabolism, and biofilm formation. In COG, the top three categories were energy production and conversion (C), amino acid transport and metabolism (E), and inorganic ion transport and metabolism (P). We also examined the top 20 KEGG pathways of the new metabolic reconstruction (Fig. S6C). The greatest number of genes in iSD1509 belong to the category “Metabolic pathways.” Interestingly, other significant pathways are associated with biosynthesis of secondary metabolites, metabolism in diverse environments, biosynthesis of antibiotics, indicating a reconstruction of a metabolically versatile organism (Fig. S6C).
Extended Materials and Methods

Initial Reconstruction

For the initial reconstruction, CarveMe\textsuperscript{47} with gap-filling function in LB medium was utilized. From the previous \textit{P. aeruginosa} model (iPau1129)\textsuperscript{17}, we added the biomass reaction along with other required reactions to simulate the growth of the model on LB medium. For this process, we first converted the ModelSEED\textsuperscript{51} metabolite identifiers of iPau1129 to BiGG identifiers. Furthermore, the bounds of non-growth associated reaction were also added from iPau1129. Then, we removed all the artificial sink and demand reactions added by CarveMe by making sure that the respective metabolites can be produced or consumed by added reactions which are supported by gene evidence.

Then, the \textit{Pseudomonas aeruginosa} PA14 reactome was inspected for the gene-protein-reaction (GPR) associations using a custom pipeline (Fig. S1). First, seventeen models from BiGG database and their respective protein FASTA files were downloaded (Table S2). Then, we performed a bidirectional blast hit (BBH) for all the reactions present in the model by prioritizing the strains that are taxonomically closer to PA14 strain. We applied a stringent method such that only the top hits in both directions were considered as the correct gene. If not, we manually inspected the top blast hits in different databases including IMG\textsuperscript{52} and KEGG\textsuperscript{53}. These GPR associations were then used as alternatives to CarveMe predictions. We performed a rigorous manual check for the reactions whose GPR associations were derived from the BBH approach from models other than those from \textit{Pseudomonas putida}. For any discrepancy between GPR associations from CarveMe and those from BBH approach, we checked IMG and/or KEGG and/or iPau1129 model to assign correct gene associations. Any reactions that did not have associated GPRs or with no evidence to be present in PA14 were discarded in this process. We also removed any unnecessary loops during this process. The list of deleted reactions is provided (Dataset S1). We iteratively improved and validated the model using substrate utilization\textsuperscript{17} and gene essentiality\textsuperscript{48} data which led to the modification of the biomass reaction and more changes to GPR associations. We also added any genes from iPau1129 that were not associated with the model at that time.

Reaction Mass and Charge Balance

For mass-and-charge balance, metabolite formula and charge were assigned using iJN1462 first, and then iML1515. Next, we manually checked the remaining metabolites on multiple databases including BiGG, MetaNetX, PubChem and ModelSEED to identify the correct formula and charge. Finally, if the information for metabolites could not be found in the aforementioned databases, we assigned the formula and charge by balancing the reactions that contain only those metabolites as the sole undetermined ones.

Manual Reconstruction

Reactions related to anaerobic metabolism, phenazine-associated metabolism, terminal oxidases and alternative terminal oxidases, thiamine metabolism, nucleotide metabolism, \textit{n-alkane} metabolism, oxalate production, anaerobic quinone production, rubredoxin-based metabolism, reactive oxygen species (ROS), and H\textsuperscript{+}-translocating NADH:quinone oxidoreductases were added using annotation in KEGG and/or extensive knowledge from literature. All the added reactions are listed (Dataset S2). This reconstruction effort led to the development of the model, iSD1509.
Substrate Utilization Screening and Gene Knockout Study

For the initial carbon source catabolic activity and gene essentiality validation, the same datasets used in Bartell et al. [17] were utilized. For substrate utilization, we simulated iSD1509 in minimal medium (Table S1) containing individual substrates and optimized for maximum biomass production. Since ModelSEED identifiers were used in Bartell et al., some of the substrate identifiers could not be converted to the BiGGG ones, and hence were removed from this analysis. Overall, 87 compounds were compared. The media composition was provided generously by Papin lab. For gene essentiality comparison, we simulated the model in LB rich medium.

For substrate utilization screening and gene essentiality assessment of iSD1509, data collected from separate experimental studies were applied. For carbon source assay, the dataset by Dunphy et al. [36] containing 190 carbon sources was collected. In the collected dataset, since only general metabolite names were provided, the metabolites whose BiGGG identifiers could be determined with high confidence were utilized for prediction. Furthermore, since the two experimental datasets ([17] and [36]) showed discrepancy in P. aeruginosa’s ability to grow on eight substrates, those compounds were removed from the analysis. Hence, the model was simulated on minimal medium containing 123 individual substrates for comparison. For assessing the gene essentiality, we used the dataset from a recent study by Poulsen et al. [35]. In this dataset, core essential genes were defined as essential genes in five media conditions and nine different strains. We analyzed these core essential genes in LB medium, SCFM (Synthetic Cystic Fibrosis Medium) and glucose minimal medium. For gene essentiality comparison in glucose minimal medium, iron had to be added even though the media used in the study presumably did not contain iron.

The metrics for the comparison of model predictions and experimental data are defined as follows:

- **Precision** = \( \frac{TP}{TP + FP} \)
- **Negative Predictive Accuracy** = \( \frac{TN}{TN + FN} \)
- **Recall** = \( \frac{TP}{EP} \)
- **Specificity** = \( \frac{TN}{EN} \)
- **Accuracy** = \( \frac{TP + TN}{EP + EN} \)


**Flux-sum Analysis**

For the metabolites of interest, we performed flux-sum analysis [54]. Briefly, flux-sum (\( \Phi_i \)) for metabolite i can be computed using the following formula,

\[ \Phi_i = \frac{1}{2} \sum_j |S_{ij}v_j| \]

where j is the reaction in which the metabolite i participates in.

For the computation of yield, the flux-sum was either divided by the growth rate (for quinone production) or by the substrate flux multiplied by the number of carbon atoms in the substrate (for proton motive force (PMF)).

**Flux Sampling**

We applied flux sampling approach [55] in the COBRApy package [28] by using optGpSampler [56] with 10000 samplings in both fumarate and glyoxylate minimal
media. The growth rate was constrained to 90% of the rate predicted in FBA simulations in the respective media condition. By using the validate method of the optGpSampler, only valid samples were kept for further analysis. Furthermore, we used the autocorrelation plots and trace plotting for convergence analysis. To compare the distributions of reaction fluxes or metabolite flux-sums between the two different media conditions, we performed rank-sum tests.

**limed-FBA**

For the simulation of dilution of cofactors, we used limed-FBA approach [27]. Briefly, all the reactions were first made irreversible. Then, for the reactions consuming or producing the desired cytosolic cofactor, a small dilution constant ($\epsilon$) was applied such that

$$(S - \epsilon S_{\text{binary}}) = 0$$

where $S_{ij}^{\text{binary}} = 1$ if $S_{ij} \neq 0$.

and $\epsilon_{ii}$ for metabolite $i$ was computed using the following formula,

$$\epsilon_{ii} = \frac{0.1}{1000 \cdot \sum_j S_{ij}^{\text{binary}}}$$

To not double-dilute biomass related metabolites, their $\epsilon_{ii}$ was made zero. Likewise, for simple transport reactions in which no chemical transformations (e.g., glucose(extracellular) $\rightarrow$ glucose (periplasm)) occur, their binary coefficients were assigned zeros to avoid any high-flux loops.

**Visualization of flux data**

We first created a preliminary Escher [57] map in the Python-based framework, and then exported the map to web-based Escher tool for appropriate changes. Default settings were applied except for the customization of color and size of the edges for better visual representation. Then, we modified the Escher maps using vector graphic tool (Affinity Designer, v. 1.9.3).

We produced the heatmaps using seaborn package (v. 0.10.1) in Python. All the respective flux or yield values were divided by their respective wild-type values. To avoid division by zeroes, all the zero fluxes were first replaced by 0.01. For color range, robust quantile computation function was applied.
Figure S1. The working pipeline for building the genome-scale reconstruction of \textit{P. aeruginosa} PA14. Both automated and semi-automated methods were applied in this pipeline, and the reconstruction was gap-filled using iPau1129 model wherever necessary. Furthermore, strain-specific reactions were added after extensive literature curation. At each stage, the model was validated and improved using gene essentiality and substrate utilization data from [17].
Figure S2. Memote scores for iSD1509. Overall, the score for the model is 88%.
Figure S3. FBA simulations agree with MFA experimental data [34]. In the figure, the width of arrows represent the percentage of glucose flux. Only absolute values are shown here. All the zero value arrows are made grey for clarity. The MFA average values are written in the blue font if their respective reactions were found in the reconstruction. Lumped reactions are represented by multiple reactions separated by commas. Since two possible reactions (ME1 and ME2) are present in the M-model for the reactions catalyzed by the malic enzyme, both are shown separated by "/". The box inset is a correlation plot that corroborates the fact that the majority of the predictions agree with the MFA results (correlation coefficient of 0.91 (p<0.001)). Grey bands denote 95% confidence interval. Within the plot, glucose import (from extracellular to periplasm) has been indicated for reference.
Figure S4. Comparison of gene essentiality between iSD1509 and iPau1129. (A) Comparing common genes between the two models for dataset retrieved Liberati et al. [48]. (B) Computation of gene essentiality for all the genes present in iSD1509 using the same dataset [48]. (C) Comparing common core essential genes (from Poulsen et al. [35]) between the two models.
Figure S5. Using iSD1509limed to demonstrate that the phenazine production is crucial for the pathogen survival. (A) Demonstration of the application of limed-FBA that predicted phenazine production is higher at lower oxygen availability condition. Normal FBA (blue) cannot predict phenazine biosynthesis as a function of oxygen availability. (B) In oxygen-limited condition (O2tex upper bound: 0.5 mmol gDW-1 hr-1), the biosynthesis of phenazine can affect the growth of the pathogen considerably. (C) A simple formulation of pyocyanin removal (i.e., a demand reaction) was added to the model, and the simulations were performed in oxygen-limited condition (O2tex upper bound: 0.5 mmol gDW-1 hr-1). As the pyocyanin removal flux increased, the model predicted that the biomass production of *P. aeruginosa* also lowered. Please note that this simulation (5C) can also be performed using the FBA model.
Figure S6. Pathway enrichment of the genes present in iSD1509. (A) The top 20 KEGG pathways of unique genes in iSD1509. (B) The top 20 COG categories of unique genes in iSD1509. (C) The top 20 KEGG pathways of all the genes present in iSD1509.
Table S1. Minimal media composition for various experiments. For model assessment and gluconate prediction study, glucose was used as the carbon substrate. For the drug potentiation/protection investigation, either fumarate or glyoxlate were used as carbon substrates.

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<th>Priority</th>
<th>Strain Name</th>
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