Expanding the results of a high throughput screen against an isochorismate-pyruvate lyase to enzymes of a similar scaffold or mechanism

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Abstract

Antibiotic resistance is a growing health concern, and new avenues of antimicrobial drug design are being actively sought. One suggested pathway to be targeted for inhibitor design is that of iron scavenging through siderophores. Here we present a high throughput screen to the isochorismate-pyruvate lyase of \textit{Pseudomonas aeruginosa}, an enzyme required for the production of the siderophore pyochelin. Compounds identified in the screen are high nanomolar to low micromolar inhibitors of the enzyme and produce growth inhibition in PAO1 \textit{P. aeruginosa} in the millimolar range under iron-limiting conditions. The identified compounds were also tested for enzymatic inhibition of \textit{E. coli} chorismate mutase, a protein of similar fold and similar chemistry, and of \textit{Y. enterocolitica} salicylate synthase, a protein of differing fold but catalyzing the same lyase reaction. In both cases, subsets of the inhibitors from the screen were found to be inhibitory to enzymatic activity (mutase or synthase) in the micromolar range and capable of growth inhibition in their respective organisms (\textit{E. coli} or \textit{Y. enterocolitica}).

Keywords
siderophore; isochorismate pyruvate lyase; chorismate mutase; salicylate synthase
1. Introduction

Most pathogens require iron for survival because this metal serves as a required cofactor in many essential biological processes including DNA biosynthesis and cellular respiration [1]. While direct mechanisms for iron uptake (for example, uptake of heme) have evolved, indirect methods of iron acquisition, such as the use of high affinity iron chelators called siderophores, allow for iron scavenging regardless of the source [1]. Siderophore biosynthesis has been linked to virulence in many pathogenic bacteria [2–10]: without the ability to scavenge iron from the host, the bacteria are incapable of causing disease. Salicylate-capped siderophores are produced by several pathogenic bacteria, such as Yersinia spp. including the causative agent of plague, Mycobacterium tuberculosis, which is the causative agent of tuberculosis, and Pseudomonas aeruginosa, a nosocomial pathogen infecting susceptible hosts, including burn victims, cancer patients with hematological malignancies and those undergoing chemotherapy, patients with immune deficiencies such as AIDS patients, and Cystic Fibrosis (CF) patients [11]. Compounds that prevent the formation of salicylate or its incorporation into siderophores may prove to be a new class of antimicrobial drugs in this age of antibiotic drug resistance.

P. aeruginosa, Yersinia spp., and M. tuberculosis make chemically related siderophores that all use a salicylate cap (Figure 1A), the hydroxyl group of which is involved in iron chelation [12]. The salicylate 2 is formed in two steps from chorismate 4 with an isochorismate 1 intermediate (Figure 1B). The first step is the general acid-general base isomerization of chorismate to form isochorismate [13–15], whereas the second is the pericyclic migration of the enolpyruvyl tail [15, 16]. Because the cyclic transition state for this reaction includes a hydrogen, the result is the elimination of the tail and the formation of salicylate and pyruvate 3. For Yersinia spp. (siderophore: yersiniabactin) and M. tuberculosis (mycobactin), both reactions are catalyzed by a salicylate synthase, Irp9 and MbtI, respectively. The generation of the siderophore pyochelin by P. aeruginosa requires two enzymes, an isochorismate synthase (PchA) and an isochorismate-pyruvate lyase (PchB). Interestingly, PchA, Irp9 and MbtI are all structural homologues in the MST (menaquinone, siderophore and tryptophan biosynthesis) family, but as yet it is unknown why PchA cannot perform the pericyclic lyase reaction [17]. There are homologues of PchA that are isochorismate synthases, which also cannot perform the pericyclic lyase reaction. E. coli MenF [18, 19] is involved in menaquinone biosynthesis whereas EntC [20–23] from E. coli and VibC [24, 25] from Vibrio cholera are found in the biosynthetic pathways for dihydroxybenzoate capped siderophores. For MenF, EntC and VibC, the inability to perform the lyase reaction is biologically logical, since isochorismate is required for the formation of the biosynthetic products whereas salicylate is not [17].

The current work was initiated as a high throughput screen for the isochorismate-pyruvate lyase from P. aeruginosa (PchB), which performs the second, pericyclic reaction in the generation of the pyochelin siderophore. PchB is easily generated, we have adopted an effective microbial production system for obtaining the substrate isochorismate [26], and the salicylate product is fluorescent, allowing for the development of an enzymatic assay suitable for automation. PchB is a structural homologue of E. coli chorismate mutase (EcCM) of the AroQ structural class (Figure 2) [27, 28]. EcCM is the N-terminal domain of...
the well-studied bifunctional P-protein [29], the terminal enzyme of the shikimate biosynthetic pathway that is the branch point for the production of aromatic amino acids [30]. Both PchB and EcCM perform pericyclic reactions with similar transition states, differing in the alignment of the enolpyruvyl tail over the ring and thus the atomic composition of the cyclic transition state (Figure 2) [17]. The shikimate pathway is also recognized as an attractive target for antimicrobial drug design, since mammals do not synthesize aromatic amino acids. Rational design of inhibitors has been conducted for the chorismate mutases [31–38], including the TSA (transition state analogue), an oxabicyclic acid generated by Bartlett [34, 35]. EcCM is refractory to screening, because it lacks a good spectrophotometric handle and is difficult to make in sufficient quantities.

While pyochelin has been implicated in virulence [4, 10], any compounds that are effective inhibitors against PchB would need to be used in combination with inhibitors to the other iron scavenging pathways in *P. aeruginosa*, including that of the high affinity siderophore pyoverdin. Salicylate synthases (for example, Irp9 and MbtI mentioned above) perform the same pericyclic reaction as PchB (Figure 2), but are found in siderophore biosynthetic pathways necessary for virulence by organisms that are more limited in variety of iron uptake systems (most particularly, in *Yersinia spp*. [2, 3, 6] and *M. tuberculosis* [5, 7–9]). While salicylate synthases perform the same reaction, they do so using a different protein scaffold (Figure 2). Salicylate synthesis and activation of salicylate by adenylation for incorporation into siderophores have been the targets of inhibition studies to produce lead compounds for antimicrobial drug design. Rational design of inhibitors to enzymes such as isochorismate synthase, salicylate synthase and salicylate adenylase has been documented [39–59], as well as high throughput screening for inhibitors of salicylate synthase [60].

We hypothesized that compounds identified as inhibitors in an enzymatic screen may also be effective against an enzyme that performs the same chemistry albeit using a differing protein fold. By the same token, an inhibitor identified by screening methods may also be effective against a protein with a homologous protein scaffold and similar transition states, in this case, allowing for a way to identify inhibitors to an enzyme that is not suitable for screening. Therefore, we report a high-throughput screen designed to identify inhibitors of the isochorismate-pyruvate lyase from *P. aeruginosa*, measuring decreased salicylate production from isochorismate in the presence of an inhibitory compound. The compounds were further tested for inhibition of salicylate production from chorismate by the *Y. enterocolitica* salicylate synthase Irp9 which performs the same pericyclic lyase reaction but using a different protein fold. An orthogonal assay was used to measure inhibition of pyruvate production for both PchB and Irp9. Due to similarities in protein fold, substrate and transition state, identified compounds were also tested for inhibition of *E. coli* chorismate mutase activity, measuring chorismate disappearance. Finally, growth inhibition was measured for *P. aeruginosa*, *E. coli* and *Y. enterocolitica* for two of the most promising compounds.
2. Results

2.1 High Throughput Screen

Our assay measures the conversion of isochorismate to salicylate and pyruvate by the isochorismate-pyruvate lyase, PchB. The assay takes advantage of the fluorescent properties of salicylate, and is amenable to standard lab and high throughput conditions. Using the lab conditions and concentrations derived from steady state kinetics \[27\] as a starting point, a high-throughput assay was developed. Optimization of the assay conditions were tested by varying enzyme and substrate concentration and incubation times. The assay was insensitive to DMSO up to concentrations of 2%.

The assay was automated at the University of Kansas High Throughput Screening Laboratory and used to screen PchB versus the ChemDiv, ChemBridge, Microsource, Prestwick and the University of Kansas Chemical Methodologies and Library Development libraries. A pilot screen identified compound 6 as a positive control. Wells in each plate were reserved for no enzyme negative controls and 10 µM compound 6 positive controls to allow for the calculation of plate-to-plate statistics and to detect irregularities in the screen. Each compound was screened at a concentration of 10 µM. Substrate and enzyme dilutions were prepared fresh each day, with 30 plates run per day. Many compounds (258) were auto-fluorescent (showed > 100% inhibition) and were eliminated from further analysis (Supplemental Figure 1A). Compounds that showed inhibition effects greater than plate median plus three standard deviations but less than/equal to 100% (422 compounds) were deemed “actives” – a hit rate of 0.4%. Z’ factors on individual plates ranged from 0.5 to 0.9 and was 0.8 overall (Supplemental Figure 1B). The signal to baseline ranged from 10 to 30. Active compounds (231 of the 422) were confirmed by rescreening from the source plates in the high-throughput assay. Based on score (IC\(_{50}\) values from dose response data), availability and price, five compounds from three structural classes (9–13) were purchased for further evaluation (Figure 3). Compounds 6–8 were no longer commercially available and so were synthesized as described in the Methods.

2.2 Inhibition of isochorismate-pyruvate lyase (PchB)

The eight selected hits from the high-throughput screen were more rigorously examined with the same conditions that were used to obtain the original kinetic constants \[27\]. Half maximal inhibitory concentration (IC\(_{50}\)) values for the prospective inhibitors were calculated from dose response curves based on percent inhibition of PchB (Table 1). All compounds were effective with IC\(_{50}\) values ranging from 100 nM to 10 µM. \(K_I\) values were determined for compounds 6, 9 and 10: 44 +/- 2 nM, 80 +/- 8 nM, and 350 +/- 40 nM, respectively, and all three were competitive inhibitors. An orthogonal assay was adopted to measure pyruvate production (as opposed to salicylate production) using an absorbance based assay. The calculated IC\(_{50}\) values for this assay were comparable to those from the original salicylate fluorescence assay.

2.3 Inhibition of salicylate synthase (Irp9)

Since Irp9 performs the same chemistry as PchB, albeit by an enzyme of a different structural class, the same assays (salicylate production and pyruvate production) were used...
to determine IC₅₀ values for the eight selected hits from the high-throughput screen. The compounds of the first structural class (6–8) were not inhibitory for Irp9 in either assay (Table 1). The remaining compounds showed IC₅₀ values ranging from 25 to 160 µM in the salicylate production assay, but these values were only recapitulated in the pyruvate production assay for compounds 10 and 12. A Kᵢ value was determined for compound 10: 0.8 +/− 0.2 µM, a competitive inhibitor.

2.4 Inhibition of chorismate mutase (EcCM)

EcCM and PchB are of the same structural class and both perform pericyclic reactions on similar substrates (chorismate and isochorismate). Therefore, the eight selected hits from the high-throughput screen were also tested for inhibition of EcCM using a chorismate disappearance absorbance assay. Compounds 6 through 13 gave IC₅₀ values between 30 and 200 µM (Table 1). Kᵢ values were determined for compounds 6 and 10: 30 +/− 10 µM, and 40 +/− 20 µM, respectively, and both were competitive inhibitors.

2.5 Bacterial Growth Inhibition

To determine if the compounds were effective at inhibiting bacterial growth, half maximal effective concentration (EC₅₀) values were determined for P. aeruginosa, Y. enterocolitica, and E. coli. The HTS and subsequent screens provided a minimal structure-activity relationship for the two compound classes represented by compounds 6 and 10. Therefore, these two compounds were chosen for EC₅₀ determination. The EC₅₀ values were determined in iron-rich (200 µM) or iron-poor (5 µM) media for both P. aeruginosa and Y. enterocolitica to determine if inhibition is related to iron uptake. The EC₅₀ values were not determined in iron-free media, because the bacteria could not be reliably grown in the absence of iron even without potential inhibitors. For P. aeruginosa, EC₅₀ values were determined for the wildtype PAO1 strain, for a pyoverdin-minus PAO1 strain, and for a pyochelin-minus PAO1 strain. The pyoverdin-minus strain is a ΔpvdA knockout strain, deficient in the ornithine hydroxylase enzyme that is the first committed step for production of the high-affinity pyoverdin siderophore. The pyochelin-minus strain is a ΔpchE knockout strain, deficient in the nonribosomal peptide synthetase required for salicylate incorporation. These data are normalized relative to bacteria grown in the absence of inhibitor and plotted as relative growth. The highest concentration of the compounds tested in these assays was 5 mM, limited by solubility and vehicle (DMSO) toxicity.

Compounds 6 and 10 are sub-micromolar inhibitors of the isochorismate-pyruvate lyase required for pyochelin production. These compounds did not cause growth inhibition in iron-rich media, regardless of P. aeruginosa, strain (Figure 4A and 4B, Table 2). This was anticipated, since the bacteria were not reliant on siderophore production for iron uptake when the media is replete with iron. Under iron-limiting conditions, if the compounds were specific for inhibition of PchB, we would expect the strongest inhibition in pyoverdin-minus strain, as this strain is more reliant on the pyochelin pathway for iron-uptake. The wildtype PAO1 strain could show some growth inhibition or this could be overcome by the functional and more efficient pyoverdin pathway. The pyochelin-deficient strain should show no inhibition in relative growth, since the pyochelin production pathway was genetically disrupted. Both the wildtype PAO1 strain and the pyoverdin-minus strain showed growth
inhibition for both compounds at concentrations that were >5 mM. The pyochelin-minus strain was not inhibited by compound 6 but was inhibited at > 5mM by compound 10. These data suggest that compound 6 maybe more specific for the pyochelin pathway than compound 10. However, these data suggest that under iron-limiting conditions there may be a second target that is also inhibited by compound 10. *P. aeruginosa* has chorismate mutases that may be inhibited. Additionally, *P. aeruginosa* is highly antibiotic resistant due in part, to the presence of efflux pumps for the export of xenobiotics. Growth inhibition would likely be augmented in the presence of an efflux pump inhibitor, such as phenylalainine-arginine-β-naphthylamide [61], a future direction of this work.

Compound 6 was not an enzymatic inhibitor of the *Y. enterocolitica* salicylate synthase Irp9, but compound 10 was a micromolar inhibitor. If our hypothesis was correct and compound 10 was specific for yersiniabactin biosynthesis, then neither should inhibit *Y. enterocolitica* bacterial growth under iron rich conditions, and only compound 10 should inhibit under iron poor conditions. If the inhibitors are non-selective, then both may inhibit bacterial growth under both iron rich and iron poor conditions. If the inhibitors work against both the yersiniabactin biosynthesis and another enzymatic process, then compound 10 should be more effective under iron poor conditions. Based on the data in Figure 4C and Table 2, both compounds give EC50 concentrations in the millimolar range under both iron rich and iron poor conditions (indeed, the differing iron conditions were within error of each other), suggesting that compound 10 is not selective for siderophore biosynthesis and that both inhibitors are causing growth inhibition by some other mechanism.

Compounds 6 and 10 are low micromolar inhibitors of *E. coli* chorismate mutase. In growth inhibition assays against Seattle strain *E. coli*, compound 6 arguably showed no growth inhibition, while compound 10 gave an EC50 value of 3 mM (Figure 4D).

3. Discussion

Two inhibitors of PchB are reported in the literature: Bartlett’s TSA (*Ki* ~2 µM) and adamantane-1-phosphonate (*Ki* >5 mM) [62]. The work reported here identifies three compounds with *Ki* values in the nanomolar range: 6 (*Ki* 44 +/- 2 nM), 9 (*Ki* = 80 +/- 8 nM) and 10 (*Ki* = 350 +/- 40 nM). In addition, compounds 6 and 10 show growth inhibition of *P. aeruginosa* at millimolar concentrations under iron-limited conditions. Therefore, these compounds represent the best known inhibitors of PchB, and may serve as initial lead compounds for the development of new antibiotics.

To our knowledge, the best known inhibitor for *E. coli* chorismate mutase is Bartlett’s TSA. The literature reports inhibition of this and similar compounds as *I50/Km*1, with a value of 0.008 for the TSA [34, 35]. Similar calculations were performed for compounds 6 and 10 versus EcCM, and values of 0.05 and 0.053 were determined, better than all of the reported compounds save the TSA. These compounds therefore represent new scaffolds suitable for

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1Bartlett and colleagues reported inhibition of chorismate mutase/prephenate dehydrogenase by their transition state analogues as *I50/Km* values [34, 35], where they defined *I50* as “the concentration of inhibitor giving 50% inhibition when the substrate concentration equals *Km*” [34]. For the sake of direct comparison, we likewise make these calculations.
optimization as chemical probes and inhibitors of chorismate mutases. Furthermore, compound 10 also was effective at growth inhibition of *E. coli* in the millimolar range.

Rational design of inhibitors against isochorismate and salicylate synthases and similar chorismate-utilizing enzymes have primarily targeted varying the pyruvylenol tail of the substrate and have produced inhibitors with low micromolar *K*ᵢ values [47, 51, 52, 63] Interestingly, in two cases, PchB was used to convert isochorismate generated by an isochorismate synthase into salicylate for detection but there is no mention as to whether the inhibitors were tested for inhibition of PchB alone [51, 52]. A high-throughput screen against the salicylate synthase from *M. tuberculosis* (MbtI) identified two structural classes of inhibitors, which showed *IC*₅₀ values in the low micromolar range [60]. The first was a diarylsulfone reminiscent of compound 9; however, compounds of this scaffold have been identified as pan assay interference compounds [64] and are therefore unlikely for further development. The second was a benzimidazole-2-thione similar to compounds 10–12. We likewise determined a micromolar *IC*₅₀ values for compound 10 with the *Y. enterocolitica* salicylate synthase Irp9, which shows growth inhibition of *Y. enterocolitica* in the low millimolar range. These fused heterocyclic ring systems may be an excellent starting point for inhibition in the MST enzyme class.

We hypothesized that an inhibitor identified through a high throughput screen may be effective against enzymes of similar scaffold or the same catalytic function. The screen against the isochorismate-pyruvate lyase identified competitive inhibitors that were found to likewise be competitive inhibitors against chorismate mutase (similar fold and active site structure) and salicylate synthase (same catalytic mechanism). While one may speculate that the ability to “target hop” may be only possible with competitive inhibitors, exploiting similarity of the active site and/or transition state, it is also possible to envision a scenario where an allosteric regulator binding site may also be similarly employed. The bacterial inhibition studies suggest that the compounds may be multi-selective. In other words, the compounds inhibit proteins of similar structure and/or function *in vitro*, and may do so *in vivo* as well. All of the bacterial cultures grown generate chemically-related siderophores with salicylate or dihydroxybenzoate caps utilizing enzymes that are structurally or functionally related the PchB and/or Irp9. These bacterial species also produce aromatic amino acids, thereby utilizing a chorismate mutase of the same structural family as EcCM. However, these studies do not rule out the possibility that these inhibitors are non-selective, inhibiting a target dissimilar and completely independent to those tested. Identification of the cellular target(s) is the next important step in this experimentation.

4. Conclusions

The HTS screen presented here identified inhibitors of the isochorismate-pyruvate lyase from *P. aeruginosa*. Lead compounds were scrutinized in an orthogonal assay and for bacterial growth inhibition. The inhibitors were similarly tested against an enzyme that performs identical chemistry in a different protein scaffold (salicylate synthase) and against an enzyme with a homologous protein scaffold that performs similar chemistry (chorismate mutase). Inhibitors were identified that are better than those previously reported in the literature, and also cause growth inhibition of *P. aeruginosa* in a seemingly iron-dependent
way. A subset of the identified inhibitors were also effective against enzymes of similar scaffold (EcCM) or identical chemistry (Irp9). The result is a series of compounds are among the most effective inhibitors to date for all three enzymes and may prove useful in the discovery of future antibiotics.

5. Methods

5.1 Preparation of substrates

Isochorismate was isolated from *Klebsiella pneumoniae* 62-1 harboring the entC plasmid pKS3-02 [26] with only minor changes, as described previously [27]. Chorismate was isolated from *Klebsiella pneumonia* 62-1 [65] with only minor changes, as previously described [14].

5.2 Preparation of Overexpression Plasmids

The PchB overexpression plasmid (without a histidine tag) was generated as previously described [28]. The Irp9 overexpression plasmid was generated as previously described [66].

*E. coli* chorismate mutase (EcCM) is a domain of a larger pheA gene. The EcCM domain portion of the pheA gene was amplified from *Escherichia coli* K-12 substrain genomic MG1655 DNA (ATCC) by polymerase chain reaction by use of Master Mix (Eppendorf) with 1.5 mM magnesium acetate. The forward primer (5′-ATC GT\_\_\_\_\_ATA TGA CAT CGG AAA ACC CGT TA-3′) includes an *Nde*I site (underlined) and the reverse primer (5′-TT\_\_\_\_\_AGC TTT CAG AGA AAA GCG ATG CGT GCT G-3′) contains a stop codon and a *Hind*III site (underlined). The amplified 352 base pair fragment was digested with *Nde*I and *Hind*III and ligated into the pET28b plasmid (Novagen) digested with the same enzymes. The resultant plasmid encodes the chorismate mutase domain of pheA with an N-terminal histidine tag and no mutations.

5.3 Protein overexpression and purification

PchB without the histidine tag was overexpressed and purified as previously described [28]. Irp9 was overexpressed and purified as previously described [66].

BL21 Star (DE3) pLysS *E. coli* containing the EcCM expression plasmid were grown in LB medium containing 50 µg/mL kanamycin at 37 °C with shaking (250 rpm) until an OD$_{600}$ of ~0.8 was reached. The culture temperature was reduced to 30 °C and the cells were harvested by centrifugation (6 000g, 10 min, 4 °C) after 4 h. The cell pellet was resuspended in 15 mL of 20 mM Tris-HCl pH 8.8, 500 mM NaCl, 5 mM imidazole (buffer A) per liter of culture. Cells were disrupted by use of a French pressure cell (35 000 psi), and cellular debris was removed by centrifugation (12 000g, 30 min, 4 °C). The supernatant was applied to a chelating Sepharose fast-flow column (Amersham Biosciences) charged with nickel chloride and pre-equilibrated in buffer A. EcCM protein eluted at 250 mM imidazole in a linear gradient of 5–500 mM imidazole in buffer A. The pooled fractions were applied to a Superdex 75 size-exclusion column (Amersham Biosciences) equilibrated with 25 mM sodium phosphate pH 8.5, 150 mM KCl, 10% glycerol. The fractions containing EcCM were pooled and concentrated by use of an Amicon stirred cell with a YM-10 membrane to
417 µM as determined by absorbance at 280 nm (ext. coeff 1,490 M$^{-1}$ cm$^{-1}$) and stored at −80 °C.

5.4 High-throughput screening

To identify inhibitors of PchB required for salicylate synthesis, high throughput screening was performed using an assay in which isochorismate is converted to salicylate by PchB. The fluorescence of salicylate was measured with an excitation wavelength of 300 nm and an emission wavelength of 400 nm. The assay was adapted to 384-well (Costar #3575) format and used to screen the compound libraries available at the University of Kansas High Throughput Screening Laboratory, Lawrence, Kansas. The compound collection was derived from both commercially available compound libraries (Prestwick (1120 compounds), Microsource Spectrum collection (2000 compounds), ChemDiv library (50,000 compounds), Chembridge library (43,736 compounds)) and the compounds (3360) synthesized at the University of Kansas Center for Chemical Methodologies and Library Development (CMLD). All enzyme and substrate stocks were prepared in reaction buffer containing 50 mM Tris-HCl, pH 8.0, 150 mM NaCl and 1 mM TCEP. The assay was initiated by adding PchB (12 nM final concentration) enzyme mixture to compounds (10 µM final concentrations) in a 384-well plate and incubated 5 mins at room temperature. The reaction was initiated by adding substrate (isochorismate, 3 µM final which is 1.5-times the $K_m$) solution to the wells containing the compounds and enzyme mixture. The fluorescence intensity was measured at 0 and 20 mins, during which time the reaction was linear at 25 °C, using a Tecan Safire 2 plate reader. Each plate had 16 wells of positive controls (DMSO with no compound) and 14 wells for minus protein negative controls. A compound identified from a previous ChemBridge library screen (ChemBridge #5325243) was included in two wells as a positive assay control (6 µM). In summary, a total number of 100,216 compounds were screened.

5.5 HTS data analysis

Relative Fluorescence Units (RFU) per minute was calculated by subtracting the fluorescence intensity from pre-reads (before the addition of isochorismate) from that of the end reads (20 mins after the addition of isochorismate). The percent inhibition was calculated from median RFU/min in all DMSO/compound containing wells. A hit rate of 0.4% was obtained with 422 compounds showing >50% inhibition in salicylate synthesis over the DMSO treated controls. An average Z’ value of 0.80 ± 0.05 was obtained across all plates screened.

5.6 Secondary screening at HTS facility

All 422 compounds were cherry picked and tested for activity in an 8-point dose-response curve with 2-fold serial dilutions starting at 30 µM. The primary screening assay was used to confirm activity of the cherry-picked compounds. A reconfirmation rate of 50% was obtained and 131 compounds were found to be dose-responsive, with IC$_{50}$ values ranging from 410 nM to >30 µM.
5.7 Preparation of compounds for confirmatory assays and minimum inhibitory concentration

Compounds 9–13 were purchased from ChemDiv. Compounds 6, 7, and 8 were no longer available from ChemBridge and were prepared by a modified procedure that was originally reported by Miller and colleagues [67].

5.7.1 General procedure—Anthranilic acid (177 mg, 1.29 mmol) was added to a solution of Na$_2$CO$_3$ (5 mL, 0.63 M) at 60°C. The appropriate sulfonyl chloride (1.55 mmol, 1.2 equiv) was added in three portions over a period of 10 min. The reaction was heated to 75°C for 30 min and then raised to 85°C and quenched with a solution of HCl (1 mL, 6 M). The solution was cooled to room temperature and the resulting precipitate collected by vacuum filtration and was dried under vacuum overnight. All materials were obtained from commercial suppliers and used without further purification. $^1$H NMR and $^{13}$C NMR were recorded on a Bruker AV-500 with cryoprobe. Chemical shifts (δ) are reported in ppm downfield from TMS and are referenced to the residual DMSO solvent peak. High resolution mass spectroscopy (HRMS) was performed using a LCT Premier (Micromass Ltd., Manchester UK) time of flight mass spectrometer with an electrospray ion source. Melting points are uncorrected and were measured on a Thomas Hoover Capillary Melting Point Apparatus.

5.7.2 5-(N-(2-carboxyphenyl)sulfamoyl)-2-hydroxybenzoic acid (Compound 6) —$^1$H NMR (500 MHz, DMSO) δ 10.99 (s, 1H), 8.14 (d, $J = 2.5$ Hz, 1H), 7.89 (dd, $J = 7.9$, 1.5 Hz, 1H), 7.84 (dd, $J = 8.8$, 2.5 Hz, 1H), 7.57 (m, 1H), 7.49 (dd, $J = 8.3$, 0.9 Hz, 1H), 7.14 (td, $J = 8.0$, 1.2 Hz, 1H), 7.08 (d, $J = 8.8$ Hz, 1H). $^{13}$C NMR (126 MHz, DMSO) δ 170.19, 169.76, 164.32, 139.62, 134.63, 133.47, 131.64, 130.06, 128.84, 123.78, 119.07, 118.66, 117.30, 113.97. HRMS: [M-H]$^-$ calcd 336.0178, found 336.0183. Melting point: 221–222°C (dec.)

5.7.3 2-(3-carboxyphenylsulfonamido)benzoic acid (Compound 7) —$^1$H NMR (500 MHz, DMSO) δ 11.21 (s, 1H), 8.26 (t, $J = 1.6$ Hz, 1H), 8.15 (m, 1H), 8.02 (ddd, $J = 7.9$, 2.0, 1.1 Hz, 1H), 7.88 (dd, $J = 7.9$, 1.5 Hz, 1H), 7.69 (dd, $J = 11.8$, 4.0 Hz, 1H), 7.57 (m, 1H), 7.50 (ddd, $J = 8.3$, 0.9 Hz, 1H), 7.15 (td, $J = 7.9$, 1.2 Hz, 1H). $^{13}$C NMR (126 MHz, DMSO) δ 169.68, 165.77, 139.33, 139.20, 134.58, 134.04, 132.03, 131.62, 130.91, 130.33, 127.31, 123.97, 119.24, 117.65. HRMS: [M-H]$^-$ calcd 320.0229 found 320.0229. Melting point: 234–236°C (dec.)

5.7.4 2-bromo-5-(N-(2-carboxyphenyl)sulfamoyl)benzoic acid (Compound 8) —$^1$H NMR (500 MHz, DMSO) δ 11.16 (s, 1H), 8.09 (d, $J = 2.4$ Hz, 1H), 7.91 (t, $J = 8.4$ Hz, 1H), 7.90 (s, 1H), 7.80 (dd, $J = 8.4$, 2.4 Hz, 1H), 7.57 (ddd, $J = 8.5$, 7.4, 1.6 Hz, 1H), 7.47 (dd, $J = 8.3$, 0.8 Hz, 1H), 7.17 (td, $J = 7.9$, 1.1 Hz, 1H). $^{13}$C NMR (126 MHz, DMSO) δ 169.59, 165.85, 138.99, 138.25, 135.53, 134.61, 134.58, 131.68, 130.25, 128.73, 126.01, 124.10, 119.30, 117.84. HRMS: [M-H]$^-$ calcd 397.9334, found 397.9333. Melting point: 219–220°C (dec.)
5.8 Salicylate fluorescence assay

For the PchB assays, 50 nM enzyme was added to 50 mM potassium phosphate, pH 8 and the reaction was initiated by the addition of isochorismate. For the Irp9 assays, 100 nM enzyme was added to 50 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 10% (v/v) glycerol and the reaction was initiated by the addition of chorismate. Initial velocities were determined at 25 °C by measuring the accumulation of salicylate by fluorescence with an excitation wavelength of 310 nm and an emission wavelength of 430 nm using a Cary Eclipse fluorescence spectrometer (Varian) with temperature controller. Initial velocities were measured for 120 seconds during which the reaction was linear. The substrate concentration was set at the \( K_m \) of 2 µM for IC\(_{50} \) determination, and the inhibitor concentrations varied. For the \( K_i \) determination for compounds 9 and 10, the PchB concentration was 50 nM and the isochorismate concentrations ranged from 0–75 µM with inhibitor concentrations ranging from 0–100 nM for compound 9 and 0–1.5 µM for compound 10. The data was collected as for the IC\(_{50} \) determination measuring the fluorescence of salicylate at 430 nm upon excitation at 310 nm.

The \( K_i \) was determined for compound 6 using a TgK Scientific SF-61DX2 stopped-flow apparatus equipped with a photomultiplier detector and a mercury-xenon lamp at 25 °C. Equal volumes of enzyme with inhibitor were mixed with isochorismate (for PchB \( K_i \) determination) or chorismate (for Irp9 \( K_i \) determination) and initial velocity measurements were collected as an increase in salicylate fluorescence upon excitation at 310 nm using a cutoff filter at 360 nm. Initial velocities determined using the stopped-flow were measured for 30 seconds during which the reaction was linear. The final concentration of PchB was 50 nM, the isochorismate concentration ranged from 0–50 µM and the inhibitor concentration ranged from 0–100 nM in a reaction buffer of 50 mM potassium phosphate, pH 8, 1 mM TCEP. The final concentration of Irp9 was 100 nM, the chorismate concentration ranged from 0–40 µM and the inhibitor concentration ranged from 0–10 µM in a reaction buffer of 50 mM Tris, pH 7.5, 10 mM MgCl₂, 10% glycerol.

5.9 Pyruvate production assay

This assay is based on the method described by Anthon and Barrett [68]. For the PchB pyruvate production assays, 50 nM enzyme was added to 50 mM potassium phosphate, pH 8 and the reaction was initiated by the addition of isochorismate. For Irp9, 200 nM enzyme was used and the reaction was initiated by the addition of chorismate. Initial velocities were determined at 25 °C by adding 70 µl of sample at 2 min time points to 70 µl 0.25 g/L 2,4-dinitrophenylhydrazine in 1 M HCl and incubated 10 min at 37 °C. The absorbance at 515 nm was read upon addition of 70 µl 1.5 M NaOH using a Cary 50 MPR microplate reader (Varian) and converted to pyruvate concentration by means of a standard curve. The substrate concentration was set above the \( K_m \) of 10 µM for IC\(_{50} \) determination, and the inhibitor concentrations varied. All experiments were performed in triplicate.

5.10 Chorismate disappearance assay

For the EcCM assays, the disappearance of chorismate was determined using a TgK Scientific SF-61DX2 stopped-flow apparatus equipped with a photomultiplier detector and a mercury-xenon lamp operated at 37 °C. Equal volumes of enzyme with inhibitor were mixed...
with chorismate in reaction buffer consisting of 50 mM Tris-HCl, pH 8, 2.5 mM EDTA, 20 mM βME, 0.01 % BSA and initial velocity measurements were collected as a decrease in absorbance at 310 nm for 30 seconds during which the reaction was linear. Pre-injection enzyme, substrate, and inhibitor concentration were twice the final concentration in the cell (post-mixing). The post-mixing concentrations were 500 nM for the enzyme, substrate chorismate concentration was set at the $K_m$ of 600 µM, and the inhibitor concentration varied from 0–500 µM. All experiments were performed in triplicate. For the $K_i$ determination of compounds 6 and 10, the final concentration of EcCM was 500 nM, the chorismate concentration ranged from 0–2 mM and the inhibitor concentration ranged from 0–50 µM.

5.11 Bacterial strains used for growth inhibition assays

Wild type strains include *E. coli* strain Seattle (ATCC, 25922), *Yersinia enterocolitica* (ATCC 9610) and *P. aeruginosa* PAO1 (provided by Dr. Jeffrey L. Urbauer, University of Georgia, Athens, GA). *P. aeruginosa* deletion mutant strains were acquired from a *P. aeruginosa* PAO1 transposon mutant library (University of Washington [69]) and include a *P. aeruginosa* PAO1 PvdA deletion mutant (PW5011) and a *P. aeruginosa* PAO1 PchE deletion mutant (PW8175).

5.12 Growth inhibition assay

*P. aeruginosa* and *Y. enterocolitica* were grown overnight in LB media and diluted to an OD$_{600}$ of 0.5 with PMH-D media [70]. Dose response curves were determined by adding 10 µl diluted cells with 90 µl of PMH-D media with 5 µM FeCl$_3$ (iron-limited) or 200 µM FeCl$_3$ (iron-rich) containing inhibitor concentrations of 0–5 mM. The plates were incubated at 37 °C with shaking (200 rpm) for 24 hrs at which time the well solution was centrifuged to pellet the cells and the media with residual inhibitor was removed. The cells were resuspended in 100 µl LB media and the absorbance at 600 nm was read using a Cary 50 MPR microplate reader (Varian). *E. coli* were grown overnight in LB media and diluted to an OD$_{600}$ of 0.2 with LB media. Dose response curves were determined by adding 10 µl diluted cells with 90 µl of tryptic soy broth containing inhibitor concentrations of 0–5 mM. The plates were incubated at 37 °C with shaking (200 rpm) for 10 hrs and the absorbance was read at 600 nm. All experiments were performed in triplicate.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

EcCM  
E. coli chorismate mutase

EntC  
isochorismate synthase from E. coli

Irp9  
salicylate synthase from Yersinia enterocolitica

MbtI  
salicylate synthase from Mycobacterium tuberculosis

MenF  
isochorismate synthase from E. coli

MST  
protein family containing proteins of menaquinone, siderophore or tryptophan biosynthesis

PchA  
isochorismate synthase from Pseudomonas aeruginosa

PchB  
isochorismate-pyruvate lyase from Pseudomonas aeruginosa

TSA  
transition state analogue

VibC  
isochorismate synthase from Vibrio cholera

Reference


Figure 1. Salicylate-capped Siderophores and the Enzymes Targeted
a. The salicylate-capped siderophores generated by *P. aeruginosa* (pyochelin), *Yersinia spp.* (yersiniabactin) and *M. tuberculosis* (mycobactin). b. The isochorismate-pyruvate lyase from *Pseudomonas aeruginosa* (PchB) is the specific enzyme of screen, and performs the pericyclic cleavage of isochorismate 1 to form salicylate 2 and pyruvate 3. Inhibitors were also tested against the salicylate synthase from *Yersinia enterocolitica* (Irp9), which is a chorismate-utilizing enzyme and a structural homologue of the isochorismate synthase PchA from *P. aeruginosa*. Irp9 and PchA catalyze the conversion of chorismate 4 to isochorismate.
1. Irp9 subsequently catalyzes the same lyase reaction as PchB from an isochorismate intermediate, whereas PchA does not. The compounds were also tested for inhibition of \textit{E. coli} chorismate mutase (EcCM), also a chorismate-utilizing enzyme, which performs a pericyclic reaction similar to that catalyzed by PchB, generating prephenate 5.
Figure 2. Enzyme scaffolds and transition states

The X-ray crystallographic structures for PchB (PBD code: 3REM) with salicylate and pyruvate bound, EcCM (1ECM) with Bartlett’s TSA inhibitor bound, and Irp9 (2FN1) with Mg$^{2+}$, salicylate and pyruvate are depicted as cartoons. Each of these enzymes is a homodimer, with one monomer shaded darker than the other, and the active sites identified by the ligands shown as sticks. PchB and EcCM share the same fold – they are AroQ enzymes, whereas Irp9 is in the MST family of enzymes. The transition states for the reactions catalyzed, isochorismate-pyruvate lyase (left) and chorismate mutase (right), are shown below. The transition states are similar, differing only in the alignment of the pyruvlenol tail over the ring to make a cycle at the 1 (mutase) or 2 (lyase) carbon. PchB and Irp9 perform the same chemistry using different scaffolds, whereas PchB and EcCM perform related reactions in the same scaffold. It should be noted that PchB has adventitious mutase activity, albeit very low.
Figure 3. Compounds selected from screen for further evaluation

6: R = OH
7: R = H
8: R = Br

10: R¹ = R³ = NO₂, R² = OCO₂CH₃
11: R¹ = Cl, R² = OCO₂CH₃, R³ = NO₂
12: R¹ = R³ = Br, R² = OH
Figure 4. Growth inhibition of *Pseudomonas aeruginosa*, *Yersinia enterocolitica* and *E. coli* growth

**a.** *P. aeruginosa* growth inhibition by compound 6. **b.** *P. aeruginosa* growth inhibition by compound 10. For both A and B, left: WT PAO1 strain; middle: Δpvda PAO1, a pyoverdin minus strain; right: ΔpchE PAO1, a pyochelin minus strain. **c.** *Y. enterocolitica* growth inhibition by compound 6 (left) and compound 10 (right). For A–C: symbols: • iron poor, ∙ iron rich. **d.** *E. coli* growth inhibition by compound 6 (•) and compound 10 (∙) under iron rich conditions.
IC$_{50}$ values determined for the isochorismate-pyruvate lyase PchB, the salicylate synthase Irp9, and the chorismate mutase EcCM.

<table>
<thead>
<tr>
<th>Compound</th>
<th>PchB IC$_{50}$ (µM)</th>
<th>Irp9 IC$_{50}$ (µM)</th>
<th>EcCM IC$_{50}$ (µM)</th>
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<td>Salicylate Production Assay</td>
<td>Pyruvate Production Assay</td>
<td>Salicylate Production Assay</td>
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<td>NI</td>
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<tr>
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<td>NI</td>
</tr>
<tr>
<td>8</td>
<td>0.24 ± 0.01</td>
<td>0.094 ± 0.001</td>
<td>NI</td>
</tr>
<tr>
<td>9</td>
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<td>0.07 ± 0.01</td>
<td>80 ± 20</td>
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<td>0.53 ± 0.09</td>
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<td>1.8 ± 0.0</td>
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<tr>
<td>12</td>
<td>10 ± 1</td>
<td>10 ± 1</td>
<td>60 ± 20</td>
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<td>13</td>
<td>1.4 ± 0.0</td>
<td>1.4 ± 0.1</td>
<td>42 ± 3</td>
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NI= not inhibitory
## Table 2

Half maximal value of bacterial growth inhibition

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<th>Compound</th>
<th>PAO1 p.a.</th>
<th>ΔpvdA p.a.</th>
<th>ΔpchE p.a.</th>
<th>Y. enterocolitica</th>
<th>E. coli</th>
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</tr>
<tr>
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<td>NI</td>
<td>NI</td>
<td>&gt; 5</td>
<td>3 ± 1</td>
</tr>
<tr>
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<td>10</td>
<td>NI</td>
<td>NI</td>
<td>3.1 ± 0.9</td>
<td>3.2 ± 0.2</td>
</tr>
</tbody>
</table>

NI= not inhibitory