Inhibition of *Mycobacterium tuberculosis* DosRST two-component regulatory system signaling by targeting response regulator DNA binding and sensor kinase heme

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Abstract

*Mycobacterium tuberculosis* (Mtb) possesses a two-component regulatory system, DosRST, that enables Mtb to sense host immune cues and establish a state of non-replicating persistence (NRP). NRP bacteria are tolerant to several anti-mycobacterial drugs and are thought to play a role in the long course of tuberculosis (TB) therapy. Therefore, small molecules that inhibit Mtb from establishing or maintaining NRP could reduce the reservoir of drug tolerant bacteria and function as an adjunct therapy to reduce treatment time. Previously, we reported the discovery of six novel chemical inhibitors of DosRST, named HC101A-106A, from a whole cell, reporter-based phenotypic high throughput screen. Here, we report functional and mechanism of action studies of HC104A and HC106A. RNAseq transcriptional profiling shows that the compounds downregulate genes of the DosRST regulon. Both compounds reduce hypoxia-induced triacylglycerol synthesis by ~50%. HC106A inhibits Mtb survival during hypoxia-induced NRP, however, HC104A did not inhibit survival during NRP. An electrophoretic mobility assay shows that HC104A inhibits DosR DNA binding in a dose-dependent manner, indicating that HC104A may function by directly targeting DosR. In contrast, UV-visible spectroscopy studies suggest HC106A directly targets the histidine kinase heme, via a mechanism that is distinct from the oxidation and alkylation of heme previously observed with artemisinin (HC101A). Synergistic interactions were observed when DosRST inhibitors were examined in pair-wise combinations with the strongest potentiation observed between artemisinin paired with HC102A, HC103A, or HC106A. Our data collectively show that the DosRST pathway can be inhibited by multiple distinct mechanisms.

Keywords: Chemical biology, two-component regulatory systems, microbial pathogenesis.
Introduction

*Mycobacterium tuberculosis* (Mtb) is the causative agent of tuberculosis (TB). Mtb is an intracellular pathogen that can latently infect the host without causing disease symptoms (1). During chronic infection, it can establish a dormant state known as non-replicating persistence (NRP) where Mtb modulates its metabolism in response to environmental and host immune cues, such as hypoxia, acidic pH, and nutrient starvation (2, 3). DosRST is a two-component regulatory system that regulates Mtb persistence (4-6). It consists of two sensor histidine kinases, DosS and DosT, and the cognate response regulator DosR, which regulates expression of about 50 genes in the DosRST regulon (6-8). The pathway can be induced by host intracellular stimuli, such as nitric oxide (NO), carbon monoxide (CO) and hypoxia, through DosS and DosT (9-11). DosS is an oxygen and redox sensor, whereas DosT acts an oxygen sensor (12-14). Both kinases sense ligands via the heme group, and are inactive when the heme exists as either the Met (Fe$^{3+}$) form (DosS) or the oxy (Fe$^{2+}$-O$_2$) form (DosT) in the presence of O$_2$ (13). However, hypoxic conditions activate the kinases by inducing the conversion of DosS to the ferrous form and DosT to the deoxy form. Therefore, DosS/T play overlapping and distinct roles in sensing the redox status and oxygen level of the environment to turn on the DosR pathway (11, 15).

Non-replicating bacilli are naturally tolerant to many anti-mycobacterial drugs, such as isoniazid (INH) that only kills replicating Mtb (16-18). During TB infection, the disease presents as a spectrum of replicating and non-replicating bacteria; the NRP population of bacteria are thought to be responsible, in part, for the 6-month long course of TB treatment. This long antibiotic regimen makes controlling the TB epidemic challenging and has likely contributed to the evolution of drug-resistant Mtb strains. Therefore, there exists an urgent need to identify new strategies to treat the disease, with a particular focus on discovering new ways to shorten the course of TB therapy. Targeting the DosRST pathway is a promising strategy, because dosRST mutants are attenuated in *in vitro* models of hypoxia-driven NRP (19) and in animal...
models that generate hypoxic granulomas, including non-human primates, guinea pigs, and C3HeB/FeJ mouse models of TB infection (4, 20-22). Furthermore, deletion of DosR-regulated gene tgs1, which is involved in triacylglycerol (TAG) synthesis, causes reduced antibiotic tolerance (23, 24). Therefore, inhibiting the DosRST pathway and killing the reservoir of NRP bacteria may function to narrow the spectrum of TB disease and shorten the course of TB therapy.

Fluorescent reporter strains can be used as tools in drug discovery to conduct targeted, whole cell phenotypic screens for pathways that play a role during pathogenesis, but have limited impact when bacteria are grown in vitro (25, 26). In an effort to discover the new chemical probes that inhibit Mtb persistence, we previously performed a whole cell phenotypic high throughput screening (HTS) of a >540,000 compound library using the DosRST regulon reporter strain CDC1551 (hspX'::GFP) (27). We discovered six compounds that inhibit the DosR-dependent, hypoxia-induced GFP fluorescence. In the previous report, we showed that the HC101, HC102 and HC103 series functioned to inhibit NRP associated physiologies, including TAG accumulation, survival during hypoxia and isoniazid tolerance. Mechanism of action studies showed that the HC101 series, composed of artemisinin and related analogs, functioned by oxidizing and alkylating the DosS and DosT heme. HC102 and HC103 did not modulate the DosS/T heme, and were instead found to inhibit sensor kinase autophosphorylation. Here, we report the characterization of two additional compounds, HC104A and HC106A. Transcriptional and biochemical analysis demonstrate that both compounds function to downregulate the DosRST pathway and inhibit persistence associated physiologies. Biochemical studies show HC104A and HC106A function by distinct mechanisms of action, with HC104A inhibiting DosR DNA binding and HC106A interacting with DosS and DosT heme to block environmental sensing. Studies examining the pair-wise interactions between the five DosRST inhibitors revealed synergistic interactions, including strong potentiating interactions of artemisinin and HC106A. Structure activity relationship studies of HC106 identified functional groups of HC106A
that are required for activity and enabled optimization of HC106 potency to nanomolar effective concentrations against whole cell Mtb.

Results

Inhibition of the DosR regulon by HC104A and HC106A

Characterization studies were undertaken with two putative DosRST regulon inhibitors, HC104A and HC106A (Fig. 1a) (27). Half-maximal effective concentration (EC$_{50}$) studies using the CDC1551 (hspX'::GFP) DosRST-dependent fluorescent reporter strain, show that HC104A and HC106A inhibit DosRST-dependent GFP fluorescence with EC$_{50}$ values of 9.8 µM and 2.48 µM, respectively (Fig. 1b and 1c).

The compounds have minimal impact on Mtb growth, suggesting they are also potential DosRST inhibitors, as DosRST is not required for survival under the conditions of mild hypoxia used in the reporter-based assay. RNA-seq-based transcriptional profiling was undertaken to determine if the DosRST regulon was inhibited by the compounds. Mtb was treated with 40 µM HC104A, HC106A or dimethyl sulfoxide (DMSO) control for 6 d in a standing flask, and following incubation RNA was extracted, sequenced and analyzed for differential gene expression relative to the DMSO control. As a control for the DosR regulon, transcriptional profiling was also previously conducted on a DMSO treated CDC1551(ΔdosR) mutant strain (27). The transcriptional profiles showed that the genes strongly repressed by HC104A and HC106A (>2-fold; q<0.05) are from the dosR regulon (Fig. 2a-c, Supplementary Dataset 1, 3).

HC106A exhibited a remarkably strong reduction of gene expression, with transcripts for tgs1 and hspX being almost undetectable by RNA-seq following HC106A treatment. Interestingly, while HC106A broadly inhibited genes of the DosRST regulon, HC104A only strongly inhibited part of the DosR regulon, with the strongest inhibition reserved for hspX, the promoter used to drive reporter fluorescence in the screen. These RNA-seq results were...
validated by semi-quantitative RT-PCR, with HC104A causing downregulation of dosR, hspX, and tgs1 in vitro by 6-, 570- and 13-fold, respectively; whereas HC106A downregulated these three genes by 49-, 1360-, and 1424-fold, respectively (Supplementary Fig. 1), with hspX and tgs1 transcripts being below the level of detection by qRT-PCR.

Comparisons of transcriptional profiles from the inhibitor treated wild type (WT) Mtb strain to a CDC1551(ΔdosR) mutant strain showed that there are a total of 26 genes and 53 genes from dosR regulon inhibited by HC104A and HC106A, respectively. Notably, HC104A and HC106A caused an additional 119 genes and 35 genes to be repressed that were not repressed in the CDC1551(ΔdosR) mutant strain (Fig. 2c). This observation suggested that these two compounds exhibit some DosR-independent activities. To confirm the specificity of the compounds, RNA-seq was also performed on CDC1551(ΔdosR) mutant background (Supplementary Dataset 2-3) treated with HC104A or HC106A. This analysis identified 171 genes and 51 genes that are downregulated (>2-fold; q<0.05) by HC104A and HC106A, respectively (Fig. 2d). This finding indicates that HC104A and HC106A impact other targets beside DosR regulon, with HC106A showing greater on-target specificity than HC104A. Based on these findings, we conclude that: 1) HC106A strongly and specifically inhibits the DosRST regulon; and 2) HC104A strongly inhibits a portion of the DosRST regulon, with several notable off-target activities.

To assess the impact of the inhibitors on the DosRST pathway on intracellular Mtb, murine bone marrow-derived mouse macrophages were infected with Mtb and treated with 40 μM HC104A and HC106A for 48 h. Total bacterial RNA was isolated and analyzed by RT-PCR for hspX and tgs1 gene differential expression. The results demonstrate that the induction of hspX and tgs1 were inhibited 185- and 10-fold by HC104A and 6- and 4-fold by HC106A, respectively (Fig. 3a). These findings confirm that HC104A and HC106A can access Mtb inside
the macrophage, however, the reduced repression of the pathway by HC106A as compared to broth culture, suggests that the molecule may not be able to efficiently target intracellular Mtb.

The DosRST pathway is also induced by redox signals such as vitamin C and NO (27). To examine whether HC104A and HC106A can repress the induction of DosRST pathway by vitamin C or NO, Mtb cells were pretreated with HC104A or HC106A for 24 h followed by vitamin C or NO induction for 2 h. The expression of DosR-regulated genes (hspX and tgs1) was examined by real time-PCR. Vitamin C and DETA-NONOate (NO donor) strongly induced hspX and tgs1 as previously reported ((27), Fig. 3b). For instance, vitamin C induced hspX and tgs1 by 2162- and 58-fold, respectively; whereas DETA-NONOate upregulated hspX and tgs1 3024- and 113-fold, respectively (Fig. 3b). Mtb cells pretreated with HC106A showed strong inhibition of hspX and tgs1 induction by vitamin C and DETA-NONOate. For example, HC106A inhibited the hspX and tgs1 transcripts by 78- and 14-fold following vitamin C treatment, respectively, and 362- and 151-fold following DETA-NONOate treatment. Following vitamin C treatment, HC104 showed inhibition of hspX by 3.4-fold and no effect on tgs1, or 302- and 6.6-fold inhibition of hspX and tgs1 following DETA-NONOate treatment. These findings show that HC104A and HC106A act as inhibitors of the DosRST pathway in response to redox signals in addition to hypoxia.

Inhibition of Mtb persistence-associated physiologies

Mtb accumulates TAG during hypoxia (27-29). DosR directly regulates tgs1, which encodes a TAG synthase that is involved in the last step of TAG biosynthesis and is required for TAG accumulation during hypoxia (30). Transcriptional profiling showed that HC104A and HC106A repress expression of tgs1. Based on this transcriptional profiling, we hypothesized these two compounds may inhibit TAG biosynthesis during NRP. To test our hypothesis, Mtb cells were radiolabeled with 14C-acetate and treated with HC104A or HC106A for 6 d. Lipids were isolated and analyzed by thin layer chromatography (TLC). As previously observed,
DMSO treated CDC1551(ΔdosR) mutant displayed a strong (87%) reduction of TAG accumulation as compared to DMSO treated WT (Fig. 3c and Supplementary Fig. 2). Mtb cells treated with HC104A or HC106A showed a ~50% reduction of TAG accumulation, supporting our hypothesis that the compounds can inhibit TAG biosynthesis.

DosRST has been previously reported to be required for survival during NRP, where deletion of DosR causes greatly reduced survival during prolonged hypoxic stress (19). The impact of HC104A and HC106A on Mtb survival during NRP was examined using the hypoxic shift-down model (27). Mtb survival was examined following 10 d of treatment with the compounds at 40 µM. The ΔdosR mutant control had 15% survival relative to DMSO, and was partially complemented, supporting the proposal that survival during hypoxia is DosR dependent. Mtb cells treated with HC106A displayed 50% survival relative to DMSO control (Fig. 3d), whereas, HC104A had no impact on Mtb survival during NRP, an observation that suggests the portion of the DosR regulon inhibited by HC104A is not essential for survival during NRP.

**HC104A inhibits DosR DNA binding**

There are several potential targets of HC104A and HC106A to directly inhibit DosRST signaling, including: 1) direct inhibition of DosS/T sensor kinase activity, 2) modulation of the heme in the sensor, or 3) inhibition of DosR binding of DNA. To investigate the biochemical mechanisms of action of HC104A and HC106A, inhibition of DosS autophosphorylation was initially evaluated. The DosS protein was treated with different concentrations of HC104A and HC106A from 10 µM to 40 µM, or with 40 µM HC103A as a positive control that was previously discovered to be a DosS/T autophosphorylation inhibitor (27). As previously observed, HC103A strongly inhibited DosS autophosphorylation, but HC104A and HC106A had no inhibitory activity.
(Supplementary Fig. 3). This suggests HC104A and HC106A are not directly inhibiting DosS/T autophosphorylation activity.

Next, a UV-visible spectroscopy assay was employed to investigate if HC104A targets to the heme of the sensor kinase DosS. Treatment of DosS protein with the reducing agent dithionite (DTN) caused a shift of the Soret peak from 403 nm to 430 nm as previously described (13, 27). Addition of HC104A to reduced DosS did not shift the peak to the oxidized position, suggesting that HC104A does no modulate DosS heme redox (Supplementary Fig. 4). Together, these data indicate that HC104A does not inhibit DosRST signaling by targeting sensor kinase autophosphoryation or heme and supported examining DosR as a potential target.

Inspection of the HC104A structure revealed it had significant similarity to the compound viristatin (31). Viristatin is an anti-virulence compound that inhibits *Vibrio cholera* cholera toxin production by targeting the transcription regulator ToxT (31). Viristatin inhibits ToxT protein dimerization and subsequently interferes with DNA binding, thereby inhibiting the transcription of downstream genes involved in toxin production. Based on the similarity of chemical structure between HC104A and viristatin, we hypothesized that HC104A may be targeting DosR, and interfering with DNA binding. To test this hypothesis, an electrophoretic mobility shift assay (EMSA) was employed to investigate the impact of HC104A on DosR DNA binding. Recombinant DosR protein, ranging from 0.5 µM to 4 µM, was treated with 40 µM HC104A or a DMSO control and tested for binding to fluorescently labeled *hspX* promoter DNA. In the DMSO treated control, DosR bound promoter DNA beginning at a concentration 2 µM DosR protein (Fig. 4a). Treating the reaction containing 2 µM DosR protein with HC104A significantly inhibited DNA binding by ~22-fold compared to DMSO control (Fig. 4b). To further characterize the impact of HC104A on DosR binding of DNA, a dose-response study was performed. Reactions containing 2 µM recombinant DosR proteins were treated with different concentrations of
HC104A or virstatin ranging from 1 – 80 µM. HC104A inhibited DosR binding of DNA beginning at 10 µM HC104A. The fraction of free DNA increased as HC104A concentration increased (Fig. 3.4c). For example, the fraction of free DNA was 72%, 89%, 92%, 96% and 100% for 20 µM, 30 µM, 40 µM, 60 µM and 80 µM HC104A, respectively, whereas DMSO control had 15% free DNA (Fig. 4d). Thus, HC104A significantly inhibits DosR-DNA binding in a dose-dependent manner. Reactions treated with virstatin had no impact on DosR binding of DNA (Supplementary Fig. 5a). Consistent with these observations, virstatin did not have any impact on DosRST signaling in the whole cell Mtb fluorescence reporter assay (Supplementary Fig. 5b). These findings support the hypothesis that HC104A may function by inhibiting DosR DNA binding activity and has an activity that is distinct from the related molecule virstatin.

**HC106A modulates DosS heme**

DosS and DosT have a channel that exposes the heme to the environment and enables interactions with gases (32, 33). This channel is an Achilles heel that can be targeted by small molecules. Previously, it was shown that the artemisinin modulates DosS/T by oxidizing and alkylating heme carried by the kinases (27). UV-visible spectroscopy studies were conducted to examine if HC106A modulated DosS heme. Recombinant DosS was purified from *E. coli*, degassed and the change of DosS heme spectrum was monitored under anaerobic conditions by UV-visible spectroscopy. Treating DosS with the reducing agent dithionite (DTN) caused the Soret peak to shift to 430 nm as shown previously (13, 27). HC106A was added to the reaction following DTN treatment to observe the impact on the DosS heme UV-visible spectrum. HC106A caused the DosS Soret peak to immediately shift to 422 nm, where the peak was stably maintained for 2 h (Fig. 5a). This spectrum shift is different from artemisinin, where under identical conditions, artemisinin causes the DosS Soret peak to gradually shift back to the
oxidized state at 403 nm (27). These findings show that HC106A may also interact with sensor kinase heme, but via a mechanism that is distinct from artemisinin-heme interactions.

The Soret peak at 422 nm is consistent with previously described spectra that are observed when DosS heme interacts with NO or CO (13). To confirm this observation, DosS was treated with 100 µM CORM-2 (a CO donor) which caused a shift of the Soret peak to 422 nm, similar to what was observed for HC106A (Fig. 5b). This finding supports a hypothesis that HC106A may also be directly binding to the heme. Notably, CO activates DosS kinase function, whereas HC106A functions to inactivate the regulon, demonstrating that the impact of heme binding by CO or HC106A has differing impacts on the sensor kinase switch.

Amino acid substitutions in the channel exposing the DosS heme to the environment, such as DosS E87L or G117L, can limit access of artemisinin to modulate heme (27). To support the hypothesis that HC106A directly targets DosS heme, we tested the impact of these amino acid substitutions on HC106A/DosS heme interactions. Treating DosS(E87L) with HC106A exhibited a profile similar to wild type DosS with the Soret peak shifting to 422 nm (Supplementary Fig. 6). However, DosS(G117L) had no change to the overall spectrum after HC106A treatment (Fig. 3.5c). This finding indicates that DosS(G117L) is resistant to HC106A, and confirms that HC106A accesses the heme via a similar mechanism as artemisinin.

To confirm DosS is a target of HC106A in Mtb, we examined the impact of overexpressing DosS protein in Mtb. If DosS is the target of HC106A, overexpression may reduce the effectiveness of HC106A. WT DosS protein was constitutively expressed from the hsp60 promoter in Mtb. The vector control showed that both hspX and tgs1 genes were downregulated by HC106A by 2331- and 3470-fold, respectively (Fig. 5d). Overexpressing DosS provided significant resistance to HC106A, with hspX and tgs1 showing 23- and 16.5-fold less inhibition, respectively, relative to the empty vector control. This observation of resistance in Mtb is consistent with the biochemical data supporting the view that DosS is a direct target of HC106A.
Synergistic interactions between DosRST inhibitors

With multiple distinct inhibitory activities of the HC101A, HC102A, HC103A, HC104A and HC106A, we sought to examine if potentiating or antagonistic interactions existed between the molecules, when targeting the DosRST pathway. To examine these interactions, checkboard assays were performed with pairwise comparisons of artemisinin (HC101A), HC102A, HC103A, HC104A and HC106A. CDC1551 (hspX'::GFP) was treated with combinations of two compounds ranging from 50 \( \mu M \) to 0.08 \( \mu M \) in 96-well plates. DosR-driven GFP fluorescence and optical density were measured following 6 d of treatment. The Combination Index (CI) was calculated for each drug pair based on the Chou-Talalay method in the CompuSyn software package (34, 35), where CI values of < 1, = 1 and >1 indicate synergistic, additive or antagonistic interactions, respectively. Among all 64 compound pairs, artemisinin combined with HC102A, HC103A, HC104A and HC106, showed 46, 49, 41, and 50 combinations that have CI <1, respectively (Fig. 3.6). Notably, some CI values are below 0.1 when artemisinin was paired with HC102A, HC103A or HC106A combinations. Example dose response curves illustrate these synergistic interactions (Fig. 6). Several other pairwise comparisons also demonstrated synergy (Supplementary Fig. 7), however, in general, these interactions had CI between 1 and 0.1, supporting weaker synergistic interactions, as compared to combinations with artemisinin. Overall, these studies provide the evidence that the inhibitors function by distinct mechanisms and may be combined to improve potency.

Structure-activity relationship study for HC104A and HC106

We conducted a catalog search for HC104 and HC106 analogs and obtained 10 commercial analogs for each series to define initial structure activity relationships (SAR). For HC104A we observed that a bromine in the 5-position is required for activity and that the R2 dimethylamine group is not required (Supplementary Table 1). For example, HC104B is
identical to HC104A except for the removal of the bromine (R1), which results in a complete loss of activity in the whole cell assay. Whereas, replacement of the R2 group with a methyl (HC104G) results in an active compound, although ~5-fold less active than HC104A. Although not highly potent, its ligand efficiency, cLogD and druglikeness are in the range of what would be considered acceptable to good as a starting point for further manipulation. For HC106A (Table 1), catalog SAR work led to new understandings of the nature of the series. We first found that the simple removal of an ortho chloro on the “A” ring of HC106A leads to ~ 2-fold enhanced activity, with an EC_{50} in the whole cell Mtb assay for DosRST inhibition of 1.33 μM (HC106F). It was also found that the use of an alternative isomer of the isoxazole had no detectable activity (HC106C).

To further understand the SAR of the HC106 series, additional analogs were synthesized to examine the need of the central urea functionality and whether modifications can be tolerated (Table 1). A pyridyl analog (MSU-41425), designed to replace the isoxazole also demonstrated no activity as was the symmetrical 4-chloroaniline derived urea (MSU-41324). However, the bis-isoxazole urea (MSU-39444) provides an EC_{50} of 1.7 μM, indicating that the isoxazole is important for function. Isoxazoles are unique among heterocycles in that they exist in multiple tautomeric forms as supported by initial NMR studies (36). We next explored the need of one of the -NHs of the urea, capping it with a methyl (MSU-39451), integrating it into a ring for conformational restriction (MSU-39453), and replacing with a methylene unit (MSU-39449). In all cases, reduced activity (0.5 - 1 log) was observed but not all activity was lost. Thus, HC106A is a potent whole cell inhibitor of the DosRST pathway, with flexibility to be improved via SAR.

To further test the SAR, we conducted a Topliss Tree evaluation of the “A-ring” aniline (Table 20)(37). To reliably prepare the derivatives, we explored and established a general preparation (Supplementary Fig. 8). This route is preferred relative to alternative approaches
for its cleanliness, yields and ease of purification, usually by trituration. It is also anticipated that it will allow access to future derivatives. Using HC106F and HC106A as starting points, we prepared the 3,4-dichlorochloro and 3-chloro derivatives (MSU-39452 and MSU-39445, respectively). Both the 3- and 4-chloro derivatives demonstrated greater activity than 3,4-dichloro (MSU-39452). We found that replacing the 4-chlorophenyl ring with pyridyl analogs (MSU-39448 and MSU-39450) lead to similar activity. Focusing on 4-position derivatives, we found that fluoro (MSU-39446), bromo (MSU-41464) and methoxy (MSU-39447), as electron p-orbital donating substituents, also lead towards incrementally increased activity. Para-t-butyl phenyl (MSU-41442), provided slightly diminished activity. Electron withdrawing substituents, such as 4-CO₂Me (MSU-4165), 4-trifluoromethyl (MSU-41463) and biphenyl (MSU-41443) saw activity similar to the 4-chlorophenyl derivative (HC106F). Overall, several analogs were discovered with significantly ~4-fold enhanced potency, with several inhibitors having whole cell DosRST inhibitory EC₅₀ below 1 µM, including EC₅₀ of 0.63 µM, 0.61 µM, 0.54 µM and 0.75 µM for MSU-33189, -39447, -39446, and -39455, respectively. The parent analog HC106A had an EC₅₀ of 2.48 µM.

There appears to be little sensitivity, positive or negative, for electron-drawing substituents other than the biphenyl derivative (MSU-41443), which is likely the result of negative steric interactions in the binding domain. Additional derivatives (replacement of dichlorophenyl) were also prepared to further probe the size and nature of the binding domain. We began by replacing the chlorophenyl ring of HC106F with benzyl- (MSU-41462), isobutyl- (MSU-41542), cyclopentyl- (MSU-41546) and cyclohexyl- (MSU-42002) analogs. All analogs demonstrated similar activities, relative to HC106F and the simple phenyl analog (MSU-33189), suggesting flexibility of fragments that could bind in this domain.

Kinetic solubility assays were conducted for selected analogs and all exhibited excellent aqueous solubility greater than >100 µM, except for MSU-41443 (Table 2). This finding shows that the urea group present in the HC106A does not have a detrimental impact.
on HC106 aqueous solubility. All of the tested derivatives also demonstrated favorable
mouse microsomal stability, including. Overall, the nanomolar whole cell potency, flexible
SAR, good microsomal stability and excellent solubility, confirm that the HC106 series is a
suitable series for continued optimization to identify a drug-like lead.

DosRST is a two-component regulatory system required for Mtb environmental sensing,
adaptation, and persistence. By using a fluorescent reporter strain CDC1551 (hspX’:GFP), we
have previously discovered small molecule inhibitors of DosRST, named HC101A-HC106A,
from the whole-cell phenotypic HTS. Here, we report the characterizations of HC104A and
HC106A as DosRST inhibitors. Both compounds downregulated genes in the DosR regulon,
and inhibited TAG biosynthesis. HC106A also reduced Mtb survival during NRP.

In a UV-visible spectroscopy assay, we observed that HC106A interacts with the heme
of sensor kinase DosS. The UV-visible spectrum of HC106A-treated DosS is similar to those of
CO- or NO-treated DosS. The overlap between the CO and HC106A spectra supports that
HC106A may also directly bind to the heme of DosS. Interestingly, CO activates sensor kinases,
whereas HC106A inhibits them. This could be due to the difference in conformational changes
induced by CO and HC106A, or binding of HC106A may lock the sensor kinases into an inactive
state. Furthermore, the DosS G117L substitution in recombinant DosS blocks the heme
exposing channel and provides resistance to HC106A. This means that, similar to artemisinin,
this channel is also important for the activity of HC106A. These findings provide additional
evidence that the heme-exposing channel in DosS/T can be exploited by small molecules to
inhibit the heme from sensing signals and to disrupt signal transduction of a two-component
regulatory system. To our knowledge, these two distinct mechanisms of actions are novel
mechanisms to inhibit the DosRST pathway.

Mechanistic studies via EMSA indicates that HC104A may function by targeting DosR
and inhibiting DosR DNA binding. Virstatin had no effect on DosR DNA binding and no impact
on the DosRST signaling in whole cells, showing that although both compounds share a similar
structure, they function by distinct mechanisms. From the catalog SAR study, we found that the
bromine group is required for the compound. Virstatin does not have the bromine, and the R-
group is butyric acid instead of dimethylamine. These two differences are enough to differentiate
the activity of the compounds. Moreover, transcriptional profiling shows that the most repressed
genes by HC104A are from the DosR regulon, providing additional evidence that HC104A is
somewhat selective for specific DosR regulated genes. Interestingly, the genes most
downregulated by HC104A, including hspX, Rv2030c, pfkB, and Rv2028c, are from the same
operon under control of hspX promoter, which is strongly induced by DosR in hypoxia. This
result suggests that HC104A is more specific to target hspX operon genes as compared to other
DosR regulated genes. This finding leads to the speculation that HC104A may be more efficient
to prevent DosR binding to the hspX promoter than the other DosR promoters. HC104A may fit
better in the pocket in the interface of DosR-hspX’ complex. This postulation also supports the
idea that HC104A may not have an impact on DosR protein dimerization, which would lead to
universal downregulation of DosR-regulated genes. Detailed characterizations of HC104A on
inhibiting DosR dimerization and promoter binding specificity is the subject of ongoing
characterizations.

This study together with the previous report reveals multiple distinct inhibitory
mechanisms for the five compounds, HC101A-104A, HC106A. Four out five compounds,
HC101A-HC103A and HC106A, are proposed to function by targeting the sensor kinases
DosRST. Furthermore, these four compounds are effective at decreasing Mtb survival during
NRP in hypoxic-shift down assay. In contrast, the compound targeting DosR, HC104A, had no
impact on Mtb survival during NRP. This result is consistent with a previous report that showed
only the ΔdosS mutant exhibits a survival defect in the C3HeB/FeJ mouse model (22). Notably,
the dosR mutant strain used in our study is also lacking dosS expression due to polar impacts of
the deletion, supporting the hypothesis that the reduced survival during NRP may be dependent
on DosS. Alternatively, we noted that HC104A only inhibits a portion of the DosR regulon with
only 26 genes being downregulated. This finding suggests that the DosR regulated genes that are inhibited by HC104A (Fig. 2, and Supplementary Dataset 1, 3) are not required for DosRST-dependent persistence during NRP or not sufficiently inhibited to have an impact on survival.

The transcriptional profiling data showed off-target impacts in the treated dosR mutant strain. These effects may be due to inhibition of DosS/T signaling that functions independent of the response regulator. Several recent studies have demonstrated the occurrence of cross-interactions between histidine kinases and response regulators in Mtb. For instance, Lee et al. shows that DosT can interact with the other non-cognate response regulators, including NarL and PrrA (38). Our transcriptional profiling of the compound treated dosR mutant strain suggests that some genes downregulated by HC101A, HC103A and HC106A may be DosS-dependent but DosR-independent (Supplementary Fig 9). In the prior study, six genes are similarly regulated between artemisinin and HC103A in the treated dosR mutant strain, including Rv0260c that encodes a putative response regulator (27). HC106A and HC103A share four differentially regulated genes in the dosR mutant, including argC, argJ, argB, and argF, which are genes involved in arginine biosynthesis. Our data presented here and in the literature point to the possibility that DosST modulate gene expression independently of DosR.

The synergy studies show significant synergistic interactions between artemisinin, HC102A, HC103A, HC104, or HC106A. Moreover, artemisinin exhibited the greatest synergistic activities with HC102A, HC103A or HC106A, indicating that inhibition of histidine kinases by a second inhibitor can lead to synergistic inhibition of the DosRST pathway. This interaction could be due to both sensor kinases being required for full induction of the DosR regulon, where DosT responds early during hypoxia and DosS further induces the regulon at later times (39), and it is possible that the inhibitors have different affinities for DosS or DosT. Thus, multiple inhibitors could inhibit both DosS and DosT better than an inhibitor alone. Interestingly, artemisinin shows the greatest synergism with HC106A. Both compounds are proposed to target the heme of
DosS/T, but through different mechanisms. This finding shows that both inhibitors can enter the channel of DosST to interact with the heme and do so without antagonizing interactions.

The identification of new antibacterial agents and tuberculosis drugs has been associated with the realization that these compounds can occupy a different region of chemical space relative to drugs in most other therapeutic areas. Series HC106 is easily in the range of the characteristics for compounds currently in use or in development as anti-TB compounds. Series HC104 needs further exploration to fully assess its suitability. The physicochemical properties of a drug, such as solubility and permeability, impact its oral bioavailability as these factors influence absorption, distribution, metabolism, and excretion. Series HC106 demonstrates excellent aqueous solubility (Table 2, with compounds generally having solubility >100 µM). The melting points of most of the compounds in the series and the corresponding cLogP are also supportive appropriate solubility. The ability of an anti-TB drug to reach its target site is greatly hampered by the highly impermeable Mtb cell envelope. Both series (HC104 and HC106) demonstrate low micromolar and nanomolar whole cell EC₅₀ values, suggesting adequate cell wall permeability. SAR studies of HC106 show that there remain significant opportunities for optimization of HC106 for potency and drug properties and future studies will be focused on such optimizations.

**Methods**

**Bacterial strains and growth conditions.** Mtb CDC1551, CDC1551 (ΔdosR) strains were used in this study. All cultures were grown at 37°C and 5% CO₂ in 7H9 Middlebrook medium supplemented with 10% OADC (oleic acid albumin dextrose catalase) and 0.05% Tween-80 in standing, vented tissue culture flasks, unless stated otherwise.

**EC₅₀ assays.** The assay was performed as previously described (27). Briefly, the (hspX'::GFP)...
reporter strain culture was diluted to an OD$_{600}$ of 0.05 in fresh 7H9 media, pH 7.0, and 200 µL of diluted culture was aliquoted in clear-bottom, black, 96-well plates (Corning). Cells were treated with an 8-point (2.5-fold) dilution series ranging from 200 µM – 0.32 µM. For the structure relationship studies for the HC106 series, a 12-point (2.5-fold) dilution series of HC106 analogs ranging from 200 µM – 8.4 nM were used. GFP fluorescence and optical density were measured following 6 d incubation. Percentage fluorescence and growth inhibitions were normalized to a rifampin-positive control (100% inhibition) and DMSO-negative control (0% inhibition). EC$_{50}$ values were calculated for each compound using GraphPad Prism software package (version 6). Each experiment was performed with two technical replicates per plate and two biological replicates, and the error bar represents the s.d. of the biological replicates. Experiments were performed twice with similar results.

**Transcriptional profiling and data analysis.** Transcriptional profiling studies were conducted as previously described in Zheng et al. (27). Briefly, CDC1551 or CDC1551 (ΔdosR) cultures were treated with 40 µM HC104A, HC106A or DMSO control for 6 d. The starting OD$_{600}$ was 0.1 in 8 mL of 7H9 medium in standing T25 vented tissue culture flasks. Bacterial growth consumes oxygen and stimulates the DosRST pathway. The total bacterial RNA from two biological replicates was isolated and prepared for sequencing as previously described (40). The RNA-seq data were processed and analyzed using the SPARTA software package (41). Sequencing data are available at the GEO Database (Accession GSE115892).

**Real time-PCR assays.** The vitamin C and NO assays were performed as previously described (27). Briefly, cultures at an OD$_{600}$ of 0.6 were pretreated with 80 µM HC104A, HC106A or a DMSO control for 24 h, and induced with 50 µM DETA-NONOate or 20 mM vitamin C for 2 h. For the HC106A resistance assays, CDC1551 was transformed with the empty replicating
plasmid pVV16 or the plasmid expressing dosS from the strong hsp60 promoter (pVV16-DosS), and treated with 20 μM HC106A for 6 d. Total bacterial RNA was isolated and differential gene expression of DosR-regulated genes, including hspX and tgs1, was quantified. The experiment was performed in three technical replicates and error bars represent the s.d from the mean. The experiment was repeated twice with similar results. To examine Mtb gene expression in macrophages, murine bone-marrow derived macrophages were isolated as previously described (42) and seeded in T75 vented, tissue culture flasks. Macrophages were infected with CDC1551 with multiplicity of infection ratio of 1:20 as previously described (42). After infection, the flasks were treated with 40 μM HC104A or HC106A or DMSO for 48 h, with three individual flasks for each treatment. Total bacterial RNA was isolated after treatment, and the transcripts of DosR-controlled genes (hspX and tgs1) were quantified in RT-PCR. The experiment was conducted with three biological replicates. The error bar represents the s.d. of the biological replicates.

**TAG biosynthesis.** The lipid labelling and TAG TLCs were performed as previously described (27). Briefly, CDC1551 was cultured at an initial OD600 of 0.1 and radiolabeled with 8 μCi of [1,2-14C] sodium acetate in T25 vented tissue culture flasks. The cultures were treated with 40 μM HC104A, HC106A or DMSO for 6 d at 37°C. CDC1551 (ΔdosR) and dosRS complement strains were also examined. Total lipids were extracted and 14C incorporation was determined by scintillation counting. 20,000 c.p.m. of total lipids were analyzed by TLC using silica gel 60 aluminum sheets (EMD Millipore). To determine TAG accumulation, the lipids were developed in hexane-diethyl ether-acetic acid (80:20:1; vol/vol/vol) solvent system. The TLC was exposed to a phosphor screen for 3 d, and imaged on a Typhoon imager and TAG was quantified using ImageJ software (43). The experiment was repeated twice with similar results, and the error bar represents the s.d. of two biological replicates.
NRP survival assays. Survival during NRP was examined using the hypoxic shift down assays as previously described (27, 44). Briefly, CDC1551 cells were treated with 40 μM HC104A, HC106A or DMSO control in a 24-well plate (1 mL/well). CDC1551 (ΔdosR) and dosRS complement strains were also examined. Plates were incubated in an anaerobic chamber (BD GasPak) for 12 d. It took 48 h for cultures to become anaerobic, as monitored by a methylene blue control. Bacterial CFUs were numerated on 7H10 agar plates following incubation. The experiment was repeated twice with similar results.

DosR protein purification. DosR full length protein was purified as previously described (45). Briefly, the dosR gene (Rv3133c) was cloned into pET15b (Novagen Darmstadt, Germany) using the primer set: forward primer 5'-TTTCATATGGTGAAGGTTTCTTCTGGTCATGAC-3'; reverse primer 5'-TTTGGATTCATGGGATCCATGCACCGGGTGG-3'. The His6-DosR protein was expressed in E. coli BL21(DE3) strain. The culture was grown to OD600 0.5-0.6, and induced with 1 mM IPTG for 6.5 h at 29°C. The cell pellet was suspended in lysis buffer (20 mM Tris-HCl, pH 8.0, 10% glycerol, 500 mM NaCl, 0.5 mg/ml lysozyme and 0.1 mg/ml PMSF), and incubated at 37°C for 30 min. The soluble fraction of lysate was collected after centrifugation and applied to a TALON metal affinity Co2+ column (Clontech). The column was washed twice with washing buffer (20 mM Tris-HCl, pH 8.0, 10% glycerol, 500 mM NaCl) without imidazole, then with 20 mM imidazole. The protein was eluted with the same buffer containing 300 mM imidazole. The fractions with the most DosR protein (as determined by SDS-PAGE) were pooled together for dialysis in 25 mM Tris-HCl, pH 8.0. The final protein concentration was determined using a Qubit kit (Invitrogen).
**Electrophoretic mobility assay.** The assay is fluorescence-based using 6-carboxyfluorescein (6FAM) labeled 385 bp probe from the *hspX* promoter. In designing the primer set, 6FAM was added to the 5' ends of forward and reverse primers. The *hspX* probe was synthesized via PCR using the primer set: forward primer 5'-6FAM-CAACTGCACCGCGCTCTTGATG-3'; reverse primer 5'-6FAM-CATCTCGTCTTCCAGCGCATCAAC-3'. The probe was purified by Qiagen PCR purification kit. The DosR protein was pre-phosphorylated in 10 µL of phosphorylation buffer (40 mM Tris-HCl, pH 8.0, 5 mM MgCl₂, 50 mM lithium potassium acetyl phosphate), and incubated at room temperature for 30 min. The protein was then transferred to binding buffer in a final volume of 20 µL (final concentration, 25 mM Tris-HCl, pH 8.0, 0.5 mM EDTA, 20 mM KCl, 6 mM MgCl₂, 10 nM probe, 1 µg poly-dl-dC (Sigma Aldrich)), and treated with HC104A or an equal volume of DMSO or virstatin (Santa Cruz Biotech). Two different assays were performed. Firstly, different DosR protein concentrations from 0.5 µM to 4 µM were treated with 40 µM. Second, dose response assays were performed with 2 µM DosR treated with different concentrations of HC104A or virstatin from 1 µM to 80 µM. After incubating on ice for 30 min, the reactions were terminated by adding 1 µL 80% glycerol, and loaded on a native 5% Tris/Borate/EDTA (TBE) polyacrylamide gel. The gel was run at 50 V, for 5-6 h at 4°C in 1X TBE buffer, and was imaged using a Typhoon scanner with appropriate filters that can detect florescence at excitation = 495 nm, emission = 520 nm. Binding of the unbound probe was quantified using ImageJ (43). The assay was repeated at least twice with similar results. The error bar represents the s.d. of two biological replicates.

**UV-visible spectroscopy assay.** DosS and mutant proteins were purified and analyzed as previously described (27). Briefly, 7.5 µM of recombinant DosS protein was deoxygenated with argon gas in a sealed cuvette. The protein was reduced with 400 µM DTN for 20 min. The reaction was then treated with 100 µM HC106A, 400 µM HC104A, 100 µM CORM-2...
(tricarbonyldichlororuthenium (II) dimer) or equal volume of DMSO. The UV-visible spectra were recorded for kinetic changes over 2 h. The experiment was repeated at least twice with similar results.

**Checkerboard synergy studies.** The reporter strain CDC1551 (hspX::GFP) was treated with pairs of DosRST inhibitors from 50 µM – 0.08 µM in 96-well plates, including HC101A-HC104A and HC106A. GFP fluorescence and OD$_{600}$ were measured after 6 d incubation. The percentage of fluorescence inhibition (FI) and growth inhibition were calculated for each drug pair, with limited growth inhibition observed. The FI data was utilized for further analysis of interactions using CompuSyn software (34). The Combination Index (CI) value was calculated for each drug pair according to the Chou-Talalay method, which is based on the Median-Effect equation derived from the Mass-Action Law principle (35, 46). The resulting CI values provide quantitative determination of drug interactions, including synergism (CI < 1), additive effect (C = 1), and antagonism (C > 1).

**Autophosphorylation assay.** The DosS autophosphorylation assay was performed as previously described (27). Recombinant DosS protein was treated with 10 µM, 20 µM or 40 µM of HC104A, or HC106A. DMSO and 40 µM HC103A were also included as positive and negative controls, respectively.

**Kinetic solubility assay.** The assay was performed with 7-point (2-fold) dilutions from 200 µM - 3.125 µM for HC106 analogs. Mebendazole, benxarotene and aspirin were also included as controls. The drug dilutions were added to PBS, pH 7.4, with the final DMSO concentration of 1%, and incubated at 37 °C for 2 h. The absorbance at 620 nm was measured for each drug.
dilution to estimate of the compound solubility. Three replicates were examined for each dilution.

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Author Contributions

H.Z., E.E. and R.B.A conceived the experiments. H.Z. performed the Mtb physiology and biochemistry experiments. B.A. synthesized the analogs. E.E. directed the medicinal chemistry optimizations. H.Z, E.E. and R.B.A wrote the manuscript.

Disclosures

RBA is the founder and owner of Tarn Biosciences, Inc., a company that is working to develop new TB drugs.
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Figure Legends

Figure 1. HC104A and HC106A inhibit DosRST reporter fluorescence. (a) Chemical structures of HC104A and HC106A. HC104A (b) and HC106A (c) inhibited DosR-driven GFP fluorescence signal in a dose-dependent manner, while having minimal impact of Mtb growth. The EC_{50} values of fluorescence inhibition for HC104A and HC106A are 9.8 µM and 2.5 µM, respectively.

Figure 2. Transcriptional profiling shows HC104A and HC106A inhibited DosR regulon during hypoxia. Magnitude-amplitude plots showing differential gene expression of Mtb cells treated with 40 µM HC104A (a) or HC106A (b). The labeled genes represent selected genes that belong to the DosRST regulon. The red dots represent genes with significant differential expression, q<0.05. (c) A Venn diagram for the downregulated genes (>2-fold; q< 0.05) of WT CDC1551 treated with HC104A or HC106A compared to that of CDC1551 (ΔdosR). (d) Venn diagram for downregulated genes (>2-fold; q< 0.05) of CDC1551 (ΔdosR) treated with HC104A or HC106A.

Figure 3. Inhibition of DosR regulon and persistence-associated physiologies by HC104A and HC106A. (a) Inhibition of DosR regulon in murine macrophages infected with Mtb and treated with HC104A and HC106A for 48 h. Bacterial RNA was isolated after incubation, and the differential gene expression of hspX and tgs1 bone-marrow derived were quantified by qRT-PCR. The error bars represent the standard derivation of three biological replicates. (b) HC104A and HC106A inhibit DosR regulon induction by vitamin C and NO. Cells were pretreated with the compounds or DMSO for 24 h, and induced with vitamin C or NO for 2 h. Total bacterial RNA was isolated, and the transcripts of DosR-regulated genes, hspX and tgs1, were quantified by qRT-PCR. The difference in the drug treated samples compared to DMSO treated samples in response to vitamin C or NO are significant with a p-value <0.001 based on a t-test, except
the one marked as non-significant (n.s.). The error bars represent the standard deviation of three replicates. The experiment was repeated twice with similar results. (c) Inhibition of TAG accumulation of Mtb treated with HC104A or HC106A. Mtb cells were treated with 40 µM of the compounds and labeled with [1,2-14C] sodium acetate for 6 d. Total lipid was isolated and analyzed by TLC. TAG accumulation was quantified from the TLC. The error bars represent the standard derivation of two biological replicates. (d) Mtb cell survival during NRP when treated with HC104A or HC106A during NRP. Mtb cells were pretreated with 40 µM of compounds for 48 h in an anaerobic chamber, and continued incubation for 10 d. Surviving bacteria were enumerated on 7H10 agar. The error bars represent the standard derivation of three biological replicates. The experiment was repeated twice with similar results.

Figure 4. Inhibition of DosR DNA-binding by HC104A. (a) DosR protein ranging from 0.5 µM to 4 µM was treated with DMSO or 40 µM HC104A and binding to the hspX promoter was examined by EMSA. HC104A inhibits DosR DNA binding at 2 µM concentration. (b) The free DNA of each reaction was quantified in ImageJ and the percentage of free DNA was normalized using reactions containing 0.5 µM DosR as 100% free DNA. Differences between reactions containing 2 µM DosR treated with DMSO or HC104A are significant (**P value <0.005 based on a t-test). The error bars represent the standard derivation of two biological replicates. (c) Dose-dependent impact of HC104A on DosR DNA binding. DosR protein at 2 µM was treated with HC104A at concentrations from 1 µM to 80 µM. (c) The free DNA of each reaction was also quantified in ImageJ with the percentage of free DNA is normalized using the reaction containing 80 µM HC104A as 100% free DNA. The differences between treated reactions as compared to DMSO control are significant. (*P value <0.05 and **P value <0.005 based on a t-test). The error bars represent the standard derivation of two biological reps.
Figure 5. Interactions between HC106A and DosS heme. WT DosS protein was treated with dithionite (DTN) and then 100 μM HC106A (a) or 100 μM pf CORM-2 (a CO donor) (b). The UV-visible spectra of the two treatments exhibited a shift of the Soret peak to a common position of 422 nm. (c) DosS with a G117L amino acid substitution, that is predicted to block the heme exposing channel, provides resistance to HC106A. The spectrum of the mutant protein did not change, after HC106A treatment, indicating resistance to HC106A. (d) Overexpression of DosS protein promotes resistance to HC106A treatment in Mtb. Mtb cells with the pVV16 empty vector or the DosS overexpression plasmid were treated with 20 μM HC106A for 6 d. Bacterial RNA was isolated for analysis of the differential gene expression of hspX and tgs1 and analyzed by qRT-PCR. Overexpression of DosS caused 23- and 16.5-fold increase of hspX and tgs1 transcripts, respectively, compared to the empty vector control (***P value <0.0001 based on a t-test). The error bar represents the standard derivation of the mean for three technical replicates. The experiments were repeated twice with a similar result.

Figure 6. Synergistic interactions between DosRST inhibitors. CDC1551 (hspX::GFP) was treated with pairwise combinations of two compounds at concentrations of 50 μM to 0.08 μM. GFP fluorescence was measured and used to calculate percentage inhibition. The data were analyzed in the CompuSyn software to determine the combination index (CI) for the panel of each drug combination, including (a) artemisinin and HC102A; (b) artemisinin and HC103A; (c) artemisinin and HC104A; and (d) artemisinin and HC106A. Example EC50 curves are presented with individual compounds or a selected synergistic combination to illustrate the potentiating interactions.

Table 1. Initial SAR studies of the HC106 series. The HC106 analogs with different R-groups were synthesized or purchased. The reporter strain CDC1551 (hspX::GFP) was
treated with across doses of each analog from 200 µM to 0.328 µM. The EC$_{50}$ values of fluorescence inhibition calculated for each analog to determine their potency.

Table 2. Early SAR studies of “A-ring” analogs of HC106. The HC106 analogs with different R-groups were synthesized. The reporter strain CDC1551 (hspX‘::GFP) was treated with across doses of each analog from 200 µM to 0.328 µM. The EC$_{50}$ values of fluorescence inhibition were calculated for each analog to determine their potency. The other chemical properties of the analogs are also included. Kinetic solubility and mouse microsomal stability were both experimentally determined.
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6
| ID#   | Compound | EC$_{50}$ (µM) | ID#   | Compound | EC$_{50}$ (µM) |
|-------|----------|----------------|-------|----------|----------------|
| HC106F | ![Compound](image1) | 1.33          | MSU-41324 | ![Compound](image2) | >200          |
| HC106A | ![Compound](image3) | 2.48          | MSU-39449 | ![Compound](image4) | 5.2           |
| HC106C | ![Compound](image5) | >200          | MSU-39451 | ![Compound](image6) | 4.14          |
| MSU-39450 | ![Compound](image7) | 1.95          | MSU-39453 | ![Compound](image8) | 16.62         |
| MSU-39444 | ![Compound](image9) | 1.7           | MSU-41422 | ![Compound](image10) | >200          |
| MSU-41425 | ![Compound](image11) | >200          |        |           |                |

Table 1
| ID#     | Compound R = | EC<sub>50</sub> (µM) | MW (g/mol) | TPSA (Å<sup>2</sup>) | Ligand Efficiency (LE) | m.p. (°C) | cLogP | Solubility (µM) @ 30 minutes | Microsomes (% remaining) |
|---------|--------------|------------------------|------------|----------------------|------------------------|-----------|-------|-------------------------------|---------------------------|
| HC106F  |              | 1.33                   | 237.6      | 63                   | 0.50                   | 183       | 2.2   | >200                          | 115%                      |
| HC106A  |              | 2.48                   | 272.1      | 63                   | 0.41                   | 164       | 2.7   | 110                           | -                         |
| MSU-33189 |            | 0.63                   | 203.1      | 63                   | 0.56                   | 183-184   | 1.6   | -                             | -                         |
| MSU-39447 |            | 0.61                   | 233.2      | 72                   | 0.49                   | 166-167   | 1.5   | >100                          | -                         |
| MSU-39448 |            | 1.42                   | 238.6      | 75                   | 0.49                   | 158-159   | 1.3   | -                             | -                         |
| MSU-39450 |            | 1.95                   | 238.6      | 75                   | 0.38                   | 122-123   | 1.3   | -                             | -                         |
| MSU-39446 |            | 0.54                   | 221.2      | 63                   | 0.52                   | 176-177   | 1.7   | >100                          | 70%                       |
| MSU-41464 |            | 1.2                    | 282.1      | 67                   | 0.51                   | 183-184   | 2.3   | -                             | 109%                      |
| MSU-39445 |            | 0.75                   | 237.6      | 63                   | 0.52                   | 159 (dec) | 2.1   | >100                          | -                         |
| MSU-39452 |            | 2.47                   | 272.1      | 63                   | 0.45                   | 164       | 2.8   | -                             | -                         |
| MSU-41442 |            | 2.08                   | 259.3      | 63                   | 0.41                   | 163       | 3.1   | >100                          | -                         |
| MSU-41443 |            | 11.2                   | 279.3      | 63                   | 0.32                   | 193       | 3.2   | 14                            | -                         |
| MSU-41463 |            | 1.67                   | 271.2      | 67                   | 0.42                   | 178       | 2.4   | -                             | 84                        |
| MSU-41465 |            | 1.16                   | 261.2      | 94                   | 0.43                   | 194       | 1.5   | -                             | 73                        |
| MSU-41462 |            | 1.12                   | 217.2      | 67                   | 0.45                   | 128-130   | 1.3   | -                             | -                         |
| MSU-41542 |            | 1.34                   | 183.2      | 67                   | 0.62                   | 187       | 0.9   | -                             | -                         |
| MSU-41545 |            | 4.75                   | 309.3      | 76                   | 0.32                   | 164       | 2.9   | -                             | -                         |
| MSU-41546 |            | 2.15                   | 195.2      | 67                   | 0.56                   | 182       | 1.1   | -                             | -                         |
| MSU-42002 |            | 1.21                   | 209.2      | 67                   | 0.54                   | 180       | 1.4   | -                             | -                         |
| MSU-42003 |            | 1.94                   | 282.1      | 67                   | 0.49                   | 157       | 2.3   | -                             | -                         |
| MSU-42004 |            | 2.36                   | 231.3      | 67                   | 0.45                   | 165       | 1.7   | -                             | -                         |

Table 2
Supplementary Methods and Figures

Inhibition of Mycobacterium tuberculosis DosRST two-component regulatory system signaling by targeting response regulator DNA binding and sensor kinase heme

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Supplemental Methods

Experimental procedures for urea formation

Formation of acyl chloride 2. To a stirred solution of isoxazole acid 1 (1 eq.) in dry tetrahydrofuran (THF, 0.4 M) under N₂ atmosphere was added oxalyl chloride (1.5 eq.) dropwise over 5-10 min followed by dimethylformamide (DMF) (cat.) and the reaction mixture was continued to stir at room temperature. Upon completion, the reaction mixture was concentrated into a residue in vacuo and the residue was dissolved in THF and concentrated again to ensure the removal of excess oxalyl chloride. The crude acyl chloride 2 was used directly in the next step without further purification.

Formation of acyl azide and rearrangement into isocyanate 3.

The crude acyl chloride 2 was dissolved in THF (0.4 M) and stirred at room temperature under N₂ atmosphere. Trimethylsilyl (TMS) azide (2 eq.) was added dropwise over 5 min and stirring was continued. Upon completion of the reaction, the mixture was diluted with ethyl acetate (0.4 M) and quenched with H₂O (0.4 M). The two layers were separated, and the organic layer was dried over anhydrous Na₂SO₄ and filtered. The ethyl acetate solvent was swapped into toluene (0.1 M) by the addition of toluene followed by removal of the ethyl acetate in vacuo. Care was taken not to concentrate the toluene. The toluene acyl azide solution was heated at reflux conditions under N₂ atmosphere for 4 h to give the desired isocyanate 3 which was used as a solution in toluene in the next step.

Formation of urea 4. The crude isocyanate solution in toluene was mixed with different amines (1.5 eq.) and stirred at room temperature overnight. Isolation of the ureas was done by diluting the reaction mixture with hexanes, stirring for few hours and filtration of the formed precipitate. The solid material was washed with hexanes and dried under high vacuum. The urea products usually do not require further purifications. All products were analyzed by ¹H NMR and high-resolution mass spectrometry (HRMS).
Synthesis of **MSU-41422** (amide) from acyl chloride 2. Acyl chloride 2 (1 eq.) was dissolved in dichloromethane (DCM, 0.2 M) and 4-chloroaniline (1.2 eq.) was added. The reaction mixture was stirred at room temperature overnight. The reaction mixture was concentrated in vacuo and the residue was purified by flash column chromatography.

**Synthesis of MSU-39449.** A slurry of 4-chlorophenyl acetic acid (0.10 g, 0.60 mmoles) in 2.0 mL of DCM was treated with oxalyl chloride (0.12 mL, 0.9 mmoles) and 1 drop of DMF. The mixture (with gas evolution) gradually became homogeneous and was stirred for 30 min. The mixture was concentrated in vacuo, diluted with 5 mL of DCM and again concentrated in vacuo, the process repeated three times. The resulting residue was again dissolved in DCM (2 mL) and treated with 5-aminoisoxazole (0.030 g., 0.36 mmoles), followed by pyridine (0.48 mL, 0.6 mmoles). The mixture was then allowed to stir overnight. The mixture was then quenched with 1.0 N HCl and extracted with DCM. The organic layers were combined, washed with saturated KHCO₃, dried with Na₂SO₄ and concentrated in vacuo. Medium pressure liquid chromatography (SiO₂, 100% DCM to 3% methanol / DCM) to provide a solid (0.023 g).

**¹H NMR and HRMS data:**

**MSU# 39444.** ¹H NMR (500 MHz, DMSO-d₆) δ 10.54 (s, 2H), 8.44 (d, J = 1.9 Hz, 2H), 6.11 (d, J = 2.0 Hz, 2H). HRMS (ESI) m/z calculated for C₇H₆N₄O₃ [M+H], 195.0513 found 195.0518.

**MSU# 39449.** ¹H NMR (500 MHz, DMSO-d₆) δ 11.91 (s, 1H), 8.41 (s, 1H), 7.39 (d, J = 2.0 Hz, 2H), 7.35 (d, J = 2.0 Hz, 2H), 6.19 (s, 1H), 3.73 (s, 2H). HRMS (ESI) m/z calculated for C₁₁H₁₀ClN₂O₂ [M-H], 235.0269; found 235.1989.

**MSU# 39445.** ¹H NMR (500 MHz, DMSO-d₆) δ 10.41 (s, 1H), 9.14 (s, 1H), 8.40 (d, J = 2.0 Hz, 1H), 7.82 – 7.58 (m, 1H), 7.31 (dd, J = 4.9, 1.8 Hz, 2H), 7.07 (dt, J = 6.4, 2.3 Hz, 1H), 6.06 (d, J = 1.9 Hz, 1H). HRMS (ESI) m/z calculated for C₁₀H₉ClN₃O₂ [M+H], 238.0378; found 238.0365.
**MSU# 39452.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 10.50 (s, 1H), 9.23 (s, 1H), 8.51 – 8.28 (m, 1H), 7.86 (d, \(J = 2.5\) Hz, 1H), 7.64 – 7.43 (m, 1H), 7.37 (dd, \(J = 8.9, 2.5\) Hz, 1H), 6.26 – 5.80 (m, 1H).

HRMS (ESI) m/z calculated for C\(_{10}\)H\(_8\)ClN\(_3\)O\(_2\) [M+H], 271.9989; found 271.9969.

**MSU# 39447.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 8.83 (s, 1H), 8.43 – 8.29 (m, 1H), 7.36 (d, \(J = 8.4\) Hz, 2H), 6.86 (d, \(J = 8.6\) Hz, 2H), 6.00 (d, \(J = 1.9\) Hz, 1H), 3.70 (s, 3H). HRMS (ESI) m/z calculated for C\(_{11}\)H\(_{12}\)N\(_3\)O\(_3\) [M+H], 234.0874; found 234.0863.

**MSU# 39451.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 10.13 (s, 1H), 8.34 (d, \(J = 1.9\) Hz, 1H), 7.56 – 7.41 (m, 2H), 7.41 – 7.22 (m, 2H), 6.04 (d, \(J = 2.0\) Hz, 1H), 3.25 (s, 3H). HRMS (ESI) m/z calculated for C\(_{11}\)H\(_{11}\)ClN\(_3\)O\(_2\) [M+H], 252.0535; found 252.0596.

**MSU# 39453.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 10.49 (s, 1H), 8.41 (d, \(J = 1.9\) Hz, 1H), 7.84 (d, \(J = 8.7\) Hz, 1H), 7.27 (dt, \(J = 2.3, 1.1\) Hz, 1H), 7.18 (dt, \(J = 8.8, 1.6\) Hz, 1H), 6.14 (d, \(J = 1.9\) Hz, 1H), 4.13 (dd, \(J = 9.1, 8.2\) Hz, 2H), 3.17 (t, \(J = 8.6\) Hz, 2H). HRMS (ESI) m/z calculated for C\(_{12}\)H\(_{11}\)ClN\(_3\)O\(_2\) [M+H], 264.0535; found 264.0552.

**MSU# 39448.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 10.55 (s, 1H), 9.25 (s, 1H), 8.64 – 8.24 (m, 2H), 7.98 (d, \(J = 8.6\) Hz, 1H), 7.45 (d, \(J = 8.6\) Hz, 1H), 6.08 (d, \(J = 18.1\) Hz, 1H). HRMS (ESI) m/z calculated for C\(_{9}\)H\(_8\)ClN\(_4\)O\(_2\) [M+H], 239.0331; found 239.0364.

**MSU# 39450.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 11.00 (s, 1H), 9.66 (s, 1H), 8.42 (d, \(J = 1.9\) Hz, 1H), 8.34 (d, \(J = 2.6\) Hz, 1H), 7.96 – 7.79 (m, 1H), 7.73 (d, \(J = 8.9\) Hz, 1H), 6.10 (q, \(J = 2.7, 2.2\) Hz, 2H). HRMS (ESI) m/z calculated for C\(_9\)H\(_8\)ClN\(_2\)O\(_2\) [M+H], 239.0331; found 239.0356.

**MSU# 39446.** \(^1\)H NMR (500 MHz, DMSO-\(d_6\)) \(\delta\) 10.24 (s, 1H), 8.26 (s, 1H), 7.54 – 7.37 (m, 2H), 7.26 – 7.02 (m, 2H), 6.03 (d, \(J = 2.0\) Hz, 1H). HRMS (ESI) m/z calculated for C\(_{10}\)H\(_8\)F\(_2\)N\(_3\)O\(_2\) [M+H], 222.0674; found 222.0675.

**MSU# 41422.** \(^1\)H NMR (500 MHz, Chloroform-\(d\)) \(\delta\) 8.41 (d, \(J = 1.8\) Hz, 1H), 8.26 (s, 1H), 7.71 – 7.52 (m, 2H), 7.44 – 7.33 (m, 2H), 7.06 (d, \(J = 1.8\) Hz, 1H). HRMS (ESI) m/z calculated for C\(_{10}\)H\(_8\)ClN\(_2\)O\(_2\) [M+H], 223.0269; found 223.0265.
\textbf{MSU# 41324.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 9.91 (s, 1H), 9.50 (s, 1H), 8.31 (d, $J = 2.6$ Hz, 1H), 7.85 (dd, $J = 9.0$, 2.7 Hz, 1H), 7.67 (d, $J = 8.9$ Hz, 1H), 7.39 – 7.30 (m, 2H). HRMS (ESI) m/z calculated for C$_{12}$H$_{10}$Cl$_2$N$_3$O [M+H], 282.0196; found 282.0182.

\textbf{MSU# 41425.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 8.83 (s, 1H), 7.53 – 7.39 (m, 2H), 7.39 – 7.16 (m, 2H). HRMS (ESI) m/z calculated for C$_{13}$H$_{11}$Cl$_2$N$_2$O [M+H], 281.0243; found 281.0258.

\textbf{MSU# 41443.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.36 (d, $J = 28.4$ Hz, 1H), 9.13 (d, $J = 45.7$ Hz, 1H), 8.40 (d, $J = 1.9$ Hz, 1H), 7.70 – 7.59 (m, 5H), 7.59 – 7.49 (m, 3H), 7.43 (t, $J = 7.7$ Hz, 3H), 7.32 (t, $J = 7.4$ Hz, 1H), 6.06 (d, $J = 1.9$ Hz, 1H). HRMS (ESI) m/z calculated for C$_{16}$H$_{14}$N$_3$O$_2$ [M+H], 280.1081; found 280.1083.

\textbf{MSU# 41442.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.15 (s, 1H), 8.77 (s, 1H), 8.38 (d, $J = 1.9$ Hz, 1H), 7.44 – 7.27 (m, 4H), 6.03 (d, $J = 2.0$ Hz, 1H), 1.25 (s, 9H). HRMS (ESI) m/z calculated for C$_{14}$H$_{18}$N$_3$O$_2$ [M+H], 260.1394; found 260.1406.

\textbf{MSU# 33189.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.20 (s, 1H), 8.85 (s, 1H), 8.39 (d, $J = 1.9$ Hz, 1H), 7.54 – 7.37 (m, 2H), 7.35 – 7.21 (m, 2H), 7.02 (t, $J = 7.4$ Hz, 1H), 6.04 (d, $J = 1.9$ Hz, 1H). HRMS (ESI) m/z calculated for C$_{10}$H$_{10}$N$_3$O$_2$ [M+H], 204.0768; found 204.0777.

\textbf{MSU# 33231.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.29 (s, 1H), 9.01 (s, 1H), 8.39 (d, $J = 1.9$ Hz, 1H), 7.56 – 7.41 (m, 2H), 7.41 – 7.27 (m, 2H), 6.05 (d, $J = 1.9$ Hz, 1H). HRMS (ESI) m/z calculated for C$_{10}$H$_{9}$ClN$_3$O$_2$ [M+H], 238.0378; found 238.0391.

\textbf{MSU# 41462.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.18 (s, 1H), 8.31 (d, $J = 2.0$ Hz, 1H), 7.43 – 7.07 (m, 3H), 6.92 (s, 1H), 5.94 (d, $J = 1.9$ Hz, 1H), 4.30 (d, $J = 6.0$ Hz, 2H). HRMS (ESI) m/z calculated for C$_{11}$H$_{12}$N$_3$O$_2$ [M+H], 218.0924 found 218.0956.

\textbf{MSU# 41463.} $^1$H NMR (500 MHz, DMSO-d$_6$) $\delta$ 10.39 (s, 1H), 9.29 (s, 1H), 8.41 (d, $J = 1.9$ Hz, 1H), 7.67 (d, $J = 1.0$ Hz, 4H), 6.08 (d, $J = 1.9$ Hz, 1H). HRMS (ESI) m/z calculated for C$_{11}$H$_{9}$F$_3$N$_3$O$_2$ [M+H], 272.0642; found 272.0653.
**MSU# 41464.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.29 (s, 1H), 9.01 (s, 1H), 8.39 (d, $J = 1.9$ Hz, 2H), 7.50 – 7.45 (m, 1H), 7.45 – 7.40 (m, 1H), 6.05 (s, 1H). HRMS (ESI) m/z calculated for C$_{10}$H$_9$BrN$_3$O$_2$ [M+H], 281.9873; found 281.9876.

**MSU# 41465.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.38 (s, 1H), 9.27 (s, 1H), 8.41 (d, $J = 2.0$ Hz, 1H), 8.00 – 7.79 (m, 2H), 7.66 – 7.48 (m, 2H), 6.08 (d, $J = 1.9$ Hz, 1H), 3.81 (s, 3H). HRMS (ESI) m/z calculated for C$_{12}$H$_{12}$N$_3$O$_4$ [M+H], 262.0823; found 262.0827.

**MSU# 41545.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.17 (s, 1H), 8.71 (s, 1H), 8.37 (d, $J = 1.9$ Hz, 1H), 7.57 – 7.14 (m, 5H), 7.11 – 6.69 (m, 2H), 6.02 (d, $J = 1.9$ Hz, 1H), 5.06 (s, 2H). HRMS (ESI) m/z calculated for C$_{17}$H$_{16}$N$_3$O$_3$ [M+H], 310.1187; found 310.1178.

**MSU# 41546.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 9.69 (s, 1H), 8.30 (d, $J = 1.9$ Hz, 1H), 6.42 (d, $J = 7.2$ Hz, 1H), 5.91 (d, $J = 1.9$ Hz, 1H), 3.92 (h, $J = 6.7$ Hz, 1H), 1.83 (dq, $J = 12.8$, 6.6, 6.0 Hz, 3H), 1.73 – 1.46 (m, 5H), 1.46 – 1.13 (m, 3H). HRMS (ESI) m/z calculated for C$_9$H$_{14}$N$_3$O$_2$ [M+H], 196.1081; found 196.1144.

**MSU# 42002.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 9.80 (s, 1H), 8.30 (d, $J = 1.9$ Hz, 1H), 6.34 (d, $J = 7.8$ Hz, 1H), 5.90 (d, $J = 1.9$ Hz, 1H), 3.58 – 3.39 (m, 1H), 1.77 (dt, $J = 11.1$, 3.7 Hz, 1H), 1.64 (dt, $J = 12.9$, 4.1 Hz, 1H), 1.52 (dd, $J = 10.4$, 6.3 Hz, 1H), 1.37 – 0.93 (m, 3H). HRMS (ESI) m/z calculated for C$_{10}$H$_{16}$N$_3$O$_2$ [M+H], 210.1238; found 210.1282.

**MSU# 42004.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.07 (s, 2H), 8.31 (d, $J = 2.0$ Hz, 1H), 7.37 – 7.06 (m, 5H), 6.40 (s, 1H), 5.93 (d, $J = 1.8$ Hz, 1H), 3.31 (t, $J = 7.2$ Hz, 2H), 2.74 (t, $J = 7.2$ Hz, 2H). HRMS (ESI) m/z calculated for C$_{12}$H$_{14}$N$_3$O$_2$ [M+H], 232.1081; found 232.1105.

**MSU# 42003.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.36 (s, 1H), 9.06 (s, 1H), 8.40 (d, $J = 1.9$ Hz, 1H), 7.83 (t, $J = 2.0$ Hz, 1H), 7.46 – 7.03 (m, 3H), 6.07 (d, $J = 2.0$ Hz, 1H). HRMS (ESI) m/z calculated for C$_{10}$H$_9$BrN$_3$O$_2$ [M+H], 281.9873; found 281.9915.

**MSU# 41542.** $^1$H NMR (500 MHz, DMSO-$d_6$) $\delta$ 10.20 – 9.49 (m, 1H), 8.30 (d, $J = 1.9$ Hz, 1H), 6.44 (t, $J = 5.9$ Hz, 1H), 5.91 (d, $J = 1.9$ Hz, 1H), 2.93 (t, $J = 6.3$ Hz, 2H), 1.68 (dh, $J = 13.3$, 6.7
Hz, 1H), 0.85 (d, J = 6.7 Hz, 6H). HRMS (ESI) m/z calculated for C₈H₁₄N₃O₂ [M+Na], 206.0899; found 206.0926.

Supplementary Dataset 1. Differential gene expression data of WT Mtb treated with inhibitors

Supplementary Dataset 2. Differential gene expression data of the DosR mutant treated with the inhibitors.

Supplementary Dataset 3. Complete gene expression tables for transcriptional profiling experiments.

Supplemental Table 1 and Figures 1-9
Supplemental Table 1. Structure and properties of HC104 series. Catalog structure activity relationship study performed for HC104 analogs with different R-groups. The reporter strain CDC1551 (hspX':::GFP) was treated across doses of each analog from 200 µM to 0.328 µM. The EC50 values of fluorescence inhibition calculated for each analog to determine their potency. The other chemical properties of the analogs are also included.

| Compound | ID#   | MW (g/mole) | EC50 (µM) | Ligand Efficiency | CLogP (cLogD at pH 7.4) | Druglikeness |
|----------|-------|-------------|-----------|-------------------|-------------------------|--------------|
|          | HC104A| 361.2       | 9.8       | 0.34              | 3.0 (0.55)              | 4.4          |
|          | HC104G| 290.1       | 43.8      | 0.36              | 2.3 (2.4)               | 0.55         |
|          | HC104B| 282.3       | >200      | 0.26              | 2.9 (-0.24)             | 6.2          |
Supplemental Figure 1. Inhibition of the DosR regulon by HC104 and HC106A during hypoxia. Mtb cells were treated with 40 µM compounds for 6 d, and total bacterial RNA was isolated. The DosR regulated genes, dosR, hspX, and tgs1 were quantified in qRT-PCR. The error bars represent the standard derivation of three replicates. The experiment was repeated at least twice with similar results.
Supplemental Figure 2. TLC of TAG reduction in Mtb treated with HC104A and HC106A. Mtb cells were treated with 40 µM of the compounds and labeled with [1,2-14C] sodium acetate in T25 vented tissue culture flasks for 6 d. Total lipid was isolated and analyzed BY TLC. The experiment was repeated twice with similar results.
Supplemental Figure 3. Autoradiograph examining the impact of HC104A and HC106A on DosS autophosphorylation. DosS protein was treated with 10 µM, 20 µM or 40 µM of the compounds, with DMSO and HC103A as positive and negative controls, respectively. The results show that HC104A and HC106A have no effect on DosS autophosphorylation.
Supplemental Figure 4. Investigation of interaction between HC104A and DosS. WT DosS treated with 400 µM HC104A shows no impact on shifting of the Soret peak in the UV-visible spectroscopy assay. The experiment was repeated at least twice with similar results.
Supplemental Figure 5. The impact of virstatin on DosR DNA-binding and DosRST signaling in Mtb. (a) Chemical structure of virstatin. (b) Dose-response curve of virstatin shows no effect on inhibition of Mtb DosR-driven GFP fluorescence. (c) DosR protein at 2 µM was treated with 9 point dose response of virstatin from 1 µM to 80 µM. The reactions were analyzed on native PAGE gel. The experiment was repeated at least twice with similar results.
Supplemental Figure 6. Investigating the interaction between HC106A and DosS heme. DosS E87L protein was treated with 100 µM HC106A after being reduced with DTN. The UV-visible spectra were recorded after each treatment, and showed no change on the overall spectrum compared to WT protein. The experiment was repeated at least twice with similar results.
Supplemental Figure 7. Checkerboard assays examining paired interactions of DosRST inhibitors. CDC1551 (hspX::GFP) was treated with different combination of two compounds from 50 µM to 0.08 µM. GFP fluorescence was measured and used to calculate percentage inhibition. The data were analyzed in the CompuSyn software to determine the combination index (CI) for the panel of each drug combination, including (a) artemisinin and HC104A; (b) HC102A and HC103A; (c) HC102A and HC104A; (d) HC102A and HC106A; (e) HC103A and HC104A; (f) HC103A and HC106A; (g) HC104A and HC106A. Selected dose response curves are presented to illustrate synergistic interactions. The experiment was repeated twice with similar results.
Supplemental Figure 8. Synthetic scheme for the HC106 analogs.
Supplemental Figure 9. Comparison between artemisinin, HC103 and HC106 for interactions. (a) Venn diagram for the downregulated genes (>2-fold; q< 0.05) of CDC1551 (ΔdosR) treated with artemisinin, HC103A, or HC106A. (b) Overlap exists between artemisinin and HC103A or HC103A and HC106A differentially expressed genes (downregulated >2X, q<0.05). The heatmaps represent the commonly downregulated genes between the two compounds.