A Novel Small-Molecule Inhibitor of the \textit{Mycobacterium tuberculosis} Demethylmenaquinone Methyltransferase MenG Is Bactericidal to Both Growing and Nutritionally Deprived Persister Cells

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ABSTRACT Active tuberculosis (TB) and latent \textit{Mycobacterium tuberculosis} infection both require lengthy treatments to achieve durable cures. This problem has partly been attributable to the existence of nonreplicating \textit{M. tuberculosis} "persisters" that are difficult to kill using conventional anti-TB treatments. Compounds that target the respiratory pathway have the potential to kill both replicating and persistent \textit{M. tuberculosis} and shorten TB treatment, as this pathway is essential in both metabolic states. We developed a novel respiratory pathway-specific whole-cell screen to identify new respiration inhibitors. This screen identified the biphenyl amide GSK1733953A (DG70) as a likely respiration inhibitor. DG70 inhibited both clinical drug-susceptible and drug-resistant \textit{M. tuberculosis} strains. Whole-genome sequencing of DG70-resistant colonies identified mutations in \textit{menG} (rv0558), which is responsible for the final step in menaquinone biosynthesis and required for respiration. Overexpression of \textit{menG} from wild-type and DG70-resistant isolates increased the DG70 MIC by 4× and 8× to 30×, respectively. Radio-labeling and high-resolution mass spectrometry studies confirmed that DG70 inhibited the final step in menaquinone biosynthesis. DG70 also inhibited oxygen utilization and ATP biosynthesis, which was reversed by external menaquinone supplementation. DG70 was bactericidal in actively replicating cultures and in a nutritionally deprived persistence model. DG70 was synergistic with the first-line TB drugs isoniazid, rifampin, and the respiratory inhibitor bedaquiline. The combination of DG70 and isoniazid completely sterilized cultures in the persistence model by day 10. These results suggest that MenG is a good therapeutic target and that compounds targeting MenG along with standard TB therapy have the potential to shorten TB treatment duration.

IMPORTANCE This study shows that MenG, which is responsible for the last enzymatic step in menaquinone biosynthesis, may be a good drug target for improving TB treatments. We describe the first small-molecule inhibitor (DG70) of \textit{Mycobacterium tuberculosis} MenG and show that DG70 has characteristics that are highly desir-
able for a new antitubercular agent, including bactericidality against both actively growing and nonreplicating mycobacteria and synergy with several first-line drugs that are currently used to treat TB.

Treatment of *Mycobacterium tuberculosis* has become increasingly challenging with the rise of drug-resistant strains. Existing treatment of drug-susceptible tuberculosis (TB) requires administration of multiple drugs for a minimum of 6 months; treatment of drug-resistant TB can extend to years and is often unsuccessful (1, 2). TB in humans probably includes several sub-populations of *M. tuberculosis* with differing metabolic states (3, 4). Some populations, for example, those that reside in well-aerated cavities, replicate quickly and are easily killed by a number of drugs, while other populations composed of nonreplicating persistent bacteria are much less drug susceptible (3, 5, 6). Persistent *M. tuberculosis* downregulates metabolic processes required for active growth such as cell wall biosynthesis, and these organisms become relatively or absolutely tolerant to drugs (5, 7–10). New anti-TB drugs with the ability to kill *M. tuberculosis* in both replicating and nonreplicating states could be used to treat drug-resistant TB and also potentially to reduce the required duration of treatment. The effectiveness of TB treatment could be further enhanced if new drugs were synergistic with current therapies (11).

Unlike many bacteria, *M. tuberculosis* cannot support its energy needs through substrate-level phosphorylation. Instead, both actively growing and nonreplicating persistent *M. tuberculosis* bacteria are dependent on respiration to synthesize adequate amounts of ATP (12). The *M. tuberculosis* respiratory chain consists of various electron donors that transfer two electrons to lipoquinone with the help of corresponding dehydrogenases. Lipoquinone then transfers these electrons to the oxidoreductases, which then reduce terminal electron acceptors (13). The electrochemical gradient generated in the process is utilized to synthesize ATP by F,F<sub>1</sub> ATP synthase (4). *M. tuberculosis* has the capacity to utilize numerous electron donors and acceptors depending on its microenvironment (13). However, menaquinone (MK9) and its saturated form [MK-9(H2)]1 are the only lipoquinones in *M. tuberculosis* that transfer electrons from dehydrogenases to the terminal electron oxidases (14, 15). Thus, menaquinone represents an essential vulnerable point in the electron transport chain and a prime target for development of new drugs. The absence of menaquinone biosynthesis in humans further supports the druggability of enzymes involved in this pathway. Menaquinone biosynthesis begins with chorismate derived from the shikimate pathway. The conversion of chorismate to isochorismate is the first committed step in menaquinone biosynthesis (16). Isochorismate is then converted to demethylmenaquinone by a cascade of at least 8 different enzymes (15). As the final step in menaquinone biosynthesis, MenG (Rv0558) catalyzes methylation of demethylmenaquinone (DMK9) using S-adenosylmethionine (SAM), resulting in formation of menaquinone (12).

Here, we searched for specific inhibitors of respiration in *M. tuberculosis* utilizing a novel respiratory pathway-specific whole-cell-based screen to test a library of known anti-TB compounds provided by GlaxoSmithKline (GSK) (17). We report a novel anti-TB chemotype, GSK1733953A (here renamed DG70), which was discovered through this screen; show that DG70 acts through inhibition of MenG; and investigate the effects and potential therapeutic utility of MenG inhibition.

**RESULTS**

**Identification of a biphenyl amide inhibitor from a compound library.** To identify novel inhibitors of mycobacterial respiration, we developed a whole-cell respiratory pathway-specific screen that combined the advantages of a whole-cell screen with those of a target-based approach (18). A *Mycobacterium bovis* BCG (BCG) reporter strain was created by fusing the putative promoter (PcydAB) for the *M. tuberculosis* cydAB operon to an mWasabi reporter. The cydAB operon has been shown to be upregulated in response to respiratory poisoning along with low oxygen tension during
the transition to anaerobiosis in *M. tuberculosis* (19, 20). We used this reporter strain to detect possible respiratory inhibitors that are also able to penetrate into live mycobacterial cells. This reporter strain was first validated by testing it against known respiratory inhibitors such as the NADH/succinate dehydrogenase inhibitor thioridazine (THZ), the ATP synthase inhibitor bedaquiline (BDQ), and PA824, which appears to have multiple targets, including the *M. tuberculosis* cytochrome *bc*$_1$ and cytochrome *bd* oxidase (21–23). Negative controls included drugs that inhibit or kill *M. tuberculosis* by unrelated mechanisms. Significant induction of the PcydAB-fused *mWasabi* reporter was seen only in cells treated with the respiratory inhibitors (Fig. 1A). We then screened a diverse library of 168 compounds already known to inhibit *M. tuberculosis* but where the mechanism of action was unknown (17). DG70, a biphenyl benzamide (Fig. 1B), was selected from a number of positive screening hits because it consistently induced the PcydAB promoter at least 1.7-fold over the dimethyl sulfoxide (DMSO) control ($P < 0.05$) (Fig. 1A). The induction was comparable to those with THZ, BDQ, and PA824 (21–23) (Fig. 1A) and showed a peak of 3.1-fold induction when a full range of DG70 concentrations were tested (see Fig. S1 in the supplemental material).

**DG70 is potent against *M. tuberculosis* complex.** We tested the activity of DG70 against a panel of clinical and laboratory drug-susceptible and drug-resistant *M. tuberculosis* strains. DG70 had a MIC of 4.8 μg/ml against *M. tuberculosis* H37Rv and 1.2 to 9.6 μg/ml against drug-resistant strains, suggesting a novel molecular target (Table 1). The MIC variation in clinical strains was similar to that observed among clinical drug-susceptible strains when tested with isoniazid (INH) and rifampin (RIF) (24). DG70 lacked activity against nontuberculous mycobacteria (NTM) and a panel of Gram-positive, Gram-negative, ESKAPE (*Enterococcus faecium*, *Staphylococcus aureus*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa*, and *Enterobacter* species), and surrogate select agent bacteria (Table S1). Consistent with these findings, an alignment of MenG amino acid sequences showed that NTM had only 64.5 to 88% identity to *M. tuberculosis* (Fig. S2), and molecular docking studies of DG70 in its predicted MenG binding sites identified significant steric clashes in the NTM MenG sequences studied (Text S1). Of the other bacteria tested, none of their MenG sequences had more than 40.6% amino acid identity to *M. tuberculosis* MenG. Cytotoxicity testing in Vero cells showed a 50% cytotoxic concentration (CC$_{50}$) of $>77$ μg/ml and a selectivity index (SI = CC$_{50}$/MIC for *M. tuberculosis*) of $>30$. DG70 also inhibited intracellular *M. tuberculosis* growth in J774A.1 macrophages at concentrations equivalent to its *in vitro* MIC (Fig. 2).
DG70 targets MenG in *M. tuberculosis*. We searched for the target of DG70 by selecting for DG70-resistant mutants in *M. tuberculosis* and BCG. Two *M. tuberculosis* isolates (70P3 and 70P7) that were 8- and 32-fold resistant to DG70, respectively, were whole genome sequenced. Each mutant had a unique single nucleotide polymorphism (SNP) in *menG* (*rv0558*), leading to amino acid substitutions F118L and V20A, respectively (Table 2). Targeted sequencing of *menG* in additional DG70-resistant *M. tuberculosis* and BCG mutants revealed other *menG* SNPs (Table 2). Moreover, docking results against homology models of MenG predicted that DG70 would interact favorably with

![Figure 2](image-url)
the four residues mutated in DG70-resistant \textit{M. tuberculosis} isolates. DG70 interacted with two to three of the residues mutated in DG70-resistant \textit{M. tuberculosis} in each top-ranked docked mode (Fig. 3A and B), and all four residues interacted with DG70 in the entire collection of top-ranked modes (Fig. 3C and D). Detailed descriptions of the

\begin{table}[h]
\centering
\caption{Single nucleotide polymorphisms detected in \textit{M. tuberculosis} and \textit{M. bovis} BCG spontaneous mutants resistant to DG70}
\begin{tabular}{llll}
\hline
Strain & DG70 MIC$_{90}$ (\mu g/ml) & Gene(s) mutated & Amino acid change \\
\hline
H37Rv & & & \\
70P3$^a$ & $>$80 & menG, sigL & V20A \\
70P7$^a$ & 40 & menG, ponA1 & F118L \\
DRM1 & $>$80 & menG & W75A \\
DRM7 & $>$80 & menG & S188A \\
\hline
BCG & & & \\
70B1 & 19.2 & menG & A60V \\
70B2 & 19.2 & menG & A60V \\
70B4 & 19.2 & menG & D7H \\
70R1 & 9.6 & menG & G99V \\
70R2 & 9.6 & menG & G99V \\
70R5 & 19.2 & menG & D7H \\
70S1 & $>$40 & menG & R43W \\
70S3 & 9.6 & menG & V136F \\
70S4 & 9.6 & menG & R190M \\
\hline
\end{tabular}
\end{table}

$^a$Whole-genome-sequenced strains.

\textbf{FIG 3} Predicted binding modes of DG70 with models of MenG. (A and B) Multitemplate-based homology model of MenG from MODELLER. (C and D) Model made by threading MenG onto the known class, architecture, topology, and homology domains in PsiPRED. (A and D) Ribbon representation of these two MenG models. The residues that mutate in \textit{M. tuberculosis} upon DG70 treatment are shown according to the Corey, Pauling, and Koltun scheme with blue carbon atoms, and residues that mutate in BCG are displayed with purple carbon atoms. Two representative docked modes of DG70 are presented in panels B and D as ball-and-stick models with dark green carbon atoms.
plausible binding modes and the sequence variability of these key positions are provided in Text S1. To further investigate whether MenG was targeted by DG70, we used the hsp60 promoter to overexpress menG in BCG on an integrative plasmid, overexpressing either wild-type menG or the DG70-resistant F118L (MenG-F118L) or V20A (MenG-V20A) menG mutants (Fig. 4). Compared to the strain containing empty vector, overexpressing wild-type menG conferred a moderate degree of DG70 resistance and overexpressing mutant menG conferred increased levels of resistance (Table 1). In fact, the strain overexpressing menG_F118L was completely resistant to even 20 \mu{g/mL} of DG70. These data strongly suggested that DG70 targets MenG in M. tuberculosis. Additionally, MenG has been shown to be essential in M. tuberculosis by Himar1-based transposon mutagenesis (25) and thus is a plausible drug target.

**DG70 inhibits menaquinone biosynthesis.** MenG is a demethylmenaquinone methyltransferase that catalyzes one of the last steps of the menaquinone (MK9) biosynthesis pathway by methylation of demethylmenaquinone (15, 25, 26). Consistent with the known function of MenG, we observed that supplementing DG70-treated cultures with 300 to 400 \mu{M} of the MK9 analog MK4 partially rescued DG70-treated cells (Table 1). To further investigate the effect of DG70 on menaquinone biosynthesis in whole cells, metabolic labeling was performed using L-[methyl-14C]methionine (16). De novo synthesis of menaquinone in M. tuberculosis was monitored by preparative thin-layer chromatography (TLC). Menaquinone [MK-9(II-H2)] was identified by comigration with an authentic MK9 standard. Cells treated with as little as 1 \times MIC of DG70 showed complete suppression of [14C]methyl incorporation into menaquinone while the untreated lane as well as INH- or BDQ-treated lanes showed no such effect, strongly suggesting that DG70 was specially inhibiting MenG (Fig. 5A and B). To confirm that the disappearing radiolabeled moiety was in fact menaquinone, an unlabeled fraction of neutral lipids from an untreated BCG culture was also chromatographed on the same TLC. We then eluted the unlabeled material with \( R_f \) corresponding to the radioactively labeled material of interest from the silica gel TLC plate and subjected it to high-resolution mass spectrometry (HRMS) with atmospheric pressure chemical ionization (APCI) as described previously (16). The resulting mass spectrum clearly showed the presence of a molecular ion peak with an \( m/z \) value of 787.6387 atomic mass units (amu), consistent with the predominant mycobacterial menaquinone [MK-9(II-H2)]. The fully unsaturated and less abundant form of menaquinone (MK9) with a peak at an \( m/z \) value of 785.6263 amu was also observed in the corresponding mass spectra. Even though BCG can be partially rescued from DG70 by the addition of 400 \mu{M} MK4, in the L-[methyl-14C]methionine incorporation assay (Fig. 5B, lane 4), external supplementation of MK4 did not circumvent inhibition of [14C]methyl incorporation in menaqui-
This suggested that although MK4 can physically substitute for MK9 and maintain the viability of mycobacteria treated with DG70, MK4 does not prevent inhibition of MK9 synthesis by DG70. To conclusively show that DG70 specifically inhibits MenG, we treated BCG with DG70 at 1/$\text{MIC}_{100}$, 5/$\text{MIC}_{100}$, and 10/$\text{MIC}_{100}$ in liquid medium for 3 days and then analyzed its neutral lipids through HRMS with APCI. The relative ratio of menaquinone [MK-9(II-H$_2$)] ($m/z$ 787.6387 amu) to demethylmenaquinone (DMK9) ($m/z$ 771.8202 amu) was at least 7-fold lower in DG70-treated cells at 1/$\text{MIC}_{100}$ than in untreated cells (Fig. 5C). In contrast, a decrease in the ratio of MK-9(II-H$_2$)/DMK9 was not observed when strain 70B1, a DG70-resistant BCG mutant, was treated with DG70 at up to 5/$\text{MIC}_{100}$ MIC of the parent strain. Finally, the ratio of MK-9(II-H$_2$)/DMK9 did decrease in strain 70B1 when it was treated with DG70 above its MIC, at 10/$\text{MIC}_{100}$ MIC of the parent strain (Fig. 5D).

**DG70 inhibits oxygen consumption and ATP synthesis.** Menaquinones are essential electron carriers in the respiratory chain of *M. tuberculosis*, and inhibitors of their biosynthesis would be expected to indirectly affect respiratory ATP synthesis and oxygen utilization. The effect on mycobacterial oxygen consumption was determined...
by monitoring methylene blue decolorization at 665 nm for 24 h in Parafilm-sealed culture tubes in the presence or absence of DG70. Decolorization of methylene blue indicates utilization of oxygen and a functional respiratory chain in bacteria (19). DG70 strongly prevented decolorization of methylene blue (Fig. 6A). Inhibition of oxygen consumption was comparable to that with the positive THZ-treated control, whereas INH, which does not inhibit respiration, as well as other negative controls, clearly decolorized methylene blue. Intracellular ATP was measured in BCG treated with DG70 or not, and a significant decrease in ATP levels was observed in DG70-treated cells compared to controls (Fig. 6B). Supplementation of the culture medium with 400 μM MK4 rescued the BCG cells from the effects of DG70 on oxygen consumption as well as ATP synthesis (Fig. 6A and B). However, external MK4 (400 μM) supplementation of cultures treated with THZ or BDQ failed to reverse the effect on oxygen consumption and ATP synthesis.

**DG70 is bactericidal against nutrient-deficient, persistent mycobacteria.** Under nutrient-deficient conditions, mycobacteria enter a nonreplicating state and have been previously shown to become tolerant to most drugs (27). However, respiratory ATP synthesis is essential to both actively growing and persistent mycobacteria (13). Furthermore, menaquinone levels have been shown to decrease in hypoxic cultures of *M. tuberculosis*, and MenA (the enzyme upstream of MenG in menaquinone biosynthesis) inhibition in BCG cultured under nutrient-deficient conditions leads to a 3-log₁₀ decrease in CFU (16, 28). These prior results suggested that nonreplicating mycobacteria might be particularly susceptible to MenG inhibition. We therefore tested the activity of DG70 on *M. tuberculosis* persisters using nutrient-deprived culture conditions as an *in vitro* model of the dormancy phenotype. In this model, DG70 resulted in 4-log₁₀ and 6.5-log₁₀ reductions of CFU after 7 days of treatment at 5× and 10× MIC, respectively (Fig. 7). This activity was far superior to those of all the other known anti-TB drugs tested. The activity of DG70 against a nonreplicating population of *M. tuberculosis* was also better than previously reported for other menaquinone biosynthesis inhibitors (16, 28, 29). Confirming the role of menaquinone inhibition in this model, the effect of DG70 was reversed by 400 μM MK4.

**DG70 interactions with known anti-TB compounds.** Single-drug TB treatment rapidly leads to resistance, and therefore, combination therapies are utilized (30). We performed synergy studies between DG70 and the first-line drugs INH, ethambutol (EMB), and RIF as well as known respiratory inhibitors BDQ, THZ, and PA824 to examine the *in vitro* effect of combined TB treatment. A checkerboard titration in microtiter plates was used as previously described (31). We observed strong synergy between DG70 and INH as well as between DG70 and RIF with ≤FIC (fractional inhibitory concentrations) of 0.4 and 0.5, respectively (Table 3; also see Fig. S4 in the supplemental material). We also examined the effect of treating cultures with DG70 and other
The combination of BDQ and DG70 showed promising synergy with a FIC of 0.5. The interactions of DG70 with THZ and PA824 were additive with FIC values of 0.7 and 1.0, respectively.

**Enhanced killing of BCG with DG70 in combination with BDQ, INH, or PA824.**

We studied whether DG70 was bactericidal and whether bactericidal activity could be increased by combining DG70 with other TB drugs. Log-phase \textit{M. tuberculosis} cultures were treated with DG70 and other antitubercular agents, both individually and in combination at $10 \times$ their MIC. DG70 was highly bactericidal against BCG \textit{in vitro}, leading to >5-log$_{10}$ killing over 21 days. This activity was comparable to that of BDQ alone and better than that of INH or PA824 alone, especially as both drugs showed regrowth after 7 and 10 days, respectively (Fig. 8A). The bactericidal activity of DG70 could be reversed by pretreating the mycobacterial cultures with supplemental MK4. Interestingly, combining DG70 with other respiratory drugs such as BDQ or PA824 resulted in enhanced killing.

**TABLE 3 Synergy between DG70 and antitubercular drugs**

| Drug combination | MIC (µM) | FIC | FIC index, category$^a$ |
|------------------|----------|-----|------------------------|
|                  | Alone    | In combination |       |
| BDQ              | 0.39     | 0.16          | 0.4    | 0.5, synergistic       |
| DG70             | 12.5     | 1.5           | 0.1    |                       |
| PA824            | 0.1      | 0.1           | 1.0    | 1.0, additive         |
| DG70             | 12.5     | 0.35          | 0.03   |                       |
| THZ              | 6.25     | 3.12          | 0.5    | 0.7, additive         |
| DG70             | 12.5     | 3.12          | 0.2    |                       |
| INH              | 0.39     | 0.09          | 0.2    | 0.4, synergistic      |
| DG70             | 12.5     | 3.12          | 0.2    |                       |
| RIF              | 0.02     | 0.005         | 0.25   | 0.5, synergistic      |
| DG70             | 12.5     | 3.1           | 0.25   |                       |
| EMB              | 6.25     | 3.12          | 0.5    | 0.6, additive         |
| DG70             | 12.5     | 1.5           | 0.12   |                       |

$^a$FIC categories: $\leq$0.5, synergistic; >0.5 to $\leq$1, additive; >1 to <4, no interaction; $\geq$4, antagonism.
resulted in more rapid killing, producing sterile cultures (no CFU detected on plating) after 21 and 10 days of treatment, respectively (Fig. 8C and D). However, the most rapid killing was seen in the INH-DG70 combination, where mycobacterial cultures were reduced by 5 log₁₀ CFU after only 2 days of treatment, compared to either INH or DG70 alone, where the same 2-day treatment reduced the cultures by only 1 log₁₀ or less (Fig. 8B). Remarkably, the combination of INH and DG70 sterilized cultures after only 10 days of incubation.

**Validation and initial profiling for DG70.** DG70 was independently synthesized, and this new preparation exhibited whole-cell efficacy and Vero cell cytotoxicity that was similar to the GlaxoSmithKline (GSK)-sourced material. The synthesis of DG70 and analogs (Table 4) centered around construction of the carboxylic acid and amine coupling partners, ultimately joined under Schotten-Baumann conditions (Text S2). The compound exhibited modest kinetic solubility in pH 7.4 aqueous phosphate-buffered saline (PBS) (S = 0.66 μg/ml) but poor stability in the presence of a mouse liver microsomal preparation (half-life [t₁/₂] = 0.753 min, CLint = 921 μl/min/mg protein). The major metabolite in this decomposition was determined to result from demethylation of both methyl ethers (Fig. S5). The MS data are consistent with the structure of bis(phenol) JSF-2950, which had significantly reduced whole-cell activity versus *M. tuberculosis* (MIC = 39 μg/ml). A focused set of DG70 analogs was prepared to further probe the antitubercular structure-activity relationships (SARs). Whole-cell efficacy depended on the presence of both methoxy groups in DG70 (cf. JSF-2907, -2971, and -2910), as well as the terminal methoxyphenyl moiety (cf. JSF-2926). The benzoyl moiety
TABLE 4 Validation and initial probing of DG70 structure-activity relationships

| Name       | Structure | MIC (µg/ml) | CC₉₀ (µg/ml) |
|------------|-----------|-------------|--------------|
| DG70 (JSF-2911) | ![Structure](image) | 2.4       | >80          |
| JSF-2912   | ![Structure](image) | 9.6       | >80          |
| JSF-2908   | ![Structure](image) | 4.8       | 50           |
| JSF-2909   | ![Structure](image) | >77       | >50          |
| JSF-2951   | ![Structure](image) | 77        | >50          |
| JSF-2949   | ![Structure](image) | 39        | 24           |
| JSF-2950   | ![Structure](image) | 39        | 50           |
| JSF-2910   | ![Structure](image) | 9.6       | >50          |
| JSF-2907   | ![Structure](image) | >77       | 50           |
| JSF-2971   | ![Structure](image) | 9.6       | >50          |
| JSF-2926   | ![Structure](image) | 9.6       | >70          |

SAR was less straightforward. Removal of either the fluoro group (cf. JSF-2908) or both fluoro and chloro groups (cf. JSF-2912) led to small losses in activity, while excision of solely the chloro group (cf. JSF-2909) forfeited all efficacy. The amide carbonyl and amide N-H were also critical to whole-cell efficacy (cf. JSF-2951 and JSF-2949). To confirm MenG specificity, a subset of analogs was tested for MIC against DG70-resistant mutants, and these mutants were resistant to active DG70 analogs (Table S2).

DISCUSSION

We developed a novel whole-cell-based screen to detect inhibitors of the respiratory pathway, using induction of PcydAB as a highly specific reporter strain. This assay allowed us to identify DG70 as a potent inhibitor of both replicating and nonreplicating mycobacteria as well as drug-resistant *M. tuberculosis*. Our studies strongly suggest that the primary mechanism of action of DG70 is through inhibition of *M. tuberculosis* MenG, an essential enzyme involved in a critical terminal step of menaquinone biosynthesis. We isolated DG70-resistant mutants with SNPs in *menG*. Homology modeling and protein threading placed most of the mutations associated with resistance in close proximity either to the MenG binding site for *S*-adenosylmethionine, a cofactor used for methylation of DMK9 by MenG, or to the substrate site. Moreover, overexpression of mutant *menG* conferred high-level resistance on otherwise DG70-susceptible BCG strains. Finally, a combination of metabolic labeling and HRMS studies conclusively showed that DG70 inhibited the metabolic step catalyzed by MenG. DG70 treatment inhibited oxygen utilization in sealed mycobacterial culture tubes and significantly...
reduced ATP levels inside treated mycobacterial cells, further supporting the role of 
DG70 in inhibiting the respiratory chain.

Although earlier work has validated menaquinone biosynthesis as a druggable 
target in M. tuberculosis, DG70 is the first chemotype that specifically inhibits MenG (16, 
29, 33), and our study is the first to show that this inhibition is highly bactericidal in M. 
tuberculosis persisters. DG70 showed several logs more activity against persisters than 
the currently licensed anti-TB drugs INH, RIF, and BDQ, and activity was also 80-fold 
higher than that of PA824. The increased effectiveness of DG70 in nonreplicating 
mycobacteria might be due to a lack of energy-generating pathways other than the 
respiratory chain in the nongrowing persistent mycobacteria (34). Menaquinone is 
known to be downregulated in persisters (16, 28), which could also result in heightened 
susceptibility to MenG inhibition. Although our data are promising and suggest that 
DG70 effectively kills nongrowing M. tuberculosis, it is important to note that the model 
that we used to measure killing in stationary phase requires culturability when the 
cultures are reexposed to conditions favorable for log-phase growth. Therefore, we 
cannot definitively determine whether the treated M. tuberculosis bacteria were killed 
as persisters under nongrowing conditions or whether they subsequently died when 
placed under growth-promoting conditions.

This work is also the first to note the highly desirable synergies between menaqui-
none biosynthesis inhibitor and several important anti-TB drugs. DG70 and BDQ 
displayed almost identical killing curves in vitro. Both compounds showed delayed 
onset of bactericidal activity with less than 1-log10 killing in 4 days. However, they 
proved to be highly bactericidal, leading to a 5-log10 drop in CFU over the course of 
21 days. Our results are consistent with previous findings that compounds which target 
the mycobacterial respiratory chain exhibit a delayed onset of bactericidal activity (23). 
Interestingly, combinations of DG70 and BDQ showed significantly better killing than 
did either compound by itself and completely sterilized cultures in 21 days. Further-
more, DG70 combined with either INH or PA824 was able to sterilize cultures in 10 days. 
This is particularly remarkable because it suggests that treatment with two drugs that 
target different enzymes can overcome both the persistence and the drug resistance 
that develop in the presence of single drug.

The activity and physiochemical and metabolic stability profiling of DG70 and its 
analogs support this biphenyl benzamide as a validated antitubercular hit. However, 
this series must be modified for improved metabolic stability, beginning by improving 
its stability in mouse liver microsomes. The identification of an inactive bis(phenol) 
metabolite and DG70 SAR and computational docking studies suggest an optimization 
route toward candidates for in vivo efficacy assessment. If developed into a drug, DG70 
or another inhibitor of MenG with similar biological effects could become a highly 
effective component of TB therapy, with activity against both replicating and nonrep-
licating mycobacteria and considerable synergy with other anti-TB drugs.

MATERIALS AND METHODS

  **Bacterial strains.** M. tuberculosis H37Rv (ATCC 27294) and BCG (ATCC 35734) were used as reference 
  strains. An avirulent ΔpanCD ΔleuCD derivative of H37Rv (mc²6206) was a kind gift of William R. Jacobs, 
  Jr., at the Albert Einstein College of Medicine, NY. The clinical M. tuberculosis strains tested were selected 
  from collections of clinical isolates established by UNICEF, UNDP, the World Bank, and WHO Special 
  Programs for Research and Training in Tropical Diseases (TDR). NTM and ESKAPE species and strains were 
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  from collections of clinical isolates established by UNICEF, UNDP, the World Bank, and WHO Special 

  **Chemical and compound library.** A library of 168 structurally diverse small molecules with known 
  whole-cell anti-M. tuberculosis activity was obtained from GlaxoSmithKline (GSK) (17). Antibiotics and 
  MK4 were purchased from Sigma-Aldrich. Stock solutions were prepared in dimethyl sulfoxide (DMSO), 

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and working stocks were diluted in the medium to 100 μM final concentrations. The MK9 standard was purchased from Santa Cruz Biotechnology.

**Construction of the CAW306 screening system.** The mWasabi gene was cloned into pMV306, a promoter-absent integrative vector, between HindIII and EcoRI restriction sites to generate pMV306mWasabi. The 489-bp sequence upstream of the translation start codon of the cydA gene was amplified from M. tuberculosis H37Rv genomic DNA using primers containing KpnI and HindIII restriction enzyme sites in the forward and reverse primers, respectively. The primer sequences were as follows: FP, 5′-GGCGTAGCTTGTGTCATCACG 3′, and RP, 5′-GACAGCTTACATGTGCTACATCCC 3′. The amplified product was then cloned between the KpnI and HindIII sites of pMV306mWasabi to generate the pMV306pCydnmWasabi construct. After confirmation by DNA sequencing, the construct was electroporated into BCG and selected on 7H10 plates containing hygromycin B (50 μg/ml) to create the recombinant BCG strain CAW306. For screening, CAW306 was cultured to an optical density at 595 nm (OD<sub>595</sub>) of 0.2 in 7H9 medium supplemented with OADC and 0.02% Tween 80. This culture was then dispensed into 96-well black clear-bottom half-area plates (Corning) at 90 μl/well. Compounds from the test library were then added to plate wells in 10-μl aliquots for a final 10 μM test concentration. Positive and negative controls were included in each plate in triplicate. Plates were incubated for 24 h at 37°C, and the fluorescence of each well was recorded at an excitation (Ex) wavelength of 488 nm and an emission (Em) wavelength of 509 nm in a SpectraMax M5 microplate reader. Compounds were scored based on fold induction over vehicle. The library was screened in singlets due to limitations in the amount of compound available; however, each positive hit was tested in triplicate and then resynthesized or synthesized. Z<sup>r</sup> factor values were calculated for each plate using the mean and standard deviation of both positive and negative controls. The results from plates with a Z<sup>r</sup> factor of ≥0.7 were deemed technically acceptable.

**MIC determination.** MICs of M. tuberculosis, BCG, NTM, and other bacterial isolates were measured in 96-well microtiter plates using a microdilution alamarBlue assay (MABA) as described previously (18). Briefly, bacterial cells grown to mid-log phase were diluted (1:100) in medium and 50 μl of this dilution was added to each well in 11 2-fold serial dilutions of test compound in the medium and then incubated for 7 days at 37°C. alamarBlue (Invitrogen) reagent was added along with 12.5 μl of 20% Tween 80 (Sigma) to evaluate bacterial cell viability. Plates were scanned after 24 h at 570-nm absorbance with a reference wavelength of 600 nm.

**Isolation of resistant mutants and their genetic characterization.** Mid-log-phase M. tuberculosis cultures (10<sup>6</sup> CFU/ml) were plated on 7H10 agar containing DG70 at 4× and 8× MIC and incubated for 3 to 4 weeks at 37°C. Isolated colonies were purified and grown in liquid culture and subjected to MIC determination. Genomic DNA was isolated from confirmed resistant mutants and subjected to whole-genome sequencing using Illumina NextSeq at >100× coverage. Variant calling was performed by CLC Genomic Workbench 7.5 using the H37Rv reference NC_000962.3 and the sequenced laboratory parent H37Rv strain as references (18).

**Construction of overexpression strains.** The menG gene was PCR amplified from the wild type as well as the DG70-resistant mutant of M. tuberculosis using primers FP (5′-agggatctagctgccgcgcgc 3′) and RP (5′-tccagctttcactgcggggtttgtttgc 3′) and cloned into pMV306H-hsp60 (18) between NdeI and HindIII. The sequence-verified plasmids were transformed into BCG to create strains BCG:pMV306PHsp60, BCG:pMV306Hsp60-menG_Wt, BCG:pMV306Hsp60-menG_F118L, and BCG:pMV306Hsp60-menG_V20A.

**Cytotoxicity testing.** Vero cells (ATCC CCL-81) were used to test cytotoxicity as described previously (18). Briefly, Vero cells were plated at a concentration of 10<sup>3</sup> cells/well in 96-well plates and incubated for 2 to 3 h to allow the cells to settle. Test compounds were serially diluted separately in Eagle’s minimal essential medium to generate test concentrations ranging from 200 to 0.1× M. tuberculosis MIC. The serial dilutions were then added to the plated cells and incubated for 48 h at 37°C. The viability of Vero cells exposed to each compound was determined using an MTT [3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide] cell viability kit (Promega).

**Determination of compound interactions by checkerboard assay.** We used a checkerboard assay to test for interactions between DG70 and drugs with known anti-TB activity using alamarBlue as a viability marker (35). Drug combinations were categorized based on a FIC of ≤0.5 as synergism, a FIC of >0.5 and ≤1 as additivity, a FIC of >1 and <4 as no interaction, and a FIC of ≥4 as antagonism (36).

**Determining bactericidal concentration in a nutritionally deprived persistence model.** Exponentially growing cultures of M. tuberculosis at an OD<sub>600</sub> of 0.4 to 0.6 were harvested by centrifugation (3,000 × g, 10 min), washed twice with PBS (Invitrogen) supplemented with 0.025% Tween 80, and incubated in this nutritionally deprived medium for 6 weeks at 37°C. The bactericidal activity of each compound and compound combination against the resulting nongrowing bacilli was determined by exposing each culture to compounds for 7 days at 37°C and then measuring CFU by plating appropriate dilutions in 7H9 medium on 7H11 agar plates (37).

**Twenty-one-day kill kinetics.** Mid-log-phase BCG cultures were diluted in fresh medium to an OD<sub>600</sub> of 0.1 to 0.2. Compounds were then added to culture aliquots at 10× MIC of each compound. Dilutions of each culture were plated at selected intervals on 7H10 agar plates to determine CFU.

**Intracellular activity in J774A.1 murine macrophages.** Intracellular activity was evaluated in J774A.1 (ATCC 113-67) cells as described previously (18). Briefly, J774A.1 cells were maintained in Dulbecco’s minimal essential medium (DMEM) supplemented with 10% heat-inactivated fetal bovine serum (FBS). The cells were placed into 96-well tissue culture white clear-bottom plates (Corning) in a 100-μl volume containing 2 × 10<sup>6</sup> cells per well and allowed to adhere overnight. The avirulent ΔapnCD ΔeluCD M. tuberculosis strain mc<sup>6206</sup> was cultured to mid-log phase; diluted in DMEM supplemented with FBS, pantothenic acid, and leucine; and added to the adherent macrophages at 10<sup>5</sup> cells/well. The
assay plates were then incubated for 4 h to allow bacteria to infect the macrophages. The remaining extracellular bacteria were then removed by replacing the medium with 100 μl of DMEM containing 50 μg/ml of gentamicin and incubating it for 1 h. The infected cells were then washed twice with 100 μl of DMEM supplemented with FBS, pantothenic acid, and leucine, and the final wash was replaced with 100 μl of complete DMEM containing the indicated concentrations of each test compound. Each compound concentration was tested in triplicates. On day 3, cells were washed with DMEM, lysed with distilled water (dH2O), serially diluted, and plated for CFU.

**ATP determination.** BCG was grown to an OD595 of 0.3. DG70 and other control drugs were added at selected concentrations. The cultures were then incubated at 37°C for 3 days. Intracellular ATP was quantified using a BacTiter Glo microbial cell viability assay kit (Promega) as previously described (23). ATP standards ranging from 0.1 to 10,000 nM were also included.

**Oxygen consumption.** The effect of DG70 on *M. tuberculosis* oxygen consumption was tested using a methylene blue decolorization assay as previously described (19). Decolorization was monitored by measuring absorbance at 6, 12, and 24 h at 665 nm. In the rescue experiment, the indicated concentration of MK4 was added prior to the inhibitor.

**Menadione analysis by whole-cell labeling.** Five-milliliter cultures of BCG were grown to an OD595 of 0.6, pelleted, and resuspended in 1/10 fresh medium, and 0.5-ml subsets were incubated with compounds for 1 h prior to the addition of 5 μCi L-[methyl-14C]methionine (American Radiolabeled Chemical) and further incubated for 4 h. Neutral lipids were then extracted (16), dissolved in hexane, applied to silica gel TLC plates, and developed using hexane-diethyl ether (95:5, vol/vol).

**Identification of menaquinone.** Ten-milliliter cultures of BCG were grown in 7H9 medium containing 0.05% Tween to mid-log phase and treated with compounds for 5 generations (3 days). Neutral lipids were then extracted as described above and analyzed by high-resolution mass spectrometry on a Bruker Daltonics mass spectrometer with atmospheric pressure chemical ionization as the ionization interface (16).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02022-16.

**TEXT S1,** DOCX file, 0.1 MB.

**TEXT S2,** DOCX file, 0.04 MB.

**FIG S1,** DOCX file, 0.1 MB.

**FIG S2,** DOCX file, 1.2 MB.

**FIG S3,** DOCX file, 2.8 MB.

**FIG S4,** DOCX file, 0.8 MB.

**FIG S5,** DOCX file, 0.3 MB.

**TABLE S1,** DOCX file, 0.01 MB.

**TABLE S2,** DOCX file, 0.01 MB.

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