Construction of an overexpression library for *Mycobacterium tuberculosis*

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Abstract

There is a pressing need to develop novel anti-tubercular drugs. High-throughput phenotypic screening yields chemical series that inhibit bacterial growth. Target identification for such series is challenging, but necessary for optimization of target engagement and the development of series into clinical drugs. We constructed a library of recombinant *Mycobacterium tuberculosis* strains each expressing a single protein from an inducible promoter as a tool for target identification. The library of 1733 clones was arrayed in 96-well plates for rapid screening and monitoring growth. The library contains the majority of the annotated essential genes as well as genes involved in cell wall and fatty acid biosynthesis, virulence factors, regulatory proteins, efflux, and respiration pathways. We evaluated the growth kinetics and plasmid stability over three passages for each clone in the library. We determined expression levels (mRNA and/or protein) in 396 selected clones. We screened the entire library and identified the Alr-expressing clone as the only recombinant strain, which grew in the presence of D-cycloserine (DCS). We confirmed that the Alr-expressing clone was resistant to DCS (7-fold shift in minimum inhibitory concentration). The library represents a new tool that can be used to screen for compound resistance and other phenotypes.

Introduction

One-quarter of the global population is infected with *Mycobacterium tuberculosis*, the causative agent of tuberculosis (TB) [1, 2]. Due to the increasing prevalence of drug-resistant tuberculosis and an ageing arsenal of anti-tuberculosis drugs, there is a pressing and continuing need to develop novel, well-tolerated anti-tubercular drugs. Current TB drug discovery has focused on phenotypic screens against chemical libraries to identify compound scaffolds that inhibit bacterial growth [3, 4]. Treated as series, these compounds are optimized to minimize mammalian cytotoxicity while maintaining antibacterial activity and improving bioavailability. Identification of the target or metabolic pathway is crucial to drug discovery and development to improve on-target activity and specificity.

While genetic and biochemical methods have been applied successfully to determine intracellular targets in *M. tuberculosis* [5–16], identification of a target is often difficult and requires multiple approaches. For example, the isolation of resistant mutants may lead to target identification in about 20% of cases [5, 14]. Additional methods for target identification will therefore increase the potential to identify relevant targets or metabolic pathways, enhancing drug development. Target overexpression

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conferring resistance to antimycobacterial compounds has been used as a diagnostic or confirmatory assay once the target has been identified; in these cases, the target was previously predicted by other methods and overexpression was used to confirm the target [12, 17–22].

Overexpression libraries have been used in Schizosaccharomyces and Saccharomyces, covering 90 and 95% of each species ORF, respectively [23, 24]. In Escherichia coli, heterologous expression of genomic libraries from Piriformospora indica and Marinobacter aquaeoli identified genes conferring salt and terpene tolerance, respectively [25, 26]. Likewise, an inducibly expressed library of E. coli transcription factors was used to identify factors that mediated beta-lactam sensitivity in antibiotic-resistant E. coli [27]. In M. tuberculosis, systematic overexpression of transcription factors was used to derive a transcription factor interaction network [28].

In order to expand the methods available for target identification, we constructed a library of M. tuberculosis strains using an inducible expression system in which each clone expresses a single M. tuberculosis protein. We evaluated each clone for growth under induced and noninduced conditions and determined plasmid stability over several passages. We also monitored expression, either by protein or mRNA levels, for a subset of clones. The library was arrayed in 96-well plates for ease of use. We confirmed the functionality of the system by screening the library for resistance to D-cycloserine (DCS) and confirmed that the clone expressing alr was selected.

Materials and methods

Mycobacterium tuberculosis culture

Mycobacterium tuberculosis was grown in Middlebrook 7H9 medium supplemented with 0.05% w/v Tween 80 and 10% v/v oleic acid, albumen, dextrose, catalase (OADC) supplement (Becton Dickinson), or on Middlebrook 7H10 agar with 10% v/v OADC. Hygromycin (Hyg) was added to 100 µg/ml and anhydrotetracycline (ATc) to 150 ng/ml where required. For long-term storage, an equal volume of 50% w/v glycerol was added to 96-well plates and stored at -80°C.

Library construction

The majority of the clones were constructed by polymerase chain reaction (PCR) amplification from H37Rv genomic deoxyribonucleic acid (DNA) using oligonucleotides incorporating the Gateway recombination sequences; products were cloned into pDONR221 using BP clonase, and subsequently into pDTNF and/or pDTCF vectors using LR clonase (Thermo Fisher). This set was augmented with Gateway Entry clones received from the Pathogen Functional Genomics Resource Center (J. Craig Venter Institute); for these genes, expression vectors were generated by Gateway cloning into pDTNF and pDTCF expression vectors [28]. Plasmids were electroporated into M. tuberculosis H37Rv [29]. Recombinant strains were grown in liquid medium and arrayed into 96-well plates.

Evaluation of library growth

Growth was measured in 96-well plates as follows; 10 µl of culture was inoculated into 90 µl 7H9-OADC-Tw-Hyg100 ± ATc in a 96-well plate and incubated at 37°C for 7 days. Growth was measured at OD590 and the growth ratio of induced to uninduced was calculated.

Determination of plasmid stability

Cultures were passaged three times in 96-well plates by inoculating 10 µl into 90 µl 7H9-OADC-Tw plus Hyg and ATc for 7 days. Cell lysates were generated from 96-well plates. Plates were placed on a heating block at 100°C for 10 min; cultures were filtered through a 96-well, 0.2 µm filter plate (Millipore) by centrifugation at 4000 rpm and collected into fresh 96-well plates. PCR amplification of the gene inserts was carried out using primers Walk-F1: 5’ GTGAGAAAGGGTCTCTGAGCAC 3’ and pDTNF-R3: 5’ CTCAGACTCGAGGTATCG 3’. Control PCR using primers to amplify the hyg gene was carried out using primers HygF2: 5’ GAAGCTGCGCGTTTCTCCG 3’ and HygR2: 5’ CTGACGGGGA ACACGGTGCTC 3’.

Determination of overexpression

Ten strains were selected at random from each library plate and inoculated into an OD590 of 0.05 in 5 ml 7H9-Tw-OADC plus Hyg and ATc (where indicated) in three 16 mm borosilicate tubes containing stirrer bars. Growth was monitored for 7 days by OD590. The three cultures were pooled and bacteria harvested by centrifugation. Bacterial pellets were washed in 10 mM Tris pH 8.0, 0.05% Tween 80 and resuspended in 10 mM Tris pH 8.0. Cell-free extracts were generated by lysis using lysing matrix B and a Fastprep (MP Biomedical) at speed 6.0 for 30 s. The lysate was centrifuged for 5 min and passed through a 0.20 µm filter. For Western analysis, 60 µg of protein was loaded and electrophoresed on a 4–12% Bis-Tris gradient gel (Invitrogen) and transferred to PVDF membrane using the iBlot system (Invitrogen). Proteins were detected using an anti-FLAG rabbit IgG primary antibody (Genscript), HRP conjugated goat anti-rabbit IgG secondary antibody and detected using SuperSignal™ West Femto Maximum Sensitivity Substrate (Thermo-Fisher). An amino-terminal FLAG-BAP (Sigma-Aldrich) fusion protein was used as the positive control.

To determine mRNA levels, total RNA was extracted as described [30]. cDNA was synthesized using the Transcriptor High Fidelity cDNA Synthesis Kit (Roche) and random hexamers. Quantitative PCR was carried out with primer/probe combinations for each gene (Supplementary Table S1) on a Roche LightCycler 480 in duplicate. Samples were also evaluated in duplicate for sigA expression. Reactions were formulated in LightCycler 480 Multiwell 384 well plates (Roche) containing 10 µl 2x LightCycler 480 Probes Master (Roche), 7.2 µl of 5 µM primer mix, 0.2 µl of 25 µM probe, 2 µl of cDNA, and 0.6 µl PCR grade water. Samples were cycled as follows; 95°C for 10 min initial denaturation, 45 cycles of 95°C for 10 s denaturation, 56°C for 1 min amplification, and 72°C for 1 s extension. To determine copy number, genomic DNA was used to generate a standard curve.

Library screen

M. tuberculosis clones were cultured in in 100 µl 7H9-Tw-OADC-Hyg in 96-well plates for 8 days. Clones were sub-cultured into 96-well plates containing 100 µl 7H9-Tw-OADC ± ATc and incubated for 6 days (mid-log phase). Assay plates were inoculated with 10 µl culture into medium ± ATc, ± 150 µM DCS (four combinations). Plates were incubated for 5 days and growth measured at OD590. The growth ratio was calculated from the OD590 at Day 5 compared to Day 0.
Determination of minimum inhibitory concentration

Minimum inhibitory concentrations (MICs) were determined in liquid culture by measuring OD₅₉₀ as described [31]. Briefly, a 2-fold dilution series of compound was prepared in medium in 96-well plates. Each plate included controls for background (medium/DMSO only, no bacterial cells), zero growth (100 µM rifampicin) and maximum growth (2% DMSO), as well as a rifampicin dose response curve. Cultures were inoculated to an OD₅₉₀ of 0.05 and incubated for 5 days before measuring growth. A dose response curve was plotted as % growth and fitted to the Gompertz model using GraphPad Prism 6. MIC was defined as the minimum concentration at which growth was completely inhibited and was calculated from the inflection point of the fitted curve to the lower asymptote (zero growth).

Results

Construction of a panel of expression plasmids for M. tuberculosis

Since overexpression of a drug target often leads to resistance, we wanted to construct a library of overexpression strains that could be used to identify the targets, or mechanisms of resistance, to novel chemical agents. Our overall aim was to use such a panel to support phenotypic screening. We aimed to construct a library of M. tuberculosis strains, each overexpressing a single gene (or operon in a few cases) which could be screened rapidly for growth in a high-throughput format. We decided to use an inducible expression system in order to minimize any issues arising from toxic effects of overexpression of individual genes. We used the expression plasmids pDTNF or pDTCF in which the gene is cloned downstream of a tetracycline-inducible promoter with either an N-terminal or C-terminal FLAG tag fused to the protein in a shuttle vector (Fig. 1) [28].

The M. tuberculosis genome has ~4000 annotated ORFs [32]. We decided to first construct an expression panel containing the in vitro essential genes identified by Sassetti et al. [33]. We constructed expression constructs for the majority of the essential genes. Donor plasmids were obtained either from the PFGRC library or PCR-amplified and cloned into a Gateway donor vector. Final expression plasmids were generated using Gateway cloning technology. A proportion of the essential genes (87) were cloned into both vectors to generate protein fusions with either N-terminal or C-terminal FLAG-tagged proteins. We were able to construct the majority of the expression plasmids with 542/614 essential genes represented in our set. Once we had constructed the essential set, we generated a second set of plasmids representing non-essential genes. These covered genes involved in lipid metabolism, cell wall synthesis, transcriptional/translational regulation, respiration, virulence, and efflux.

Plasmids were transformed into M. tuberculosis and recombinants selected on hygromycin-containing plates. Transformants for each plasmid were cultured in liquid medium and then transferred into 96-well plates and numbered (Table 1). The library was organized in several groups containing a number of plates termed POETs = Plate Of Over-Expressing TB and organized into smaller panels. The first panel, (POET 21-30) contained the majority of the essential genes. Panel 2 (POET 31-36) contained expression clones from genes involved in lipid metabolism along with other pathways involved in virulence, detoxification, and adaptation. Non-essential genes involved in metabolism and respiration were arranged in POET 37-41. The full library consists of a total of 1733 recombinant strains of M. tuberculosis generated from 1725 plasmids (a small number of plasmids were transformed more than once) each carrying a plasmid overexpressing a single gene or operon (Supplementary Table S2).

Determination of growth kinetics for each clone

We anticipated that overexpression of certain proteins might have a negative effect on bacterial growth; therefore we determined the growth rate for every clone in the presence and absence of ATc (induced and non-induced). Strains were cultured in 96-well plates and growth measured at Day 0 and Day 5 (Table 2 and Fig. 2). For each strain we calculated the ratio of growth under induced/uninduced conditions, since we expected that overexpression would be most likely to show a toxic effect. The majority of strains (97%) showed a growth rate comparable to the wild-type strain; using a cutoff of 1.5-fold change only 55 strains showed reduced growth (Supplementary Table S3). Only two strains showed increased growth, and that was only ~1.5-fold over the wild-type strains (Supplementary Table S3).

Figure 1: Vectors used to generate expression plasmids. pDTNF and pDTCF contain mycobacterial (MycORI) and E.coli replication origins (ori), hygromycin resistance, and an inducible promoter (Uv15tet0). Gene inserts are cloned into the attR1 and attR2 sites thereby eliminating the chloramphenicol and ccdB toxin selection markers with FLAG tags either N-terminal or C-terminal to the inserted gene.
Table 1: Composition of the overexpression library

| Category                  | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 | 31 | 32 | 33 | 34 | 35 | 36 | 37 | 38 | 39 | 40 | 41 | Total |
|---------------------------|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|-------|
| Insertion sequences and phages | 2  | 0  | 1  | 0  | 0  | 0  | 3  | 0  | 1  | 0  | 0  | 0  | 1  | 0  | 0  | 1  | 0  | 0  | 0  | 0  | 0  | 8    |
| Intermediary metabolism and respiration | 31 | 38 | 45 | 21 | 31 | 14 | 14 | 18 | 29 | 0  | 3  | 4  | 4  | 8  | 11 | 88 | 85 | 87 | 88 | 88 | 707  |
| Information pathways       | 17 | 8  | 10 | 8  | 25 | 49 | 44 | 42 | 48 | 20 | 4  | 0  | 0  | 0  | 3  | 6  | 0  | 0  | 0  | 0  | 0  | 240  |
| Cell wall and cell processes | 12 | 22 | 12 | 21 | 14 | 9  | 12 | 11 | 14 | 1  | 3  | 4  | 1  | 5  | 6  | 0  | 1  | 0  | 0  | 0  | 0  | 148  |
| Conserved hypothetical     | 17 | 6  | 8  | 2  | 8  | 8  | 7  | 3  | 8  | 1  | 2  | 0  | 2  | 1  | 2  | 0  | 2  | 0  | 0  | 0  | 77   |
| Virulence, detoxification, adaptation | 3  | 6  | 3  | 1  | 2  | 1  | 1  | 3  | 2  | 3  | 3  | 23 | 23 | 25 | 24 | 29 | 22 | 0  | 0  | 0  | 0  | 188  |
| Lipid metabolism           | 3  | 7  | 8  | 7  | 3  | 2  | 5  | 1  | 7  | 26 | 31 | 24 | 38 | 22 | 20 | 0  | 0  | 0  | 0  | 0  | 0  | 204  |
| PE/PPE                    | 1  | 0  | 0  | 1  | 1  | 2  | 1  | 0  | 3  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 9    |
| Regulatory proteins        | 2  | 1  | 1  | 1  | 4  | 3  | 3  | 4  | 23 | 26 | 21 | 18 | 20 | 21 | 0  | 0  | 0  | 0  | 0  | 0  | 152  |
| Total                     | 88 | 88 | 88 | 62 | 88 | 88 | 88 | 88 | 88 | 88 | 88 | 88 | 88 | 87 | 88 | 88 | 1733|

Gene classifications were taken from Tuberculist (now Mycobrowser) https://mycobrowser.epfl.ch. The composition of each plate (POET 21–41) is given.

Table 2: Growth of overexpression library strains

| Method                  | 96-well plate | 5 ml culture |
|-------------------------|---------------|--------------|
| Strains evaluated       | 1733          | 155          |
| Increased growth        | 2             | 16           |
| Normal growth           | 1676          | 130          |
| Reduced growth          | 55            | 9            |

Strains were grown in 5 ml stirred cultures or in 96-well plates ± 150 ng/ml ATc for 7 days and growth measured by OD$_{590}$. Increased growth is defined as >1.5-fold growth in the presence of ATc. Reduced growth is defined as <0.5-fold growth in the presence ATc. Normal growth is defined 0.5- to 1.5-fold growth in the presence of ATc.

Determination of plasmid stability

A small fraction of strains had reduced growth, which we assume is due to toxicity from the expressed protein. Plasmid instability in *M. tuberculosis* has been seen, and is related to toxicity of protein expression [34]. Therefore, we expected some plasmids might be unstable due to protein toxicity [35–37], and the fact that the inducible system has leaky expression [28, 36].

We evaluated plasmid stability over three passages under selective pressure under both induced and uninduced conditions (Table 3 and Fig. 3). Strains were passaged three times and then tested for the presence of the correct insert size in the plasmid using primers which flanked the gene insert. Plasmids were considered stable where an amplified product of the correct size was obtained. Where no insert was amplified, we used primers for the hygromycin gene to confirm the presence of a plasmid; we were able to amplify the hygromycin resistance cassette for all these strains and they grew in the presence of hygromycin confirming that a plasmid was present. For strains in which the third passage lacked a product of the correct size, we tested the second passage, and if no correct amplicon was obtained, we tested the first passage.

We evaluated the majority of the library and found that loss of the gene from the plasmid occurred only in a small subset of strains (~7%) after three passages. After one passage, only 26 transformants (1.5%) of the entire library did not have a plasmid with the correct sized insert. Thus, we confirmed that for the majority of the library, maintenance of the plasmid even under induced conditions was stable. Of interest, the plasmids showing the most instability contained a number of essential genes, namely embB, hycQ, ctpH, Rv1002c, rocA, hsl, pkS7, f0E, ftsK, fmt, and rplC.

Confirmation of protein expression in *M. tuberculosis* recombinant strains

We wanted to confirm that expression of proteins was being maintained in an ATc-inducible fashion in our recombinant clones. We selected a subset of strains in which we determined either mRNA and/or protein levels. A subset of strains was selected randomly from the library and grown in 5 ml cultures with ATc ranging from 0 to 200 ng/ml in order to determine the expression levels in response to ATc. We also monitored growth over time for each strain. The wild-type strain showed no reduction in growth in the presence of ATc up to 200 ng/ml. The majority of strains we selected showed no significant reduction in growth up to 200 ng/ml ATc (Fig. 4). We generated cell-free extracts from these cultures and measured protein levels by Western blot using an anti-FLAG tag antibody. We also measured mRNA levels for each gene. For proteins, only the plasmid-encoded protein would be detected, whereas for mRNA we determined the total levels from both plasmid and chromosomal copies of the genes. The majority of strains demonstrated overexpression (mRNA and protein) at 100 or 200 ng/ml ATc (Table 4A). Protein expression is dependent on gene codon use, protein sequence and structure, and level of inducer; with some genes requiring high concentrations of inducer to achieve detectable levels of protein. For example, for RpoZ (Rv1390) protein...
expression levels plateaued at 50 ng/ml of ATc (Fig. 5C and D), whereas DnaJ2 (Rv2373c) had maximal expression at 100 ng/ml ATc and robust protein expression was observed even at 10 ng/ml ATc (Fig. 5A and B). For both of these genes, we saw inducer-dependent increases in mRNA levels (Fig. 5C and D) which were similar to protein levels, but they did not correlate exactly, either due to post-transcriptional factors or due to the fact that mRNA measures the total of native and recombinant protein.

Once we had determined that 100–200 ng/ml was optimal for inducing protein expression, we selected a larger set of clones for evaluation. We randomly selected 10 strains from each plate (396 in total) and measured protein and mRNA levels. Since we saw the majority of protein expression detected at 100–200 ng/ml ATC and the strains had no growth defect, we selected 150 ng/ml ATC for subsequent work. We measured protein levels from bacteria cultured in the presence of 150 ng/ml ATC (Table 4B). We detected protein expression in 116 of the 396 strains tested by Western. Since we used cell-free extracts, we would not expect to detect membrane or secreted proteins, which applies to 53 of our clones (selected at random).

Validation screen

We wanted to confirm that the library could be used to identify targets whose overexpression leads to resistance. To establish proof of concept, we selected DCS as our candidate drug. DCS is an established second-line antitubercular compound with known psychoactive properties [38]. DCS prevents proper cell wall construction by interfering with peptidoglycan biosynthesis; specifically the production of a mycolyl–arabinogalactan–peptidoglycan complex [39]. DCS is proposed to have two essential primary in cell targets; D-alanine racemase (Rv3423c—alr) and D-alanine: D-alanine ligase (Rv2981c—ddlA) and overexpression of these genes in M. smegmatis confers resistance to DCS [40, 41].

A section of the library containing 326 clones (POET 21-24) was grown in 96-well plates under uninduced and induced conditions and 65X MIC (150 μM) DCS (MIC = 31 ± 0.4 μM). Growth was measured after 5 days (Fig. 6A and B). In the absence of DCS, robust growth was measured, with an average OD590 of 0.34 ± 0.044 (no ATc) and 0.31 ± 0.057 (plus ATc). In contrast, in the presence of DCS, minimal growth was seen, with an average OD590 of 0.022 ± 0.008 (no ATc) and 0.017 ± 0.006 (plus ATc). A single strain grew in the presence of DCS reaching an OD590 of 0.37 (plus ATc). This strain also grew in the absence of ATc induction (OD590 = 0.38) (Fig. 6A). The strain exhibiting robust growth contained the alanine racemase gene (Rv3423c).

We monitored the kinetics of growth in the presence of DCS for the entire 96-well plate containing the alr expression strain. Growth of the air strain alone was observable as early as Day 3 in induced conditions. By Day 5, growth was observed for both induced and uninduced (Fig. 6B). This confirmed both that expression of Alr was leading to resistance, but also that ATc induction of expression was effective. In order to confirm resistance; we generated a fresh transformant with the air expression plasmid and determined the MIC. The strain
**Figure 4:** Growth of RpoZ and DnaJ2 expression strains. *M. tuberculosis* recombinant strains expressing DnaJ2 or RpoZ gene were cultured in 5 ml medium plus increasing concentrations of ATc (0–200 ng/ml). Growth was measured over time. (A) Wild type (no vector). (B) Rv2373c (DnaJ2). (C) Empty vector. (D) Rv1390 (RpoZ). Data are the mean ± SD of three independent cultures.

**Figure 5:** Expression of recombinant proteins in *M. tuberculosis*. *M. tuberculosis* recombinant strains expressing DnaJ2 (Rv2373c) or RpoZ (Rv1390) were cultured in 5 ml medium plus increasing concentrations of ATc (0–200 ng/ml). (A and C) Western blot using α-FLAG antibody. *M* = ECL Rainbow Marker; *M2* = Novex Sharp Protein Standards; The amino-terminal Flag control - FLAG-BAP fusion protein (2.5–20 ng). (B and D) RNA was isolated from cultures. cDNA was synthesized and subjected to qPCR. Copy number for each genes was determined using a standard curve generated using genomic DNA and normalized to sigA transcripts. Data are the mean ± SD of three independent cultures.
demonstrated significant resistance, with a 6–7-fold shift in MIC from the wild-type (Fig. 6D and F). The presence of the alr gene and overexpression of the protein was confirmed by PCR and Western, respectively confirming that protein expression was evident in both uninduced and induced states (Fig. 6E).

Table 4: Expression from recombinant plasmids in M. tuberculosis

| ATc (ng/ml) | Protein | mRNA |
|------------|---------|------|
| (A) Strains evaluated at variable ATc |
| 0          | 0       | 10   |
| 1          | 2       | 8    |
| 10         | 5       | 10   |
| 50         | 9       | 12   |
| 100        | 11      | 13   |
| 200        | 11      | 15   |
| Total strains tested | 18 | 17  |
| (B) Strains evaluated at single concentration |
| 0          | 9       |      |
| 150        | 108     |      |
| Total strains tested | 396 |      |

Recombinant strains were randomly selected from the library and evaluated for (A) mRNA and protein levels or (B) protein levels only. ATc was added at the concentrations indicated. The number of strains in which protein was detected or in which mRNA levels were increased in a dose-dependent fashion is indicated.

Discussion

As new compound series pass through the drug development pipeline, identification of their biological target(s) and further optimization of those compound series on the target becomes essential. We have constructed a library of M. tuberculosis containing 1733 recombinant strains each overexpressing an individual gene or operon as a complement to existing target identification methods. The library was constructed to prioritize essential genes, as well as genes involved in cell wall and fatty acid biosynthesis, virulence factors, regulatory proteins, efflux, and aerobic and anaerobic respiration metabolic pathways; the library covers ~40% of the protein coding genes in the M. tuberculosis genome [42].

The majority of the library behaved well demonstrating normal growth, stable plasmid maintenance and inducible expression of the required protein. Most of the recombinant strains tested had increased protein expression at 100–200 ng/ml of ATc which is consistent with previous studies using this expression system [28, 36]. A small number of clones were unstable, as determined by loss of the correct plasmid insert on passaging. Overexpression of these genes may be metabolically disruptive or burdensome. These genes, particularly the essential genes, may warrant further investigation as potential drug targets since modulation of their expression levels is deleterious.

When confirming the resistance of the air overexpression from the screen, we noted that Alr was expressed under both the induced and uninduced conditions. Expression in the

Figure 6: Use of the overexpression library to identify the target of a compound. M. tuberculosis recombinant strains were inoculated with 10 μl culture in 90 μl medium ± 150 ng/ml ATc ± 150 μM α-cycloserine. The growth ratio of each strain was calculated as OD590 at Day 5/OD590 at Day 0. (A) No α-cycloserine. (B) 150 μM α-cycloserine C. (C) Growth kinetics of plate containing Alr-expressing recombinant strain. A single 96-well plate containing 80 independent recombinant clones was incubated for 5 days in presence of 150 μM D ± 150 ng/ml ATc. The growth kinetics for all 80 strains are shown. The Alr-expressing strain which demonstrated growth is indicated with arrows (Alr + ATc and Alr–Atc). (D) Determination of MICs for α-cycloserine. MICs were determined in liquid culture by measuring OD590 after 5 days growth at 37 °C using the Gompertz algorithm [31]. (E) Gene integrity within the overexpression plasmid was confirmed by amplification of the gene from cell lysates using plasmid-specific primers followed by sequencing. Protein overexpression was confirmed by anti-FLAG Western.
absence of inducer was also observed in a small proportion of strains evaluated by Western (9 out of 396), consistent with low level basal expression from of this ATc expression system [28, 36]. These genes (alr, dnaJ1, hisF, argF, murD, Rv2215, acpM, topA, Rv0372c) are all annotated as essential and it is possible that the higher expression level reflects low protein degradation and turnover rates.

We screened DCS against the overexpression library as a proof of concept; we identified alr overexpression strain as a target for DCS. Interestingly, the ddaA overexpression strains, present in the portion of the strain library screened, did not show resistance to DCS, suggesting differences between M. smegmatis and M. tuberculosis.

In conclusion, we have constructed a library of 1733 recombinant expression strains of M. tuberculosis for use in drug target identification and to determine the mechanisms of resistance. The library can also be used for growth on alternate media, e.g., solid medium, alternate nutrient sources, restrictive media, under different conditions, e.g., anaerobic, intra-macrophage, and will be a useful method for future drug discovery studies.

Availability
The complete library and individual expression plasmids are available from the authors on request.

Supplementary data
Supplementary data are available at Biology Methods and Protocols online.

Conflict of interest statement. None declared.

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